

ORIGINAL ARTICLE

Bacterial gene abundances as indicators of greenhouse gas emission in soils

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Nitrogen fixing and denitrifying bacteria, respectively, control bulk inputs and outputs of nitrogen in soils, thereby mediating nitrogen-based greenhouse gas emissions in an ecosystem. Molecular techniques were used to evaluate the relative abundances of nitrogen fixing, denitrifying and two numerically dominant ribotypes (based on the $\geq 97\%$ sequence similarity at the 16S rRNA gene) of bacteria in plots representing 10 agricultural and other land-use practices at the Kellogg biological station long-term ecological research site. Quantification of nitrogen-related functional genes (nitrite reductase, *nirS*; nitrous oxide reductase, *nosZ*; and nitrogenase, *nifH*) as well as two dominant 16S ribotypes (belonging to the phyla *Acidobacteria*, *Thermomicrobia*) allowed us to evaluate the hypothesis that microbial community differences are linked to greenhouse gas emissions under different land management practices. Our results suggest that the successional stages of the ecosystem are strongly linked to bacterial functional group abundance, and that the legacy of agricultural practices can be sustained over decades. We also link greenhouse gas emissions with specific compositional responses in the soil bacterial community and assess the use of denitrifying gene abundances as proxies for determining nitrous oxide emissions from soils.

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Introduction

The impact of agricultural practices on the environment has been studied extensively, leading to changes in land management policy worldwide (Tilman *et al.*, 2002). Yet, surprisingly little is known about the interactions between agroecosystem management practices and the soil microbial community, which has a key role in nutrient transformation and chemical cycling (Staley and Reysenbach, 2002). The Kellogg biological station long-term ecological research (KBS-LTER) site has hosted numerous microbiological studies (Bruns *et al.*, 1998, 1999; Broughton and Gross, 2000; Phillips *et al.*, 2000a,b; Buckley and Schmidt, 2001; Blackwood and Paul, 2003), but few studies have focused on quantitative analysis of bacterial community composition in relation to nitrogen turnover rates, specifically those related to greenhouse gas emissions. In addition, comparative quantitative analysis of specific functional or

phylogenetic groups within the soil community is still limited. To date, genes encoding enzymes involved in nitrogen cycling have been targets of choice for studies focusing on functional groups (that is, guilds) of bacteria (Leininger *et al.*, 2006; Henry *et al.*, 2008). This focus is well founded as nitrogen is essential for plant growth, along with phosphate, carbon, hydrogen and oxygen. However, simultaneous comparison of the abundance of multiple N-cycle-related genes across multiple treatments and land-use types has not yet been conducted, especially when framed around an ecosystem-level process, such as, greenhouse gas emissions from soils.

The United States Department of Agriculture tracks emissions of multiple greenhouse gases related to agricultural activities and ranks them based on their global warming potential (GWP). Of all the sources of GWP in cropping systems, including CO₂ and CH₄, none are more poorly quantified than N₂O production (Robertson and Grace, 2004). This represents a tremendous knowledge gap regarding the role of N₂O in global warming, especially considering the fact that its GWP is 296-fold greater than that of CO₂ and it is frequently the major source of GWP in agricultural systems (Robertson *et al.*, 2000; Robertson and Grace, 2004; EIA, 2008). Poor quantification of

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N₂O is to a large extent linked to the challenges of measuring N₂O fluxes in the field, requiring numerous measurements with inherently high variability. This constraint has limited data collection, and, until refined, represents a rudimentary measurement of a globally important activity. The ability to construct more informative models, predictions and mitigation strategies related to greenhouse gas emissions depends on the development of new analytical approaches that are more efficient and accurate than currently available ones. As microbial populations control natural production and consumption of nitrous oxide, their abundance and activities represent a potential method for predicting gas emissions.

Using real-time quantitative PCR (qPCR), we examined the abundance of key soil microbial guilds and taxa, including nitrogen-fixing bacteria (through the *nifH* gene) and denitrifying bacteria (through the *nirS* and *nosZ* genes). Although nitrification represents a key step in the conversion of ammonia nitrogen into its gaseous forms (NO₂ and NO₃), this process is less relevant to N₂O emissions. Further, nitrifier numbers are typically low in soils and are challenging to quantify using direct qPCR as it is applied to all other genes analyzed making nitrification measures beyond the scope of this study. Quantification of two numerically predominant operational taxonomic units (OTUs) belonging to the phyla *Acidobacteria* and *Thermomicrobia* obtained at KBS (Morales and Holben, 2009) was also performed for comparison with data obtained using function-based primers. These analyses were performed across 10 different treatments based on land-use types at the KBS-LTER. Analyses in other soils using 16S rRNA gene-based PCR denaturing gradient gel electrophoresis have shown that soil type may be the strongest selector of soil microbial community structure (Wakelin *et al.*, 2008), but that study focused on large-scale rearrangements in community composition measured using a broad-scale technique. However, changes in functional or phylogenetic group abundance may go undetected when using such 16S rRNA gene-based approaches, as poor resolution between taxa due to gene conservation, nonspecific primers and other factors (for example, see Morales and Holben, 2009) can lead to mistaken conclusions about functional group abundance or population dynamics. To examine the extent to which this occurs in this system, the current study employed qPCR targets directly related to functional traits, as well as two 16S rRNA gene-based ribotypes, to compare the patterns observed based on the abundance of genes encoding key enzymatic activities with those observed using 'OTU-based primers'.

