Grass rhizosheaths: associated bacterial communities and potential for nitrogen fixation

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Grasses (Poaceae) include about 10,000 species and inhabit many varied habitats (Campbell 1985). About 70% of cultivated land is planted in crop grasses (Judd et al. 1999). Most grasses are not hosts to nitrogen-fixing bacteria, and growth of grasses may be inhibited due to a deficiency of nitrogen (Hopkins 2000). Grasses growing on relatively nutrient-poor soils are at a particular disadvantage, and growth may be greatly enhanced by soil bacteria (Kuske et al. 2002).

Although dinitrogen gas (N₂) comprises 78% of earth’s atmosphere, nitrogen is often limiting for organisms in many terrestrial ecosystems because plants and animals cannot utilize dinitrogen directly as a nitrogen source. During nitrogen fixation, dinitrogen gas is reduced to ammonia (NH₃), making nitrogen available for uptake by various organisms. Microbial nitrogen fixation accounts for 53.8% of terrestrial nitrogen fixation, which is greater than anthropogenic nitrogen fixation due to ammonia fertilizer production (38.5%) and abiotic (7.7%) nitrogen fixation (Schlesinger 1991). Diverse groups of Bacteria and some Archaea fix nitrogen, including many free-living soil bacteria and plant-associated symbionts such as the Rhizobiaceae, which inhabit the root nodules of legumes.

Microbial nitrogen fixation is catalyzed by the nitrogenase enzyme complex, which consists of a 2-subunit dinitrogenase component, encoded by the genes nifD and nifK, and a dinitrogenase reductase subunit, encoded by the nifH gene (Wang et al. 1988, Madigan and Martinko 2006). This nitrogenase complex also reduces acetylene to ethylene. Hence, the potential for nitrogen fixation may be assayed by incubating an anaerobic sample with acetylene and analyzing the sample for conversion of acetylene to ethylene (Knowles and O’Toole 1975, Turner and Gibson 1980, Boddey 1987, Capone 1993, Weaver and Danso 1994).

Some species of grasses possess rhizosheaths (Wullstein et al. 1979, Wullstein 1980, Buckley 1982). The rhizosheath consists of a layer of sand grains tightly bound to the root by mucilage, a mucopolysaccharide. The function of rhizosheaths in grasses is not well understood. Price (1911) suggested that the rhizosheath is important in water conservation, and this idea was supported by Young (1995), who stated that mucigel within the rhizosheath increases water-holding capacity. Other suggestions of rhizosheath function include harboring bacteria that fix nitrogen, thereby supplying the plant with ammonia in soils where nitrogen is limiting (Wullstein et al. 1979, Wullstein 1980, Buckley 1982).

The rhizosphere, or region of soil adjacent to a plant root, has been reported to have higher overall numbers of bacteria as well as
differences in the composition of bacterial species when compared to the bulk soil surrounding it (van Diepeningen et al. 2005). Gochnauer et al. (1989) determined bacterial numbers associated with rhizosheaths of maize. They reported similar numbers of viable bacteria in bare root regions and in roots with rhizosheaths, but they isolated more spore-forming bacteria and bacteria capable of growth in nitrogen-free media from sheathed regions. Evidence for nitrogen fixation in soil has been measured using an acetylene reduction assay (Turner and Gibson 1980). Andrews et al. (2003) reviewed nitrogen fixation in cultivated grasses and noted that few bacteria-grass interactions have been studied and those that have been studied have produced inconsistent levels of usable nitrogen.

The difficulty of studying organisms in the rhizosphere has been noted by Hawkes et al. (2007). Most work with rhizosphere microbes has been done on their relationships with agronomically important plant species (Hawkes et al. 2007). One recent study reported microbe-root interaction on 3 grass species in an arid grassland of Utah (Kuske et al. 2002). Bacterial communities in the soil are essential for biogeochemical cycling of elements such as carbon and nitrogen, especially in aridlands (Kuske et al. 2002). The study of bacterial communities in the rhizosphere of plants is complicated by the fact that perhaps 90% of the bacterial species present in the soil are not readily culturable using standard bacteriological techniques (Ward et al. 1990). Despite the pitfalls of culture-based techniques, the composition and diversity of soil (and rhizosphere) microbial communities may be examined by extraction of total microbial DNA and PCR of selected genes.

