A quantitative molecular technique was developed for rapid analysis of microbial community diversity in various environments. The technique employed PCR in which one of the two primers used was fluorescently labeled at the 5' end and was used to amplify a selected region of bacterial genes encoding 16S rRNA from total community DNA. The PCR product was digested with restriction enzymes, and the fluorescently labeled terminal restriction fragment was precisely measured by using an automated DNA sequencer. Computer-simulated analysis of terminal restriction fragment length polymorphisms (T-RFLP) for 1,002 eubacterial sequences showed that with proper selection of PCR primers and restriction enzymes, 686 sequences could be PCR amplified and classified into 233 unique terminal restriction fragment lengths or “ribotypes.” Using T-RFLP, we were able to distinguish all bacterial strains in a model bacterial community, and the pattern was consistent with the predicted outcome. Analysis of complex bacterial communities with T-RFLP revealed high species diversity in activated sludge, bioreactor sludge, aquifer sand, and termite guts; as many as 72 unique ribotypes were found in these communities, with 36 ribotypes observed in the termite guts. The community T-RFLP patterns were numerically analyzed and hierarchically clustered. The pattern derived from termite guts was found to be distinctly different from the patterns derived from the other three communities. Overall, our results demonstrated that T-RFLP is a powerful tool for assessing the diversity of complex bacterial communities and for rapidly comparing the community structure and diversity of different ecosystems.

Species richness (the number of species within a community) and species evenness (the sizes of species populations within a community) are two essential parameters for defining community structure and diversity. Quantitation of these two parameters is severely limited by conventional culture-dependent methods by virtue of the fact that large fractions (>85 to 99.999%) of the organisms existing in nature appear to be refractory to cultivation (1, 3, 11, 30, 35). Another limitation of these approaches is that any departure from the original environmental parameters during cultivation can alter the community structure through the imposition of new selective conditions (8). In effect, a new community structure evolves, which may not accurately reflect the original structure.

Several molecular approaches now provide powerful adjuncts to the culture-dependent techniques. One approach in particular that couples PCR and rRNA-based phylogeny has been effective in the exploration of microbial environments and the identification of uncultured organisms. The stepwise strategy of this approach is to isolate total community DNA and use this DNA as a template for PCR amplification of 16S rRNA genes with universal or domain-specific primers. This is usually followed by construction of a clone library for genes encoding rRNA (rDNAs) and rapid screening of the library based on sequence differences (4) or by determining restriction fragment length polymorphisms (RFLPs) of the rDNAs (19). The richness and evenness of a community are qualitatively estimated based on the number of unique clones and the relative frequencies of the various “ribotypes” detected. While this approach obviates the requirement for cultivation, construction and screening of clone libraries are laborious. More recently, techniques dependent on DNA melting behavior (20, 21) or single-strand DNA conformation (14) have been developed as a means of circumventing library construction. These methods have been employed to assess community structure and to provide relatively crude estimates of species diversity, but are limited by the comparatively insensitive staining technologies employed and do not provide information concerning the specific phylogenetic groups that comprise a microbial community.

Here we report on a quantitative molecular method for rapid analysis of complex microbial communities in which an existing technology was extended. RFLP analysis of 16S rDNA, otherwise known as amplified rDNA restriction analysis (ARDRA), has been used for several years as a method for rapid comparison of rDNAs (13, 19). Briefly, rDNAs are obtained by PCR amplification by using universal primers, and the product is digested with restriction enzymes with 4-bp recognition sites. The typical analysis of restriction digests for isolates or clones is performed on relatively low-resolution agarose gels. For community analysis, the potentially large number of fragments can be resolved by using polyacrylamide gels to produce a community-specific pattern (17, 18). However, ARDRA is of limited use for demonstrating the presence of specific phylogenetic groups or for estimating species richness and evenness. In the technique reported on here, the initial steps of DNA isolation, PCR amplification, and restriction were similar to those used for ARDRA. However, one of the primers used was labeled with a fluorescent dye so that when the preparation was analyzed with an automated DNA sequencer, the sizes of only the terminal restriction fragment (T-RF) could be determined and the amount could be quan-
**TABLE 1. Sequences of primers used in this study**

