Soil denitrifier community size changes with land use change to perennial bioenergy cropping systems

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Abstract. Dedicated biomass crops are required for future bioenergy production. However, the effects of large-scale land use change (LUC) from traditional annual crops, such as corn–soybean rotations to the perennial grasses (PGs) switchgrass and miscanthus, on soil microbial community functioning is largely unknown. Specifically, ecologically significant denitrifying communities, which regulate N₂O production and consumption in soils, may respond differently to LUC due to differences in carbon (C) and nitrogen (N) inputs between crop types and management systems. Our objective was to quantify bacterial denitrifying gene abundances as influenced by corn–soybean crop production compared to PG biomass production. A field trial was established in 2008 at the Elora Research Station in Ontario, Canada (n = 30), with miscanthus and switchgrass grown alongside corn–soybean rotations at different N rates (0 and 160 kg N ha⁻¹) and biomass harvest dates within PG plots. Soil was collected on four dates from 2011 to 2012 and quantitative PCR was used to enumerate the total bacterial community (16S rRNA) and communities of bacterial denitrifiers by targeting nitrite reductase (nirS) and N₂O reductase (nosZ) genes. Miscanthus produced significantly larger yields and supported larger nosZ denitrifying communities than corn–soybean rotations regardless of management, indicating large-scale LUC from corn–soybean to miscanthus may be suitable in variable Ontario climatic conditions and under varied management, while potentially mitigating soil N₂O emissions. Harvesting switchgrass in the spring decreased yields in N-fertilized plots, but did not affect gene abundances. Standing miscanthus overwinter resulted in higher 16S rRNA and nirS gene copies than in fall-harvested crops. However, the size of the total (16S rRNA) and denitrifying bacterial communities changed differently over time and in response to LUC, indicating varying controls on these communities.

1 Introduction

Future energy needs require dedicated biomass crop production for bioethanol and combustion-based electricity generation. Corn (Zea mays L.)–soybean (Glycine max L.) rotations currently dominate the landscape across Ontario and the northern US Corn Belt (Gaudin et al., 2015), and corn grain is currently the primary feedstock for bioethanol production in Canada (Jayasundara et al., 2014). The C₄ perennial grasses (PGs) switchgrass (Panicum virgatum L.) and miscanthus (Miscanthus spp.) have been proposed as alternate feedstock crops to corn for biomass-based bioenergy production due to their large biomass yields, reduced nitrogen (N) and water requirements, decreased nutrient leaching, and potential for increased soil carbon (C) storage (Blanco-Canqui and Lal, 2009; Foster et al., 2013). Large-scale production of C₄ PGs in Ontario and the northern Corn Belt would require land use change (LUC) from existing corn–soybean rotations to PG biomass cropping systems (Deen et al., 2011; Kludze et al., 2013; Liang et al., 2012; Sanscartier et al., 2014).
Few studies have assessed how this LUC may influence soil microbial community functioning. In particular, soil denitrifying communities represent an ideal subset of the soil microbial community to target to assess changes in ecosystem functioning due to agricultural management and LUC. Denitrifying bacteria represent approximately 5% of the total soil microbial biomass (Braker and Conrad, 2011) and have been identified in over 60 genera (Philippot, 2006), encompassing a wide range of phylogenetic and functional diversity. Multiple studies have linked changes in denitrifier communities with plant types or development stage (Bremer et al., 2007; Hai et al., 2009; Petersen et al., 2012), N fertilization (Hallin et al., 2009; Yin et al., 2014), organic or conventional crop management (Reeve et al., 2010), perennial vs. annual crop land use (Bissett et al., 2011), and C and N inputs (Bastian et al., 2009). These studies suggest that LUC from corn–soybean rotations to PG species may influence the soil bacterial communities which drive soil N₂O production and consumption.

N₂O is a potent greenhouse gas with a global warming potential 296 × that of CO₂ (IPCC, 2007). However, measuring N₂O directly in the field is often difficult with chamber methods in cropping systems that produce large above-ground biomass. Additionally, including multiple field treatments (e.g. as in a randomized complete block design) makes micrometeorological methods of N₂O flux impossible to obtain. Instead, relative abundances of denitrifier genes can be used to assess a soil’s potential to produce (e.g. nirS or nirK) and consume (e.g. nosZ) N₂O via denitrification, representing a qualitative proxy of relative N₂O emission potential of a soil (Butterbach-Bahl et al., 2013; Hallin et al., 2009; Morales et al., 2010; Petersen et al., 2012; Philippot, 2002). Denitrifier community size has been correlated with denitrification process rates (Hallin et al., 2009; Wu et al., 2012) and denitrification potential (Attard et al., 2011; Cuhel et al., 2010; Enwall et al., 2010). Potential denitrifying activity and denitrifying community size have also been shown to be correlated with each other in some studies (Hallin et al., 2009; Morales et al., 2010; Szukics et al., 2010; Throbäck et al., 2007), suggesting community size may indicate potential differences in soil N processes after LUC. Particularly, the nosZ-bearing community may act as a N₂O sink and counter high N₂O production rates (Braker and Conrad, 2011; Philippot et al., 2011), therefore influencing N₂O emissions (Cuhel et al., 2010; Morales et al., 2010; Philippot et al., 2011).