Four general hypotheses were tested. The first stated that 16S-based taxon abundance estimates would be higher than those observed for the functional genes as the former can detect multiple phylogenetic subgroups that might each harbor

multiple different bacterial functional groups (for example, denitrifiers and nitrogen fixers can both be found within the same genus). Second, we hypothesized that, due to the high number of leguminous and other nitrogen-fixing symbiotic plants in certain land-use treatments at the KBS-LTER site, nitrogenase reductase gene (*nifH*) numbers would be consistently higher in all treatments involving leguminous cover crops (that is, treatments 1–4 (T1–4), soybean; and T6, alfalfa). The third hypothesis predicted that abundance of denitrification bacterial genes (*nosZ* and *nirS*) would be relatively uniform across all treatments, given the widespread distribution of this metabolic activity across the breadth of bacterial phylogenetic groups. Finally, as the balance between input and output of nitrogen gas in soils is respectively controlled by nitrogen fixers and denitrifiers, we hypothesized that differences in bacterial gene abundances of these key nitrogen cycling genes between annual, perennial and successional sites would correspond to those observed for greenhouse gas emission rates for these sites.

Materials and methods

Study site and sample collection

Samples were collected from the KBS-LTER Row-Crop Agriculture site in mid-Michigan (for an overview of that project see <http://lter.kbs.msu.edu/>). Our study examined the bacterial community in the replicate plots of Treatment 1–8 (T 1–8) of the main experimental site, as well as two additional successional and forest sites (Table 1). Four of the eight main site treatments are annual crop rotations (T1–4), two are perennial (T5, poplars; and T6, alfalfa), and two are successional systems under native vegetation

Table 1 Kellogg biological station LTER treatment and successional regimes

Treatment	Crop cover	Notes
T1	WCS	STD, chisel plowed
T2	WCS	STD, no-till
T3	WCS	Org red, N at planting, WCCC, H, PPC
T4	WCS	Rotary hoed, WCCC, PCC
T5	Continuous poplar	
T6	Continuous alfalfa	
T7	Native successional	Last plowed on spring of 1989
T8	Native mid-successional	Never plowed, soil organic matter historical control
DFR	Deciduous forest	Late successional site
SFR	Successional	40 to 60-year-old former agricultural field

Abbreviations: DFR, deciduous forest; LTER, long-term ecological research; N, nitrogen added; Org Red, reduced input organic; PCC, post planting cultivation; SFR, successional forest; STD, standard; WCCC, winter clover cover crop; WCS, wheat corn soy rotation.

(T7, former agricultural site left fallow following spring plowing in 1989; and T8, never plowed or cultivated). The two forested sites comprised of a set of 40 to 60-year-old successional sites (former agricultural fields, SFR (successional forest) and DFR (a deciduous forest site that has never been cut)). All soils in these treatments are classified as fine loamy, mixed and mesic Typic Hapludalfs. Sampling was carried out on 2 May 2007 by collecting five randomly positioned, 0–20 cm soil cores using a soil probe from each of the treatment replicates as is standard at the LTER site (<http://lter.kbs.msu.edu/protocols/112>). Each set of five samples was sieved through 2 mm mesh and then mixed thoroughly in equal proportions, providing a single composite sample for each replicate treatment plot (that is, there were either three or six replicated plots per treatment, and one composited sample was developed and examined for each replicate plot). All soil samples were stored in Whirl-Pak bags on dry ice or at -70°C immediately after sieving and mixing until processed for bacterial community DNA extraction. All samples were processed within 20 days of sampling.

DNA extraction

Total community DNA was extracted from 0.25 g of each soil sample, in triplicate, using the MoBio PowerSoil DNA Isolation Kit (MoBio, Solana Beach, CA, USA) according to the manufacturers instructions and using sterile MilliQ water in the final elution step. All DNA samples were stored at -20°C until used in downstream analyses.

Real-time qPCR assays

Real-time qPCR was performed using an iCycler iQ thermocycler (Bio-Rad, Hercules, CA, USA) with an ABsolute QPCR SYBR green mix (AbGene, Epsom, UK) using primers and conditions previously described (nitrogenase reductase (*nifH* gene) (Rösch and Bothe, 2005; Yergeau *et al.*, 2007); nitrite reductase (*nirS* gene) (Throback *et al.*, 2004; Yergeau *et al.*, 2007); nitrous oxide reductase (*nosZ*) (Henry *et al.*, 2006); *Thermomicrobia* group 4 (OTU-specific based on $\geq 97\%$ sequence similarity at the 16S rRNA gene) (Morales and Holben, 2009); *Acidobacteria* group 6 (OTU-specific based on $\geq 97\%$ sequence similarity at the 16S rRNA gene) (Morales and Holben, 2009)) and are summarized in Appendix Table S1. Although no function- or 16S-based primer sets are necessarily comprehensive across the spectrum of microbial diversity, they have been widely used to good effect in comparative studies (for example, between treatments), as is the case in the current study. Known template standards were made from cloned PCR products amplified from whole-genome extracts of pure bacterial isolates (see Appendix Table S1), and each standard was sequenced to confirm target identity.