<table>
<thead>
<tr>
<th>Primer</th>
<th>Specificity</th>
<th>Sequence (5’→3’)</th>
<th>Reference(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>8f</td>
<td>Domain</td>
<td>AGAGTTTGATCTGGCTCAG</td>
<td></td>
</tr>
<tr>
<td>341f</td>
<td>Domain</td>
<td>CCTACGGGAGGCAGCA</td>
<td></td>
</tr>
<tr>
<td>536r</td>
<td>Universal</td>
<td>CAGCMGCCGCGATATG</td>
<td></td>
</tr>
<tr>
<td>527f</td>
<td>Universal</td>
<td>ACGCCGCGCTGGTC</td>
<td>This study</td>
</tr>
<tr>
<td>926r</td>
<td>Domain</td>
<td>CGTCAATTCCTTATGTT</td>
<td></td>
</tr>
<tr>
<td>1406r</td>
<td>Universal</td>
<td>AGCCGGCCTGTGRC</td>
<td></td>
</tr>
</tbody>
</table>

**FIG. 1. Primer pairing and target sites used in this study.** The fraction of the database that could be matched by a particular primer pair (values in parentheses) was determined by using pattern-matching software (PatScan). The settings for mismatch/insertion/deletion in PatScan that determine primer specificity were as follows: 0/0/0 at the last four nucleotides from the 3’ end and 3/0/0 at any other nucleotide.

**MATERIALS AND METHODS**

16S rRNA sequence database. The Ribosomal Database Project (RDP) database (release 5.0) contains almost 2,900 aligned 16S rRNA sequences (16). For the analysis of T-RFLPs, a subset of sequences containing at least 1,420 bp (Escherichia coli numbering) was selected from the RDP database; this subset included all of the sequences with fewer than two contiguous unassigned positions and with primer sites complementary to the primers selected. The final subset included chosen 1,007 eubacterial sequences and 95 archaeabacterial sequences.

**Computation of T-RFLP primer specificity and terminal fragment length from the RDP database.** A pattern-matching program, PatScan, developed by R. Overbeek (Argonne National Laboratory, Chicago, Ill.), was used for computing 16S rDNA primer specificity and for predicting the theoretical lengths of T-RFs. Primer specificity was estimated by considering the specificity of forward and reverse primer sequences, the individual primer annealing specificity, and the length of the sequence between the forward and reverse primers. The sequences and annealing sites of the forward and reverse primers considered in this study are shown in Fig. 1 and Table 1. The specificity of each pair of primers was expressed as the percentage of the total number of sequences in the database that were matched. The matched sequences were then used as a subset of the database for computing the lengths of T-RFs that would be obtained after digestion with a given restriction enzyme. The following 10 restriction enzymes were used in this simulation: AciI (CGGTAG), AluI (CCCGG), BglII (GGGAT), BlnIII (GGGCC), EcoRI (GGATCC), HaeIII (GCGG), HinfI (GATC), HpaI (CCCGG), MboI (GGATCC), and TaqI (GCGG).

**Model and natural microbial communities.** The six bacteria in a model microbial community were Alcaligenes faecalis ALCI, Raoultia eutrophus, Arthrobacter globiformis ATCC 8010, Bacillus subtilis BAC11, Escherichia coli HB101, and Pseudomonas aeruginosa ATCC 27853. All of these bacteria were obtained from the American Type Culture Collection, Rockville, Md. (ATCC). The strains were maintained on nutrient agar slants at room temperature and were subcultured every 2 weeks. The bacteria were grown at 37°C in Luria-Bertani medium (pH 7.0) and harvested by centrifugation at 7,000 × g for 10 min. The supernatants were discarded, and the pellets were resuspended in deionized water and serially diluted (1:10) with sterile 0.85% NaCl solution. Serial dilutions were plated onto nutrient agar plates and incubated at 37°C for 24 h. The plates were chosen for the isolation of single colonies, and the purity of cultures was determined by microscopic examination. The bacteria were also cultured in Trypticase soy broth (pH 7.0) at 37°C for 24 h. The cultures were then plated onto nutrient agar plates, and the purity of cultures was determined by microscopic examination. The bacteria were also cultured in Trypticase soy broth (pH 7.0) at 37°C for 24 h. The cultures were then plated onto nutrient agar plates, and the purity of cultures was determined by microscopic examination.