Denitrification nirS and nosZ gene targets represent the two most important steps in the denitrification pathway that produce gaseous by-products, and account for a large proportion of functional N genes in some studies (Stone et al., 2015). The first step in denitrification that produces a gaseous N product is the reduction of nitrite (NO₃⁻) to nitric oxide (NO), catalyzed by nitrite reductases either encoded by the cytochrome cd₁ (nirS) or copper-containing (nirK) genes, which are equivalent but have not been detected within the same species (Zumft, 1997). We chose to quantify nirS because three-quarters of cultured denitrifiers possess the nirS gene (Zumft, 1997) and some molecular reports indicate nirS may dominate in abundance over nirK in some natural environments (Deslille et al., 2014; Nogales et al., 2002), indicating it may be a better-suited target for relative characterization of potential nitrite-reducing communities than nirK. Additionally, nirK has been recently identified in autotrophic ammonia-oxidizing species (Cantera and Stein, 2007; Casciotti and Ward, 2001), calling into question its utility in specifically targeting denitrifying communities. The nosZ target codes for nitrous oxide reductase, which catalyzes the reduction of N₂O to N₂ in the denitrification pathway, indicating nosZ-bearing communities help to complete the N cycle and determine the N₂O:N₂ balance. 16S rRNA was chosen as a molecular target for the total bacterial community size; although 16S rRNA gene copies vary from 1 to 15 copies per genome, its use has continued to be the “gold standard” for microbial ecology (Case et al., 2007; Vos et al., 2012). Although an alternate target, such as rpoB, which is a single copy gene would be valuable if assessing phylogenetic diversity, there are no universal primers for it (Adékambi et al., 2009) as rpoB is not conserved enough to be of use as a universal marker and only a subset of the microbial community can be targeted (Vos et al., 2012). Many studies have used 16S rRNA gene copy numbers as a proxy for the total bacterial community size, and some have found that the total estimated numbers of proteobacteria species was not significantly different if using rpoB or 16S rRNA markers (Vos et al., 2012). As this study has not assessed phylogenetic relationships of the microbial communities, 16S rRNA is an appropriate target for the relative comparison of the overall bacterial community size between environmental treatments/variables.

LUC resulting from displacement of corn–soybean rotations by PG production may alter soil microhabitats and therefore soil microbial N-cycling due both to extensive root and rhizome biomass and to large leaf litter inputs to soils in perennial vs. annual systems (Dohleman et al., 2012). Within studies targeting soil microbial communities in biomass cropping systems to date (Hedenec et al., 2014; Liang et al., 2012; Mao et al., 2013, 2011; Orr et al., 2015), the effects of various management practices (e.g. N fertilization and harvest) on soil microbial community functioning have not been an area of focus. The effect this type of LUC may have on soil microbial communities may depend on PG management practices in these systems.

Currently, there is no consensus regarding optimal N fertilization practices for increased yields in PG production as yield responses can be highly variable depending on environmental conditions and crop species (Deen et al., 2011). Depending on downstream use, miscanthus can be harvested in the fall pre-frost, harvested post-frost kill, or left to overwinter as standing biomass for harvest in the spring. Switchgrass is commonly harvested in the fall, and is often windrowed (cut, swathed, and left on soil) over winter due to produc-
ers' limitations in collecting and storing harvested biomass in winter (REAP, 2008; Sokhansanj et al., 2009). Differences in N fertilizer requirements and harvest regimes may alter C and N inputs (Attard et al., 2011) and may influence LUC impacts on soil denitrifier community sizes.

Our objective was to compare the effects of LUC from corn–soybean to PG biomass production on the relative abundances of total (16S rRNA gene target) and denitrifier (nirS and nosZ gene targets) soil bacterial communities 3–4 years after PG planting. Soil was collected on four dates from 2011 to 2012 from a field trial established in Ontario in 2008. This study is unique in that it consists of two PG biomass crops produced in parallel with the existing common land use of corn–soybean rotation within the same field trial site. It also includes unfertilized and fertilized plots in both annual and perennial systems, and varied harvest practices within PG plots. We hypothesized that soils from PG plots would support larger total bacterial and denitrifier communities than soils from corn–soybean plots due to increased shoot residue return and root inputs to soils in PG systems, as well as that soils from PG plots with biomass harvested in the spring would support larger total bacterial and denitrifier communities than supported by soils from PGs harvested in the fall due to increased root inputs and leaf loss to soil over winter.