The purpose of this study was to determine the abundance of bacterial communities on the rhizosheaths of grasses growing in sand dune environments. We hypothesized that there would be more bacteria in the root/rhizosheath complex than in the surrounding bulk soil. Secondly, we hypothesized that at least some of the bacteria associated with the root/rhizosheath complex would be capable of nitrogen fixation.

**METHODS**

**Study Site**

The study site (45°33.4′N, 103°29.6′W) is located in an area with a continental climate with large seasonal and daily temperature fluctuations. Elevation at the site is approximately 900 m. The nearest weather-reporting station (approximately 15 km ESE) recorded an average annual precipitation of 37.6 cm, with the greatest amount of rain in June (7.6 cm). Temperature extremes recorded from 1993 to 2005 ranged from 42 °C in July to −39 °C in January. The average wind velocity was 19.0 km · h⁻¹ (South Dakota Office of Climatology 2006). The soil present at the study site is characterized as Zeona loamy fine sand. Organic or mineral horizons were not observed in the active dunes. In flat areas the sand is approximately 150 cm deep, while some dunes are greater than 6 m in height (Johnson 1988). Vegetation across the region is a mixed grassland, with dominant plants on the dune tops including *Calamovilfa longifolia* (Hook.) Scribn., *Achnatherum hymenoides* (R. & S.) Barkw., *Psoralidium lanceolatum* (Pursh) Rydb., and *Rumex venosus* Pursh.

**Field Sampling**

Two transects of 100 × 2 m were established longitudinally along 2 dune tops in a sand dune area in Harding County in northwestern South Dakota. Due to the prevailing northwest winds, dunes (and thus, transects) run in a northwest to southeast direction. Plant specimens were obtained 7 June 2006. Entire plants were collected to ensure that the roots being studied were properly identified. The plants collected for this study were *A. hymenoides* (R. & S.) Barkw., *C. longifolia* (Hook.) Scribn., *Hesperostipa comata* (Trin. & Rupr.) Barkw., and *Pascopyrum smithii* (Ryd.) A. Löve. Due to the sparse vegetation on the dune tops, percent cover was not measured. For enumeration of bacteria by scanning electron microscopy (SEM) and by culturing, rhizosheaths of 6 plants of each species as well as ten 20-g samples of bulk soil were collected. Bulk soil samples used for determination of soil bacteria were collected from approximately 5–10 cm below the surface of the dune tops at least 1 m away from the nearest visible plant stem.

Fourteen samples were collected from each transect for soil analysis, 7 at 0–15 cm and 7 at 15–46 cm. Samples from each transect were pooled for determination of percent organic matter (determined by weight loss upon ignition) and nitrate (determined from 0.01 M
Al₂(SO₄)₃ 0.02 M H₃BO₃), phosphorus (determined by 0.5 M NaHCO₃), and potassium concentrations (determined by 1 N NH₄OAc) by the Soil Laboratory at South Dakota State University. Soil particle surface area was estimated by 2 methods to allow comparisons of our analysis to published values of bacterial abundance. Soil particles in 2 samples of sand were weighed. The soil was transferred to a microscope slide with 2-sided adhesive tape, and the number of particles and longest diameter were determined by visual observation under a dissecting microscope with an ocular micrometer. Calculations were done for both spherical and cuboidal particle shapes. In the second method, the total mass of the sample was divided by the number of particles to find the average mass per particle. This mass was then divided by 2.65 g · cm⁻³, the density of quartz, to determine an average particle volume. The surface of a particle was again calculated by assuming the particle was either a simple cube or a sphere. The averaged area was multiplied by the total number of particles in the sample and divided by the total mass of the sample to arrive at average surface area per gram of soil.

**Root and Rhizosheath Measurements**

Using a dissecting microscope for observation, we measured rhizosheath diameters with a micrometer for 5 samples from each of 3 grass species with the largest rhizosheaths (A. hymenoides, C. longifolia, and H. comata). The soil particles of the rhizosheath were carefully removed, and the root diameters (excluding root hairs) were also measured with a micrometer.

