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Antibiotic Effects on Microbial Communities Responsible for Denitrification and N$_2$O Production in Grassland Soils

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Antibiotics in soils may affect the structure and function of microbial communities. In this study, we investigated the acute effects of tetracycline on soil microbial community composition and production of nitrous oxide (N$_2$O) and dinitrogen (N$_2$) as the end-products of denitrification. Grassland soils were pre-incubated with and without tetracycline for 1-week prior to measurements of N$_2$O and N$_2$ production in soil slurries along with the analysis of prokaryotic and fungal communities by quantitative polymerase chain reaction (qPCR) and next-generation sequencing. Abundance and taxonomic composition of bacteria carrying two genotypes of N$_2$O reductase genes (nosZ-I and nosZ-II) were evaluated through qPCR and metabolic inference. Soil samples treated with tetracycline generated 12 times more N$_2$O, but N$_2$ production was reduced by 84% compared to the control. In parallel with greater N$_2$O production, we observed an increase in the fungi:bacteria ratio and a significant decrease in the abundance of nosZ-II carrying bacteria; nosZ-I abundance was not affected. NosZ-II-carrying Bacillus spp. (Firmicutes) and Anaeromyxobacter spp. (Deltaproteobacteria) were particularly susceptible to tetracycline and may serve as a crucial N$_2$O sink in grassland soils. Our study indicates that the introduction of antibiotics to agroecosystems may promote higher N$_2$O production due to the inhibitory effects on nosZ-II-carrying communities.

Keywords: denitrification, nitrous oxide, tetracycline, nosZ, bacteria, fungi

INTRODUCTION

Microbial denitrification is a dominant respiratory pathway for reactive N-removal in terrestrial and aquatic ecosystems. Diverse microorganisms belonging to several genera of bacteria, archaea, and fungi perform denitrification (Stein and Klotz, 2017). Dinitrogen gas (N$_2$) is the end-product of complete denitrification: the stepwise reduction of nitrate (NO$_3^-$) and nitrite (NO$_2^-$) to nitric oxide (NO), nitrous oxide (N$_2$O), and dinitrogen (N$_2$). However, many microorganisms do not carry the necessary genes to perform complete denitrification and may instead release N$_2$O as the end-product of incomplete denitrification. Since N$_2$ is radiatively inert and N$_2$O is a potent greenhouse gas and the dominant source of stratospheric ozone depletion, the atmospheric impacts of complete and incomplete denitrification are dramatically different (Ravishankara et al., 2009; Neubauer and Megonigal, 2015). Denitrifiers capable of complete denitrification to N$_2$ rely on the presence and expression of the nosZ gene, which encodes N$_2$O reductase, the enzyme that converts N$_2$O to N$_2$.
(Thomson et al., 2012). NosZ-carrying prokaryotes, found in both Archaea and Bacteria, consume N₂O and therefore are an important biological sink for N₂O in soils (Ma et al., 2011; Jones et al., 2014; Domeignoz-Horta et al., 2015; Samad et al., 2016). Because the nosZ gene is missing in fungal genomes, fungal denitrifiers have an incomplete denitrification pathway and are a source of N₂O (Shoun et al., 2012; Maeda et al., 2015; Mohapo et al., 2015).

Recent studies of nosZ gene diversity have revealed the presence of two nosZ genotypes, clade I (nosZ-I) and clade II (nosZ-II), that have different phylogenetic, physiologic, and ecological properties (Hallin et al., 2017). The taxonomic and genetic diversity of nosZ-II carrying prokaryotes is greater than nosZ-I and is positively correlated to soil N₂O sink capacity (Jones et al., 2013, 2014; Domeignoz-Horta et al., 2015). Additionally, a larger percent of nosZ-II carrying prokaryotes do not possess the genes that encode for nitrite reductase (Nir) and nitric oxide reductase (Nor) enzymes, which are necessary to produce N₂O (Graf et al., 2014; Jones et al., 2014). This indicates that nosZ-II carrying bacteria are mostly N₂O consumers, rather than N₂O producers. The abundance and diversity of the two clades also appear to respond differently to environmental parameters including pH, moisture, and nutrient concentrations (Jones et al., 2014; Domeignoz-Horta et al., 2015; Samad et al., 2016). When considering responses to environmental contaminants, such as antibiotics, the differential responses of the two nosZ genotypes have not been studied yet.

