

## Community Composition and Functioning of Denitrifying Bacteria from Adjacent Meadow and Forest Soils

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**We investigated communities of denitrifying bacteria from adjacent meadow and forest soils. Our objectives were to explore spatial gradients in denitrifier communities from meadow to forest, examine whether community composition was related to ecological properties (such as vegetation type and process rates), and determine phylogenetic relationships among denitrifiers. *nosZ*, a key gene in the denitrification pathway for nitrous oxide reductase, served as a marker for denitrifying bacteria. Denitrifying enzyme activity (DEA) was measured as a proxy for function. Other variables, such as nitrification potential and soil C/N ratio, were also measured. Soil samples were taken along transects that spanned meadow-forest boundaries at two sites in the H. J. Andrews Experimental Forest in the Western Cascade Mountains of Oregon. Results indicated strong functional and structural community differences between the meadow and forest soils. Levels of DEA were an order of magnitude higher in the meadow soils. Denitrifying community composition was related to process rates and vegetation type as determined on the basis of multivariate analyses of *nosZ* terminal restriction fragment length polymorphism profiles. Denitrifier communities formed distinct groups according to vegetation type and site. Screening 225 *nosZ* clones yielded 47 unique denitrifying genotypes; the most dominant genotype occurred 31 times, and half the genotypes occurred once. Several dominant and less-dominant denitrifying genotypes were more characteristic of either meadow or forest soils. The majority of *nosZ* fragments sequenced from meadow or forest soils were most similar to *nosZ* from the *Rhizobiaceae* group in  $\alpha$ -*Proteobacteria* species. Denitrifying community composition, as well as environmental factors, may contribute to the variability of denitrification rates in these systems.**

In addition to ameliorating eutrophication and removing excess nitrate ( $\text{NO}_3^-$ ) from drinking water, the process of denitrification can limit primary productivity and contribute to global warming through the formation of nitrous oxide ( $\text{N}_2\text{O}$ ). Despite the essential role of denitrification in the global nitrogen (N) cycle, little is known about how denitrifying community structure relates to denitrification rates in natural environments. Examining this relationship may improve understanding of denitrification and provide insights into the functional role of microbial diversity.

Denitrifying bacteria are defined solely by the ability to reduce  $\text{NO}_3^-$  or nitrite ( $\text{NO}_2^-$ ) to  $\text{N}_2\text{O}$  or  $\text{N}_2$  in the absence of oxygen. Denitrification is not specific to any one phylogenetic group; the trait is found in about 50 genera, mostly in the *Proteobacteria* species (48). Therefore, to analyze denitrifier diversity, functional genes (such as those coding for  $\text{NO}_2^-$  reductase [*nirK* and *nirS*] and  $\text{N}_2\text{O}$  reductase [*nosZ*]) in the denitrification pathway have been retrieved (using PCR and cloning and sequencing) from environmental samples (4, 36, 39). Denitrifier community profiles have been generated using PCR-coupled terminal restriction fragment length polymorphisms (T-RFLPs) (1, 3, 40). Avrahami et al. (1) recently found that denitrifying community composition was related to ammonium ( $\text{NH}_4^+$ ) addition and enhanced  $\text{N}_2\text{O}$  emissions from soil.

Plant communities presumably have strong effects on below-ground microbial community structure (18, 32). Therefore, we investigated denitrifying soil communities from two broadly contrasting plant communities of meadow vegetation and adjacent coniferous forest in the Western Cascade Mountains of Oregon. We hypothesized that because these vegetation types likely differed in soil N cycling processes, denitrifier communities would also differ. Differences in N cycling were assumed because the net production of  $\text{NO}_3^-$  is higher in analogous meadow soils than in coniferous forests (5, 14). Although factors leading to vegetation differences are complex (21, 25), annual turnover and higher concentrations of N in meadow plant tissue likely contribute to differences in N cycling (12, 28).

The objective of this study was to examine spatial variation in denitrifier communities and N cycling processes from adjacent meadow and forest soils. We used standard activity assays to confirm that N cycling processes differed between meadow and forest soils. Community composition was measured with T-RFLP profiles of the *nosZ* fragment analyzed by Rösch et al. (36). We assessed relationships between denitrifier communities and functioning with multivariate techniques, and we investigated phylogenetic relationships among *nosZ* clones. Recently, Mintie et al. (26) examined nitrifying populations from the same soils.

### MATERIALS AND METHODS

**Site characteristics and sampling.** Study sites were located at the H. J. Andrews Experimental Forest (44.2°N, 122.2°W) in the Western Cascade Mountains of Oregon. We examined two montane sites (referred to here as Carpenter

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TABLE 1. Properties of the mineral soil from the H. J. Andrews meadow-to-forest transects (mean of 0- to 10-cm depth)

Measurement	Value for Carpenter site		Value for Lookout site		Reference or source
	Meadow	Forest	Meadow	Forest	
pH	5.8	5.2	5.6	5.1	26
Total C (g of C kg <sup>-1</sup> of dry soil)	94.7	131.8	115.9	142.3	This study
Total N (g of N kg <sup>-1</sup> of dry soil)	6.8	5.8	9.3	9.2	This study
C:N	13.8	22.8	12.7	15.7	This study
Nitrate (mg of N kg <sup>-1</sup> of dry soil) <sup>a</sup>	2.9	1.5	3.5	1.9	This study
Ammonium (mg of N kg <sup>-1</sup> of dry soil)	4.6	1.4	2.6	1.7	This study
Net nitrification (mg of N kg <sup>-1</sup> of dry soil day <sup>-1</sup> ) <sup>b</sup>	2.60	0.04	2.15	0.20	This study
Net ammonification (mg of N kg <sup>-1</sup> of dry soil day <sup>-1</sup> )	-0.42	0.43	-0.20	0.79	This study
Net N mineralization (mg of N kg <sup>-1</sup> of dry soil day <sup>-1</sup> )	2.18	0.47	1.96	0.99	26
Nitrification potential (mg of N kg <sup>-1</sup> of dry soil day <sup>-1</sup> )	8.90	0.26	8.67	0.74	26
DEA (mg of N kg <sup>-1</sup> of dry soil day <sup>-1</sup> )	0.78	0.04	1.58	0.17	This study

<sup>a</sup> Nitrate and ammonium were measured with an autoanalyzer, as reported for N mineralization (26).

<sup>b</sup> Net rates were measured over a 10-day laboratory incubation (26).

and Lookout) with adjacent meadow and forest vegetation and similar soils. Details of the study sites have been reported elsewhere (26, 35).