**DNA extraction.** Genomic DNA of each bacterium and total DNA from environmental samples were isolated by using a previously described protocol (27), with minor modifications. Approximately 10 mg (dry weight) of biomass or 5 g of aquifer sand was homogenized with a tissue grinder or a sterilized mortar and pestle in 9.0 ml of lysis buffer (100 mM Tris-HCl, 100 mM NaCl, 0.5% Sarkosyl, and 2.5% SDS). The mixture was then subjected to three freeze-thaw cycles. Hexadecyltrimethyl ammonium bromide and sodium chloride were added to final concentrations of 1.0% (w/v) and 0.7 M, respectively, and the mixtures were incubated for 20 min at 65°C to precipitate polysaccharides and cellulolytic proteins. Each precipitate was removed by extraction with 2 volumes of phenol-chloroform-isomyl alcohol (25:24:1), followed by extraction with chloroform-isomyl alcohol (24:1). DNA was recovered by the addition of 1 volume of isopropanol followed by centrifugation at 10,000 × g for 10 min. The pellet was washed with 70% ethanol, dried, and resuspended in 200 to 400 µl of distilled water. Additional purification, if necessary, was by electrophoresis in a low-melting-point agarose gel (26).

**PCR conditions.** Reaction mixtures for PCR contained 1× PCR buffer, each deoxynucleoside triphosphate at a concentration of 200 µM, 1.5 mM MgCl2, each primer at a concentration of 0.1 µM, and 2.5 U of Taq DNA polymerase (Gibco BRL, Gaithersburg, Md.) in a final volume of 100 µl. The primers used, 8f-Hex and 926r-TET, were labeled at the 5’ end with the phosphoramide dyes 5-hexachlorofluorescein and 5-tetrachlorofluorescein (Operon, Inc., Alameda, Calif.), DNA amplification was performed with a model 9600 thermal cycler (Perkin-Elmer, Norwalk, Conn.) by using the following program: a 3-min hot start at 94°C, followed by 35 cycles consisting of denaturation (30 s at 94°C), annealing (45 s at 56°C), and extension (2 min at 72°C) and a final extension at 72°C for 5 min. Amplified DNA was verified by electrophoresis of aliquots of PCR mixtures (2 µl) in 1% agarose in 1× TAE buffer.

**16S rDNA T-RFLP.** Fluorescently labeled PCR products (100 µl) were purified by using WIZARD PCR purification columns (Promega, Madison, Wis.) and were eluted in a final volume of 50 µl. Aliquots (10 µl) of amplified 16S rDNAs of six bacterial strains were separately digested with HhaI (Boehringer Mannheim Biochemicals [BMB] Indianapolis, Ind.) according to the manufacturer's instructions. Aliquots (5 µl) of each digested product or a mixture of the six digested products were mixed with 2 µl of 5× loading dye buffer and resolved by electrophoresis through an 8% nondenaturing acrylamide gel (acrylamide-N,N′-methylenebisacrylamide, 29:1 [17]). The RFLPs were visualized by staining with SYBR green I dye according to the instructions of the manufacturer (FMC Bioproducts, Rockland, Maine) and then photographed. For natural communities, 6-µl aliquots of purified PCR products were digested with 20 U of HhaI, MspI, Rsal, or HhaI plus MspI (BMB), and the T-RFLP fingerprint of each community was determined as described below.