Primer validation analyses were performed before use as previously described (Morales and Holben, 2009).

Variance in gene abundance measurements was determined between replicate plots of the same treatment by pooling three individual DNA preparations from each replicate plot in equimolar amounts to provide a representative sample for that replicate plot. A total of 5 ng of DNA was used to compare gene abundance in each plot. For comparing the effect of different treatments on the overall abundance of each gene, DNA preparations from all replicate plots and extractions within a treatment were combined in equimolar amounts to provide a representative sample for that treatment. All qPCR reactions for any single sample were run at least in triplicate, as described above.

Correct target amplification from soil DNA was confirmed by cloning PCR fragments from T1. Triplicate standard PCR reactions were performed separately as described above for each primer pair using total community DNA from T1, which represents the canonical treatment practice for the KBS-LTER site. The resultant PCR products were purified and cloned as previously described (Morales and Holben, 2009) to confirm that specific amplification of the corresponding target had occurred.

Statistical analysis

Relationships between microbial gene abundance, successional stage, greenhouse gas flux and other environmental parameters, were determined by principal components analysis (PCA) with data matrices composed of chemical and bacterial qPCR data (Supplementary Table S2) for T1–8, SFR and DFR of the KBS-LTER, collectively representing annual, perennial and successional sites. Chemical data were extracted from Robertson *et al.* (2000) and represented gas fluxes and their respective greenhouse warming potentials, aboveground net primary productivity, $\text{NO}_3\text{-N}$, N mineralization potentials and soil carbon concentrations over an 8-year period. The chemical metadata set used was from 1991 to 1999 (as reported in Robertson *et al.*, 2000) because this timeframe maximized the number of variables available for analysis, as several of the measurements were not continued beyond that point. However, more recent gas emission data through 2007 (presented in Supplementary Figure S6) strongly support the suggestion that the same soil processes and relationships persisted at KBS through our sampling time and beyond, as observed differences in these parameters are consistent across treatments and the relative relationships between treatments remain constant. Further, the classification of treatments as a sink or source of gases on the basis of current or past mean gas fluxes for any treatment remains the same. Data were organized with rows representing treat-

ments and columns representing individual variables. Principal component scores were plotted by site, abbreviated as shown in the site description and in Table 1, for the first two principal components. Parameters driving the distribution of the PCA plots were determined by querying all variables against the first three principal components. Individual chemical factors were also independently queried to qPCR results on a per gene basis by conducting pairwise correlations.

To show co-trends between genetic and chemical variables, data were also analyzed using co-inertia analysis (CIA) (Dray *et al.*, 2003), computed with the MADE4 package in the R statistical software environment (Culhane *et al.*, 2005). CIA is a dimensional reduction procedure designed to measure the similarity of two sets of variables (measurements), as they are associated with a single set of cases. Due to very strong correlations (>0.99) between some chemistry measurements, we were able to unclutter the CIA plot by using CH₄, N₂O and Organic C as proxies for CH₄-C, N₂O-N and Organic C (kg), respectively.

Results

Bacterial gene abundance and diversity

The efficiencies of real-time PCR assays for all targets averaged 92% (s.d. $\pm 4\%$), allowing for direct comparison of results for all targets. Statistically significant differences in gene target abundances were observed between replicate plots under the same treatment (T1) for all tested genes (Table 2). As anticipated, qPCR results obtained from pooling DNA extracted from each individual replicate plot and run as a representative sample for that treatment resulted in mean values within the standard deviation of the true replicates (Figure 1). Two 16S rRNA gene targets representing the numerically dominant bacterial groups *Thermomicrobia* and *Acidobacteria* based on sequences generated from this site (Morales *et al.*, 2009) were the most abundant of all targets tested (Figure 1), supporting the first hypothesis regarding higher abundances based on phylogenetic targets (that is, 'genus-level' 16S rRNA genes) compared with functional gene (that is, enzyme coding) targets.

By contrast, the measured abundances of the nitrogenase gene (*nifH*) did not support our second

hypothesis that predicted higher abundance in treatments containing crops with known symbiotic nitrogen fixation associations (for example, soybean rotations). Indeed, the abundance of *nifH* was generally higher in the successional treatments (SFR and DFR) than in traditional agricultural sites (Figure 1). Denitrifier abundance, as indicated by *nirS* and *nosZ* gene abundance, varied significantly between treatments, which did not support the third hypothesis predicting comparable denitrifier numbers between treatments.