**Enumeration of Bacteria**

**SCANNING ELECTRON MICROSCOPY.**—Roots and rhizosheaths of the 4 different species of grasses collected from the sand dune sites were prepared for observation with SEM. Portions of roots with rhizosheaths were preserved in 70% ethanol and dehydrated in an ethanol series. Liquid carbon dioxide was substituted for ethanol, and in a critical point dryer (E3000 series II, Polaron Equipment Ltd., Watford Hertfordshire, U.K.) the temperature and pressure were increased to exceed 31 °C and 7.39 MPa (critical point of carbon dioxide), respectively, to prevent damage by dehydration at ambient temperature and pressure.

The specimens were then coated with gold in a sputter coater (Effacoater 18930, E. F Fullan Co, Clifton Park, NY) and examined using a scanning electron microscope (5600LV, JEOL USA, Peabody, MA) to observe rhizosheath structure and to quantify numbers of bacteria found within the rhizosheath complexes. We compared rhizosheaths of different grass species; the overall number of bacteria found; and the size, shape, and location of bacteria within the rhizosheath (on sand grains, on root hairs, or other). Bacteria counts from bulk soil collected from areas without observable plant roots were also measured by counting cells on soil particles on scanning electron micrographs. These counts were compared to counts from the rhizosheath samples. Bacterial numbers were quantified following the procedure outlined in Barnes et al. (2005). We recognize that SEM provides images of only surface structures but is valuable as it can provide images of microorganisms that cannot be cultured on standard media.

**BACTERIAL CULTURE.**—One gram of bulk soil or rhizosheath (including root) from a grass was vortexed at high speed for one minute in 10 mL of autoclaved tap water, diluted in autoclaved water, and plated onto R2A agar (Rea-soner and Geldreich 1985) containing 40 mg · mL⁻¹ cycloheximide to inhibit fungal growth. Plates were incubated at 25 °C for 4 days.

**Acetylene Reduction to Ethylene**

The nitrogenase enzyme reduces acetylene to ethylene (Turner and Gibson 1980). Therefore, we evaluated the nitrogen fixation potential of the rhizosheath complex by assaying for this activity. We incubated rhizosheath samples in an atmosphere of nitrogen and 10% acetylene and detected the percentage of acetylene that was converted to ethylene as an assay for potential nitrogenase activity. In this assay the rhizosheath and root material could not be easily separated, so 4.0 g of material containing both root tissue and rhizosheath were sealed in vials along with 600 μL of sterile deionized water. Since Boddey (1987) indicated that nitrogenase activity associated with grasses is sensitive to O₂, the vials were evacuated and filled with N₂ gas at ambient pressure 3 times to completely remove O₂. After the final purge and refill, 0.10 volume of acetylene was added to the vial to provide root material with a 90% N₂:10% acetylene atmosphere, which is sufficient to achieve maximum reduction in grass.
species (Boddey 1987). Ethylene production undergoes an unexplained lag period (Boddey 1987), so samples were incubated at room temperature for 8 days. Two additional vials containing only N$_2$ and acetylene were also incubated during this time to act as controls.

After the incubation period a 100-μL sample of gas from each vial was injected into a GS-Q column (J&W Scientific, Folsom, CA) in a gas chromatograph (Agilent 6890N, Agilent Technologies, Inc., Santa Clara, CA) equipped with a flame ionization detector (FID). In this column any ethylene is cleanly separated from the relative amounts of the 2 gases can be determined from the output of the FID detector.

**Extraction of Microbial DNA and Amplification of nifH**

The presence of the genes for dinitrogenase reductase (nifH) in bacteria associated with rhizosheaths was tested by extraction of microbial DNA and PCR amplification of nifH. Genomic DNA was extracted from 4.0 g of soil from rhizosheaths (grass roots and adherent soil particles) of C. longifolia, P. smithii, A. hymenoides, and H. comata, as well as from 2 samples of bulk soil. DNA was extracted using the methods of Zhou et al. (1996), except that the incubations of soil with the detergent/proteinase K extraction buffer were 30 minutes instead of 2 hours. Crude DNA was then further purified with a 0.6-mL spin-column containing Sepharose 4-B, followed by a spin-column containing polyvinyl polypyrrolidone.