2 Materials and methods

2.1 Site description and experimental design

A field trial was established in 2008 at the University of Guelph Research Station in Elora, ON (43°38'46.73" N, 80°24'6.66" W). The field site was cultivated on 16 May and 6 June 2008. Switchgrass (Panicum virgatum L. “Shelter”) was planted on 6 June 2008. Miscanthus (M. sinensis × M. sacchariflorus “Nagara”, M116) was planted on 12 June 2008, and soybean (Glycine max L.) was planted on 24 June 2008 and annually rotated with corn (Zea mays L.). Corn was planted on 5 May 2010, soy was planted on 3 June 2011, and corn was re-planted on 18 May 2012, with annual light cultivation to prepare seedbeds for planting. In 2007, prior to trial establishment, the experimental area was planted to barley (Hordeum vulgare L.). The soil type is a London silt loam (Gray Brown Luvisol).

The field trial was a split–split–strip plot design with three replications. The main plot factor was PG crop or an-
annual rotation (miscanthus, switchgrass, and corn–soybean). Main treatment plots measured 6.2 m × 26.0 m. Nitrogen fertilizer (0 or 160 kg N ha⁻¹) was applied in strips randomly within replicates. 160 kg N ha⁻¹ subplots received hand-broadcast urea fertilizer (46-0-0; N-P-K) in May 2011 or hand-broadcast ammonium nitrate fertilizer (34-0-0; N-P-K) in May 2012, after soil sampling procedures described below. Main treatments were split into two harvest timings (fall or spring) within the PG fertilizer strips only. Miscanthus plots were either harvested in the late fall season after

post-frost kill or left standing to overwinter until spring harvest. Switchgrass plots were harvested in the fall or cut and assembled into windrows in the field for biomass removal in the spring. Spring harvest of PGs occurred before N fertilizer was applied. Harvest methods of PG yields (dry harvested biomass (tonnes) ha⁻¹) are described in Deen et al. (2011).

Figure 1 illustrates the seasonal and annual variation in daily average air temperature (°C) and daily precipitation (mm) measured at the Elora Research Station.

2.2 Soil sampling and analysis

Baseline bulk density and carbon measurements were measured for each main plot on 23 October 2010. For bulk density, two soil cores per plot were collected at 0–5 cm depth using 2.5 cm diameter cylindrical aluminum cores. Cores were weighed before and after drying for 24 h at 105 °C (Maynard and Curran, 2007). For soil carbon analysis, 10 soil cores per plot were collected from both 0–15 and 15–30 cm depths using a 5 cm diameter soil corer on a Z-shaped transect, and then composited per treatment plot for each depth. Total soil carbon and inorganic carbon were analyzed with a Leco® Carbon Determinator CR-12 (model no. 781-700, Leco Instruments Ltd.) following the dry combustion technique (Périé and Ouimet, 2008) on approximately 0.300 g of dried, ground, and homogenized soil (Table 1).

For molecular analyses, soil was sampled on four dates (9 May 2011, 30 October 2011, 2 May 2012, and 20 October 2012). October sampling dates occurred before fall harvest of PG crops, while May sampling dates occurred before N fertilizer application and after spring PG biomass removal (Fig. 1). Ten soil cores per plot were sampled aseptically to 15 cm depth using a 5 cm diameter soil corer on a Z-shaped transect, composited and kept on ice until transport back to the laboratory. The transect shape was chosen to encompass plot heterogeneity; at a pre-trial study date, initial analysis indicated gene abundances were not significantly different between bulk or rhizosphere soils in corn–soybean or PG plots, possibly due to the large root biomass/leaf loss to soils in perennial plots and residual soy/corn residue cover on soil in corn–soybean plots. Soil samples were divided for storage at 4 and −20 °C.

Mean values of gravimetric soil moisture (g g⁻¹) are shown in Fig. 1. Soil exchangeable NO₃⁻-N and NH₄⁺-N were determined for each of the soil samples by KCl extraction. Soil samples (10.0 g) were placed into 125 mL flasks and 100 mL of 2.0 M KCl was added to each flask. Flasks were stoppered and shaken for 1 h at 160 strokes per minute; solutions were allowed to settle and were then filtered through Whatman no. 42 filter paper (Whatman plc, ME, USA). Extractable NO₃⁻-N and NH₄⁺-N were determined colorimetrically with segmented flow analyses (AA3, SEAL Analytical, Wisconsin, USA) via a cadmium reduction (US Environmental Protection Agency, 1974) and a Berthelot reaction, respectively (Fig. 2).
Table 1. Mean soil properties measured at the Elora Research Station.