**16S rDNA T-RFLP.** The precise lengths of the T-RFs from the amplified rDNA products were determined by electrophoresis with a model 373A automated sequencer (Applied Biosystems Instruments [ABI], Foster City, Calif.), as follows. For the model community, 2 µl of digested DNA from each strain or a combination of the digests was mixed with 2 µl of distilled water, 0.5 µl of 5× loading buffer (ABI), and 0.5 µl of DNA fragment length standard (Rox 2500 or Tamara 2500; ABI). The sizes of the DNA fragments in these mixtures were 37, 94, 109, 116, 172, 186, 222, 233, 258, 269, 358, 381, 470, 490, 536, 827, 21, 011, 181, 172, 2208, 2162, 2465, 2481, 2860, 4529, 4771, 5099, and 14,079 bp. This mixture was then denatured at 94°C for 5 min and immediately chilled on ice prior to electrophoresis with the automated DNA sequencer in the GeneScan mode. Aliquots (2.5 µl) of the mixtures were loaded onto a 36-cm 6% denaturing polyacrylamide gel. Electrophoresis was done for up to 20 h with limits of 2,500 V and 40 mA. After electrophoresis, the lengths of fluorescently labeled T-RFs were determined by comparison with internal standards using GeneScan software (ABI).

**Species richness, community signature, and community similarity.** The similarity of communities could be estimated by visual comparison of T-RFLP patterns or by numerically analyzing the pattern of T-RFLPs in gel images with GelCompare software (version 3.1; Applied Maths, Kortrijk, Belgium). Following the normalization (no background subtraction) and image analysis steps of the software, a stacked band pattern of three T-RFLPs from the same community was generated for each community. Two different band-matching coefficients were used to produce a similarity matrix for the pattern fragments in the samples. The Jaccard coefficient considered the presence or absence of bands and the number of T-RFs in common in communities, as well as the total number of T-RFs observed. The area-sensitive coefficient took the last step into account, as well as the difference in the relative abundance (peak area) of each T-RF. The unweighted pair group method using average linkages was applied to cluster patterns to obtain a similarity dendrogram for each coefficient. For both coefficients a value of 0 means that the community fingerprints are completely different from one another, and a value of 1 indicates that they are identical.
TABLE 2. Potential number of 5'- and 3'-T-RF lengths or ribotypes generated for sequences that could be annealed from the 1,102 complete 16S rRNA sequences in the RDP database with four different primer pairs

<table>
<thead>
<tr>
<th>Restriction enzyme</th>
<th>5' end</th>
<th>3' end</th>
<th>5' end</th>
<th>3' end</th>
<th>5' end</th>
<th>3' end</th>
<th>5' end</th>
<th>3' end</th>
</tr>
</thead>
<tbody>
<tr>
<td>AflI</td>
<td>129</td>
<td>92</td>
<td>155</td>
<td>108</td>
<td>129</td>
<td>84</td>
<td>86</td>
<td>56</td>
</tr>
<tr>
<td>BstUI</td>
<td>159</td>
<td>132</td>
<td>173</td>
<td>72</td>
<td>144</td>
<td>69</td>
<td>67</td>
<td>61</td>
</tr>
<tr>
<td>DdeI</td>
<td>180</td>
<td>74</td>
<td>159</td>
<td>109</td>
<td>93</td>
<td>80</td>
<td>84</td>
<td>45</td>
</tr>
<tr>
<td>HaeIII</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>HhaI</td>
<td>136</td>
<td>92</td>
<td>194</td>
<td>127</td>
<td>103</td>
<td>99</td>
<td>87</td>
<td>62</td>
</tr>
<tr>
<td>HinII</td>
<td>124</td>
<td>73</td>
<td>158</td>
<td>105</td>
<td>75</td>
<td>82</td>
<td>111</td>
<td>20</td>
</tr>
<tr>
<td>MboI</td>
<td>170</td>
<td>126</td>
<td>129</td>
<td>105</td>
<td>100</td>
<td>93</td>
<td>138</td>
<td>76</td>
</tr>
<tr>
<td>MspI</td>
<td>133</td>
<td>35</td>
<td>194</td>
<td>27</td>
<td>106</td>
<td>32</td>
<td>72</td>
<td>79</td>
</tr>
<tr>
<td>RsaI</td>
<td>18</td>
<td>140</td>
<td>36</td>
<td>113</td>
<td>110</td>
<td>83</td>
<td>73</td>
<td>66</td>
</tr>
</tbody>
</table>

