

Diversity of Nitrogen-Fixing Bacteria Associated with Switchgrass in the Native Tallgrass Prairie of Northern Oklahoma

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Switchgrass ($Panicum\ virgatum\ L$.) is a perennial C_4 grass native to North America that is being developed as a feedstock for cellulosic ethanol production. Industrial nitrogen fertilizers enhance switchgrass biomass production but add to production and environmental costs. A potential sustainable alternative source of nitrogen is biological nitrogen fixation. As a step in this direction, we studied the diversity of nitrogen-fixing bacteria (NFB) associated with native switchgrass plants from the tallgrass prairie of northern Oklahoma (United States), using a culture-independent approach. DNA sequences from the nitrogenase structural gene, nifH, revealed over 20 putative diazotrophs from the alpha-, beta-, delta-, and gammaproteobacteria and the firmicutes associated with roots and shoots of switchgrass. Alphaproteobacteria, especially rhizobia, predominated. Sequences derived from nifH RNA indicated expression of this gene in several bacteria of the alpha-, beta-, delta-, and gammaproteobacterial groups associated with roots. Prominent among these were *Rhizobium* and *Methylobacterium* species of the alphaproteobacteria, Burkholderia and Azoarcus species of the betaproteobacteria, and Besulfuromonas and Besul

Switchgrass (*Panicum virgatum* L.) is a warm-season C_4 grass that is native to the tallgrass prairies of North America, and it has been targeted for development as a bioenergy crop by the U.S. Department of Energy (1). Its features, such as perenniality, adaptation to diverse edaphic conditions, wide geographic distribution, growth on acidic soils, and mutualistic associations with soil microorganisms, make it an attractive choice for cultivation on marginal lands that are poorly suited to food crops (2-4).

Although switchgrass is thrifty in its use of N to produce biomass compared to other crops (2, 5), addition of N fertilizer enhances growth and generally is needed to maintain switchgrass productivity over multiple years (6, 7). However, application of N fertilizer increases production costs, reduces the energy balance of biomass production, and can be harmful to the environment through release of N-containing gases to the atmosphere and soluble N compounds to groundwater, which can lead to eutrophication (8). Biological nitrogen fixation has the potential to reduce the use and negative consequences of industrial N fertilizer and to put the nascent biofuels industry on a more sustainable path. Biological nitrogen fixation has been demonstrated in association with switchgrass (9, 10). However, to the best of our knowledge, the bacteria responsible for such activity are unknown, as is the diversity of potentially nitrogen-fixing bacteria associated with this species.

Nitrogen-fixing bacteria (NFB) associate with many grasses, including maize (11–13), rice (14), sugarcane (15), and *Miscanthus* species (16). These diazotrophs are endophytic, living between plant cells, and/or epiphytic, living on the surface of plant organs, and most do not elicit plant defense responses (17). However, the extent to which diazotrophic bacteria contribute to the N economy of grasses remains unclear.

Most soil microbes remain uncultured and largely uncharacterized, in part because appropriate culture conditions have not been found. Therefore, we used a culture-independent approach, based on PCR amplification and sequencing of *nifH* DNA (18, 19), to assess the diversity of potential diazotrophs associated with switchgrass. *nifH* has been used by many researchers to study total

and functional diversity of bacteria in various environments (15, 20–22), and phylogenies based on *nifH* and 16S rRNA generally are congruent (23, 24). Additionally, PCR amplification of cDNA derived from *nifH* RNA was used to determine whether any bacteria express this gene in association with switchgrass. The results of this work indicate diverse and abundant NFB associated with switchgrass and active expression of the *nifH* nitrogen fixation gene in some of these bacteria.

MATERIALS AND METHODS

Plant material. Switchgrass (*Panicum virgatum* L.) plants were harvested from separate locations in the tallgrass prairie (Oklahoma, USA) in April and July 2010. GPS coordinates of the sampling sites were between 36°38′38″N to 36°48′48″N latitude and 96°10′26″W to 96°26′23″W longitude (see Table S1 in the supplemental material). Plants were uprooted, and loosely attached soil was removed from roots by vigorous shaking. Roots and shoots were rinsed with tap water at the collection site to remove soil particles and were transferred to the laboratory on dry ice. Samples were stored at -80°C within 24 h of collection.