Antibiotics are introduced into soils from multiple anthropogenic sources, such as in applications of animal manure and biosolids, inappropriate disposal of unused medicines, and in wastewater treatment effluents (Boxall, 2004). Animals excrete between 17 and 90% of antibiotics administered during livestock production in their feces and urine (Massé et al., 2014), and soil concentrations can range from a few microgram to gram per kilogram of soil (Thiele-Bruhn, 2003). Animal manures with antibiotics are frequently applied to agricultural soils as fertilizer (Sengelov et al., 2003; Zhu et al., 2013) where they can affect microbial denitrification and N₂O production. To achieve this goal, we conducted a laboratory experiment with grassland soil samples treated with tetracycline, an inhibitor of bacterial protein synthesis that belongs to one of the largest classes (tetracyclines) of antimicrobials used in the United States livestock industry (USEPA, 2013). Tetracycline compounds are ranked second in production and usage of antibiotics worldwide (Daghir and Droguí, 2013). We hypothesized that tetracycline would cause a shift in microbial community structure, leading to a greater relative abundance of fungi to bacteria, lower abundance of nosZ-carrying bacteria, lower N₂ production, and greater net production of N₂O. This is the first report simultaneously evaluating the impacts of antibiotic exposure on prokaryotic and fungal communities as well as the two clades of nosZ-carrying bacteria associated with soil N₂O emission.

**MATERIALS AND METHODS**

**Soil Collection**

Soils were collected from a managed grassland farm (60 ha) in Emmons County, ND, United States (46°24′22″; 100°23′16″), where there had been no history of antibiotic or other agro-chemical application (including fertilizer), based on 80 years of farm records. Since 1990, the farm has been enrolled in the Conservation Reserve Program (CRP) because the soils were classified as highly erodible and not suitable for crop production by the United States Department of Agriculture Farm Services Agency (USDA-FSA). The CRP allows only limited grassland harvest (every 3–5 years) and no grazing (Phillips et al., 2015). A total of 16 plots (1 m²) were established within a 0.2 ha area to collect soil core samples following spring thaw on March 19, 2014. Air temperature at the time of coring was 2°C.

Two sets of surface soil samples (2.5 dia. × 10 cm depth) were collected using a hand auger at random locations within each plot and stored at 4°C. One set was collected to determine background bulk density, gravimetric soil moisture, total organic carbon (TOC), total nitrogen (TN), pH, and soil texture (sand, silt, and clay). Porosity and percent water-filled pore space (%WFPS) at the time of soil collection were calculated based on bulk density, gravimetric soil moisture and using a particle size density of 2.65 g cm⁻³. Soil sampling and measurement protocols are detailed in Phillips et al. (2012). The second set of soil samples were collected for tetracycline exposure incubations (see below). Before coring each
plot, the auger was carefully cleaned and rinsed with alcohol. Three small diameter cores were collected per plot and mixed together to form one composited sample per plot. These were immediately shipped to the Virginia Institute of Marine Science (VIMS) for the antibiotic exposure work.

**Exposure of Soil to Tetracycline**

Each of the 16 soil samples was split into two groups and 15 g of soil were amended with either 7.5 ml of autoclaved DI water (control group) or tetracycline (tetracycline group) delivered in 7.5 ml of autoclaved DI water (final tetracycline concentration of 0.1 mg g⁻¹ soil). The tetracycline concentration used here is within the range of antibiotic concentrations known to repress microbial iron (III) reduction and denitrification (Thiele-Bruhn, 2005; Long et al., 2013). Each paired sample, control and tetracycline, was incubated in the dark at room temperature for 1 week. After incubation, samples were split into two groups, one for N₂ production rate measurements and one, which was stored at −80°C, for microbial community analysis and soil properties measurements.

