Microbial diversity along a transect of agronomic zones

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Abstract

The diversity of microbial communities constitutes a critical component of good soil-management practices. To characterize the effects of different management practices, molecular indicators such as phospholipid fatty acid (PLFA), denaturing gradient gel electrophoresis (DGGE) and composition of ammonia-oxidizing bacteria were used to analyze bacterial community structure and diversity from four eastern Washington State soils. Samples from four sites were collected representing a transect of high-precipitation to low-precipitation areas that covered different agronomic zones with different management and cropping practices. Biomass amounts estimated from extractable PLFA were significantly higher in the no-till (NT) soil than in the conventional-till (CT) soil. Similarities among the different 16S rDNA DGGE band profiles were analyzed quantitatively using correspondence analysis and this confirmed that the CT soil was the most dissimilar soil. DGGE analysis of 16S rDNA ammonia-oxidizing bacteria from the four soils revealed two identical bands, indicating little effect of agronomic practices and precipitation on these species. A second set of primers, specific for amoA (ammonia monoxygenase) genes, was used to examine ammonia oxidizers in the samples. Six banding patterns (clusters) from amplified rDNA restriction analysis of 16S rDNA fragments were observed after restriction analysis with Hinfl. Sequencing of these clones revealed the presence of only Nitrosospira-like sequences. Analysis of the sequences showed that ammonia oxidizers from the NT soil were more diverse compared to those from the CT and conservation reserve program soils. Our data showed that management and agronomic practices had more impact on bacterial community structure than annual precipitation. © 2002 Published by Elsevier Science B.V. on behalf of the Federation of European Microbiological Societies.

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1. Introduction

Microbial communities are critical components of soil and may be the earliest predictors of soil quality changes [1]. A recent definition of soil quality is the ability of soil to sustain biological functions and promote plant and animal health [2]. The diversity of microorganisms in soil is critical to the maintenance of good soil health because microorganisms are involved in many important soil functions such as carbon and nitrogen cycling, soil tilth and structure [3], and organic-matter transformation [4]. Several microbial methods including the traditional approaches have been used to study microbial diversity in agricultural soils [5,6]. Recent developments in molecular biology and biochemical assays for soil microbial community structure analysis have provided new tools to conduct experiments on soil microbial components without using the traditional approaches.

The quantitative description of microbial communities is one of the most difficult tasks facing microbial ecologists, as the classical technique of viable counting recovers only a small proportion of cells [7]. At present, only two methods have overcome the problem of selective culturing and provide an unbiased view of the structure of complex microbial communities. These are (1) analysis of microbial populations using ribosomal RNA (rDNA) and (2) phospholipid fatty acid (PLFA) analysis [8]. Microbial community analyses using rRNA can detect and identify community members with high specificity to the species and
strain level, and can also detect and suggest phylogenetic affinities of uncultured organisms [9]. PLFA can be used to quantify community structure and biomass without relying upon the cultivation of microorganisms. Unfortunately, this approach does not have the capability of identifying microorganisms at the species or strain level, but rather produces descriptions of microbial communities based on functional groupings of fatty acid profiles. This concept has resulted in the identification and quantification of viable biomass and community structure in sediments [10–12] and in agricultural soils [13,14].

Use of the 16S RNA gene as an indicator for species diversity is providing a fresh approach for investigating the structure of microbial communities. Such methods include thermal and denaturing gradient gel electrophoresis (DGGE) [15,16], terminal restriction fragment analysis [17–19], and 16S rDNA sequence analysis [20,21].

One of the critical functions of soil microorganisms in maintaining good soil health is the cycling of nitrogen. This process is carried out in part by ammonia-oxidizing bacteria. This group of bacteria is responsible for the conversion of ammonia to nitrite which is the form of nitrogen readily available to plants. Therefore, it is important to understand the community structure of ammonia oxidizers as they play a very important function in soil.