Nucleic acid extraction. Shoot and root pieces were pulverized in liquid nitrogen using a cryo-mill (6870 freezer mill; SPEX SamplePrep, USA). DNA was extracted from 0.5 g pulverized tissue using a modified cetyltrimethylammonium bromide (CTAB) method (25). Total RNA was extracted from 100 mg of pulverized tissue using an RNeasy plant minikit (Qiagen, USA) by following the manufacturer's instructions. Genomic DNA was removed by DNaseI treatment (Turbo DNase; Ambion, USA), followed by column purification of RNA using an RNeasy MinElute cleanup kit (Qiagen, USA). RNA was quantified using a Nanodrop spec-

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trophotometer (ND-100; NanoDrop Technologies, Willington, DE) and evaluated for purity with a Bioanalyzer 2100 (Agilent, Santa Clara, CA, USA). Checks for genomic DNA contamination of RNA were performed using *nifH* primers (mentioned below), 16S rRNA gene 27F (26), and 1492R primers (27).

PCR conditions for *nifH* **amplification.** A nested-PCR approach was used to amplify nifH gene fragments as described previously (28). Two degenerate primers, nifH 3 (5'-ATR TTR TTN GCN GCR TA-3') and nifH 4 (5'-TTY TAY GGN AAR GGN GG-3'), were used to amplify an ~460-bp region. One microliter of the resultant PCR product was used as the template to amplify an \sim 362-bp region using degenerate primers *nifH* 1 (5'-TGY GAY CCN AAR GCN GA-3') and nifH 2 (5'-AND GCC ATC ATY TCN CC-3'). All sample manipulations were performed under laminar flow to avoid airborne contamination. PCRs were initiated with denaturation at 95°C (5 min); 30 cycles of denaturation at 95°C (30 s), annealing at 55°C (1 min), and extension at 72°C (1 min); and a final extension at 72°C (10 min). Reverse transcription-PCR (RT-PCR) was done as described by Zani et al. (28), and cDNA synthesis was carried out using 1 µg of DNase-treated total RNA with primer nifH 3 and Super-Script III reverse transcriptase according to the manufacturer's instructions (Invitrogen, USA). Nested PCR was performed with the nifH 3 and nifH 4 primer combination, followed by the nifH 1 and nifH 2 combination. No-RT (i.e., no cDNA) controls were included for all RNA samples analyzed to check for genomic DNA contamination.

nifH amplification products were separated by electrophoresis through 2.0% agarose gels and visualized with Sybr green. Amplified fragments were excised from gels and purified using a Promega gel extraction kit (Promega, USA) and then cloned into the pGEM-T Easy vector (Promega, USA). Bacterial transformation was done using Z-competent cells (Zymo Research, USA). Forty-eight white colonies per sample were picked, and colonies were grown in 96-well plates.

nifH sequencing and phylogenetic analysis. Recombinant pGEM-T plasmids containing nifH fragments were sequenced using an ABI Prism 3100 Genetic Analyzer (Applied Biosystems, USA). DNA sequences were edited to remove vector sequences using GeneDoc (29), and identical sequences were grouped into operational taxonomic units (OTUs), using the FastGroupII tool (30) with the method of percent sequence identity with gaps of 100% identity. DNA sequences were translated into protein amino acid sequences using BioEdit software (31). Protein sequences were subjected to BLAST analysis (32) against GenBank, and nifH protein sequences from cultivated bacteria that showed the closest match to query sequences were included for phylogenetic analysis. OTUs representing 4 or more sequences were subjected to phylogenetic analysis. The MAFFT web interface was used for alignment (33).

Phylogenetic trees were constructed using MEGA (ver. 5.0) with the maximum likelihood method based on a Poisson correction model (34). The bootstrap consensus tree was produced from 1,000 replicates, and *Methanothermococcus okinawensis* NifH (accession no. NZ_AEDA01000001) was used as an outgroup.

Data analysis. *nifH* sequences from root DNA, root RNA (cDNA), and shoot DNA of individual plants were subjected to rarefaction analysis, and Shannon-Weaver and Simpson diversity indices were calculated using PAST statistical software (35). The percentage of total sequence diversity captured in sequenced clones of each organ for each plant was estimated by Good's method using the formula $[1 - (n/N)] \times 100$, where n is the number of clones appearing only once in a library and N is the total number of clones sequenced (36). Venn diagrams were drawn using Excel (Microsoft).