a The numbers of sequences which could be annealed from the 1,102 complete 16S rRNA sequences with primer sets 8f-536r, 8f-926r, 341f-926r, and 527f-1406r were 587, 686, 931, and 772.

b —, a restriction enzyme site was found in the primer sequence.

RESULTS AND DISCUSSION

T-RFLP in the 16S rRNA database. The ability of T-RFLP analysis to distinguish phylogenetic groups of bacteria was theoretically evaluated through a computer simulation of the T-RF size distribution for 1,102 complete bacterial sequences deposited in the RDP database. Four different pairs of domain-specific or universal PCR primers were selected to anneal to as many 16S rDNA sequences in the RDP database as possible. After digestion with any of 10 different restriction enzymes, the sizes of the terminal fragments amplified with the PCR primers were determined. The combination of PCR primers and enzyme(s) that gave the largest number of terminal fragments with unique sizes because of the high conservation of restriction site positions.

Based on the simulations described above, we concluded that amplification with primer pair 8f-926r followed by digestion with HhaI or MspI resulted in the production of 194 or 233 5'T-RFs, respectively, from the 16S rDNAs amplified (Fig. 2). The number of these 16S rDNAs with HhaI restriction sites in common ranged from 0 (unique to 23. In the most extreme case, 23 species belonging to four genera of the Bacteroides group and the genus Campylobacter and 23 species belonging to 12 different genera of the Neisseria group (4 genera) and the Vibrio group (8 genera) had T-RFs that were 102 and 374 bp long, respectively. As a consequence of the variable conservation of restriction site positions in 16S rDNA, the resolution of T-RFLP analysis is often reduced from the level of species (unique restriction site position in each species) to that of higher-order groups. This effectively genes from phylogenetically related organisms can result in the production of T-RFs of identical size (2, 31). For example, amplification with primer pair 8f-926r followed by digestion with either HhaI or HhaI plus MspI resulted in the production of 194 or 233 5'T-RFs, respectively, from the 16S rDNAs amplified (Fig. 2).
reduces the complexity of the profile of T-RFs obtained from a community so that estimates of diversity within communities can readily be made.

**Analysis of species diversity in a model bacterial community by T-RFLP.** The computer-simulated analysis suggested that polymorphisms among T-RF lengths could be used to characterize microbial communities. To test this approach, a model community containing six known bacterial strains was constructed. PCR amplification in which primers 8f and 926r were used. The terminal location of the fluorescent tag on the PCR products ensured that only terminal fragments were measured. The predicted outcome based on a computer-simulated restriction analysis of sequences from the RDP database indicated that the six strains used should have different RFLP (ARDRA) patterns and unique 5' and 3' T-RFs. The ARDRA patterns of the six strains were indeed different (Fig. 3, lanes 2 to 7). However, if the DNA fragments derived from the six strains were mixed together (lane 10), the pattern became too complex to estimate the species diversity, let alone derive phylogenetic information. In contrast, T-RFLP analysis readily resolved the six-member community by limiting the analysis to T-RFs (Fig. 4). Six unique T-RFs were observed for both the 5' and 3' termini of the rDNA following HhaI digestion, which led us to conclude there were at least six bacterial strains in the model community.