Previous studies [22] using soils from the same area as in this study showed a shift in microbial community structure between a grass prairie and a cultivated field when biomass C, phosphatase, and substrate utilization were measured. We used phospholipids to assess microbial biomass and community structure from the four eastern Washington State soils because PLFAs are rapidly turned over and thus represent the current living community, both quantitatively and qualitatively. DGGE of the 16S rDNA fragment amplified directly from the soil DNA using general bacterial primers was used to quantify the level of bacterial community diversity. Diversity of ammonia-oxidizing bacteria, which comprises mainly of small grains with minimal tillage (MT), conservation reserve program (CRP), legumes with no-tillage (NT), and dry-land and irrigation systems with conventional tillage (CT) (Table 1). The Natural Resources Inventory database of primary sampling unit for Washington State was used [23], along with county surveys, to select sites from the two counties, based upon the different agricultural-management histories, current farming practices, and annual precipitation. To locate each of the four sampling sites, a Magellan NAV 5000 pro Global Positioning System was used. One sample was taken from each of the sampling sites and four additional satellite samples were taken at distances of 100 m in the four cardinal compass directions. The satellite samples were collected to account for local variability, in a 100-m radius around the center sample. All samples were kept in sealed plastic bags and mixed immediately. Samples were then placed on dry ice and stored at −70°C prior to DNA extraction.

2. Materials and methods

2.1. Site selection and sampling

Soil samples (0–10 cm) were collected along a transect from representative sites in Columbia and Adams counties in eastern Washington State. Farming practices were comprised mainly of small grains with minimal tillage (MT), conservation reserve program (CRP), legumes with no-tillage (NT), and dry-land and irrigation systems with conventional tillage (CT) (Table 1). The Natural Resources Inventory database of primary sampling unit for Washington State was used [23], along with county surveys, to select sites from the two counties, based upon the different agricultural-management histories, current farming practices, and annual precipitation. To locate each of the four sampling sites, a Magellan NAV 5000 pro Global Positioning System was used. One sample was taken from each of the sampling sites and four additional satellite samples were taken at distances of 100 m in the four cardinal compass directions. The satellite samples were collected to account for local variability, in a 100-m radius around the center sample. All samples were kept in sealed plastic bags and mixed immediately. Samples were then placed on dry ice and stored at −70°C prior to DNA extraction.

2.2. Phospholipid extraction, separation, and analysis

Soil samples (5 g, n = 5) were extracted by using the modified method of Bligh and Dyer as described by Petersen and Klug [24]. The total lipid extract was fractionated into glyco-, neutral, and polar lipids [25]. The polar-lipid fraction was transesterified with a mild alkali to recover the PLFA as methyl esters in hexane. The PLFAs were separated, quantified and identified by GC-flame ion-
2.3. Fatty acid nomenclature

The suffixes ‘-c’ for cis and ‘-t’ for trans refer to geometric isomers. The prefixes ‘i-’, ‘a-’, and ‘me-’ refer to iso-, anteisomethyl, and mid-chain methyl branching, respectively, with cyclopropyl rings indicated by ‘cy’ [27].

2.4. DNA extraction, PCR primers, and DGGE analysis

Total bacterial community DNA was extracted by placing about 500 mg of soil in FastPrep tubes (BIO 101, Vista, CA, USA) containing lysis matrix. Isolation of total DNA was accomplished with a FastPrep DNA isolation kit according to the manufacturer’s protocol (BIO 101). PCR was first performed using 20–80 ng template DNA with the primers, PRBA338f and PRUN518r, located at the V3 region of the 16S rRNA genes of bacterio-iplankton [28], to assess bacterial community diversity. PCR mixtures for the bacterial 16S rDNA sequence amplification contained 10 pmol of each primer, 4 μg of bovine serum albumin, and sterile distilled water in a final volume of 25 μl. The PCR cycles used for amplification were as follows: 92°C for 2 min, followed by 30 cycles of 92°C for 1 min, 55°C for 30 s, 72°C for 1 min, and a single final extension at 72°C for 6 min.