Accession numbers. Representative sequences from each OTU were deposited in GenBank under accession numbers KF541058 to KF541088, and the protein accession numbers are AIE38835 to AIE38865.

RESULTS

Diversity of *nifH* **sequences.** A total of 28 clone libraries were produced from *nifH* PCR amplicons derived from DNA of roots

and shoots separately from 10 independent plants and RNA from roots of 8 of these plants. Up to 48 clones were sequenced from each library, yielding a total of 1,087 high-quality *nifH* sequences. Based upon shared sequence identity, these sequences formed 52 distinct OTUs, which were tentatively associated with known species based on the highest sequence similarity (Table 1).

Estimations of species coverage, diversity, and dominance were calculated for each library separately (Table 2). Good's coverage averaged over 90% for root DNA libraries, over 95% for root RNA, and over 98% for shoot DNA libraries, indicating that the majority of nifH-containing and nifH-expressing bacteria associated with switchgrass plants were represented in the sequences obtained. Based on the number of OTUs represented by sequences in each library, bacteria containing *nifH* were significantly more diverse in roots than in shoots (P = 0.0145) of switchgrass plants, with an average of approximately 10 and 6 OTUs per plant, respectively (Table 2). About one-third of nifH-containing bacteria were found to express the gene in association with switchgrass roots (Table 1). Simpson's and Shannon's indices supported these conclusions about *nifH* bacterial diversity in the different sample types (Table 2). Overall dominance of one or a few OTUs was highest in root RNA, followed by shoot DNA and root DNA (Table 2). In three of the eight plants analyzed for RNA, only a single OTU was represented in the *nifH* sequences (i.e., dominance = 1).

Seven OTUs were common to all sample types, while 22 were specific to roots (DNA plus RNA), and 10 OTUs were specific to shoots (Fig. 1).

OTUs were matched to the most closely related known nitrogen-fixing species based on *nifH* sequence comparisons using the BLAST algorithm. A total of 23 OTUs representing 704 sequences appeared to derive from alphaproteobacteria, 9 OTUs (177 sequences) from deltaproteobacteria, 8 from betaproteobacteria (146 sequences), 6 from gammaproteobacteria (37 sequences), 2 from cyanobacteria (3 sequences), and 1 OTU each from *Firmicutes* (15 sequences) and *Verrucomicrobia* (3 sequences) (Table 1 and Fig. 2).

nifH diversity and expression in roots. Among the nifH sequences derived from root-associated DNA, more than 65% originated from alphaproteobacteria, 15% from betaproteobacteria, and 12% from deltaproteobacteria. The largest number of sequences in a single OTU (OTU-8; 81 sequences) were most similar (95 to 99% identity) to nifH from Sphingomonas azotifigens (accession no. BAE71134). Approximately one-third of all sequences (33%) were affiliated with various Bradyrhizobium species and distributed into 3 OTUs (1, 2, and 10) (Table 1 and Fig. 3). At the individual plant level, OTUs 1, 2, 4, 5, 8, and 10 (all alphaproteobacteria) were present in 7 to 8 out of 10 plants (see Table S2 in the supplemental material).

Among the *nifH* sequences derived from root-associated RNA, 58% matched sequences from alphaproteobacteria, 20% from betaproteobacteria, and 19% from deltaproteobacteria (Table 1 and Fig. 2). Over one-third of transcribed *nifH* sequences (88) fell into OTU-14, which is affiliated with *Rhizobium helanshanense* (ADP37388; 95 to 100% identity) or *Sinorhizobium meliloti* (CCH40450), while 38 sequences fell into OTU-12, affiliated with *Methylobacterium nodulans* (AAQ82902). Five OTUs contained sequences derived from root RNA but not root DNA (OTU-11, affiliated with *Rubrivivax gelatinosus*; OTU-13, affiliated with *Amorphomonas oryzae*; OTU-14, affiliated with *Rhizobium helanshanense/S. meliloti*; OTU-21, affiliated with *Desulfuromonas ace*