We amplified microbial nifH genes to facilitate nifH cloning and sequencing. Polymerase chain reaction (PCR) was performed with 1–5 ng of target DNA; 50 pmoles of each primer; dATP, dCTP, dGTP, and dTTP at 0.2 mM each; and 0.3 U Taq DNA polymerase (Eppendorf Inc., Hamburg, Germany) in a reaction buffer with 50 mM KCl, 10 mM Tris-Cl (pH 8.3), 0.2 mg · mL$^{-1}$ BSA, and 2.0–2.5 mM magnesium acetate, in a total volume of 25 μL. PCR reactions were performed in a Minicycler (MJ Research, Inc., Beverly, MA) or in a GeneAmp 9600 (Perkin-Elmer, Norwalk, CT) thermal cycler. A set of primers, nifH-g1-f (5’GGT-TGTGACCCGAAAAGCTGA3’) and nifH-g1-r (5’GGCTACATGGCCTACATCTC3’), was used (Buergmann et al. 2004) for PCR. After initial denaturation at 94 °C for 4 minutes, 50 cycles of 94 °C (denaturation) for 25 seconds, 55 °C (annealing temperature) for 30 seconds, and 72 °C (extension) for 40 seconds were performed before a final incubation at 72 °C for 10 minutes.

**Cloning and Sequence Analysis**

After extraction of microbial DNA and amplification of bacterial nifH genes, nifH PCR products were cloned into plasmid vectors so that the nifH PCR products from different bacterial species could be separated from each other and sequenced. Microbial nifH PCR products were purified and cloned into a plasmid vector to facilitate DNA sequencing. PCR products (350 bp) amplified from DNA of bulk soil and rhizosheaths of A. hymenoides were isolated using agarose gel electrophoresis. The PCR products were extracted from excised gel slices using a Qiagen QIA II kit (Qiagen, Inc., Valencia, CA) and the products were cloned into the pCR2.1 plasmid vector by using the TOPO-TA cloning kit and then transformed into Escherichia coli strain TOP-10 as directed by the manufacturer (Invitrogen, Inc., Carlsbad, CA). Transformed E. coli cells were plated out on McConkey agar with 0.1 mg · mL$^{-1}$ ampicillin so that cells transformed with the pCR2.1 plasmid vector could be selected by disruption of the plasmid β-galactosidase gene. Individual clones of E. coli were grown overnight at 37 °C in 3.0 mL LB medium with 0.1 mg · mL$^{-1}$ ampicillin and plasmid DNA harvested with a commercial miniprep kit (Qiagen, Inc., Valencia, CA). The cloned PCR products were sequenced using the M13–20 primer and the Big Dye version 3.1 kit (ABI, Inc., Foster City, CA). Sequencing reactions were analyzed on an Applied Biosystems ABI Prism 310 Genetic Analyzer at the Center for Conservation of Biological Resources at Black Hills State University, Spearfish, SD. Sequences were edited and aligned using the CLUSTAL W program at the University of California San Diego supercomputing center (http://workbench.sdsc.edu/). NifH sequences from soil and rhizosheaths were analyzed using the BLAST program (Altschul et al. 1990) at NCBI (http://www.ncbi.nlm.nih.gov/BLAST/).

**Statistical Analyses**

Root and rhizosheath comparisons were analyzed using one-way ANOVA. Root diameters were compared as well as rhizosheath diameters less root diameters. Bacterial enumeration was evaluated using two-way ANOVA.
to compare counts obtained for cultured bacteria (number \cdot mm^{-2}), and counts obtained using SEM observations (number \cdot mm^{-2}), from both rhizosheaths and bulk soil samples from interspaces. Tukey-Kramer post hoc tests were completed after the ANOVA. Acetylene reduction was analyzed using natural log-transformed data in a one-way ANOVA for comparisons between each of the grass species tested and the control vials.

RESULTS

Soil Characteristics and Root Diameters

Bulk soils from dune tops from the 2 transects in the study have an average pH of 7.9 and 8.0, respectively, 1.3% and 1.2% organic matter; 3.3 and 1.7 ppm NO₃, 3 and 4 ppm P, and 140 and 123 ppm K. One gram of dune topsoil had an average surface area of 1.9 \times 10^{4} \text{ mm}^{2}.