DGGE was performed with 8% (w/v) acrylamide gels containing a linear chemical gradient ranging from 30 to 70% denaturant with 100% defined as 7 M urea and 40% formamide. Gels were run for 3 h at 200 V with a Dcode Universal Mutation System (Bio-Rad Laboratories, Hercules, CA, USA). DNA was visualized after ethidium bromide staining by UV transillumination and photographed with a Polaroid camera.

2.5. DGGE, blotting, and sequencing of ammonia oxidizers

To determine the diversity of ammonia-oxidizing bacteria in the four soils, a nested PCR assay was first performed with bacterial-specific primers 27f and 1492r [29]. A second PCR assay was performed with the CTO 189f-GC and CTO 645r [30]. The first PCR assay was performed with 50 ng of template DNA using the following parameters: 94°C for 2 min, followed by 30 cycles of 92°C for 30 s, 55°C for 60 s, and 72°C at 72°C (with the time increasing 1 s/cycle), and a final single extension at 72°C for 5 min with a reaction volume of 25 μl. Products with the expected size of 1.5 kb were excised from the gel, melted by heating at 65°C for 5 min and 1 μl was used as template in a second PCR with the CTO primers. PCR products recovered with CTO primers were subjected to DGGE as described above, except a gradient of 30–60 denaturant was used with a run time of 4 h.

A second PCR was performed with the CTO primers but without the GC-clamp for Southern blot analysis. Aliquots (10 μl) of the CTO PCR products were separated in 1% agarose gels. The DNA fragments were transferred to a nylon membrane [31] and were fixed by UV cross-linking. Hybridization analyses were conducted by using the oligonucleotide probes (Nsp436 for all Nitrosospira and Nmo254a for all Nitrosomonas) and hybridization conditions described by Stephens et al. [32]. Probe labeling and immunological detection were done by random primed incorporation of DIG Oligonucleotide 3’-end labeling kit (Boehringer Mannheim, Bedford, MA, USA) as recommended by the manufacturer.

A third PCR was performed with amoA primers that target a partial stretch of genes that encode the active-site polypeptide of amoA [33]. Amplification was performed in a total volume of 50 μl. The reaction conditions were 94°C for 2 min, followed by 30 cycles of 92°C for 60 s, 55°C for 60 s, and 68°C for 45 s, and a final single extension at 72°C for 5 min with 10 pmol of each primer. PCR products were gel-purified and extracted with Qaex-II gel extraction kit (Qiagen, Valencia, CA, USA). Purified fragments were cloned with the pGEM-T Easy vector (Promega, Madison, WI, USA) and transformed into Escherichia coli JM109. Plasmid isolations were performed using the Qiagen plasmid mini kit as described by the manufacturer. For each soil, 40 clones with the correct size of insert were selected and amplified rDNA restriction analysis (ARDRA) technique was performed to assess the genetic diversity of ammonia-oxidizing bacteria from the different soils [34]. After PCR reamplification of the clones with the vector primers M13 and T7, 5 μl of the PCR products was digested with the restriction endonuclease HinfI and analyzed by separation of fragments on a 2% low-melting agarose gel in Tris–acetate–EDTA. Representative clones from each digestive pattern were selected for sequencing and data analyses. The purified DNA was sequenced with the ABI PRISM Dye Terminator Cycle Sequencing kit with AmpliTaq DNA Polymerase, FS (Perkin Elmer).

2.6. Phylogenetic analysis

Sequence analyses were done using the BLAST database (National Center for Biotechnology Information, www.ncbi.nlm.nih.gov). Partial amoA gene sequences were aligned with parts from the complete amoA gene sequences of ammonia-oxidizing bacteria obtained from the BLAST gene bank. Sequence alignments were performed using the PILEUP program from the University of California-Riverside Genetics Computer Group (GCG programs). Matrices of evolutionary distances were computed using the Phylip program with the Jukes–Cantor model [35]. Phylogenetic trees were constructed and
checked by bootstrap analysis (100 data sets) using the program SEQBOOT. Bootstrap values represent the frequency of resampling that supports a specific branching pattern.