TABLE 1 Phylotypes of nifH associated with switchgrass

	nylotypes of <i>mj11</i> associated with switchgra	Accession no.	Similarity (%)	No. of	sequences			
Phylotype				Root		ol.		
	Closest relative			DNA	RNA	Shoot DNA	Total	Taxonomic description
OTU-01	Bradyrhizobium sp. strain BTAi 1	BAC07281	98-100	35	0	15	50	Alphaproteobacteria
OTU-02	Bradyrhizobium sp. strain MAFF 210318	BAC07283	94-98	46	2	71	119	Alphaproteobacteria
OTU-03	Mesorhizobium loti	BAF95636	100	0	0	4	4	Alphaproteobacteria
OTU-04	Methylocystis sp. strain LW2	AAK97418	97-98	24	2	8	34	Alphaproteobacteria
OTU-05	Burkholderia sp. strain WSM3937	ABX80637	93-98	58	17	1	76	Betaproteobacteria
OTU-06	Burkholderia sp. strain JPY105	AFM28513	97	7	7	3	17	Betaproteobacteria
OTU-07	Burkholderia sp. strain PTK47	AAU85620	96-100	3	0	20	23	Betaproteobacteria
OTU-08	Sphingomonas azotifigens	BAE71134	95-99	81	0	55	136	Alphaproteobacteria
OTU-09	Novosphingobium sp. strain Rr 2-17	ZP_10360790	98-99	29	0	0	29	Alphaproteobacteria
OTU-10	Bradyrhizobium japonicum	ACT67985	92-98	71	0	2	73	Alphaproteobacteria
OTU-11	Rubrivivax gelatinosus	BAE15985	90-94	0	5	0	5	Betaproteobacteria
OTU-12	Methylobacterium nodulans	AAQ82902	94-97	6	38	4	48	Alphaproteobacteria
OTU-13	Amorphomonas oryzae	BAF48338	96	0	1	7	8	Alphaproteobacteria
OTU-14	Rhizobium helanshanense	ADP37388	95-100	0	88	40	128	Alphaproteobacteria
OTU-15	Sinorhizobium sp. strain SCAU224	AFH96094	98	3	0	1	4	Alphaproteobacteria
OTU-16	Azospirillum lipoferum	ABG88868	96-100	3	2	22	27	Alphaproteobacteria
OTU-17	Methylomonas sp. strain MG30	CCH22595	93	5	0	0	5	Gammaproteobacteria
OTU-18	Pseudomonas stutzeri	CAC03734	97	7	4	0	11	Gammaproteobacteria
OTU-19	Klebsiella sp. strain AL060224_04	ACM68399	99–100	0	0	18	18	Gammaproteobacteria
OTU-20	Azoarcus sp. strain BH72	YP_932042	96–98	1	21	0	22	Betaproteobacteria
OTU-21	Desulfuromonas acetoxidans	ZP_01312343	94–99	0	32	31	63	Deltaproteobacteria
OTU-22	Anaeromyxobacter sp. strain Fw109-5	YP_001380211	93–97	12	0	15	27	Deltaproteobacteria
OTU-23	Geobacter uraniireducens Rf4	YP_001229952	92	1	0	5	6	Deltaproteobacteria
OTU-24	Geobacter sp. strain M21	YP_003021955	93–96	4	3	0	7	Deltaproteobacteria