At each site, soil samples were taken along three replicate transects, which were distributed throughout sites at a distance of at least 20 m from each another. Transects ran perpendicularly to the meadow-forest boundary and consisted of eight evenly spaced sampling points, four in the forest and four in the meadow. Sampling points were spaced at 20-m intervals at Carpenter and 10-m intervals at Lookout, with larger spacing at Carpenter to account for the wider meadow-to-forest transition at this site. We sampled the mineral soil under the organic humus layer (at a depth of 0 to 10 cm) by compositing five replicate soil cores (inner diameter, 6 cm) at each sampling point within a 0.5-m radius. A total of 48 field replicate samples (i.e., 48 sample units) were taken on 26 June 2000. The soil was brought back to the laboratory on ice, stored at 4°C, and sieved (4.75-mm-pore-size mesh) about 24 h after sampling. Soil for DNA extractions was immediately frozen at -20°C after sieving. Soil for the denitrification potential measurement was kept at 4°C until analysis was performed (within 72 h after sampling). Basic characteristics of the transect soil samples are shown in Table 1. Total soil carbon (C) and N were measured on a dry-weight basis by combustion.

**Denitrification potentials.** Denitrification potential (or denitrifying enzyme activity [DEA]) measurements were carried out following the method of Tiedje (45). Fresh soil (10 g) was added to 125-ml Erlenmeyer flasks with 25 ml of a solution containing glucose (10 mM), NO<sub>3</sub><sup>-</sup> (5 mM), and phosphate buffer (pH 7.0) (50 mM). The flasks were sealed with neoprene stoppers and made anaerobic by repeated evacuation and flushing with Ar. Standard-grade acetylene was purified through an acid trap (13) and added to the headspace (10% [vol/vol]). The flasks were incubated at 250 rpm and 21°C on a rotary shaker. Gas samples (4 ml) were taken at 15 and 75 min and stored in Vacutainers (3 ml) until gas chromatography analysis of N<sub>2</sub>O levels was performed using a <sup>63</sup>Ni electron capture detector (27). Production of N<sub>2</sub>O was linear (as determined on the basis of analysis of a subset of flasks sampled every 15 min) during the 15- to 75-min interval (data not shown). Statistical differences in DEA were assessed using two-factor analysis of variance (ANOVA) for site and vegetation type (SAS software, version 7.0).

**nosZ PCR and T-RFLPs.** PCR was used to generate 700-bp *nosZ* fragments from the soil-denitrifying community. Designed in 1999 on the basis of available sequence data and the work of Scala and Kerkhof (41), our *nosZ* primers are almost identical to those independently developed by Rösch et al. (36). Our primer sequences were as follows: 5'-CGCTGTTTCITCGACAGYCAC-3' (bold characters represent nucleotides that differ from those in the sequence presented in reference 36) for the forward primer (*nosZ*-F-1181) and 5'-ATGTGCAKIG CRTGGCAGAA-3' for the reverse primer (*nosZ*-R-1880) (I, inosine; Y, T and C; K, T and G; R, A and G). Numbers included in primer designations indicate nucleotide positions at the ends of the *Pseudomonas stutzeri* 700-bp *nosZ* fragment (GenBank accession no. M22628).

DNA was extracted from soil samples (fresh weight, 0.3 g) or pure cultures using a FastDNA kit (Bio 101, Inc.) for bacterial cells or soil according to the manufacturer's instructions. DNA extracts were checked on agarose gels (1.1%) stained with ethidium bromide (0.5 µg ml<sup>-1</sup>). The DNA was quantified using a DNA TKO 100 fluorometer (Hoefer Scientific) with a calf thymus DNA standard. Reaction mixtures (50 µl) contained soil DNA (100 ng), AmpliTaq DNA polymerase (2.5 U), GeneAmp PCR buffer (1×), MgCl<sub>2</sub> (2 mM), deoxynucleo-

side triphosphates (0.2 mM each), forward and reverse primers (0.2 µM each), and bovine serum albumin (0.064 g ml<sup>-1</sup>). The forward primer was fluorescently labeled with 6-Fam (6-carboxyfluorescein). For PCR amplification of soil DNA, the temperature profile on a PTC-100 hot bonnet thermocycler (MJ Research, Inc.) was 94°C for 3 min and 25 cycles of 94°C (45 s), 56°C (1 min), and 72°C (2 min) followed by a final extension of 72°C for 7 min. For each sample, the PCR products of three reactions were pooled and purified using a QIAquick PCR purification kit (Qiagen) according to the manufacturer's instructions. PCR products were checked using agarose gels stained with ethidium bromide. Fluorescently labeled PCR products were quantified on an ABI Prism 3100 Genetic Analyzer (Applied Biosystems, Inc.) with a known concentration of a 6-Fam-labeled *nosZ* fragment obtained from the amplification of *P. stutzeri* DNA.

Samples for T-RFLP analysis consisted of approximately 1 ng of fluorescently labeled *nosZ* PCR products (digested at 37°C in 25-µl aliquots with 6.25 U of restriction endonuclease for 3 h) followed by heat inactivation at 65°C for 15 min. Three T-RFLP profiles were generated per sample unit in separate reactions with the endonucleases *CfoI* (an isoschizomer of *HhaI*), *MspI*, and *RsaI* (Roche Co.). Restriction digests (20 µl) were purified using a Sephadex G-50 column (Amersham Bioscience) and dried as pellets in a SpeedVac centrifuge. Dried pellets were rehydrated with Hi-Di formamide (9.92 µl) and an X-rhodamine MapMarker 1000 (0.08 µl) internal lane size standard. The mixture was heated at 96°C for 3 min and cooled on ice for 5 min prior to running of the samples in 96-well microtiter plates on an ABI Prism 3100 Genetic Analyzer with an injection time of 22 s, a run time of 54 min, and a 36-cm-long capillary array containing ABI Prism 3100 POP-4 polymer.

**Analysis of T-RFLP data.** Sizes and relative abundances of terminal restriction fragments (T-RFs) were quantified using GeneScan version 3.5 software (Applied Biosystems, Inc.). T-RFs with signals above a detection limit of 10 relative fluorescent units were summed for each profile; only fragments with a signal above 1.5% of the sum of all peaks were included in the analyses. The peak areas of T-RFs that differed in size by ≤1.5 bp in an individual profile were summed and considered as one fragment. Sequencing of clones indicated that almost all individual T-RFs with this variation were the same size (in base pairs). Each fragment was expressed as a percentage of the total peak area for all peaks in a profile (i.e., proportional abundance) or scored as either present or absent.