2.7. Statistical analysis of PLFA and DGGE bands

Data analyses were performed using SAS [36]. Means and standard deviations for the individual fatty acids in triplicate-sample PLFA profiles were determined to compare the composition of the PLFA profile in samples from different sites. Average mole percent ratios of selected fatty acids were used to estimate microbial biomass using selected PLFAs as biomarkers. Principle component analysis (PCA) was used to compare PLFA profiles among the four sites and the data were presented as a 3D plot for better understanding of the relationship. PCA explains the variance–covariance structure through a few linear combinations of the original variance (PLFA) with coefficients equal to the eigenvectors of the correlation matrix [37].

DNA fingerprints obtained from the 16S rDNA DGGE banding patterns on the gels were photographed and digitized using ImageMaster Labscan (Amersham-Pharmacia Biotech, Uppsala, Sweden). The lanes were normalized to contain the same amount of total signal after background subtraction. The gel images were straightened and aligned using ImageMaster 1D Elite 3.01 (Amersham-Pharmacia Biotech, Uppsala, Sweden) and analyzed to give a densitometric curve for each gel. Band positions were converted to \( R_f \) values between 0 and 1 and profile similarity was calculated by determining Dice’s coefficient for the total number of lane patterns. Dendrograms were constructed by using the unweighted pair group method with mathematical averages (UPGMA). Dice’s similarity coefficients were generated, converted into \( x/y \) line plots, and transferred to Excel files. Community similarities based on peak areas from the Excel files for the different bacterial groups (16S rDNA bands) were analyzed by correspondence analysis (CANACO 4.0, Microcomputer Power, Ithaca, NY, USA) to generate an ordination diagram as described by Ibekwe et al. [25].

3. Results and discussion

3.1. Chemical and biological properties of soils

Results of biological and chemical data from the four soils are presented in Table 1. Organic carbon and CEC were significantly higher in the Palouse silt loam soil with MT. These parameters were lowest in the Shano silt loam soil with CT. The Shano soil also had the lowest amount of annual precipitation in all of the sites studied. There were no pH differences among the four soils. No significant differences were found in dehydrogenase or biomass for the MT, CRP, or NT soils. However, the CT soil had significantly lower dehydrogenase and biomass (Table 1).

3.2. PLFA analysis for microbial community structure and biomass

Principal component analyses of PLFA data of soil communities (Fig. 1) were conducted to determine the microbial community structure of the four soils. PCA was conducted with 49 PLFAs that were present in the soil samples. The PCA plot (Fig. 1) for the four soil microbial communities accounted for 36% of the variance on the first component, while the second component accounted for 22%, and the third component accounted for 14%. This explained 72% of the total variance in the first three dimensions of the plot. PCA of the samples revealed separation of the four soils into four distinct clusters (Fig. 1). The MT, CRP, and NT soils formed distinct clusters along principle components (PCs) 1 and 2, while the CT soil clustered along PC3. All analyses were done using the correlation matrix and the Kaiser’s rule [37], which states that variables with eigenvalues greater than 1 should be used for further analysis. A second PCA was conducted on matrices with fewer PLFAs. Eighteen PLFAs with the most significant contributions to the PCA (\( r = 0.55 \)) were selected. The 3D plot constructed from the reduced matrix did not show different clustering (data not shown). PLFAs contributing most significantly to each of the principal component scores were the same for both the reduced data set (18 PLFAs) and the unreduced set (49 PLFAs). Branched, monounsaturated, and cyclopropane PLFAs