There were discrepancies between the results expected based on database analyses and the empirical data obtained. Two of the 12 T-RFs observed did not have the predicted lengths (Table 3). The observed length for the T-RF of *Alcaligenes faecalis* was found to correspond to the length from the terminus to the second HhaI site in the RDP sequence. The amount of rrm sequence variation among strains of the same species has been shown to be 0 to 5%, and thus differences in T-RFs, while not common, are also not unexpected (7, 28).

This is problematic for T-RFLP analysis and several other molecular techniques (4, 14, 17–21) if species richness in a community is to be estimated correctly but should be partially overcome as more 16S rDNA sequences are included in the database. Similarly, the observed size of the 3' T-RF of *B. subtilis* was different than the size predicted, but this was most likely due to difficulty in precisely determining the sizes of fragments larger than 600 bp with an automated sequencer. The sizing accuracy could be improved by increasing the size range of fragments used as standards. However, even in our analysis, T-RFs from 37 to 600 bp long were accurately sized to ±2 bp.

**Analysis of microbial diversity of complex bacterial communities by T-RFLP.** The species richness of natural communities was estimated by determining the number of unique T-RFs or ribotypes observed in digests of 16S rDNAs amplified by PCR from total community DNA. The T-RFLP pattern observed (referred to as the "community fingerprint") is a composite of the number of fragments with unique lengths and the relative abundance of each fragment as reflected by the size of each peak in the electropherogram. Samples of natural microbial communities, including ELAS and MITI samples and samples from a contaminated aquifer site and termite guts (TGs), were studied by T-RFLP analysis. Based on an analysis of electro-

<table>
<thead>
<tr>
<th>Terminal end</th>
<th>5'</th>
<th>3'</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Ralstonia eutropha</em></td>
<td>208 (207)</td>
<td>291 (292)</td>
</tr>
<tr>
<td><em>Alcaligenes faecalis</em></td>
<td>569 (67)</td>
<td>72 (70)</td>
</tr>
<tr>
<td><em>Arthrobacter globiformis</em></td>
<td>474 (472)</td>
<td>76 (74)</td>
</tr>
<tr>
<td><em>Bacillus subtilis</em></td>
<td>259 (240)</td>
<td>632 (690)</td>
</tr>
<tr>
<td><em>Escherichia coli</em></td>
<td>373 (373)</td>
<td>350 (352)</td>
</tr>
<tr>
<td><em>Pseudomonas aeruginosa</em></td>
<td>154 (155)</td>
<td>346 (348)</td>
</tr>
</tbody>
</table>
could have been due to differences in inocula or growth conditions.

For example, the dominant T-RFs of the ELAS and MITI samples were either absent or present in low amounts in the MITI sample. Since the MITI sample was originally activated sludge, it was probably seeded with activated sludge from different wastewater treatment processes and subsequently acclimated in a bioreactor under defined growth and substrate feed conditions for more than 3 months. The extent to which T-RFLP analysis can take advantage of rRNA-based phylogenies is dependent on the actual universality or specificity of primer pairs. Universal primers, by definition, should be complementary to all known sequences. However, none of the known universal primers for 16S rDNA (e.g., 8f and 926r) could hybridize to or amplify all sequences from spirochetes and related organisms that could theoretically be amplified with primers 8f and 926r and yield a 38-bp 5′ T-RFLP pattern following digestion with HhaI. Moreover, the electropherograms of the TG sample following digestion with MspI or RsaI also had 5′ and 3′ T-RFs that were predicted based on the sequences of 16S rRNA genes of spirochetes (data not shown). Thus, the T-RFLP fingerprints of TGs suggest that spirochetes are a dominant ribotype in this community and this suggestion is consistent with results of previous studies that have shown that spirochetes with distinctive morphological features, metabolic functions, and 16S rRNA sequences are among the most abundant groups of bacteria in TGs (5, 22). However, it should be noted that nine other 16S rRNA gene sequences in the database would theoretically also yield a 38-bp 5′ T-RF following digestion of their 16S rRNA genes with HhaI plus MspI; these sequences include those of the 16S rRNA genes of _Spirochaeta paucivorans_, _Atopobium minutum_, _Polyangium cellulosum_, _Eubacterium spp._, and _Chondromyces spp._ However, by determining the expected sizes of the 5′ T-RFs following simulated digestion of these sequences with MspI or RsaI, it was possible to conclude that the 38-bp T-RF in the electropherogram of the TG sample following digestion with HhaI was probably not derived from these organisms. These data illustrate the utility of T-RFLP analyses for presumptively identifying bacterial groups within a microbial community.