<p>| Cropping system/ | N rate  | Bulk | % organic carbon | % total carbon | Yield         |</p>
<table>
<thead>
<tr>
<th>harvest</th>
<th>(kg ha(^{-1}))</th>
<th>density(^1)</th>
<th>0–15 cm</th>
<th>15–30 cm</th>
<th>0–15 cm</th>
<th>15–30 cm</th>
<th>2011</th>
<th>2012</th>
<th>Mean(^2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Corn–soybean</td>
<td>Fall</td>
<td>0</td>
<td>1.21 AB</td>
<td>1.88</td>
<td>1.06</td>
<td>2.22</td>
<td>2.16</td>
<td>5.341</td>
<td>2.912</td>
</tr>
<tr>
<td></td>
<td>Fall</td>
<td>160</td>
<td>1.27 A</td>
<td>1.79</td>
<td>1.47</td>
<td>2.25</td>
<td>2.11</td>
<td>9.92</td>
<td>7.882</td>
</tr>
<tr>
<td>Miscanthus</td>
<td>Fall</td>
<td>0</td>
<td>1.10 B</td>
<td>2.06</td>
<td>1.44</td>
<td>2.27</td>
<td>1.72</td>
<td>17.62</td>
<td>12.77</td>
</tr>
<tr>
<td></td>
<td>Fall</td>
<td>160</td>
<td>1.10 B</td>
<td>2.13</td>
<td>1.63</td>
<td>2.36</td>
<td>1.84</td>
<td>17.43</td>
<td>18.32</td>
</tr>
<tr>
<td>Miscanthus</td>
<td>Spring</td>
<td>0</td>
<td>1.13 AB</td>
<td>2.09</td>
<td>1.53</td>
<td>2.31</td>
<td>1.69</td>
<td>12.66</td>
<td>13.38</td>
</tr>
<tr>
<td></td>
<td>Spring</td>
<td>160</td>
<td>1.13 AB</td>
<td>2.24</td>
<td>1.42</td>
<td>2.47</td>
<td>1.89</td>
<td>14.33</td>
<td>14.56</td>
</tr>
<tr>
<td>Switchgrass</td>
<td>Fall</td>
<td>0</td>
<td>1.11 B</td>
<td>2.12</td>
<td>1.43</td>
<td>2.33</td>
<td>1.61</td>
<td>7.648</td>
<td>6.458</td>
</tr>
<tr>
<td></td>
<td>Fall</td>
<td>160</td>
<td>1.09 B</td>
<td>2.12</td>
<td>1.34</td>
<td>2.32</td>
<td>1.73</td>
<td>11.1</td>
<td>10.45</td>
</tr>
<tr>
<td>Switchgrass</td>
<td>Spring</td>
<td>0</td>
<td>1.11 B</td>
<td>2.09</td>
<td>1.23</td>
<td>2.32</td>
<td>1.55</td>
<td>6.33</td>
<td>4.146</td>
</tr>
<tr>
<td></td>
<td>Spring</td>
<td>160</td>
<td>1.21 AB</td>
<td>1.92</td>
<td>1.33</td>
<td>2.23</td>
<td>1.7</td>
<td>6.905</td>
<td>6.441</td>
</tr>
</tbody>
</table>

\(^{1}\) Means of bulk density (\(n = 6\)) followed by the same letter within one column are not significantly different according to a post hoc Tukey’s means comparison (\(p < 0.05\)); carbon measurements (\(n = 3\)) were not significantly different between treatments. \(^{2}\) Mean yields (\(n = 3\)) followed by the same letter are not significantly different according to a post hoc Tukey’s means comparison (\(p < 0.05\)).

Figure 1. Mean daily air temperature (\(^\circ\)C) and daily precipitation (mm) at the Elora Research Station from January 2011 to November 2012. Soil gravimetric H\(_2\)O was measured on a per-sample basis and is shown as crop means (±SE) for each sampling date (9 May 2011, 30 October 2011, 2 May 2012, and 20 October 2012) (\(n = 12\) in perennial grasses, \(n = 6\) in corn–soybean rotation).

2.3 Soil DNA extraction

Total DNA was extracted from field-moist soil sampled from each plot (three field replicates, \(n = 3\); total plots \(n = 30\)). DNA was extracted in duplicate (\(~0.250\) g) within 48 h of sampling as per manufacturer’s protocol using a PowerSoil DNA isolation kit (Mobio, Carlsbad, USA). Duplicate extracts were then pooled, separated into aliquots, and stored at −80 °C until use in downstream analyses.

2.4 Quantification of total bacteria and functional genes

Quantitative PCR (qPCR) assays were used to enumerate the total bacterial communities (16S rRNA gene) and communities of denitrifiers by targeting nitrite reductase (nir\(S\)) and nitrous oxide reductase (nos\(Z\)) genes, using primer pairs 338f/518r (16S rRNA; Fierer et al., 2005), Cd3a/R3Cd (nir\(S\); Throbäck et al., 2004), and 1F/1R (nos\(Z\); Henry et al., 2006).

For each gene target analyzed, duplicate replicates were run in parallel on an IQ5 thermocycler (Bio-Rad Laborato-
for 60 s (nirS 30 s, annealing at 57 °C for 2 min, followed by 35 cycles of denaturing at 94 °C) with a 100 bp ladder, which resulted in clean bands and have been cited as having 89–91 % match-efficiency to all bacteria (Bergmark et al., 2012). The primers amplify one of two conserved regions in V3 of the SSU rRNA gene, resulting in a 200 bp amplicon that is within small enough to amplify via qPCR methodology and amplifies for most bacterial taxa (Bakke et al., 2011).