Fig. 1. PCA of microbial community structure from four soils in eastern Washington State as determined by their PLFA profiles. •: MT with small grain and annual rainfall greater than 400 mm, ○: CRP (annual rainfall of 350–400 mm), ▼: NT under alfalfa pasture and annual rainfall between 250–300 mm, ▲: CT with potato, and annual rainfall below 200 mm. PCA was conducted with 49 PLFAs that explained 72% of the total variance in the first three dimensions.
were heavily weighted on PC1 (a15:0, i15:0, i16:0, 16:1T7c, a17:0, i17:0, cy17:0, 18:1o9c, cy19:0). PC2 and PC3 were heavily weighted by straight-chain PLFA (PC2=12:0, a13:0, 15:0, 17:0, 20:0; PC3=14:0 and 16:0). Most of the fatty acids in PC1 are primarily of bacterial origin [38]. The clusters obtained from the soil communities indicate differences in the biomarker compositions in the four soils.

PLFA content in the four soils, as determined by microbial biomass per gram of dry weight, showed marked significant variations ranging from 67.6 nmol g⁻¹ dry wt in the Couse silt loam soil with NT to 56.3 nmol g⁻¹ dry wt in the Shano silt loam soil. The variation in biomass composition from the four soils was in agreement with the clusters obtained in Fig. 1. In a complex environment such as soil, the different fatty acid compositions can be divided into different metabolic groups for proper understanding of the community composition (Fig. 2). Hydroxy PLFA compositions in the four soils were lower than cyclopropane PLFA, and these PLFAs are characteristic of Gram-negative bacteria [38]. Monounsaturated fatty acids showed some differences among the four sites, but this did not follow any pattern associated with precipitation. Straight-chain PLFAs had the highest mole percent concentration in all the samples, with 16:0 having the highest percent concentration. Branched PLFAs, characteristic of Gram-positive bacteria [39,40], were about three- to five-fold higher in soils than the hydroxy group. The polyunsaturated PLFAs, mostly 18:2 w6c, characteristic of fungi [41], were higher in the MT and CT soils.

PLFA composition and biomass observed in this study provided some evidence of how microbial communities are shaped due to the influence of different agricultural practices and environmental conditions. These results showed that there were significant differences in microbial biomass among the four soils. For example, total PLFA contents which reflect active microbial biomass [42,43] declined significantly from 67.6 nmol g⁻¹ dry wt in Couse silt loam soil under NT and alfalfa to 56.3 nmol g⁻¹ dry wt in Shano silt loam soil under CT and potato. Organic carbon was four times greater in the Couse silt loam than the Shano soil, and this factor has been shown to have strong correlation with microbial biomass carbon [2]. Biomass from PLFA may be a strong indicator of bacterial composition since it represents all microbial biomass and indicates active living microbial components. Biomass from phospholipid has been found to be more representative of the microbial biomass than that obtained from the chloroform-fumigation method [5,44].

3.3. Analysis of microbial community diversity by DGGE

Examination of the band profile in the DGGE gel revealed that the communities consisted of five predominant bacterial DGGE groups that were present in all the soils and some minor groups that co-migrated in the DGGE gel. To compare DGGE patterns from the four soils, Dice’s indices were determined for comparisons of all profiles, and UPGMA was used to create a dendrogram describing pattern similarities (Fig. 3). The bands in the DGGE profile correspond to the 16S rDNA fragments that differ in nucleotide sequences. This represents the dominant microbial population in the four soils pooled from the center sample and the four satellite samples. Based on the analysis of the DGGE banding patterns, 17, 18, 20, and 17 major bands were observed from Palouse, Walla Walla, Couse, and Shona silt loam soils, respectively (Table 1).

Eichner et al. [45] described the community structure in activated sludge and noted that the number and intensity of bands did not equal the number and abundance of species within the microbial community due to features of 16S rDNA-based phylogeny and bias inherent to PCR amplification from complex template mixtures. The reasons for most of the limitations are that DGGE banding patterns are subjected to PCR bias due to DNA ex-
traction methods, potential preferential amplification, and the formation of chimera [46]. Other problems may result from one organism producing more than one DGGE band because of multiple, heterogeneous rRNA operons [47–49]. Also, for some phylogenetic groups of bacteria, 16S rDNA sequences do not allow discrimination between species, so one DGGE band may represent several species with identical rDNA sequences [50].