Correlating bacterial gene abundance to environmental variables

Principal component analysis of annual ecosystem averages for key environmental parameters, global warming potential (GWP), bacterial qPCR results and a combined data set comprised of all variables was employed to assess relationships between bacterial gene abundance and process-level measurements. Strong clustering of samples based on aboveground plant cover type (that is, annual, perennial, successional) was observed (Figure 2), supporting the fourth hypothesis predicting differences in greenhouse gas emissions as being correlated to differences in the balance between nitrogen fixing and denitrifying bacteria. PCA based solely on qPCR results from bacterial gene targets accounted for the most variance within the first two components ($\sim 89\%$). Lower combined principal component 1 and 2 scores were observed for annual ecosystem averages and GWP values ($\sim 75\%$) and for the combined data set ($\sim 68\%$) (Figure 2). All three PCA plots showed general clustering of sites based on land-use type, with the exception of T7 (the early-successional site which was previously a woodpile). Loadings for principal components 1 and 2 (that is, loading vectors) were plotted to show independent variable contributions to variance between the treatments (Figure 2). Successional sites were generally associated with higher organic carbon levels and *nifH* gene abundance. Perennial plant-based treatments, as well as the early native successional plot T7, exhibited increased nitrous oxide reducer abundance (*nosZ*), total carbon levels and abundance of *Thermomicrobia* and *Acidobacteria*. Aboveground annual productivity, nitrite reducers (*nirS*) and GWP were all strongly

Table 2 *t*-Test analysis of replicate plot qPCR values for each gene target within T1.

Target	Test value	Count	Mean	Std dev	Std Error of mean	<i>t</i>	df	P-level
AB#6	10	24	2.63E + 05	1.36E + 05	2.79E + 04	9.44	23	<0.001
TM#4	10	23	9.97E + 04	6.91E + 04	1.44E + 04	6.92	22	<0.001
<i>nifH</i>	10	21	3.59E + 03	1.75E + 03	3.82E + 02	9.37	20	<0.001
<i>nirS</i>	10	19	4.63E + 04	1.98E + 04	4.53E + 03	10.2	18	<0.001
<i>nosZ</i>	10	20	8.16E + 03	4.71E + 03	1.05E + 03	7.74	19	<0.001

Abbreviations: *t*, test score; df, degrees of freedom; qPCR, quantitative PCR; Std dev, standard deviation.