PC-ORD version 4.01 software (B. McCune and M. J. Mefford, PC-ORD for Windows: multivariate analysis of ecological data, 4.01 ed, MjM Software, Gleneden Beach, Oreg., 1999) was used for all multivariate statistical analyses of the *nosZ* T-RFLP profiles. Differences in community composition were assessed graphically using the ordination method of nonmetric multidimensional scaling (NMS) calculated on the basis of Sørensen's distance (19, 23). NMS axes summarize community relatedness among samples on the basis of distance measure and rank transformation (23). NMS was constrained to two ordination axes, with a random starting configuration, 300 iterations, an instability criterion of 0.0001, and 40 runs with the real data and 50 runs with randomized data; Monte Carlo tests of the real data versus the randomized data were used to assess axis significance. Functional variables (e.g., DEA and nitrification potential) were correlated with the NMS axes to evaluate relationships between overall community composition and functioning. Individual T-RFs were also correlated with NMS axes (35). Nitrification potential and DEA values were log<sub>10</sub> transformed for the correlation analysis, because the rates differed by an order of magnitude. Correlation coefficients between functional variables and NMS axes were dis-

played as vectors radiating from the center of the plot. Each vector was calculated as the hypotenuse of a right triangle, with each side of the triangle representing the correlation coefficient of the activity represented by axes one and two (23). The vector angle was a function of the relative proportions between the correlation coefficients (23). A multiresponse permutation procedure (MRPP) using Sørensen's distance and rank transformation was used to test for significant differences in community composition between groups defined by vegetation type and site (23, 24). The MRPP *A* statistic represents effect size or within-group relatedness relative to that expected by chance alone and is somewhat analogous to an  $r^2$  value (23). To identify T-RFs that differentiated denitrifier communities by vegetation type and site, indicator species analysis (7) was run with 1,000 randomizations in the Monte Carlo test.

**nosZ cloning, screening, and sequencing.** *nosZ* fragments generated from the PCR of soil DNA were cloned using a pGEM-T Easy Vector system (Promega) according to the manufacturer's instructions. Template DNAs for the PCR were pooled from two representative samples of each vegetation type. The PCR was run as described above except with 30 cycles. Prior to cloning, the 700-bp *nosZ* fragments were excised from an agarose gel and purified (Qiagen). Clones were screened using *CfoI*, *MspI*, and *RsaI* T-RFLPs as described above for direct soil profiles (defined here as T-RFLP profiling). Genotypic diversity was analyzed using species area curve analysis (treating a unique *nosZ* clone as a species) as implemented in PC-ORD version 4.01 software (McCune and Mefford, PC-ORD for Windows). Species area curve analysis consisted of randomly subsampling the clone library data set 500 times at each level of clone screening, which is analogous to generating rarefaction curves.

Plasmids were prepared using QIAprep Minipreps (Qiagen) according to the manufacturer's instructions. Both strands of the *nosZ* fragment were sequenced (using the SP6 and T7 promoter primers flanking the PCR insert in the plasmid) for 34 clones. Sequencing reactions were done with ABI Prism BigDye terminator cycle sequencing and the samples were analyzed on an ABI Prism 3100 Genetic Analyzer.

**nosZ phylogenetic analyses.** Analyses of *nosZ* sequences were carried out for the translated amino acid sequences by using PHYLIP version 3.573 software (J. Felsenstein, PHYLIP [phylogeny inference package], Department of Genetics, University of Washington, Seattle, 1993 [distributed by the author]) and TREE-PUZZLE version 5.0 software (42). With the use of the method of Friedrich (8), *nosZ* sequences (with the exception of that of *Ralstonia eutropha* ATCC 17699 [9]) were selected from cultured strains if their phylogenetic position could be confirmed using publicly available strain-specific small-subunit (SSU) rRNA sequence data and the Ribosomal Database Project's hierarchy browser and sequence matching tool (22). Amino acid sequences of *nosZ* were aligned using Clustal\_X (44), excluding residues corresponding to the primer sequences. Distance, parsimony, and maximum likelihood methods were used to infer phylogenetic relationships. Dayhoff's 001 substitution matrix (6) was used to measure evolutionary distance with PROTDIST (PHYLIP) software, and trees were constructed using the neighbor-joining algorithm (PHYLIP). PROTPARS (PHYLIP) software was used for parsimony and TREE-PUZZLE software was applied for maximum likelihood analysis, using the JTT (15) amino acid substitution matrix for evolutionary distance. Bootstrap analysis of 1,000 replicates was carried out for neighbor-joining trees and of 500 replicates was carried out for parsimony trees (PHYLIP).

**Nucleotide sequence accession numbers.** The GenBank accession numbers of the H. J. Andrews *nosZ* sequences are AY259180 to AY259212.

## RESULTS

**Denitrification potentials.** Strong functional differences were found between meadow and forest soils at the two sites (Table 1 and Fig. 1). DEA was generally an order of magnitude higher in the meadow than in the forest. This difference was highly significant at both sites ( $P < 0.0001$  for vegetation type [two-factor ANOVA]). The DEA levels (in units of nanograms of N gram<sup>-1</sup> of dry soil h<sup>-1</sup> [mean  $\pm$  1 standard error;  $n = 12$ ]) of the meadow soils were  $32.4 \pm 7.1$  for Carpenter and  $66.0 \pm 19.4$  for Lookout; the forest soils were  $1.6 \pm 0.5$  for Carpenter and  $7.0 \pm 1.4$  for Lookout. DEA showed a weak tendency to be higher at Lookout ( $P = 0.07$  for site [two-factor ANOVA]). The mean activity along the transects only changed significantly after the meadow-forest boundary (Fig. 1) was crossed.

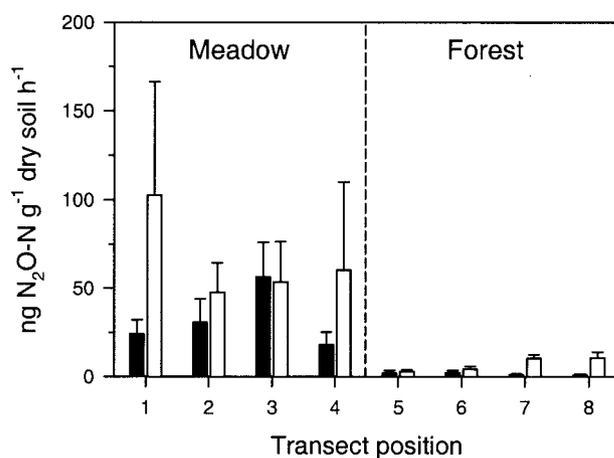


FIG. 1. Spatial variation of DEA in adjacent meadow and forest soils in the H. J. Andrews Experimental Forest. Bars represent the mean  $\pm$  1 standard error for each position along replicate transects at the Carpenter site (closed columns) and Lookout site (open columns) ( $n = 3$ ). The distance between transect positions was 20 m at Carpenter and 10 m at Lookout (see Materials and Methods).