Known template standards were made from cloned PCR products from pure culture genomic DNA (Clostridium thermocellum (16S), Pseudomonas aeruginosa (nirS), and Pseudomonas fluorescens (nosZ)) and transformed into Escherichia coli plasmids (TOPO TA cloning kit); plasmids were sequenced to confirm successful cloning and transformation of the target genes. Amplicon specificity was screened by running PCR products on an ethidium bromide-stained gel (1 % agarose, 80 V for 20 min) with a 100 bp ladder. PCR amplicons of cloned gene targets were sequenced by the Laboratory Services Department at the University of Guelph using an ABI Prism 3720 (Applied Biosystems, Foster City, CA, USA) to confirm target identity. In all qPCR assays, all unknown samples were amplified in parallel with a triplicate serial dilution (10^1–10^8 gene copies per reaction) of control plasmids. PCR assays were optimized to ensure efficiencies ranging from 93.0 to 106.4 %, with R^2's ranging from 0.990 to 0.999 and standard curve slopes of −3.177 to −3.408 by testing serial dilutions of DNA extracts in order to minimize inhibition of amplification due to humic and fulvic contaminants. Duplicate no-template controls were run for each qPCR assay, which gave null or negligible values. Melt curve analysis was used to confirm amplicon specificity. Normalization of DNA concentrations to grams of dry soil was used to give results on a biologically significant scale, which assumes similar DNA isolation efficiency across samples.

2.5 Statistical analysis

Analysis of variance was conducted in SAS 9.3 (Carlsbad, NC, USA) using a generalized linear mixed model (PROC GLIMMIX). The Shapiro–Wilks test was used to test for normality of data; studentized test for residuals confirmed the absence of outliers. The probability distributions of gene abundance data sets were log-normal or highly skewed and were analyzed using an overdispersed Poisson distribution for count data (Ver Hoef and Boveng, 2007). Bulk density, organic carbon, total carbon, nitrate, and ammonium data were log-transformed when required and fitted to the normal distribution. Within each data set, sampling time was a repeated measure; independent and interactive fixed effects were associated with crop/crop rotation, nitrogen application rate, and harvest timing within perennial grasses, while field replicate and its associated interactions were random effects. The residual maximum likelihood method was employed to fit the model for all data sets. Several covariance structures were entertained before the variance components structure was chosen based on convergence and model fitting criteria. Individual treatment means within data sets were compared using a post hoc Tukey’s test for all pairwise comparisons, with significance denoted at p < 0.05.

Correlation analysis was used to assess nonparametric measures of statistical dependence between gene abundances.
and H₂O, NO₃⁻-N and NH₄⁺-N measured over time (Table S1 in the Supplement). Correlation analysis resulted in multiple significant correlations between variables; as such a principal component analysis was conducted in SAS (PROC FACTOR) on 120 samples using a VARIMAX rotation.

3 Results

3.1 Environmental and soil conditions

Environmental conditions varied during the periods prior to the four soil sampling dates (Fig. 1). Average air temperatures over the growing seasons (May–October) were 16.9 and 17.3 °C in 2011 and 2012, respectively (Roy et al., 2014); average air temperatures in spring 2012 were warmer than normal and resulted in earlier emergence of PG crops compared to 2011. Cumulative monthly precipitation was above average prior to the May 2011 sampling date (101 mm vs. 72 mm 30-year average in April 2011 and 113 mm vs. 82 mm 30-year average in May 2011) (Roy et al., 2014). In comparison, southern Ontario received very low cumulative precipitation in April 2012 (30 mm vs. 72 mm 30-year average) and May 2012 (28 mm vs. 82 mm 30-year average) (Roy et al., 2014). Cumulative precipitation levels were lower in 2012 compared to 2011 from May to August (391 mm in 2011 vs. 186 mm in 2012); however, higher than normal precipitation levels occurred in October of 2011 (129 mm vs. 77 mm 30-year average) and both September (106 mm vs. 77 mm 30-year average) and October (127 mm vs. 77 mm 30-year average) of 2012 (Roy et al., 2014). Environmental conditions prior to soil sampling directly impact soil gravimetric content measured at the time of sampling (Fig. 1) and could also impact soil N and soil bacterial communities.

Soil physical and chemical properties were assessed in October 2010, after only 2 years of contrasting management since crop establishment in 2008. The corn–soybean rotation had higher soil bulk density than soils from both miscanthus and switchgrass plots harvested in the fall. No differences in total or organic soil carbon were detected between the corn–soybean rotation and the PG treatments at either the 0–15 or 15–30 cm depth (Table 1). Soil NH₄-N levels did not differ significantly between the corn–soybean rotation and the PG soils; however, N fertilization significantly increased NH₄-N levels in soils from fall-harvested miscanthus plots (p < 0.05) (Fig. 2a). N fertilization also significantly increased NO₃-N levels in spring-harvested switchgrass (p < 0.05) (Fig. 2b). From May to October 2011, soil NH₄-N levels increased significantly and soil NO₃-N levels decreased significantly in PG soils (data not shown); a similar trend was not observed in 2012 or for soils from the corn–soybean rotation in either year.