**Soil Properties of Incubated Samples**

Soil pH was determined using a glass electrode with a soil:deionized water mixture of 1:1 (w/v). Percent organics was determined by loss of ignition (500°C, 4 h). TOC and TN were quantified using the Exeter CHN model 440 CE analyzer. Soil NO₃⁻ and NH₄⁺ were measured in a Lachat QC8000 FIA after extraction with 2 M KCl (2:1 KCl to sediment ratio).

**Activity Measurements: N₂O and N₂ Production Rates**

All 16 pairs of samples were used for soil slurry incubation experiments to measure potential rates of denitrification (moles of added ¹⁵N-labeled NO₃⁻ transformed to ³⁰N₂) as well as total N₂O production, following the method described by Long et al. (2013). One gram of each soil sample was pre-incubated anaerobically, after flushing with He gas, in 12-ml Exetainer tubes overnight. Two sets of tubes were prepared to measure N₂ and N₂O production from each sample. Pre-incubation served to deplete the resident soil NO₃⁻ and NO₂⁻ pools (NOx) prior to spiking with 200 nmol of potassium nitrate (K¹⁵NO₃, 99%). Before the addition of potassium nitrate, all tubes were reflushed with He for 5 min to remove background N₂, CO₂, and other atmospheric gases. Time course incubations (time points 0 and 1 h after ¹⁵NO₃ spike) were carried out in duplicate at room temperature. A 0.2 ml of potassium hydroxide (KOH) solution was added at each time point to stop microbial activity. The ³⁰N₂ gas in the headspace was measured on a continuous-flow isotope ratio mass spectrometer (Thermo Delta V Advantage, Thermo Scientific) in line with an automated gas bench interface (Thermo Finnigan GasBench II, Thermo Scientific). The N₂O gas in the headspace was measured using a gas chromatograph fitted with an electron capture detector (Shimadzu). The gas chromatograph was calibrated with commercial N₂O standards and the coefficient of variation for three to five replicate injections of low and high concentration standards was consistently <3%.

Potential rates of denitrification and N₂O production were calculated based on the amounts of ³⁰N₂ and N₂O, respectively, measured at T0 and T1 (1 h) after ¹⁵NO₃ addition following the method described in Long et al. (2013). The samples with high N₂ and N₂O measured at T0 (three for N₂ and six for N₂O) were excluded in the rate calculation due to incomplete killing of microbial activities, which may be resulted from insufficient amount of KOH application.