**Amplification of *nosZ* from pure cultures and soil DNA.** The *nosZ* primers amplified the expected fragment from five positive controls possessing *nosZ*, including *Paracoccus denitrificans* ATCC 17741, *R. eutropha* ATCC 17699, "*Achromobacter cycloclastes*" ATCC 21921, *P. stutzeri* ATCC 14405, and *Pseudomonas aeruginosa*. The primers did not amplify DNA from two negative controls lacking *nosZ* (*Serratia marcescens* and *Burkholderia cepacia* cep31<sup>T</sup> [group I]). Yields of DNA extracted from the H. J. Andrews soil ranged from 7 to 35  $\mu$ g of DNA g<sup>-1</sup> of dry soil. The PCR products that were amplified from the soil DNA were of the expected size (700 bp). Of the 48 field replicates, 3 were excluded from subsequent analyses because of poor PCR amplification.

***nosZ* T-RFLP profiles.** Each *nosZ* T-RFLP profile from the H. J. Andrews soil consisted of one or two dominant T-RFs and several less-dominant T-RFs. Overall, a total of 61 T-RFs were detected, with 14 to 24 T-RFs for each endonuclease (Table 2). Of the 61 T-RFs, 4 (i.e., a 357-bp T-RF detected using *CfoI*, a 111- to 112-bp T-RF detected using *MspI*, and 666- and 700-bp T-RFs detected using *RsaI*), comprising 58% of the mean total fragment abundance of each profile, were dominant. These four were the only T-RFs that occurred in all samples. The rest of the T-RFs comprised a mean of 2 to 12% of the total fragment abundance of each profile and occurred in 2 to 98% of the samples; of these, 21 T-RFs occurred only once. Each endonuclease produced a mean of 2.7 to 7.1 T-RFs per profile, depending on the endonuclease, vegetation type, and site (Table 2). *CfoI* and *RsaI* tended to produce more fragments than *MspI* (Table 2). Meadow samples tended to have slightly more fragments in each profile on average, but this result was inconsistent among endonucleases and sites (Table 2).

**Differences in *nosZ* distribution and functional relationships.** Denitrifying community composition differed significantly according to vegetation type and site (Fig. 2 and Table 3). MRPP *A* statistic values of about 0.3 (Table 3) reflect strong differences between groups (23). Differences deter-

TABLE 2. Number of T-RFs in the *nosZ* T-RFLP profiles of H. J. Andrews soils

Site and vegetation type	<i>n</i> <sup>a</sup>	No. of <i>CfoI</i> T-RFs		No. of <i>MspI</i> T-RFs		No. of <i>RsaI</i> T-RFs		No. of combined T-RFLPs	
		Total	Mean (SE)	Total	Mean (SE)	Total	Mean (SE)	Total	Mean (SE)
Carpenter									
Meadow	12	12	7.1 (0.4)	6	4.2 (0.2)	13	4.9 (0.3)	31	16.2 (0.7)
Forest	10	11	5.7 (0.5)	10	4.2 (0.5)	10	5.0 (0.2)	31	14.9 (1.1)
Lookout									
Meadow	12	15	6.9 (0.6)	8	3.3 (0.4)	15	6.0 (0.7)	38	16.2 (1.4)
Forest	11	11	5.1 (0.2)	5	2.7 (0.2)	7	4.5 (0.2)	23	12.4 (0.4)
Total	45	24	6.2 (0.3)	14	3.6 (0.2)	23	5.1 (0.2)	61	15.0 (0.5)

<sup>a</sup> *n*, total number of field replicates analyzed.

mined on the basis of the proportional abundance of or the presence or absence of *nosZ* T-RFs were evident (Table 3). Four groups (corresponding to vegetation type and site) were determined on the basis of proportional abundance levels (Fig. 2). Differences by site (as determined on the basis of the presence or absence of *nosZ* T-RFs) were less apparent for the forest samples (Table 3). NMS values for axes one and two were statistically significant ( $P = 0.02$ ). Differences between vegetation type and site were also found by analyzing separate *CfoI*, *MspI*, or *RsaI* T-RFLP profiles, but these differences were not as strong as those seen for the combined profiles and relationships between vegetation type and site differed somewhat depending on the endonuclease used (data not shown). Ordination plots of *CfoI* profiles appeared most similar to the combined profile plots.

Functional variables were related to denitrifying community composition by plotting the relative magnitudes and directions of correlation coefficients ( $r^2$ ) between NMS axes and functional variables (Fig. 2). Examples of the strongest correlates are shown in Fig. 2. Nitrification potential and DEA were correlated with axis one in the direction of meadow samples (Fig. 2). Nitrification potentials and DEA levels were substantially higher in meadow than in forest soils (Table 1). DEA was also correlated with axis two in the direction of the Lookout meadow (Fig. 2). Net nitrification gave correlation coefficients almost identical to those for nitrification potentials (data not shown). Accumulation of  $\text{NH}_4^+$  (i.e., net ammonification) was correlated with axis one in the direction of forest samples (Fig. 2). Soil C/N ratios were correlated with axis one and two in the direction of Carpenter forest, where C/N ratios were highest (Fig. 2 and Table 1).

**Individual *nosZ* T-RFs.** The *RsaI* 666-bp T-RF level declined in proportional abundance from meadow to forest, whereas 700-bp *RsaI* T-RF levels increased from meadow to forest (Fig. 3). *RsaI* 666- and 700-bp T-RF levels were most strongly correlated with axis one (shown in the NMS plot in Fig. 2). *CfoI* 454-bp T-RF was found in every meadow sample but in none of the forest samples except for those taken near the boundary at Lookout (Fig. 3). Conversely, *RsaI* 451-bp T-RF was found preferentially in the forest (Fig. 3). In NMS plots calculated on the basis of the presence or absence of *nosZ* T-RFs (data not shown), *CfoI* 454-bp and *RsaI* 451-bp T-RF levels were most strongly correlated with axis one.

Indicator species analysis was used to further identify the T-RFs that were more prevalent according to either vegetation type or site (Fig. 4). This procedure examines the relative

abundance and presence or absence of individual T-RFs in a predefined group compared to those expected to occur by chance (23). A total of 29 significant indicator T-RFs ( $P < 0.05$ ) were found. In recognition of the several-orders-in-magnitude range in proportional abundance, T-RFs are shown in Fig. 4 on a  $\log_{10}$  scale to allow for inspection of less-abundant T-RFs. On this scale, relatively large differences in the proportional abundance of more dominant fragments appear smaller (Fig. 4).