A comparison of bare-root diameters of the 3 plant species showed no significant differences among the root diameters, but rhizosheath diameters less the root diameters were significantly larger (ANOVA: \( F_{5, 24} = 8.4, P = 0.0001 \)). Rhizosheath diameters in A. hymenoides (\( \bar{x} = 1.51 \text{ mm}, s_{\bar{x}} = 0.16 \)) were significantly larger than they were in H. comata (\( \bar{x} = 0.69 \text{ mm}, s_{\bar{x}} = 0.06 \)) and C. longifolia (\( \bar{x} = 0.80 \text{ mm}, s_{\bar{x}} = 0.13 \)); the latter two were not significantly different (\( P > 0.05 \)).

Bacterial Enumeration

Enumeration of bacteria on rhizosheaths (example in Fig. 1) of the 4 grass species observed with SEM showed a range of bacterial density. The species with the most bacteria was C. longifolia with 529.1 bacteria \cdot mm^{-2} (\( s_{\bar{x}} = 135.1 \)), while soil from interspaces between plants averaged only 45.9 bacteria \cdot mm^{-2} (\( s_{\bar{x}} = 7.3 \)). Numbers of culturable bacteria from rhizosheaths, as estimated by bacterial colonies recovered on R2A media, were about 10-fold higher in the rhizosheaths of grasses than in bulk soil. The highest number of culturable bacteria, \( 9.9 \times 10^{7} \text{ CFU} \cdot g^{-1} \) or 5178 CFU \cdot mm^{-2} (\( s_{\bar{x}} = 1583 \)) was found in rhizosheaths

Fig. 1. Paspopyrum smithii: Scanning electron micrograph of root cross section and surrounding rhizosheath with attached soil particles.
of *C. longifolia*, while bulk soil contained $0.3 \times 10^7$ CFU $\cdot$ g$^{-1}$ or 171 CFU mm$^{-2}$ ($s_\chi = 66.8$).

A two-way ANOVA showed that cultured bacterial cell numbers were significantly larger than numbers observed using SEM ($F_{1,1119} = 117, P \ll 0.0001$), and that bacterial abundance varied by grass species and bulk soil ($F_{3,4119} = 16.3, P \ll 0.0001$). Tukey-Kramer post hoc tests showed that all 4 grass species had significantly more bacteria than the bulk soil. In addition, *C. longifolia* rhizosheaths had more bacteria than *A. hymenoides*, *H. comata*, or *P. smithii*, and the latter 3 grasses were similar in bacterial numbers. We believe that a significant interaction between grass species and observational type ($F = 11.2, P \ll 0.0001$) was observed because SEM grossly underestimated bacterial abundance on rhizosheaths, causing an inability to distinguish amongst the abundance of bacteria on rhizosheaths of various species of grasses.

Percentages of bacterial morphotypes observed by SEM varied among grass species (Table 1). Most species of grass had predominantly bacillus-type bacteria (Fig. 2), with the exception of *P. smithii* and bulk soil, which had higher percentages of coccus-type bacteria (Fig. 2). Actinomycetes (Fig. 3) were uncommon except in *A. hymenoides* rhizosheaths.

### Table 1. Percent frequency of bacterial morphotypes, as visualized by SEM, observed in rhizosheaths of 4 grass species and in bulk soil $\geq 1$ m from any plants.

<table>
<thead>
<tr>
<th>Grass species</th>
<th>Samples (n)</th>
<th>Bacillus (%)</th>
<th>Coccus (%)</th>
<th>Actinomycete (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Achnatherum hymenoides</em></td>
<td>11</td>
<td>49</td>
<td>23</td>
<td>27</td>
</tr>
<tr>
<td><em>Calamovilfa longifolia</em></td>
<td>22</td>
<td>95</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td><em>Hesperostipa comata</em></td>
<td>14</td>
<td>70</td>
<td>26</td>
<td>5</td>
</tr>
<tr>
<td><em>Pascopyrum smithii</em></td>
<td>22</td>
<td>35</td>
<td>59</td>
<td>6</td>
</tr>
<tr>
<td>Soil</td>
<td>38</td>
<td>16</td>
<td>84</td>
<td>0</td>
</tr>
</tbody>
</table>