In a community DNA mixture such as soil, the maximum number of different rDNA fragments separated by DGGE may be vastly underestimated. Torsvik et al. [51] found that there might be as many as \(10^4\) different genomes present in soil samples. This shows that DGGE can not separate all of the 16S rDNA fragments obtained from soil microorganisms, but only the dominant species [15]. Therefore, the banding patterns obtained in this study reflect the most abundant rDNA types in the community.

The second method for the calculation of community diversity was done by first calculating the similarities of three sampling points from each soil by correspondence analysis (Fig. 4). The statistical procedure uses matrix algebra to describe banding pattern data in a multidimensional space. The data are graphed using eigenvectors that locate within the communities a set of coordinates for factors that describe the data set, with corresponding eigenvalues that describe the amount of variation that is explained by each factor. Sixty seven percent of the variance in the community structure was explained by three factors, with factor one explaining 38% and factor two 19%. The difference between the communities with respect to the first and the second eigenvectors was the distance between the two points representing the communities on the ordination diagram. A comparison of the distances showed that the microbial communities from the CT soils had the most dissimilar structure (Fig. 4). The three CT samples were separated by about 1 standard deviation on the x-axis and 3 standard deviations on the y-axis (Fig. 4). The communities associated with NT, MT, and CRP were separated on the average among and within themselves with less than 0.5 standard deviation (Fig. 4).

The use of PLFA and DGGE methods provided useful information on the composition of soil microbial communities. PLFA analysis of soil microbial community structure by PCA (Fig. 1) and the analysis of DGGE patterns (Figs. 3 and 4) revealed a common pattern of clustering from the four soils. From the three analyses, NT and CRP soils clustered closer to each other than CT and MT soils. These results suggest that soil microbial community responds more to management than precipitation. Based on a previous study [22] and this study it is quite clear that microbial community structure is very sensitive to changes in soil management. Therefore, microbial community structure sets the potential for functional response to management changes, which in turn affect soil processes.

### 3.4. Diversity of ammonia-oxidizing bacteria recovered from soils

The relationship between community structure and function was further explored by examining the functional gene encoding the \(\alpha\) subunit of amoA. Results obtained from the DGGE analysis of ammonia-oxidizing bacteria were very reproducible across the four soils (Fig. 5). DGGE patterns from the sampling points showed two similar dominant bands from the four sites that were closely related to *Nitrosospira*. Previous results from se-
quencing [30,32,52,53] using the same primers as in this study showed that the two bands contained sequences that were closely related to 

\textit{Nitrosospira}-like sequence of the \(\beta\)-subdivision ammonia oxidizers. These authors also stated that bands one and two may be doublets with a slight difference in sequence composition. The DGGE patterns obtained from our four soils and the satellite samples were very similar, even though soils one and four were tilled plots with different crops (Fig. 5). In our study, two dominant bands co-migrated in all of the soils, irrespective of management practices or precipitation. This suggests that there is a high degree of stability of ammonia oxidizers in most agricultural soils. Bruns et al. [53] observed similar DGGE patterns in agricultural soils with different crops over a 2-year period.