***nosZ* clones.** Among 225 clones, 47 *nosZ* genotypes were identified on the basis of the *CfoI*, *MspI*, and *RsaI* T-RFLP profiles of each clone. Two-thirds of the library consisted of clones from meadow soils, and one-third consisted of clones from forest soils. A total of 24 genotypes occurred once and 8 occurred twice; the most dominant genotype occurred 31 times. On the basis of species area curve analysis, a mean of 21 genotypes was found after screening 50 *nosZ* clones, a mean of

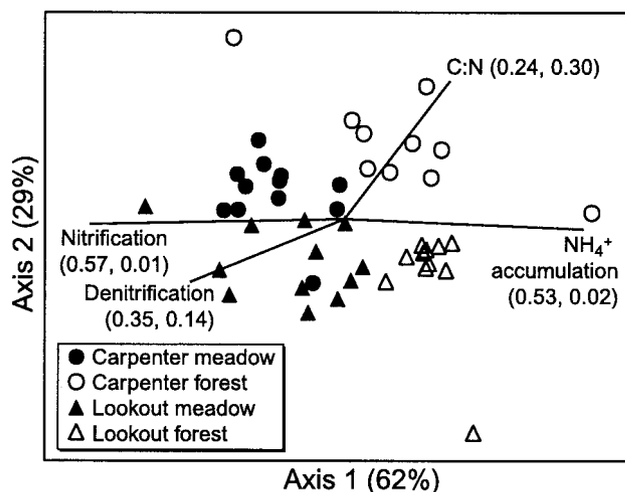


FIG. 2. NMS ordination of denitrifying community composition determined on the basis of proportional abundance of T-RFs in combined *CfoI*, *MspI*, and *RsaI* *nosZ* T-RFLP profiles. Points represent meadow-to-forest transect samples. The percentage of variation represented by each axis is shown in parentheses next to the axis name. Vectors show the directions and relative magnitudes of correlation coefficients ( $r^2$ ) between NMS axes and functional variables;  $r^2$  values (in the order  $r^2$  of axis 1,  $r^2$  of axis 2) for the correlation between the functional variable and axis one or two are shown in parentheses. Nitrification,  $\log_{10}$  of nitrification potential; denitrification,  $\log_{10}$  of DEA; C:N, soil carbon-to-nitrogen ratio;  $\text{NH}_4^+$  accumulation, net  $\text{NH}_4^+$  production (i.e., ammonification).

TABLE 3. Results of MRPP testing for significant differences between predefined groups determined on the basis of *nosZ* T-RFLP profiles (*CfoI*, *MspI*, and *RsaI* results have been combined)

Test and sites	<i>A</i> statistic <sup>a</sup>	
	Difference in proportional abundance	Difference in presence vs absence
Comparison by vegetation type		
Carpenter meadow vs Carpenter forest	0.29 ( $P < 10^{-5}$ )	0.23 ( $P < 10^{-5}$ )
Lookout meadow vs Lookout forest	0.26 ( $P < 10^{-5}$ )	0.34 ( $P < 10^{-3}$ )
Comparison by site		
Carpenter meadow vs Lookout meadow	0.18 ( $P < 0.001$ )	0.17 ( $P < 0.001$ )
Carpenter forest vs Lookout forest	0.24 ( $P < 10^{-5}$ )	0.05 ( $P = 0.02$ )
Comparison of randomized data <sup>b</sup>		
Carpenter meadow vs Carpenter forest	-0.04 ( $P = 1.0$ )	-0.01 ( $P = 0.7$ )

<sup>a</sup> The *A* statistic measures within-group relatedness compared to that expected by chance; *A* statistic values of about 0.3 reflect strong differences between groups (23).

<sup>b</sup> T-RFLP profiles from the Carpenter site were randomized among sample units.

31 genotypes was found after screening 100 clones, and a mean of 44 genotypes was found after screening 200 clones. Abundances of T-RFs in the clone library generally agreed with proportional abundances of T-RFs in direct soil profiles. Of the 29 indicator T-RFs shown in Fig. 4, 21 were present in sequenced *nosZ* clones. Of the 34 sequenced clones, 1 was nonspecific for *nosZ* (i.e., not identifiable using GenBank's BLAST search) but the T-RFs for this clone were not detected in direct soil profiles.

***nosZ* fragment size.** Actual T-RF size was determined by sequencing *nosZ* clones. Determining fragment size on the basis of T-RFLP profiling of the same clones was less precise and less accurate, but the results agreed within a few base pairs or less. In terms of accuracy, individual T-RFs >100 bp in size differed by less than 1 to 3 bp from the results obtained by sequencing (absolute mean difference, 0.9 bp). T-RFs less than 100 bp in size were 3 to 7 bp smaller than expected. In terms of precision, individual T-RFs differed in size by an average of