The extent to which T-RFLP analysis can take advantage of rRNA-based phylogenies is dependent on the actual universality or specificity of primer pairs. Universal primers, by definition, should be complementary to all known sequences. However, none of the known universal primers for 16S rDNA have been shown to hybridize to or amplify all sequences from the eukaryal, bacterial, and archaeal domains (6, 36). Further-

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**FIG. 5.** Electropherograms of the 5′ T-RFLPs of HhaI-digested 16S rDNAs amplified from four different complex communities. (A) ELAS sample. (B) Aquifer sand sample. (C) TG sample. (D) MITI sample.
more, the universal primers that are currently used were designed by using the existing 16S rRNA database, which may represent only a portion of the total species diversity of the microbial world (1, 30). The primers that we used for community analysis were eubacterium-specific primers that were previously described, and our analysis was limited to this domain. A search of the subset of RDP sequences used for our database analysis clearly shows that primer pair 8f-926r, which we used for most of the T-RFLP community analyses, excludes the Archaea. This finding is based on a search that permitted no mismatches in the five residues at the 3′ terminus of the primer but permitted up to three mismatches in the remainder of the primer. This primer pair appears to exhibit a potential bias within the bacterial domain. A search of the bacterial domain by using the criteria described above indicated that chloroplasts, the α group, the Aeromonas subgroup, and the Bifidobacterium subgroup are recognized at levels of 23, 5.6, 21, and 1%, respectively, by the 8f-926r primer pair. However, the extent to which these phylogenetic groups are amplified under our PCR conditions is not known. The remainder of the database is recognized by this primer pair at levels of greater than 75% under the conditions specified above.

Data obtained by using T-RFLPs should be cautiously interpreted. For example, the number of populations represented in the fingerprint of any given community depends on the rank abundance of each population. Microbial populations that are not numerically dominant are not represented, because the template DNAs from these populations represent a small fraction of the total community DNA. Consequently, the species diversity of the microbial community is underestimated. Moreover, differences in gene copy number between species (10) and biases introduced during cell lysis, DNA extraction, and PCR amplification (12, 15, 23–25, 29) may yield a mixture of products that do not accurately reflect the rank abundance of the original community DNA template, thus skewing the apparent abundance of different populations.

In summary, the protocol developed here was shown to be useful for estimating the phylogenetic diversity and composition of communities in various environments. It provides a means to determine the number and abundance of numerically dominant ribotypes within a community of PCR-amplified 16S rDNAs. Using universal or domain-specific primers, one can draw only very general phylogenetic inferences from the results. The degree to which detailed phylogenetic information can be obtained with this approach reflects the level of phylogenetic specificity that can be imparted to the PCR primer. The method does produce distinct community signatures that can be used to assess the similarity of different communities. This approach should provide a facile means to assess microbial diversity and to assess changes in microbial community structure that occur on temporal or spatial scales or that occur in response to environmental perturbations.

ACKNOWLEDGMENTS

We thank Ross Overbeek (Argonne National Laboratory, Chicago, Ill.) for assistance with the Pat Scan program, Bonnie Maidak (University of Illinois, Urbana-Champaign) for providing the RDP database, and Timothy Lilburn and John Breznak (Michigan State University, East Lansing) for supplying the TG sample. Much gratitude is also given to Mary Hutcheson and Bobbie Okimoto (USDA-ARS Avian Disease & Oncology Laboratory, East Lansing, Mich.) for the analysis of T-RFLP samples.

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REFERENCES