**Molecular Analysis: Bacterial and Fungal Abundance**

A subset of the soils stored at −80°C (four pairs: 4 control and 4 tetracycline samples) was selected for microbial community analysis according to the measured activity rates to include the full range of N₂ production inhibition (59 to 100% inhibition) by the tetracycline treatment. Genomic DNA was extracted from 0.5 g of soil using the PowerSoil DNA Isolation kit (MoBio). The DNA quality was assessed using a NanoDrop spectrometer (Thermo Scientific) and quantified with a Qubit™ fluorometer (Invitrogen) and the dsDNA high-sensitivity kit. The abundance of bacteria and fungi was quantified by quantitative polymerase chain reaction (qPCR) of 16S rRNA and internal transcribed spacer (ITS) genes, respectively, using the QuantStudio 6 Flex (Thermo Scientific). Standards were prepared through a serial dilution of plasmids carrying the target genes and quantified using an Agilent 220 TapeStation System (Agilent Technologies). The primers EU341F (5′-CCT ACG GGA GGC AGC AG-3′) and 685R (5′-ATC TAC GGA TTT CAC TCC TAC A-3′) were used to generate 344 bp amplicons of bacterial 16S rRNA genes. The fungal ITS region was amplified using the primers ITS1F and ITS2 (Buee et al., 2009), generating 300–400 bp fragments. The 20 µL qPCR reactions for 16S rRNA and ITS quantification consisted of 10 µL of SYBR green Go-Taq qPCR Master Mix (Promega), 0.05 µL of CRX dye, 1 µL of each primer (10 µM), 2 ng of template DNA, and were adjusted to final volume with nuclease-free H₂O. The qPCR conditions for 16S quantification were as follows: 10 min at 95°C, followed by 40 cycles of 15 s at 95°C, 30 s at 55°C, and 30 s at 72°C. Efficiency and R² values for the 16S qPCR reaction were 63% and 0.99, respectively. The detection limit was 2400 gene copies per sample. For ITS quantification, the qPCR conditions were as follows: 10 min at 95°C, followed by 35 cycles of 15 s at 95°C, 30 s at 50°C, and 1 min at 72°C. Efficiency and R² values for the ITS qPCR reaction were 55% and 0.97, respectively. The detection limit was 1170 gene copies per sample. All reactions were performed in 96 well plates with two negative controls, which contained no template DNA, to exclude any potential contamination. Reaction specificity was confirmed using gel electrophoresis in comparison with standards and monitored by analysis of dissociation curves. Gene copy number per PCR well was calculated from the standard curve according to the following equation:

\[
\text{copy number}_{\text{well}} = 10^{\frac{Ct - b}{a}}
\]

Where Ct corresponds to the threshold cycle of the sample, and a and b correspond to the y-intercept and slope of the logarithmic standard curve, respectively. Copy numbers per well were then
converted to copy number per gram of soil according to the following equation, assuming 100% DNA extraction efficiency:

\[
\text{gene copies g soil} = \left( \frac{\text{copy number}}{\text{ng DNA}} \right)_\text{well} \times \left( \frac{\text{ng DNA}}{\mu L} \right)_\text{sample} \times \left( \frac{\mu L \text{ DNA}}{g \text{ soil}} \right)_\text{extracted}
\]

**Molecular Analysis: Microbial Community Composition and Diversity**

Next-generation sequencing of prokaryotic 16S rRNA and fungal ITS genes was used to examine the composition and diversity of prokaryotic and fungal communities in soil and tetracycline samples. The communities were analyzed through barcode pyrosequencing using the Ion Torrent PGM sequencer. The variable V4 region of the 16S rRNA gene was amplified through PCR, using the forward primer 515F and a modified, barcoded reverse primer 806R (Caporaso et al., 2011). The ITS1 variable region of fungal ITS was amplified using a modified, barcoded forward primer ITS1F and the reverse primer ITS2 (White et al., 1990; Gardes and Bruns, 1993; Bellemain et al., 2010). The PCR mixture for 16S rRNA amplification contained 10 µL of Go-Taq mix, 1 µL of primers at 10 µM, 1 µL of template DNA (10–30 ng/µL), and nuclease-free H2O up to 25 µL. The PCR conditions for 16S rRNA amplification were as follows: 3 min at 95°C, followed by 25 cycles of 30 s at 95°C, 1 min at 55°C, and 1 min at 72°C, followed by 5 min at 72°C. The PCR mixture for ITS amplification contained 0.2 µL of Taq Polymerase (Invitrogen), 1 µL of primers at 10 µM, 1 µL of template DNA (10–30 ng/µL), 2.5 µL of buffer (Invitrogen), 0.75 µL of dNTPs mix (1 mM), 1.0 µL of MgCl2 (50 mM), 0.25 µL of bovine serum albumin (BSA) at 10 mg/mL, and nuclease-free H2O up to 25 µL. The PCR conditions for ITS amplification were as follows: 3 min at 95°C, followed by 25 cycles of 30 s at 95°C, 1 min at 55°C, and 1 min at 72°C, followed by 5 min at 72°C. The fragment size of the 16S rRNA (354 bp) and ITS (363 bp) amplicons and negative control amplification were checked by agarose gel electrophoresis. PCR products from each sample were pooled into a homogeneous mixture and a 2% agarose gel was run in duplicate to extract the amplicons, which were purified using an UltraClean GelSpin DNA Purification Kit (Promega). The concentration of purified amplicons was measured using a 2200 TapeStation instrument, following the manufacturer’s instruction. Pyrosequencing was conducted on the Ion Torrent PGM sequencer with barcode samples pooled on Ion 316 chips, following the Ion PGM Hi-Q Sequencing Kit protocol (Thermo Scientific).