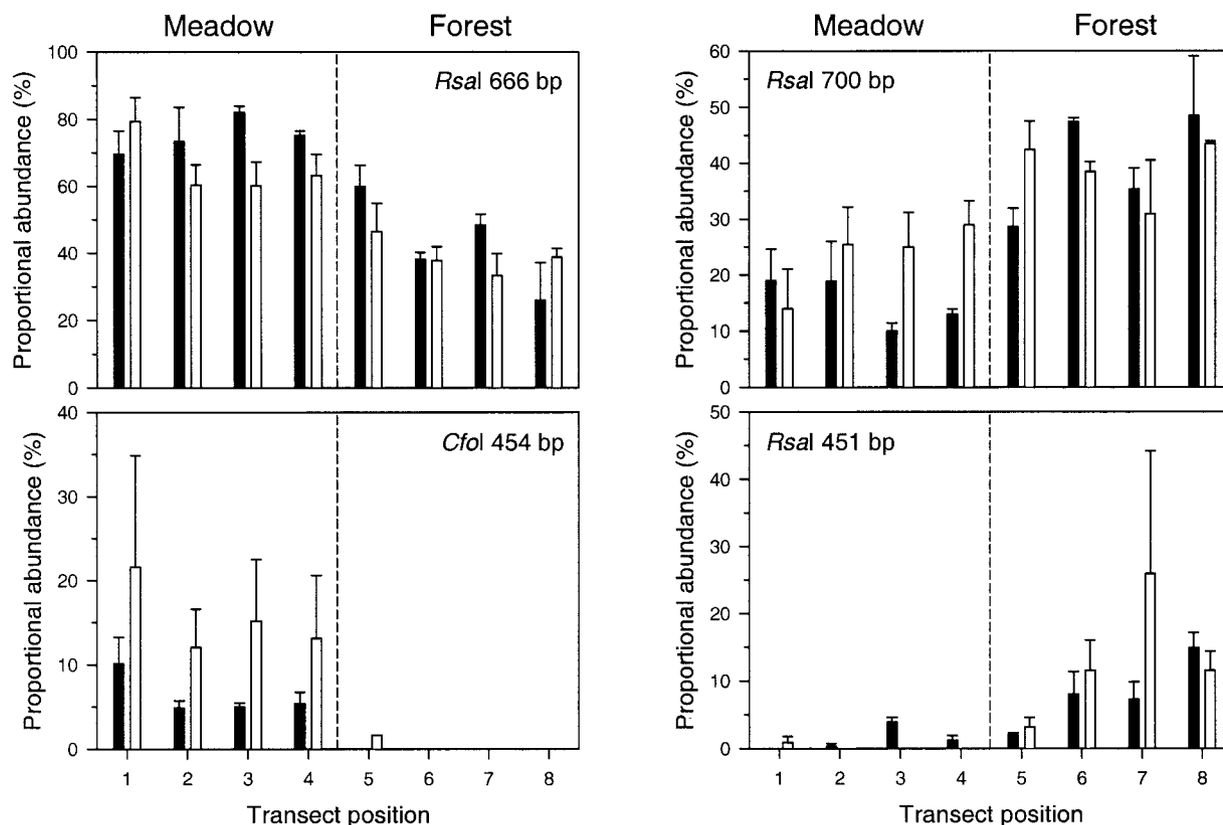


FIG. 3. Spatial variations of some key *nosZ* T-RFs from adjacent meadow and forest soils in the H. J. Andrews Experimental Forest. Bars represent the mean  $\pm$  1 standard error for each position along replicate transects at the Carpenter site (closed columns) and the Lookout site (open columns) ( $n = 3$  for all positions except  $n = 2$  for position 5 at Lookout and positions 6 and 8 at Carpenter). Each panel shows the results for an individual T-RF (identified by the corresponding restriction endonuclease and fragment size). The distance between transect positions was 20 m at Carpenter and 10 m at Lookout (see Materials and Methods).

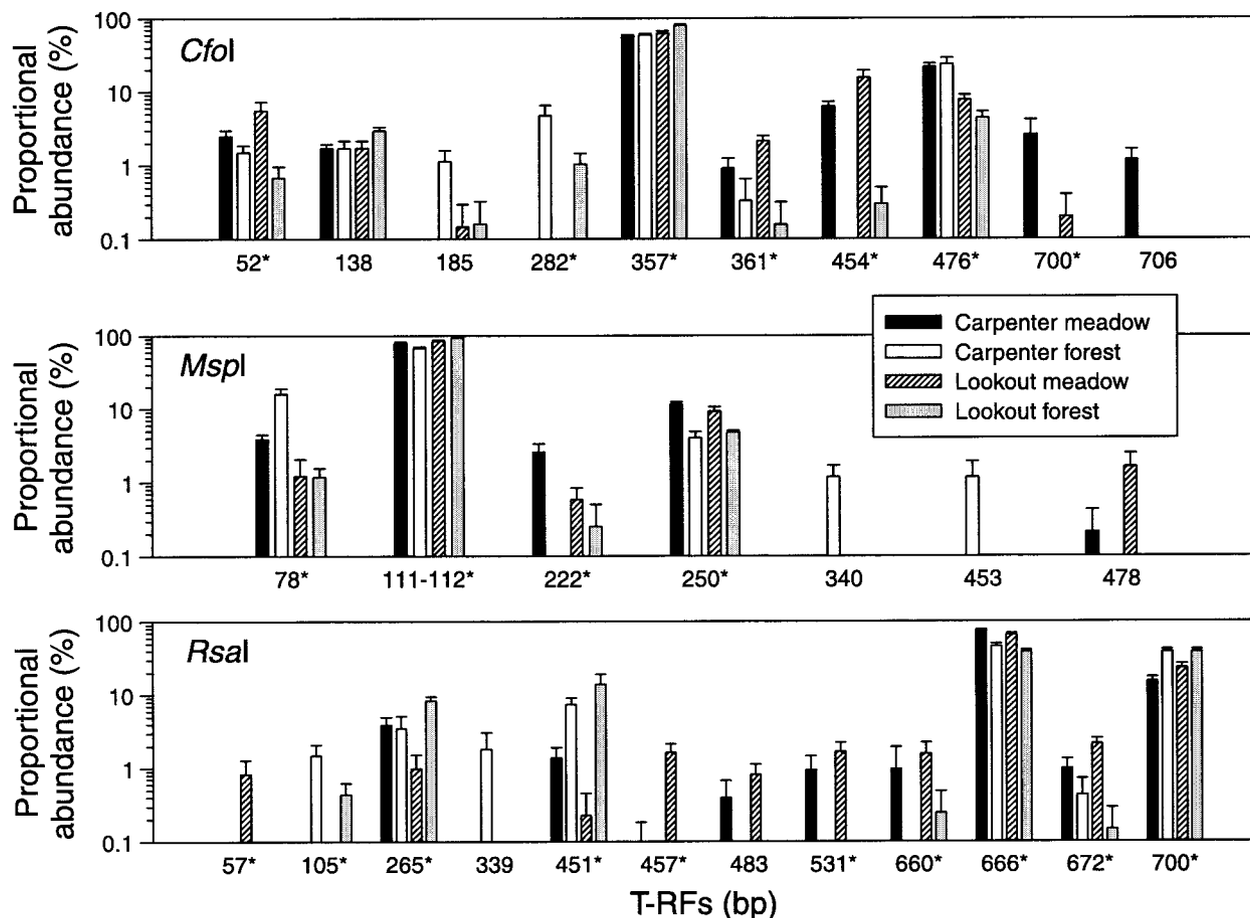


FIG. 4. *nosZ* T-RFs that differed significantly according to vegetation type or site as determined on the basis of indicator species analysis ( $P < 0.05$ ). Each panel is labeled with the name of the restriction endonuclease used to generate the T-RFs. T-RFs marked with an asterisk (\*) were present in sequenced *nosZ* clones.