3.2 Biomass yields

Despite significant differences in precipitation between 2011 and 2012, biomass yields of miscanthus and switchgrass did not differ between years. In comparison, corn grain yields were higher in 2011 vs. 2012 (Table 1). Miscanthus produced higher yields (12.7–18.3 dry t ha⁻¹) than either switchgrass or corn grain, regardless of N fertilization rate or harvest timing (Table 1). When harvested in the fall and N-fertilized, switchgrass yields were not significantly lower (10.5–11.1 dry t ha⁻¹) than miscanthus yields. Switchgrass yields from unfertilized plots were not significantly different if harvested in the fall or spring; however, switchgrass yields from fertilized plots were significantly higher (∼3–4 dry t ha⁻¹) when harvested in the fall compared to yields obtained when switchgrass was windrowed over winter.

3.3 Bacterial responses to annual and perennial crops and their management

There was no statistically significant effect of N fertilization or any significant interactions between cropping system and sampling time on any of the targeted gene abundances. Therefore, we analyzed the impact of each biomass crop under specific harvest management on soil bacterial gene abundances (Table 2). Denitrifying (nosZ) gene copy abundances were affected by LUC; regardless of harvest or N management, mean nosZ gene copies were higher in miscanthus plots than in the corn–soybean rotation, and nirS:nosZ ratios were higher in the corn–soybean soils than in miscanthus or switchgrass soils (p < 0.05) (Table 2). Under fall harvesting management, biomass crop had no impact on total bacterial 16S rRNA gene copies or nirS gene copies. However, leaving miscanthus biomass standing over winter until spring resulted in significantly higher 16S rRNA gene copies than observed in soils from fall-harvested biomass crops and significantly higher nirS gene copies than in fall-harvested switchgrass or the corn–soybean rotation (Table 2).

3.4 Temporal changes in bacterial gene abundances

Sampling date had a significant impact on gene abundances for all genes quantified (Fig. 3). Over both sampling years, 16S rRNA gene copies were significantly higher (5.2–5.4 × 10⁶ gene copies g⁻¹ dry soil) at fall (October) sampling dates compared to the ~5.5–6.4 × 10⁶ gene copies g⁻¹ dry soil quantified at spring (May) sampling dates (Fig. 3). Populations of nirS and nosZ denitrifiers represented ~1.58 and 0.26% on a gene-to-gene basis (nirS or nosZ to 16S) of the total bacterial community (data not shown) and did not follow similar trends with time of sampling (Fig. 3). The abundance of nirS gene copies was significantly higher in 2012 (4.0 × 10⁶–1.6 × 10⁷ gene copies g⁻¹ dry soil) compared to 2011 (2.5–6.3 × 10⁵ gene copies g⁻¹ dry soil), with no significant differences between May and October sampling dates within each year (Fig. 3). The abundance of nosZ gene
Table 2. Mean gene abundance responses to crop and harvest management, averaged over nitrogen application rate and time at the Elora Research Station.

<table>
<thead>
<tr>
<th>Cropping system</th>
<th>Management</th>
<th>Total soil bacteria (gene copy g(^{-1}) soil)(^{*})</th>
<th>Soil denitrifying bacteria (gene copy g(^{-1}) soil)(^{*})</th>
<th>(nirS):(nosZ) ((\times 10^{-2}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Corn–soybean</td>
<td>Fall harvest</td>
<td>(1.35 \times 10^9)(^b)</td>
<td>(1.95 \times 10^6)(^b)</td>
<td>(2.63 \times 10^5)(^b)</td>
</tr>
<tr>
<td>Miscanthus</td>
<td>Fall harvest</td>
<td>(1.38 \times 10^9)(^b)</td>
<td>(2.30 \times 10^6)(^{ab})</td>
<td>(4.47 \times 10^5)(^a)</td>
</tr>
<tr>
<td>Miscanthus</td>
<td>Spring harvest</td>
<td>(1.91 \times 10^9)(^a)</td>
<td>(3.02 \times 10^6)(^a)</td>
<td>(5.25 \times 10^5)(^a)</td>
</tr>
<tr>
<td>Switchgrass</td>
<td>Fall harvest</td>
<td>(1.41 \times 10^9)(^b)</td>
<td>(2.19 \times 10^6)(^b)</td>
<td>(3.55 \times 10^5)(^{ab})</td>
</tr>
<tr>
<td>Switchgrass</td>
<td>Spring windrow</td>
<td>(1.48 \times 10^9)(^{ab})</td>
<td>(2.46 \times 10^6)(^{ab})</td>
<td>(3.98 \times 10^5)(^{ab})</td>
</tr>
</tbody>
</table>

* Means followed by the same letter within one column are not significantly different according to post hoc Tukey’s means comparison at \(p < 0.05\) (\(n = 24\)).