Fig. 2. *Calamovilfa longifolia*: Scanning electron micrograph of rhizosheath of showing (A) rod- and (B) coccus-shaped bacteria.
Acetylene Reduction to Ethylene

The percent of acetylene converted to ethylene for samples from rhizosheaths of a particular grass species varied up to 3 orders of magnitude (Fig. 4), but due to large variance, differences in conversion among the grass rhizosheaths and controls were not significant (ln transformed, $F_{5,16} = 1.5, P = 0.25$). Individual sample values varied over 3 orders of magnitude, and the variances of these distributions were large. An additional control experiment was done where soil from a site with no plants growing within 1 m was also tested for acetylene reduction. After 24 hours the ethylene peak from the soil-only sample was 0.006% for the acetylene peak, and after 12 days the ethylene peak was 5.8% of the acetylene peak. Given these results, it was highly likely that all samples except one P. smithii sample (0.08%) and one A. hymenoides sample (0.09%) reduced acetylene to ethylene (Fig. 4). Interestingly, 2 other P. smithii samples also gave the largest values for acetylene-to-ethylene conversion.

PCR and Analysis of \textit{nifH} Genes

Bulk soil samples yielded about 50 ng DNA per gram soil, while rhizosheath samples yielded about 500 ng of combined plant and microbial DNA per gram soil. Using universal bacterial 16S rDNA primers as described by
Barnes et al. (2005), we successfully amplified fragments of 16S rRNA genes from 1.0 μL of undiluted purified DNA samples from bulk soil and rhizosheaths. Our success indicated that PCR inhibitors had been largely removed from the samples.

PCR of bulk soil with primers nifH-g1-f and nifH-g1-r produced a small amount of a 350 base-pair (bp) product only. PCR of rhizosheaths of C. longifolia, A. hymenoides, and P. smithii with the same primers yielded a mixture of PCR products of different sizes, including the expected 350 bp nifH product, but only A. hymenoides yielded enough 350 bp PCR product for cloning. No PCR products were obtained from H. comata rhizosheaths. Gel-purified 350 bp PCR products from bulk soil and A. hymenoides rhizosheaths were cloned into the pCR2.1 plasmid vector and the nifH inserts of plasmids were sequenced. A total of 19 separate nifH clones from bulk soil were sequenced. All 19 soil nifH sequences were identical (Sequence 1).

Among nifH sequences from cultured bacteria in GenBank, Sequence 1 (GenBank accession number EF158037) was most similar to nifH from Alcaligenes latus (gi|71040569|dbj|AB188122.1) (94.3% identity over 230 bp) and Ideonella sp. Long 7 (gi|33286718|gb|AY231580.1) (93.0% over 324 bp). For DNA from A. hymenoides rhizosheaths, a total of 16 clones with nifH sequences were obtained: 15 appeared identical to Sequence 1, while a second, Sequence 2, (GenBank accession number EF158038) was most similar (when compared with cultured isolates in GenBank) to nifH from Mesorhizobium loti MAFF303099 (gi|47118328|dbj|BA000012.4) (96.5% identity over 315 bp).

**Discussion**

In an attempt to better understand the occurrence of several species of grasses in relatively inhospitable environments of low-moisture and low-nutrient sand dunes, we hypothesized that plant occurrence may have been mediated by nitrogen-fixing bacteria associated with moisture-retaining mucilaginous rhizosheaths. We found strong support for this hypothesis, as the lowest numbers of bacteria found by both methods (SEM and culturing) were in the bulk soil. Furthermore, Alcaligenes latus (Malik et al. 1981) and Mesorhizobium loti (Kaneko et al. 2000) isolated from rhizosheaths are capable of nitrogen fixation.

Higher bacterial numbers in rhizosheaths are probably due to bacteria using root exudates as a source of carbon and to higher water potentials in these systems. The numbers of CFUs we observed in rhizosheaths were similar to those reported by Gochnauer et al. (1989) for maize. The number of cultured bacteria we determined from the soil (3.3 × 10^6 CFU g⁻¹) was similar to the averaged values reported by Kuske et al. (2002; 3.8 × 10^6 CFU g⁻¹) for an arid (21.4 cm annual precipitation) grassland system. Achnatherum hymenoides, the sole grass species included in both our study and in Kuske et al. (2002), had >10 times more CFUs per gram in the rhizosheaths we studied than it had in the rhizosheaths studied by Kuske et al. (2002; 44 × 10⁶ vs. 3.5 × 10⁶). Kuske et al. (2002) collected specimens from drier habitats with less organic matter (0.28%) and collected soil only via cores near plants.