1.5 bp over a range of 0.3 to 5.1 bp. The dominant *RsaI* 666-bp T-RF was identified at exactly that size in all 15 cases of sequencing, whereas with T-RFLP profiling the T-RF size differed over a range of 5.1 bp (from 664.0 to 669.1 bp). Six T-RFs (i.e., those obtained with *CfoI* [52 bp, 454 bp, and 476 bp], *MspI* [111 to 112 bp], and *RsaI* [666 bp and 700 bp]) and the two undigested PCR products (i.e., those 700 and 706 bp in size) consistently differed over a range of more than 1 bp in T-RFLP profiling. T-RFs of soil clones and direct soil T-RFLP profiles determined on the basis of T-RFLP profiling were in close agreement, differing by an absolute mean of 0.4 bp (data not shown).

***nosZ* phylogeny.** Phylogenetic analysis of *nosZ* sequences revealed five major clusters of denitrifying bacteria (Fig. 5). With the exception of those strains in cluster X, phylogenetic relationships among the cultured strains as determined on the basis of *nosZ* results were similar to taxonomic relationships determined on the basis of SSU rRNA results (Fig. 5). Tree topology was supported by all the phylogenetic analyses, with some differences in bootstrap support (Fig. 5). All the H. J. Andrews *nosZ* sequences, along with those of  $\alpha$ -Proteobacteria species (except those represented in cluster X) and most clones from a German forest soil and representatives of uncultured

marine bacteria, grouped in cluster A (Fig. 5). The sequences of the majority of the H. J. Andrews clones were most closely related to *nosZ* sequences of members of the *Rhizobiaceae* group and the clones from the German forest soil (Fig. 5). Separate from the grouping determined by *Rhizobiaceae nosZ* sequences, two minor branches of H. J. Andrews clones formed meadow- or forest-specific groups (Fig. 5). These clones had sequences with T-RFs found preferentially in meadow or forest in the direct soil T-RFLP profiles (Fig. 5). Two clones from the German forest soil formed a major cluster without any pure-culture sequences (Fig. 5, cluster Y), and the sequence of another clone grouped with the sequence of *Azospirillum irakense* (Fig. 5, cluster X). The majority of meadow and forest *nosZ* clones from the H. J. Andrews grouped with *Bradyrhizobium japonicum* USDA110 and *Rhodopseudomonas palustris* CGA009. These clones had T-RFs indicative of forest or meadow origin but were not distinguishable phylogenetically (Fig. 5).

## DISCUSSION

Shifts in the relative abundance of dominant organisms within prokaryotic functional groups, and not complete turn-

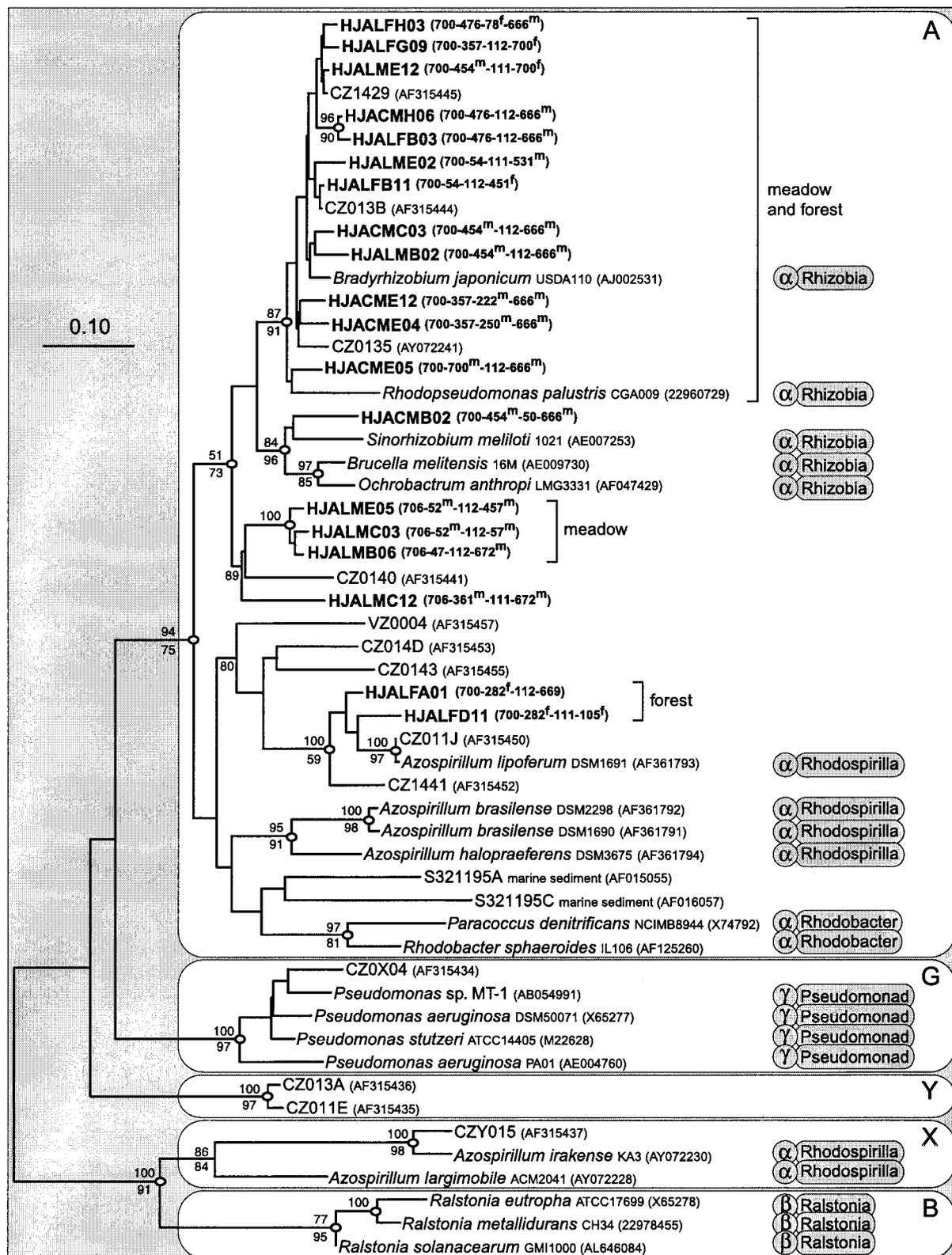


FIG. 5. Phylogenetic tree determined on the basis of evolutionary distances among *nosZ* amino acid sequences (208 positions) from soils, marine sediment, and cultured bacteria. The names of clones from the H. J. Andrews soils are shown in boldface characters, and the designations CZ and VZ identify clones from a German forest soil (36). Except for the H. J. Andrews clones, accession numbers are shown in parentheses. Taxonomic groups of cultured strains are identified to the subdivision and family level of *Proteobacteria* on the basis of SSU rRNA sequence analysis with the Ribosomal Database Project. Note that the suffix “-aceae” was dropped from the family name. Nodes with open ovals had >90% distance bootstrap support. Numbers above the branches are parsimony bootstrap values; numbers below the branches are maximum likelihood quartet-puzzling support values (analogous to bootstrap values). Sizes of the PCR products and T-RFs of the H. J. Andrews *nosZ* clones are shown in parentheses (in the order PCR product-*CfoI*-*MspI*-*RsaI*). T-RFs from direct soil T-RFLP profiles that were more prevalent in the meadow or forest are indicated with a superscript m for “meadow” and a superscript f for “forest.” Bar, evolutionary distance of 0.1.