OTU-25	Geobacter sp. strain M21	YP_003021955	97–98	35	15	5	55	Deltaproteobacteria
OTU-26	Geobacter uraniireducens Rf4	YP_001229952	95–97	3	0	6	9	Deltaproteobacteria
OTU-27	Anaeromyxobacter sp. strain Fw109-5	YP_001380211	88–91	0	5	2	7	Deltaproteobacteria
OTU-28	Desulfotomaculum gibsoniae	ZP_09101754	93–95	0	0	15	15	Firmicutes
OTU-29	Geobacter sp. strain M21	YP_003021955	89–92	13	0	0	13	Deltaproteobacteria
OTU-30	Desulfovibrio magneticus RS-1	YP_002953433	89	4	0	0	4	Deltaproteobacteria
OTU-31	Syntrophobacter fumaroxidans	YP_845148	90–93	0	0	16	16	Deltaproteobacteria
OTU-32	Methylocystis sp. strain LW5	AAK97419	98	2	0	0	2	Alphaproteobacteria
OTU-33	Coraliomargarita akajimensis	YP_003550022	89	1	0	1	2	Verrucomicrobia
OTU-34	Cupriavidus sp. strain pp2.75	ADM25241	92	1	0	0	1	Betaproteobacteria
OTU-35	Rhizobium sp. strain SMF 466_6	CCN27451	98	0	0	2	2	Alphaproteobacteria
OTU-36	Paenibacillus graminis	BAH23271	94	0	0	2	2	Firmicutes
OTU-37	Rhizobium sp. strain SMF 466_6	CCN27451	92	0	2	0	2	Alphaproteobacteria
OTU-38	Nostoc sp. strain PCC 7120	NP_484917	99	0	0	2	2	Cyanobacteria
OTU-39	Calothrix sp. strain LEGE 06100	AGG40738	97	0	0	1	1	Cyanobacteria
OTU-40	Ectothiorhodospira haloalkaliphila	ABN10975	88	1	0	0	1	Gammaproteobacteria
OTU-40 OTU-41	Desulfatibacillum alkenivorans	YP_002430688	93	1	0	0	1	Deltaproteobacteria
OTU-41	*	AEP33459	96	1	0	0	1	*
	Bradyrhizobium sp. strain cmy11			0	0			Alphaproteobacteria
OTU-43	Rhizobium sp. strain SMF 466_6	CCN27451	97		0	1	1	Alphaproteobacteria
OTU-44	Cupriavidus sp. strain pp2.75	ADM25241	98	1		0	1	Betaproteobacteria
OTU-45	Dechloromonas aromatica	YP_284634	98	1	0	0	1	Betaproteobacteria
OTU-46	Ectothiorhodospira haloalkaliphila	ABN10975	90	1		0	1	Gammaproteobacteria
OTU-47	Heliorestis baculata	BAD80875	88	0	1	0	1	Firmicutes
OTU-48	Mesorhizobium tianshanense	CAR57837	96	0	0	1	1	Alphaproteobacteria
OTU-49	Methylocystis echinoides	AAO49390	89	1	0	0	1	Alphaproteobacteria
OTU-50	Pelobacter carbinolicus	YP_006717849	93	0	0	1	1	Deltaproteobacteria
OTU-51	Geobacter bemidjiensis	YP_002138883	95	1	0	0	1	Deltaproteobacteria
OTU-52	Halorhodospira halophila	ABN10970	94	0	1	0	1	Gammaproteobacteria