Figure 3. Mean log gene copies (g\(^{-1}\) dry soil ± SE) in annual and perennial biomass cropping systems under varied harvest management at the Elora Research Station (\(n = 6\)) over time. Different letters within panels indicate significant differences according to a post hoc Tukey’s test (\(p < 0.05\)).

Copies was approximately 1.3–3.2 \(\times 10^5\) gene copies g\(^{-1}\) dry soil, but this increased significantly in May 2012 to approximately 3.2 \(\times 10^6\) gene copies g\(^{-1}\) dry soil and dropped back to previous levels by October 2012 (Fig. 3). Higher relative proportions of denitrifiers (\(nirS\) or \(nosZ\) to 16S) were observed at spring sampling dates, when total bacterial 16S rRNA gene abundances decreased in comparison to fall sampling dates (Fig. 3).

Two factors were selected in the principal components analysis, which accounted for 67.73% cumulative variance. A scree plot was examined for breaks, and factors were retained for eigenvalues \(\geq 1\). Soil NH\(_4\)-N\(^+\), soil NO\(_3\)-N, \(nirS\), and \(nosZ\) loaded on factor 1, which accounted for 43.89% variance, while soil gravimetric H\(_2\)O and 16S rRNA loaded on factor 2, which accounted for 23.84% variance (Fig. 4a and b). Differences in soil NO\(_3\)-N and NH\(_4\)-N were strongly related to differences in \(nirS\) and \(nosZ\) gene abundances observed between May 2011 and May 2012 sampling dates (Figs. 3 and 4), while the size of the total bacterial com-
systems is to produce crops with high yields, such as PGs (Sanscartier et al., 2014), which offset the amount of land required for crop production (Kludze et al., 2013). However, if PG biomass production negatively affects soil health as indicated by changes in the potential functioning of microbial communities, large-scale LUC from annual to perennial biomass production may not be as sustainable as originally proposed. As such it is necessary to identify biomass cropping systems that not only result in large biomass yields but also ensure agroecosystem sustainability by maintaining or improving ecosystem services (Orr et al., 2015), such as soil N cycling.

4 Discussion

Denitrification is an important process contributing to the production and consumption of N\textsubscript{2}O in soils, and mitigation of greenhouse gases such as N\textsubscript{2}O is required to create sustainable biomass cropping systems (Miller et al., 2008; Schlesinger, 2013). Changes in the potential functional abilities of the soil microbial community may reflect changes in LUC or agricultural management and should be considered to assess the ecological impact of biomass crop production (Hedenec et al., 2014). Currently, few studies have assessed soil microbial community responses to PG biomass production systems (Hedenec et al., 2014; Liang et al., 2012; Mao et al., 2013, 2011; Orr et al., 2015). The highest potential to reduce greenhouse gas emissions from biomass cropping community (16S rRNA) was related to soil gravimetric moisture levels (Fig. 4).

4.1 Biomass yields of annual and perennial crops

Miscanthus and switchgrass biomass yields were within the typical range of values reported previously in Ontario (Kludze et al., 2013; REAP, 2008) and Europe (Christian et al., 2008; Himken et al., 1997), despite differences in temperature and precipitation between the two study years. Corn grain yields were within the lower range for reported Ontario yields (Munkholm et al., 2013), potentially due to wetter (2011) and drier (2012) field conditions than normal over the two growing seasons (Roy et al., 2014). Deen et al. (2011) showed increases in PG biomass yields between the second and third years after PG planting at our site, whereas we measured similar yields in 2011 and 2012, indicating the PGs may have reached maximum yield potential.

Nitrogen fertilization significantly increased corn grain yields and fall-harvested switchgrass biomass yields; however, no significant increases due to N fertilization were observed in miscanthus or spring-harvested switchgrass biomass yields. Potential yield increases from N fertilization in spring-harvested switchgrass may have been offset due to leaf loss over the winter season, as increases in switchgrass yields to N fertilization have been previously observed (Nikièma et al., 2011; Vogel et al., 2002). Similar to the present study, European and US field trials have also found no response of miscanthus yields to N (Lewandowski et al., 2000, 2003; Behnke et al., 2012; Christian et al., 2008), and PG yields were minimally impacted by differences in growing season conditions compared to corn grain yields (Table 1).

Despite significant differences in biomass yields between miscanthus and corn–soybean systems, there were no significant differences in either total or organic soil carbon between any of cropping systems assessed (Table 1). Sampling of soil carbon occurred only two years after PG planting; PGs are expected to be productive for 20+ years, indicating future changes in soil carbon levels may occur. Additionally, Ontario-based land conversion modelling scenarios have estimated a soil carbon decrease of 2.5% upon miscanthus establishment (Sanscartier et al., 2014), which may have negated potential increases in soil organic carbon. However, high miscanthus yields most likely resulted in increases
in above- and below-ground plant residue return to soils (Mutegi et al., 2010; Soil Quality National, 2006); therefore, our carbon measures may not have reflected short-term changes in labile carbon sources that had occurred. Regardless of management or climatic conditions, miscanthus consistently produced large yields, emphasizing its potential as a bioenergy crop suitable for production in variable Ontario conditions.