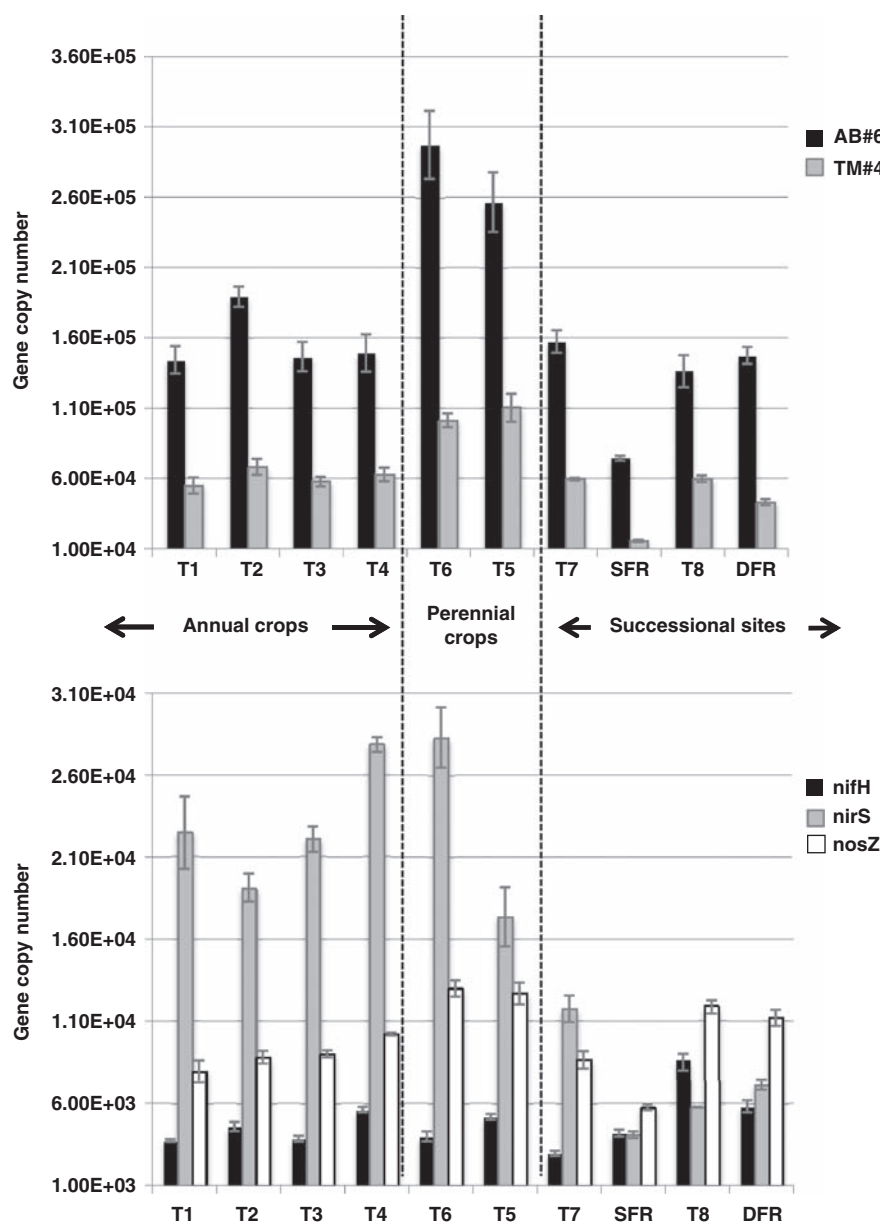


Figure 1 Specific detection of bacterial groups by quantitative PCR (qPCR). Values indicate gene copy numbers, as determined from 5 ng of DNA extracted from soil using a dilution curve with known standards. Error bars are one standard error of the mean (s.e.) of at least triplicate qPCR reactions ($n \geq 3$). AB#6 and TM#4: operational taxonomic units based on $\geq 97\%$ sequence similarity to the 16S rRNA gene representing *Acidobacteria* group 6 and *Thermomicrobia* group 4, respectively. *nifH*, nitrogenase gene; *nirS*, nitrite reductase gene; *nosZ*, nitrous oxide reductase gene.

correlated with PC1, which was responsible for clustering of sites into annual, perennial and successional treatments, with annual sites showing the highest levels of nitrite reducers and GWP.

Although plotting the first two principal components accounted for much of the variance in the data, the third component significantly increased the percentage of variance accounted for. On the basis of the first three principal components for the qPCR data alone, $\sim 98\%$ of the variance was accounted for, whereas the number decreased to $\sim 86\%$ when only the annual ecosystem averages and GWP values were analyzed. The combined data

set of all measured variables accounted for $\sim 82\%$ of the variance within the first three components, with *nosZ* gene abundance being significantly and negatively correlated (-0.85 , $P \leq 0.05$) to the third component. Correlation values of individual variables with each principal component are summarized in Supplementary Table S3.

An alternative way of analyzing the data based on CIA showed a pattern similar to that in the PCA analysis (Appendix S4). The CIA gives an RV coefficient, a global measure of similarity between data sets based on a multivariate extension of the Pearson correlation coefficient, scaled from 0 (no

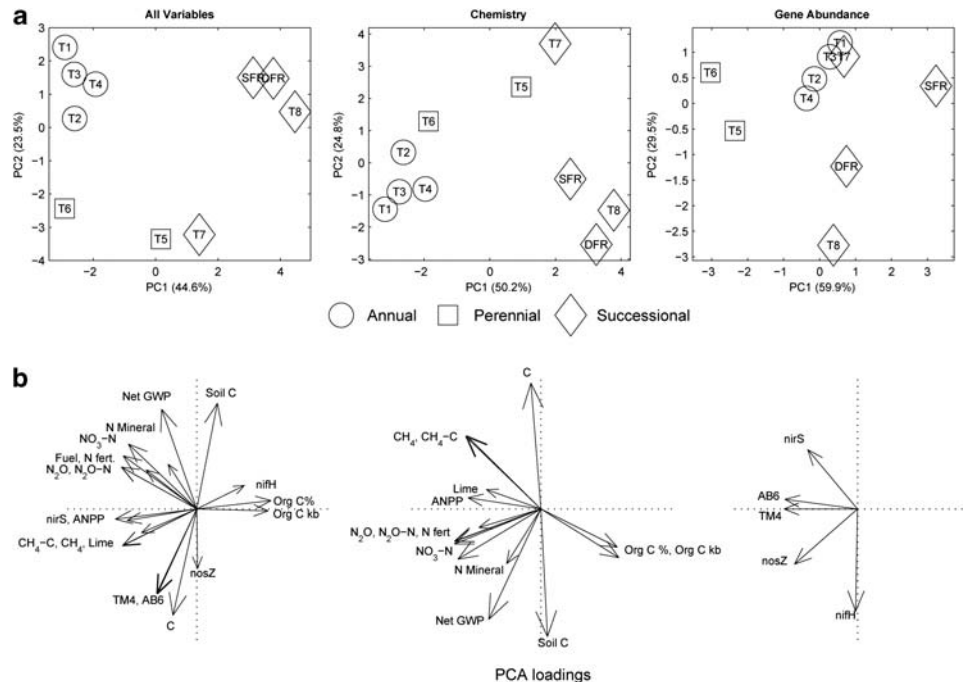


Figure 2 Principal component analysis (a) and factor loadings plot (b) for gene abundances (far right), ecosystem chemistry (center) and combined data sets (far left) summarized in Appendix S1 for different management systems at the KBS-LTER site (described in Table 1). The percentage of the variation in the samples described by the plotted principle components is indicated on the axis.