toxidans; and OTU-27, affiliated with *Bradyrhizobium japonicum*), indicating measurable *nifH* gene expression in several apparently low-abundance root bacteria (Table 1 and Fig. 3). Overall, 3 OTUs contained sequences derived from root DNA and RNA, 14 OTUs contained sequences from root DNA but not RNA, and 5

OTUs contained sequences from root RNA but not DNA (Table 1 and Fig. 1). At the individual plant level, OTU-4 contained *nifH* RNA sequences from the most plants (only 4 of 8 plants), indicating that no single *nifH*-expressing species was common to all switchgrass plants (see Table S2 in the supplemental material).

TABLE 2 nifH diversity and coverage estimates for root DNA and RNA and shoot DNA

	Value for plant no.:										
DNA/RNA type and parameter	E-01	E-04	E-06	E-08	E-13	E-18	E-28	E-31	E-36	E-40	Avg
Root DNA											
No. of taxa	11	12	14	15	8	10	5	5	9	9	9.8
No. of clones	48	48	48	44	48	46	46	44	44	48	46.4
Good's estimator (%)	89.58	83.33	93.75	79.55	95.83	93.48	97.83	93.18	93.18	95.83	91.55
Dominance	0.201	0.237	0.133	0.137	0.248	0.158	0.306	0.717	0.185	0.170	0.249
1 — Simpson diversity index	0.799	0.763	0.867	0.863	0.752	0.842	0.694	0.283	0.815	0.829	0.750
Shannon's diversity index	1.897	1.8	2.311	2.266	1.647	2.002	1.333	0.622	1.86	1.929	1.766
Evenness	0.606	0.50411	0.721	0.643	0.649	0.740	0.759	0.372	0.711	0.765	0.646
Root RNA											
No. of taxa	5	6	3		1	1		3	1	6	3.25
No. of clones	41	31	32		33	25		19	32	34	30.87
Good's estimator (%)	97.56	93.33	100.00		100.00	100.00		100.00	100.00	100.00	98.86
Dominance	0.327	0.611	0.639		1	1		0.424	1	0.279	0.659
1 — Simpson diversity index	0.673	0.389	0.361		0	0		0.576	0	0.721	0.340
Shannon's diversity index	1.271	0.884	0.656		0	0		0.972	0	1.492	0.659
Evenness	0.713	0.404	0.643		1	1		0.882	1	0.741	0.798
Shoot DNA											
No. of taxa	3	4	5	3	6	7	8	9	8	9	6.2
No. of clones	38	38	44	41	47	32	31	42	28	35	37.6
Good's estimator (%)	100.00	97.37	95.45	100.00	100.00	90.63	93.55	97.62	89.29	91.43	95.53
Dominance	0.367	0.540	0.371	0.411	0.231	0.320	0.213	0.194	0.212	0.211	0.307
1 – Simpson diversity index	0.633	0.459	0.629	0.589	0.769	0.679	0.787	0.806	0.788	0.789	0.693
Shannon's diversity index	1.043	0.872	1.155	0.967	1.579	1.437	1.769	1.874	1.753	1.827	1.428
Evenness	0.946	0.598	0.635	0.877	0.809	0.601	0.733	0.724	0.722	0.691	0.733

nifH sequences from shoot bacteria. BLAST results indicated that 63% of sequences derived from shoot-associated DNA were affiliated with alphaproteobacteria and 20% with deltaproteobacteria. Sequences affiliated with firmicutes were associated only with shoots (OTU-28) (Fig. 2). Eighteen sequences affiliated

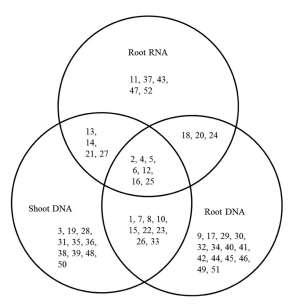


FIG 1 Venn diagram showing the number of shared and unique OTUs among the three sample types (root DNA, root RNA, and shoot DNA). Numbers indicated in the diagram are OTU identity numbers (details of each OTU are given in Table 1).

with gammaproteobacteria (*Klebsiella* species accession no. ACM68399) were represented by OTU-19 (Table 1 and Fig. 3). OTU-2 contained the largest number (71 sequences) of *nifH* sequences from shoots, which were affiliated with an uncultured bacterium related to *Bradyrhizobium* sp. strain MAFF 210318 (94 to 98% identity). Ten OTUs were unique to shoots (OTUs 3, 19, 28, 31, 35, 36, 38, 39, 48, and 50; Fig. 1), among them sequences affiliated with *Mesorhizobium loti* (OTU-3; 100% identity) and with *Syntrophobacter fumaroxidans* (OTU-31; 90 to 93% identity) (Table 1 and Fig. 3). Half or more of the plants contained *nifH* DNA sequences affiliated with *Bradyrhizobium* sp. strain BTAi 1 (OTU-1), *Burkholderia* sp. strain PTK47 (OTU-7), and *Sphingomonas azotifigens* (OTU-8). A small number of *nifH* sequences from shoots were affiliated with cyanobacteria (Table 1 and Fig. 3).

DISCUSSION

We used a culture-independent, *nifH* sequence-based approach to survey the diversity of potential nitrogen-fixing bacteria associated with switchgrass. To our knowledge, this is the first study of its kind for switchgrass, and as such it lays a foundation for future work on the isolation and characterization of associative diazotrophic bacteria from switchgrass and their potential use as a nitrogen source for this biofuel crop.

Over 1,000 *nifH* PCR amplicons derived from root and shoot DNA and root RNA of 10 plants were cloned and sequenced in this study. Roots of switchgrass plants harbored a greater diversity of *nifH*-containing bacteria than did shoots (Table 2). Similar results have been obtained for maize (37). However, only a small fraction of root-associated *nifH*-containing bacteria expressed this gene

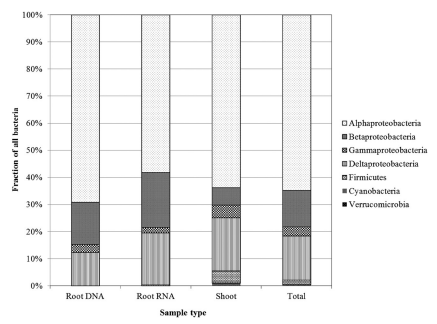


FIG 2 Class-level composition of *nifH* clone libraries. Percent abundances per clone library from root RNA (242 sequences), root DNA (451 sequences), shoot DNA (397 sequences), and overall sequences (1,060) are shown.