4.2 Bacterial responses to annual and perennial crops and their management

Some studies in biomass cropping systems have not observed differences in soil microbial responses between perennial and annual crop types (Mao et al., 2011), while others have measured significant differences in microbial abundance, diversity and community structure between these cropping types (Liang et al., 2012; Morales et al., 2010; Watrud et al., 2013). Currently, we observed significantly higher nosZ gene copies in miscanthus soils compared to corn–soybean soils, illustrating a distinct effect of LUC from corn–soybean to miscanthus production on soil N cycling (Table 2).

Due to the large biomass produced by miscanthus compared to corn, a large amount of plant residues are returned to the soil; these residues provide surface cover, decrease soil bulk density, increase water retention, and regulate temperatures (Blanco-Canqui and Lal, 2009). Previous work at the Elora Research Station found an inverse correlation between field-scale N₂O fluxes and nosZ transcript abundance in conventionally tilled corn plots with residues returned to soils (Németh et al., 2014), and increased nosZ activity after residue amendment has also been observed in lab studies (Henderson et al., 2010). High C:N plant residues have been negatively correlated with cumulative N₂O emissions (Huang, 2004) and may encourage complete reduction of N₂O to N₂ as soil available NO₃-N is limiting, so bacterial populations with the ability to reduce N₂O to N₂ are favoured (Miller et al., 2008). Presently, the primers used for nosZ gene target amplification provided good coverage of γ-Proteobacteria (Henry et al., 2006), which are stimulated by surface-applied residues (Pascault et al., 2010). Increased residue return in miscanthus plots may have selected for bacterial populations harbouring enhanced catabolic capabilities, such as N₂O reduction (Pascault et al., 2010). This implies that producing biomass crops with large yields may indirectly alter soil N cycling and potentially mitigate soil N₂O emissions due to increased residue return influencing the soil microbial community. It is likely that differences in environmental conditions (e.g. temperature, H₂O and O₂ availability) and resource quality and availability between corn–soybean and miscanthus soils related to differences in microbial community structure (Cusack et al., 2011) and selected for different dominant taxa that filled different ecological niches (Stone et al., 2015).

N fertilization did not affect targeted gene abundances; however, studies in other cropping systems have found that N fertilization affected the size of denitrifying communities (Hallin et al., 2009), nitriﬁying communities (He et al., 2007), and proportions of nirS to nirK communities (ratio of nirS:nirK genes) (Hai et al., 2009). Elevated 16S rRNA and nirS gene copies were observed in soils from spring-harvested miscanthus and windrowed switchgrass (Table 2). Increased N return via senescent leaf loss in PG plots over winter contributes to the soil organic matter pool (Heaton et al., 2009) and may have contributed to elevated total (16S rRNA) bacterial populations in these soils, concomitantly increasing nirS abundances (Huang et al., 2011).

4.3 Temporal changes in bacterial gene abundances

Total soil bacterial communities (16S rRNA) followed a seasonal trend, with elevated 16S rRNA gene copies at fall (October) compared to spring (May) sampling dates, possibly due to an increase in the availability and diversity of resources for microbial metabolism and growth over the growing season (Habekost et al., 2008). Denitrifying abundances changed differently than the total bacterial community, suggesting denitriﬁers were influenced by different proximal regulators than the total bacterial community (Figs. 3 and 4). Seasonal dynamics of N-cycling microbial communities have been previously characterized (Boyer et al., 2006; Németh et al., 2014; Wolsing and Priemé, 2004; Dandie et al., 2008; Bremer et al., 2007) and are tightly coupled with seasonal changes in labile C and N pools, temperature, and soil H₂O (Butterbach-Bahl et al., 2013; Rasche et al., 2011), indicating that local edaphic drivers may often take precedence over crop-specific drivers (Mao et al., 2013).

5 Conclusions

Miscanthus consistently produced large yields and supported larger nosZ-bearing communities than the corn–soybean rotation, emphasizing its influence on soil N cycling and its potential to mitigate soil N₂O emissions while being suitable for production in variable Ontario conditions. Additionally, miscanthus yields were not increased with N fertilization, indicating a lower N input requirement for biomass production compared to switchgrass. Higher 16S rRNA and nirS gene copies were associated with reduced yields in spring-harvested PGs, indicating that harvesting PGs in the spring may increase populations of denitriﬁers capable of producing N₂O emissions while simultaneously decreasing biomass yields. The size of both denitrifying (nirS and nosZ) and total bacterial (16S rRNA) communities changed over the sampling period; however, changes in denitrifying gene abundances did not parallel changes in the total soil bacterial community, indicating denitriﬁying communities were regulated differently than the total bacterial community. Future work measuring N₂O emissions and denitriﬁer activity (mRNA)
and community structure in these systems is required to link the effects of LUC on these communities directly with \( \text{N}_2\text{O} \) fluxes.

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