similarity) to 1.0 (identical). The genetic and chemical variables show strong similarity with an $RV = 0.76$. The significance of the similarity is underscored by a permutation test (Culhane *et al.*, 2003), in which total co-inertia (the measure of co-variability of the two data sets) is computed after permutations to one of the two data sets (randomly chosen before each permutation). This shuffles the data, disassociating the soil sites from the genetic and chemical sample values. The test yielded 100 000 permutation-based values of total co-inertia. Under the null hypothesis that the data sets are independent, only five of these 100 000 values were as large or larger than that observed for the genetic and chemical data sets, for a P -value of 0.00005. The plots also provide a view of the relative strengths of relationships between genetic and chemical variables with respect to each other and to the different soil environments. Genetic and chemical variables show the strongest overall co-trend (shortest arrows) in the group of annuals (Appendix S4). This figure also indicates that, although chemical profiles in transition treatments (former agricultural sites left to undergo natural succession; namely T7 and SFR) closely resembled expected values for successional sites, genetic variables were slower to change, retaining their original signature for longer periods. This is to say, the original bacterial community signature associated to agricultural treatments is seemingly persistent (that is, apparent) after >40 years, but this cannot be unequivocally confirmed with our data as no temporal comparisons are available.

Pairwise correlations between all of the variables showed strong (>0.65) positive correlation between bacterial gene numbers and either greenhouse gas fluxes, GWP, annual productivity or carbon levels (Appendix S5). Nitrite reductase (*nirS*) gene abundance was positively correlated to greenhouse gas (N_2O -N, NO_3 -N and CH_4 -C) emission values, GWP values and aboveground net primary productivity, while being negatively correlated to organic carbon. Nitrogenase (*nifH*) gene abundance was positively correlated to organic carbon levels. The abundance of the *nifH* gene shared a weak negative correlation (-0.358) with the abundance of the *nirS* gene (nitrite reductase) (Appendix S5). A second denitrification gene (*nosZ*), responsible for the reduction of nitrous oxide to dinitrogen, did not exhibit the same trend as *nirS* (Figure 1). The two target genes corresponding to the numerically dominant OTUs belonging to *Thermomicrobia* and *Acidobacteria* (Morales *et al.*, 2009) were the most abundant of all targets measured (Figure 1). These genes also showed a strong positive correlation (Appendix S5) across all treatments, with the highest values found in single cultivar perennial treatment plots (Figure 1).

nirS–*nosZ* gene abundance as proxy for greenhouse gas (N_2O) emissions

A simple regression analysis was conducted to compare direct measurements of nitrous oxide emissions from soils with the abundance of *nirS* gene targets minus *nosZ* gene targets (Figure 3),

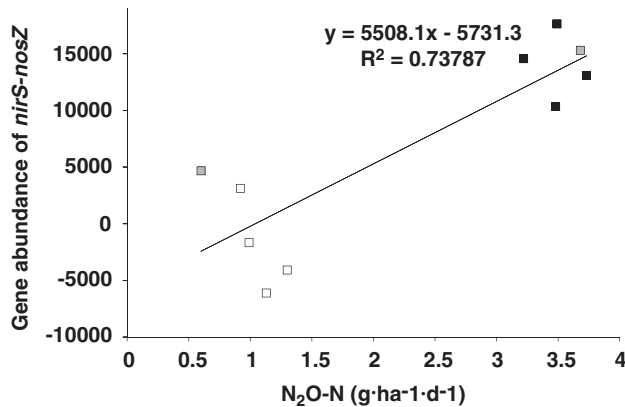


Figure 3 Simple regression analysis of nitrous oxide emission and the *nirS* gene abundance minus *nosZ* gene abundance. Annual crops (black fill); Perennial crops (gray fill); Successional sites (white fill).

which showed a strong correlation ($r^2 = 0.74$). Two discrete data clusters were observed related to direct N₂O measurements, whereas a more incremental relationship was found between site successional stage and *nirS*–*nosZ* gene abundance (Figure 3).

Discussion

Multivariate analysis of annual ecosystem parameter averages, global warming potential and bacterial gene abundances in the current study support the hypothesis that the legacy of agriculture (that is, resilience of treatment effects) has a stronger influence on soil biogeochemistry than current environmental parameters (that is, real-time soil conditions), as previously suggested by other studies (Buckley and Schmidt, 2001, 2003). Strong correlations were obtained that suggest a key role for bacterial activities in controlling responses between agricultural practice or land-use regimes and greenhouse gas emissions. The data also support the interpretation of long-term repercussions at the microbial community level to certain land-use practices. However, given that our study does not include a temporal sampling sequence, these interpretations are based on hypothesized ecosystem successions, as observed on successional treatment plots at the KBS-LTER site after 40–60 years of cessation of agricultural management. Those plots represent more advanced successional stages of the current agricultural treatments, and can be used as references for comparisons.