In our study, entire plants were excavated and bacteria were cultured directly from the root/rhizosheath complex. One could conclude that higher bacterial concentrations are present in the root/ rhizosheath complex than are present in the surrounding soil.

We observed fewer bacteria per unit surface area by SEM than by culturing. Previous reports (Barnes et al. 2005) comparing SEM observations with culture of bacteria have shown greater numbers of bacteria observed with SEM than were observed in culture. In this study the greater number of bacteria determined by culturing may be due to our inability to observe the inner surfaces of the intact rhizosheath with SEM. Only exposed surfaces of sand grains, root hairs, or root surfaces were observed. One can assume that concentrations of bacteria are higher in the internal portions of the rhizosheath or that many bacteria may be obscured by mucilage or present in biofilms. Bacteria in rhizosheath specimens were probably at least partially obscured due to an obvious mucilaginous coating present on the rhizosheaths. Support for the protective nature of the root/rhizosheath complex comes from Pohlman and McColl (1982) who noted that even after surface sterilization in 1% sodium hypochloride for 5 minutes, roots were still capable of acetylene reduction. It seems likely that some of the bacteria contained within the rhizosheath are protected from
surface sterilization or that some bacteria are endophytic.

Acetylene reduction analysis indicates that nitrogen-fixing bacteria are associated with the rhizosheaths of several species of grasses we examined. We also observed that rhizosheaths from *P. smithii* produced more ethylene than those from *C. longifolia*. Some bacteria observed by SEM or by culturing are probably not capable of nitrogen fixation. The extreme sample-to-sample variation limits the utility of statistical comparisons among species. This high sample variation is not a problem unique to this study. Knowles and O’Toole (1975) indicated a large variation of acetylene reduction associated in replicate soil core assays. Whiting et al. (1986) reported a seasonal variation in acetylene reduction and noted that reduction is correlated with CO₂ and photosynthesis levels. Pohlman and McColl (1982) showed that higher rates of acetylene reduction are associated with moisture levels close to saturation. Weaver and Danso (1994) have noted that in grasses, the nitrogen-fixation system inevitably changes due to changes in gas concentrations with collection, changes in the microbial community, and changes in the production of ethylene that may occur independently of acetylene reduction.

We successfully amplified *nifH* gene sequences from both bulk soil and from rhizosheaths of *A. hymenoides* by using the *nifH*-g1-f and *nifH*-g1-r primers designed to amplify a fragment of *nifH* from the γ-subdivision of Proteobacteria. In both bulk soil and rhizosheaths of *A. hymenoides*, the community of nitrogen-fixing bacteria appears to be dominated by a single species, possibly belonging to a species close to *Alcaligenes latus* (β-subdivision of Proteobacteria). Another species, possibly *Mesorhizobium loti* (α-subdivision of Proteobacteria), was detected in rhizosheaths of *A. hymenoides*. We noted that PCR did not yield *nifH* products from *H. comata* rhizosheaths. This may be due to inhibitors present in soil and plant materials such as phenolic compounds, which prohibit amplification of *nifH* genes, or may indicate that there are too few nitrogen-fixing bacteria present to be detectable by ordinary methods. Despite the successful PCR amplification of 16S genes from rhizosheath DNA samples, the samples had spectrophotometric absorbance ratios (260 nm / 280 nm) of 1.44 to 1.85, indicating that some contaminants may still have been present.

This study enumerates the number of bacteria present on rhizosheaths of grasses growing in dry sandy areas of the north central Great Plains by SEM observation and bacterial culturing techniques. We found that there were more bacteria in the root/rhizosheath complex than in bulk soil. Also demonstrated is the ability of some of the microorganisms to reduce acetylene to ethylene, indicating their potential to fix nitrogen in the rhizosheath. *NifH* genes have been isolated from the bacteria, indicating the ability of the organisms to fix nitrogen.

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**LITERATURE CITED**


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