(Table 2). Similar results have been found in sugarcane, spruce (15), and rice (14). Shannon's diversity indices for *nifH* transcripts associated with individual plants ranged from 0.142 to 1.245 for sugar cane, and for one spruce tree it was 0.184 (15), similar to what was found here for switchgrass.

Alphaproteobacteria were the main source of *nifH* sequences in both roots and shoots, accounting for 62% of all DNA and RNA sequences combined. Within this group, 58% of the sequences were related to *nifH* in bacteria from the order *Rhizobiales* and were present in all sample types. More specifically, these sequences were affiliated with the genera *Bradyrhizobium*, *Mesorhizobium*, *Methylobacterium*, *Rhizobium*, and *Sinorhizobium*, all of which contain species that are known to fix nitrogen in symbioses with legumes (38–45).

Based on the frequency of nifH recovered from DNA, Bradyrhizobium was the dominant genus associated with switchgrass roots and shoots, with 22% (243 sequences) of all clones distributed in 3 OTUs (Table 1). OTU-10 was closely related to Bradyrhizobium japonicum sequences. B. japonicum is a functionally diverse species (38), and some strains can fix nitrogen under freeliving conditions (46). Bradyrhizobium sp. strain MAFF210318, affiliated with OTU-2, is a nonphotosynthetic bacterium that can fix nitrogen under free-living conditions (47). Bradyrhizobium sp. strain BTAi 1, affiliated with phylotype 1, is a photosynthetic rhizobium that also can fix nitrogen under free-living conditions (8, 48). Bradyrhizobium species have been found as endophytes in rice (49) and sugarcane (15, 50). Despite the abundance of Bradyrhizobium bacteria associated with switchgrass, however, the paucity of Bradyrhizobium-like nifH sequences derived from root RNA suggests that these bacteria fix little or no nitrogen in association with native switchgrass.

Evidence was obtained for *nifH* gene expression in a variety of bacterial genera associated with switchgrass, including affiliates of *Rhizobium/S. meliloti* (OTU-14), *Methylobacterium* (OTU-12), *Desulfuromonas* (OTU-21), *Burkholderia* (OTU-5 and OTU-6),

Azoarcus (OTU-20), Geobacter (OTU-25), Bradyrhizobium (OTU-2, OTU-24, and OTU-27), Rubrivivax (OTU-11), Pseudomonas (OTU-18), Methylocystis (OTU-4), Azospirillum (OTU-16), and Amorphomonas (OTU-13), in descending order of prevalence based on numbers of sequenced clones. Half of these genera belong to the alphaproteobacteria, while the remainder partitioned into the beta-, gamma-, and deltaproteobacteria. Rhizobium helanshanense/S. meliloti (ADP37388/CCH40450)-affiliated sequences (alphaproteobacteria) were prominent among root nifH RNA, accounting for 36% of all root RNA-derived sequences (Table 1). However, only one sequence was detected from root DNA, and many (44 sequences) were identified from shoot DNA. OTU-14 showed over 97% identity with sequences from S. meliloti and various Rhizobium species, such as Rhizobium sp. strain CCN-WSX0878 (accession no. AEB96238), R. yanglingense (AFD62623), and R. undicola (AEP04095). These results indicate active *nifH* gene expression in a very small population of specific rhizobia, a conclusion that warrants confirmation in future work designed to identify the best potential nitrogen-fixing bacteria for switchgrass.

Practical application of nitrogen-fixing bacteria for N supply to switchgrass will depend not only on active nitrogenase genes but also on the ability of the plant to host a sufficiently large population of the bacteria. Given the low number of rhizobial *nifH* DNA sequences that match *nifH* RNA sequences from switchgrass roots, the latter would seem to be an impediment to the development of effective nitrogen fixation in the roots of switchgrass.