Previous research at KBS showed differences in microbial community structure between treatments (for example, indicating effects on bacterial community composition based on 16S rRNA sequences (Buckley and Schmidt, 2001, 2003); denitrifiers (Cavigelli and Robertson, 2001; Stres *et al.*, 2004); and ammonia oxidizers (Bruns *et al.*, 1999)). In addition, treatment-based differences at KBS have

been reported for greenhouse gas emission and have suggested possible mitigating properties of certain treatments (Robertson *et al.*, 2000; Suwanwaree and Robertson, 2005). Related findings have also been reported for other systems, with plant species identity affecting denitrifying communities (Bremer *et al.*, 2007), and vegetation type driving separation of community structure (Chim Chan *et al.*, 2008).

We found carbon levels to be strongly correlated to the clustering of treatments based on cropping system, suggesting a strong role for carbon as a driver of bacterial community structure. Although higher emission values of the greenhouse gas nitrous oxide were positively correlated to traditional annual crop rotations, previous work has suggested that nitrate, as applied in fertilizer, does not select for denitrifiers (Tiedje, 1988). Instead, it has been proposed that denitrifiers are generally functioning as aerobic competitors for carbon, using their denitrification capabilities only under metabolically advantageous conditions (Tiedje, 1988). Although it appears that carbon has a major role in community structure in these treatments, the activity of an established community can be significantly altered by real-time events such as nitrogen deposition (Suwanwaree and Robertson, 2005). This leads us to suggest that community composition measurements (for example, DNA-based measurements of gene abundance in treatments) are good indicators of how treatment practices shape the community in the long run, whereas rRNA or mRNA measurements would be more useful to illustrate the response of the community to changing parameters in the short term (for example, diel cycles, rainfall, fertilizer application).

Although we included qPCR analysis of two predominant taxa (at approximately the sub-phylum or ‘genus’ level) based on 16S rRNA gene quantification to assess their ubiquity and abundance, data derived from those groups are hard to interpret in the context of biogeochemical cycling, given the lack of correlation between a specific 16S ribotype and its metabolic or catabolic capabilities. Where a direct link to a given biogeochemical reaction is desired, specific tracking of relevant functional genes is likely to be more productive. Thus, in the current study, we rely on quantification of functional genes for nitrogen cycling for correlation with process-level greenhouse gas emissions from KBS soils.

Contrary, perhaps, to common assumptions, the numbers of nitrogen fixers (as determined by *nifH* gene quantification) were found to be higher in forested or successional sites than in the agricultural fields, including those with regular soybean rotations. Although leguminous plants, which include beans, clover, alfalfa, lupine and peanuts, are among the best studied systems for nitrogen-fixing symbioses (Young and Haukka, 1996; van der Heijden *et al.*, 2006; Nandasena *et al.*, 2007; Houlton *et al.*, 2008), other non-leguminous plants including grasses

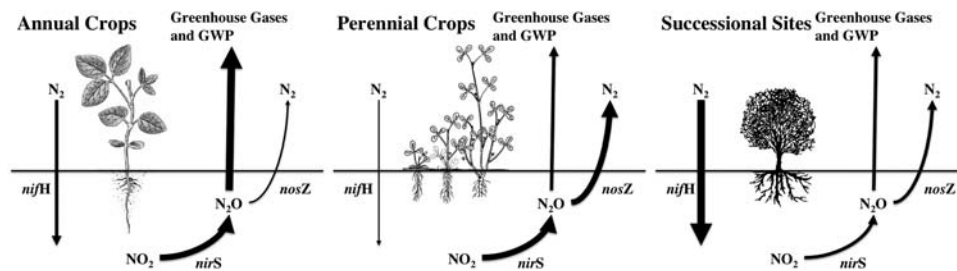


Figure 4 Generalized schematic model showing predicted changes in nitrogen flux, N-based greenhouse gas emissions and greenhouse warming potential in soils as a function of plant community differences and bacterial group abundance. Line thickness represents relative contribution of a given gene to nitrogen compound turnover rate based on data from this study and from a study by Robertson *et al.* (2000), and also summarized in Appendix S2.

(Minamisawa *et al.*, 2004; Coelho *et al.*, 2008), pine (Izumi *et al.*, 2006), wheat (Iniguez *et al.*, 2004) and alder (Ridgway *et al.*, 2004) have also been shown to possess significant populations of endophytic nitrogen-fixing bacteria. This suggests that the contribution of non-leguminous nitrogen-fixing plant symbioses and free-living nitrogen fixers is not just significant, but likely essential, in ecosystem development.