To confirm whether \textit{Nitrosospira}-like sequences of \(\beta\)-subdivision ammonia oxidizers were the dominant genera in these soils, PCR amplification of the soil samples with CTO primers were used for Southern hybridization with non-radioactive biotin-labeled oligonucleotide probes Nsp436 and very weak signals obtained with the Nmo254a probe. Hybridization patterns indicated that bands from DNA of the four soils were derived mainly from \textit{Nitrosospira} sp. with non-radioactive biotin-labeled oligonucleotide probes Nsp436 and Nmo254a for all \textit{Nitrosospira} and Nmo254a for all \textit{Nitrosomonas} (data not shown). This confirmed the identity of the PCR products. Strong hybridization signals were obtained from the four soils with probe Nsp436 and very weak signals obtained with the Nmo254a probe. Hybridization patterns indicated that bands from DNA of the four soils were derived mainly from \textit{Nitrosospira} DNA, suggesting that \textit{Nitrosospira} is more dominant in these agricultural soils than \textit{Nitrosomonas}.

To determine the genetic heterogeneity of ammonia oxidizers, the gene encoding the \(\alpha\) subunit of \textit{amoA} was selected as a functional marker for ammonia oxidizers. PCR amplification of the four soils produced DNA fragments of the expected size (ca. 491 bp). The diversity of recovered rDNA PCR fragments in the soil samples was examined by comparative ARDRA. ARDRA with \textit{Hin}I digestion revealed a total of six major banding patterns from the four soils (data not shown). Soil 3, under alfalfa and NT, had 5 ARDRA banding patterns. Soil 2, under CRP had two ARDRA banding patterns. Both soil 1 (after harvest of small grain under minimum tillage) and soil 4 (under potato and conservation tillage) had four ARDRA banding patterns. Soils 1 and 4 are from areas with the highest and lowest annual precipitation, respectively. These observations suggest that precipitation rate may not have any significant effect on the natural population of ammonia oxidizers. However, management and crop types may play a more significant role. Two cloned representatives from each pattern and from each soil were selected for sequence analysis. BLAST searches of the GenBank database confirmed that all sequenced clones represented \textit{amoA}-like sequences. The identity values for the \textit{amoA} sequences from these soil samples varied between 93.0 and 100%. Two of the cloned sequences were identified as \textit{Nitrosospira briensis} and \textit{N. briensis} C-57 (Fig. 6) and the other four were unidentified bacterium \textit{amoA} sequences of the \textit{Nitrosospira} genus as described by Rothauwe et al. [33].

The use of the functional gene target of \textit{amoA} sequences provided an example of diversity of ammonia oxidizers that is very crucial in soil-management assessment. NT soil with alfalfa had the highest diversity (five) of \textit{amoA} sequences, while CRP had the lowest (two). One possible explanation for the high diversity in the NT plot may be the continuous presence of the rhizosphere from alfalfa plants that provided a special niche for the proliferation of different strains of ammonia oxidizers compared to the CRP that was covered with grass. The CT and MT soils had four \textit{amoA} sequences each. This finding is in agreement with Bruns et al. [53] who found a greater genetic diversity of autotrophic ammonia-oxidizer sequences in soils from native, undisturbed soils than in tilled soils. \textit{N. briensis} clone A1K and unidentified \textit{amoA} clone A1N sequences were found only in soils 1 and 4 (Fig. 6). These soils were under some form of tillage regime and nitrogen fertilization. This suggests that these strains were either more adapted to these soils or were below detection limit for this technique in the other soils. On the other hand, \textit{N. briensis} C-57 clone A30 and unidentified \textit{amoA} clone A40 sequences were present in all the four soils, and may be more abundant in most agricultural soils. The presence of only two clones (A40, A30) in the CRP soil suggests that this soil contained very low numbers of ammonia oxidizers.

In this study, we have provided information on microbial community structure from four soils with contrasting soil management and environmental conditions. The use of biochemical and molecular biology tools to assess the impact of management indicated that NT practices improved the biological condition of the soil in eastern Washington State compared to CT practice. Two indica-
tors to explain this conclusion are (1) the high microbial biomass as determined by PLFA analysis and (2) the greater diversity of amoA sequences associated with the NT soil. The use of different molecular methods for the analyses of environmental and management effects is a new approach that will greatly assist in understanding changes in soil health over small time scales since microbiological parameters may be the first to indicate changes or stress.

References