**Bioinformatic Analysis: 16S rRNA and ITS sequences**

Bioinformatic analysis of the 16S rRNA sequences was performed using the mothur program (Schloss et al., 2009). Primer sequences were trimmed, and all sequences shorter than 200 bp and with a quality score lower than 25 were removed. Acacia was used to de-noise the trimmed sequences (Bragg et al., 2012). The remaining sequences were then processed using mothur (Schloss et al., 2009). Unique sequences were found after alignment with the Silva SEED database (Quast et al., 2013). Badly aligned sequences were removed, unique sequences were pre-clustered, and chimeras were removed using UCHIME (Edgar et al., 2011). Sequences were classified using the SILVA v119 taxonomy, and unknown taxa were removed (Quast et al., 2013). Operational taxonomic units (OTUs) were clustered at 97% identity using the opticlust algorithm. Bacterial and archaeal OTUs were extracted for separate analysis of each community’s richness and diversity; samples were subsampled to the lowest number of sequences to normalize the diversity estimates. Chao and Ace indexes were calculated to estimate species richness, and Shannon was calculated to estimate α-diversity and community evenness. β-Diversity among samples was estimated using the Bray–Curtis dissimilarity calculator.

Fungal ITS sequence analysis was carried out using mothur and UPARSE (Schloss et al., 2009; Edgar, 2013). After initial processing of the FASTQ files using Acacia as described above, sequences were clustered into OTUs at 97% identity, using the UPARSE pipeline (Edgar, 2013). The samples were then subsampled to the lowest number of sequences and analyzed for species richness, diversity, and evenness, as described above for the prokaryotic community. Parallel to the OTU analysis, unique ITS sequences were classified using the UNITE v6_sh_97 ITS database in mothur (Abarenkov et al., 2010).

**Bioinformatic Analysis: Inference of NosZ-Carrying Bacteria Community Composition**

Bacterial taxa carrying nosZ were identified based on a denitrification gene inference analysis on the rarefied bacterial 16S rRNA sequences (22,388 sequences per sample) using PAthway PRediction by phylogenetic plAcement (PAPRICA) (Bowman and Ducklow, 2015; Arfen et al., 2017). A customized PAPRICA database was constructed with 8,513 complete and 785 draft bacterial genomes. The nosZ genes in the reference genomes were identified based on the KEGG database and used for gene prediction as described in Arfen et al. (2017). The estimated abundances of nosZ-carrying taxa were normalized to the number of 16S rRNA gene copies predicted for each taxon. Based on the taxonomy, the taxa carrying nosZ-I or nosZ-II were identified. Inferred abundances of nosZ clades per gram of soil were also calculated by multiplying relative abundances obtained through PAPRICA with the qPCR 16S copy numbers.