over in community structure, can be associated with relatively large differences in biogeochemical processes (17, 20, 34). We found denitrification activity an order of magnitude higher in meadow than in forest soils and a significant shift in the proportional abundances of dominant denitrifying genotypes (i.e., *nosZ* *RsaI* 666- and 700-bp T-RFs) from meadow to forest soils. Several less-abundant denitrifying genotypes (e.g., a *CfoI* 454-bp T-RF and a *RsaI* 451-bp T-RF) differed in distribution between vegetation types. Denitrifying community composition was related to potential denitrification and nitrification activities and environmental factors, including soil C/N ratios.

As determined on the basis of a cross-section of studies (2, 10, 30, 47), mean DEA rates for temperate soils differ by 4 orders of magnitude (i.e., <1 to 3,000 ng of N g<sup>-1</sup> of soil h<sup>-1</sup>), with some of the lowest rates in mature coniferous forests in the Western Cascades (47) and highest rates in wet agricultural systems receiving N inputs (30). Although higher than those for the H. J. Andrews forest, our meadow values are in the low-to-medium range of DEA (i.e., 10 to 100 ng of N g<sup>-1</sup> h<sup>-1</sup>). DEA and net nitrification values were similar in magnitude (Table 1), suggesting that denitrification can serve as a sink for net nitrification in these soils.

Denitrifying communities in the meadow were much more active under denitrifying conditions compared to those in the forest (Fig. 1). Similar differences between meadow and forest soils were determined on the basis of potential N<sub>2</sub>O reductase activity (35). Because NO<sub>3</sub><sup>-</sup> or an intermediate in the denitrification pathway is necessary for sustained denitrification activity (11, 37), lack of available NO<sub>3</sub><sup>-</sup> could explain low denitrification activity in the forest soils. Nitrate availability in the forest appeared limited for several reasons, including very low nitrification potential rates and accumulation of primarily NH<sub>4</sub><sup>+</sup> in N mineralization incubations (26) (Table 1). Denitrifiers were present in the forest, probably because most denitrifiers are facultative aerobic heterotrophs which do not depend on denitrification activity for growth (46).

As determined on the basis of multivariate statistical analyses of *nosZ* T-RFLP profiles, denitrifying community composition differed among habitats. NMS and MRPP are valid methods for analyzing differences in community composition on the basis of T-RFLP data (23). Our statistical approach is further justified because it provides a quantitative framework for analyzing microbial community data and environmental interrelationships. For example, about 50% of the variation in functional variables was explained by *nosZ* distribution determined on the basis of cumulative *r*<sup>2</sup> values between process rates (or C/N ratios) and NMS axes (Fig. 2). Environmental factors (such as soil water and NO<sub>3</sub><sup>-</sup> content) often explain less than 50% of DEA variation or in situ denitrification (27, 30).

We amplified *nosZ* from pure cultures that represented a cross-spectrum of *nosZ* sequences from *Proteobacteria* species. In addition, we found divergent *nosZ* sequences in a poorly drained grassland soil that were most similar to those of *A. irakense* (53% similarity as determined on the basis of amino acid sequences), *Ralstonia solanacearum* (64% similarity), and *P. stutzeri* (88% similarity) (35). In analogy to the work of Friedrich (8), we investigated the lateral gene transfer of *nosZ* by comparing phylogenetic relationships on the basis of *nosZ* data and taxonomic relationships on the basis of SSU rRNA

data (Fig. 5). Although we detected a major discrepancy for two *Azospirillum* spp. (Fig. 5, cluster X), taxonomic and *nosZ* phylogenetic relationships were in reasonable agreement for the rest of the strains.

Therefore, the majority of denitrifiers detected in the H. J. Andrews soils may share taxonomic and functional similarities to representatives of *Rhizobiaceae* in  $\alpha$ -*Proteobacteria* species. Other studies have found numerous denitrifying sequences from soils that group closely with representatives of *Rhizobiaceae* (32, 33, 36). The *Rhizobiaceae* are metabolically diverse. Symbiotic N<sub>2</sub>-fixing *Rhizobium* and *Bradyrhizobium* spp. are known to denitrify (46), and legumes are common in the meadow at each site. However, some *Rhizobium* spp. from soil lack symbiotic genes and behave as free-living saprophytes (43). The *Rhizobiaceae* strains shown in Fig. 5 utilize a wide range of organic substrates, such as aromatic constituents present in polyphenols and lignin (29, 31). Furthermore, 2,4-D-degrading and oligotrophic bacteria that are closely phylogenetically related to *Bradyrhizobium* spp. have been isolated from soil (16, 38). None of the classical types of denitrifiers, such as pseudomonads or paracocci, were detected among sequenced clones from the H. J. Andrews soils or prevalent in other soils or marine sediments (32, 36, 39).

H. J. Andrews *nosZ* clones with T-RFs indicative of meadow or forest (i.e., *CfoI* 454-bp T-RFs and *RsaI* 451-, 666-, and 700-bp T-RFs) did not necessarily form separate phylogenetic groups (Fig. 5). Other studies comparing different habitats or treatments have resulted in similar findings (32, 33). In contrast, a few H. J. Andrews *nosZ* clones with T-RFs indicative of habitat (i.e., *CfoI* 52- and 282-bp T-RFs and *RsaI* 57-, 105-, 457-, and 672-bp T-RFs) formed monospecific phylogenetic groups differentiated according to location in meadow or forest (Fig. 5). The role of these specific groups in differentiating meadow and forest denitrification processes remains unclear.

In conclusion, denitrifying community composition and functioning appeared linked across meadow and forest soils. Denitrification activity was an order of magnitude higher in meadow than in forest soils, and denitrifying community composition differed between the vegetation types. As determined on the basis of *nosZ* phylogenetic analysis and T-RFLP profiles, organisms related to *Rhizobiaceae* may be prevalent denitrifiers in these mountain ecosystems.

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