A large number of *nifH* clones (38 in OTU-12) derived from root RNA were affiliated with *Methylobacterium nodulans* ORS 2060 (AAQ82902), another alphaproteobacteria able to form nitrogen-fixing nodules on legumes (51), as well as *Desulfuromonas acetoxidans* (32 in OTU-21) and *Azoarcus* sp. strain BH72 (21 in OTU-20). Here, *nifH* expression for these three bacteria outweighed their DNA abundance in the samples investigated.

Within the betaproteobacterial group, Burkholderia species ap-

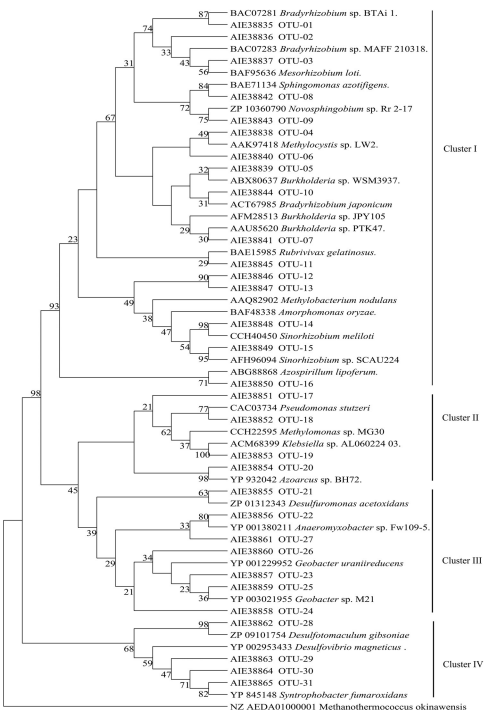


FIG 3 Phylogeny and composition of *nifH* phylotypes representing four or more sequences. OTUs from this study are represented by OTU number and the associated protein accession number. The evolutionary history was inferred by using the maximum likelihood method based on the Poisson correction model. The bootstrap consensus tree was inferred from 1,000 replicates. There were a total of 113 positions in the final data set, and analyses were conducted in MEGA5. Sequences from known bacteria are indicated by name and NCBI protein accession numbers. This tree was rooted with the *nifH* gene from the archaeon *Methanothermococcus okinawensis*.

peared to be the source of some *nifH* sequences. *Burkholderia* species have been found to nodulate and fix nitrogen in mimosoid legumes like *Mimosa pigra* (52), and some are known to associate with maize and sugarcane (53, 54). Expressed *nifH* sequences also affiliated with the betaproteobacterium *Azoarcus* sp. strain BH72

(YP_932042), which was isolated from Kallar grass and can colonize rice roots and express high levels of *nifH* transcripts there (55, 56).

Finally, the deltaproteobacteria *nifH* transcripts identified in switchgrass roots were affiliated with *Geobacter* sp. strain M21 and

D. acetoxidans (ZP_01312343). Geobacter nifH transcripts have been found previously in roots and stems of rice (22, 57).

We are aware that taxonomic inferences based on single genes, such as *nifH*, can be complicated by horizontal gene transfer. For this reason, the tentative assignments of bacterial genus and species associated with switchgrass made here should be substantiated by further work with isolated, cultured bacteria in the future. However, an independent, 16S rRNA gene-based metagenomic study of bacteria associated with switchgrass from the same sites in Oklahoma, harvested at different times of year, identified 14 of the 19 genera found in the present study (S. R. Chaluvadi and J. L. Bennetzen, personal communication).

In summary, we have identified some of the natural diversity of diazotrophic bacteria associated with switchgrass from the Oklahoma prairie. Evidence was presented that *nifH* genes of many bacterial species from the alpha-, beta-, gamma-, and deltaproteo-bacterial groups are expressed in switchgrass roots. Prominent among these were *Rhizobium* and *Methylobacterium* species of the alphaproteobacteria, *Burkholderia* and *Azoarcus* species of the betaproteobacteria, and *Desulfuromonas* and *Geobacter* species of the deltaproteobacteria. This work provides a basis for future work on the isolation, culture, and functional characterization of nitrogen-fixing endophytes of switchgrass and their possible use as nitrogen sources for cultivated switchgrass.

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