Two other nitrogen cycle-related genes encoding the denitrification enzymes nitrous oxide reductase (*nosZ*) and cytochrome cd1 nitrite reductase (*nirS*) did not exhibit similar trends in abundance across the system. Instead, what was observed was an apparent balance in the relative abundance of the two genes that can be used to predict greenhouse gas emissions and global warming potential (refer to Figures 3 and 4). To maximally relate our measurements to available metadata, we have used the same 1991–1999 data set as described by Robertson *et al.* (2000) for the KBS treatments. Whereas we note that gas emission data from 2000 to 2007 for both CH₄ and N₂O showed that emissions of these gases rose somewhat compared with the 1991–1999 timeframe (Robertson *et al.*, unpublished data (<http://lter.kbs.msu.edu/datatables/28>)), the relationship of these parameters between treatments has not changed. As noted above and in Supplementary Figure S6, this suggests that the same soil processes and relationships between treatments have persisted through to our sampling time, and that classification of treatments as a sink or source of greenhouse gas emissions remains the same. As the KBS is an LTER site with same conditions carefully maintained for more than two decades, it is well supported that ecosystem processes should remain comparable within and between treatments.

In the schematic model that we have developed to explain this behavior (Figure 4), the rate of greenhouse gas emissions is controlled by the interplay between different guilds within the local bacterial community. In this initial study, we focused on nitrogen cycling and show how the prevalence of bacteria involved in key steps in the cycle are related to the overall outcome in terms of key environmental parameters, and also how differences

in relative abundance of individual functional groups control, or at least relate to, whether a system is a net sink or a net source of greenhouse gases.

The data presented herein showed the presence of all targeted genes, and also illustrated how soil management practices have altered the relative abundance of two predominant ribotypes and several functional genes involved in different stages of the nitrogen cycle. These were linked, or at least correlated to, measured differences in greenhouse gas emissions and global warming potential resulting from various land-use practices. Although the general roles of nitrogen fixers and denitrifiers in the nitrogen cycle have been known for decades, the application of a quantitative approach based on functional gene abundance to provide predictive power regarding the fate of nitrogenous compounds in soils is novel.

Two major observations arise from these findings. The first is that monitoring bacterial community response variables, which can exhibit greater sensitivity to environmental change than other commonly used measures (Feris *et al.*, 2009), represents a robust way to monitor geochemical dynamics. Second, the possibility, perhaps probability, that bacteria can respond rapidly to environmental change without major changes in community composition through altered patterns of gene expression suggests the importance of considering the contributions and responses of microbial populations in global biogeochemical cycles to such phenomena as climate change. The latter point further suggests that there is a role for both DNA- and RNA-based approaches in modern molecular microbial ecology depending on whether the investigator is looking at long-term drivers that shape community composition or at short-term response to perturbation and change, respectively.

The variability observed with our approach, as indicated by plot-to-plot replicate variability within a treatment, likely reflects an inherent property of small- or mid-scale heterogeneities in the soil environment that affect bacterial populations locally. Thorough sampling of study sites can compensate for such variability. This is readily achieved when using molecular methods by collect-

ing multiple, small soil samples that can be pooled (composited), creating a representative sample for a site or treatment and thus its greenhouse gas production potential.

We note that our analyses focused on a single nitrite reductase gene, *nirS*. The copper-containing nitrite reductase (encoded by the *nirK* gene) was not analyzed in this study and might contribute to some of the variation not accounted for in our data. However, it has been shown that three-quarters of cultured denitrifying bacteria contain *nirS* rather than *nirK* (Zumft, 1997) and it has been found to predominate in most environments (Bothe *et al.*, 2000). It is also important to note that these experiments targeted copy numbers of genes of interest, which represent the standing community and its potential for activity rather than an actual measure of real-time gene expression levels or the corresponding enzymatic activity. Ongoing methods development to directly measure actual gene expression levels will likely enhance the resolution and accuracy of studying microbial contributions or response to key environmental functions and activities. Although our data link key gene abundance data with measured greenhouse gas potential, changes in gene expression levels could transiently change a given soil or treatment from a greenhouse gas source to a sink. Thus, making more direct molecular measurements of flux in microbial community gene expression patterns (for example, through environmental transcriptomics) is indeed highly desirable for future work.

This study provides the first quantitative assessment of the effect of land management practice on multiple microbial community constituents at both the functional and phylogenetic level. We showed that microbial assemblages do not readily return to a native or baseline community state following agricultural disturbance, consistent with previous findings that soil nutrient levels require decades or more to recover after agriculture (Robertson *et al.*, 1988, 1993; Drinkwater *et al.*, 1998; Knops and Tilman, 2000). We also present the first quantitative study illustrating interactions between different bacterial activities and their role in controlling nitrogen flux as a response to ecosystem changes.

In conclusion, we note that this initial analysis linking bacterial gene abundance data to process-level greenhouse gas emission rates represents an early step in integrating key bacterial activities to larger-scale biogeochemical cycles. Additional research in this area will extend such capabilities and allow us to assess microbial contributions and responses to ecosystem, and even to global-scale ecological phenomena such as climate change. This is particularly important as it is widely acknowledged that microorganisms govern or at least contribute to global biogeochemical cycles, yet their roles and activities are generally not even considered in current large-scale models for climate change and other global phenomena.

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