**Molecular Analysis: NosZ-I and NosZ-II Abundance**

Quantitative PCR of nosZ-I and nosZ-II genes was also performed to measure the abundance of microorganisms responsible for the reduction of N2O to N2, using the QuantStudio 6 Flex (Thermo Scientific). Standards were prepared through a serial dilution of plasmids carrying the target genes and quantified using an Agilent 220 TapeStation System (Agilent Technologies). The primers used for nosZ-I genes were nosZ1F and nosZ1R and generated 300 bp amplicons (Henry et al., 2006).
The nosZ-II genes were amplified using nosZIIF and nosZIIR primers that generated 690–720 bp amplicons (Jones et al., 2013). The 20 µL qPCR reactions for nosZ-I and nosZ-II quantification consisted of 10 µL of SYBR green Go-Tag qPCR Master Mix (Promega), 0.05 µL of CRX dye, 1 µL (nosZ-I) or 4 µL (nosZ-II) of each primer (10 µM), 2 ng of template DNA, and were adjusted to final volume with nuclease-free H₂O. The thermal cycling conditions were the following: 10 min at 95°C, followed by 50 (nosZ-I) or 55 (nosZ-II) cycles of 15 s at 95°C, 45 s at 55°C (nosZ-I) or 30 s at 54°C (nosZ-II), 30 s at 72°C, and 35 s at 80°C for fluorescence detection. Amplification efficiencies were 46 and 40% for the nosZ-I and nosZ-II genes, respectively. The detection limit was 4,600 and 1,240 gene copies per sample for the nosZ-I and nosZ-II genes, respectively. All reactions were performed in 96 well plates with two negative controls, which contained no template DNA, to exclude any potential contamination. Reaction specificity was confirmed using gel electrophoresis in comparison with standards and monitored by analysis of dissociation curves during quantitative amplification. The gene copy numbers per gram of soil were calculated as described above for 16S rRNA gene and ITS.

Statistical Analysis

Normality of all variables was assessed with Q–Q plots. Variables with large departures from normality were analyzed with non-parametric tests. A paired two-sample Mann–Whitney–Wilcoxon test (non-parametric) was used to identify significant differences between activity rates of control and tetracycline samples. Paired two-sample t-tests (parametric) were used to test for significant differences in 16S rRNA gene and ITS abundances, taxa relative abundances in prokaryotic and fungal communities, diversity estimators, and nosZ-I and nosZ-II gene abundances between control and tetracycline samples. Simple linear regressions were used to assess the relationship between nosZ abundances inferred by PAPRICA and determined by qPCR. Significant relationships for all tests were considered at α < 0.05. These statistical analyses were conducted in R (version 3.2.2. Copyright 2015 The R Foundation for Statistical Computing). Significant differences between control and tetracycline treatments in the abundance of nosZ-carrying taxa from PAPRICA analysis were tested by fitting the data into a generalized linear model (GLM) based on the negative binomial distribution using the DESeq function of the DESeq2 package in R (Love et al., 2014). A principal coordinate analysis (PCoA) was also performed to evaluate the β-diversity of bacterial, archaeal, and fungal communities using the phyloseq package in R (McMurdie and Holmes, 2013). Significant effects of the treatment in OTU dissimilarity among samples were tested by permutational multivariate ANOVA (PERMANOVA) using the adonis function of the vegan package in R (Oksanen et al., 2017).

RESULTS AND DISCUSSION

Soil Properties

Average (standard deviation) bulk density, sand, silt, and clay contents for the 0–10 cm soil depth increment were 1.19 (0.04) g cm⁻³, and 730 (13), 230 (38) and 40 (30) g kg⁻¹, respectively. At the time of soil collection, soil porosity was 55 (1.7) % and %WFPS was 38 (4). TOC and TN were 1.7 (0.2) and 0.2 (0.02)% dry weight (dw), respectively, and soil pH was 6.2 (0.1). Physicochemical factors potentially affecting microbial communities and denitrification activities were examined after the 1-week incubation period (Table 1). The only factor that was significantly different between the two groups was pH (paired t-test, p < 0.05). The tetracycline group (average pH = 6.9) had a significantly lower pH than the control group (average pH = 7.4). Despite this difference, the pH for both groups remained within the neutral region, where the 0.5 difference is not likely to significantly affect the denitrification end-products (Rochester, 2003; McMillan et al., 2016).

Effects of Tetracycline on Denitrification: N₂O and N₂ Production

Rates of potential N₂O and N₂ production were significantly affected by tetracycline treatment, as shown in Figure 1 (paired Mann–Whitney-Wilcoxon test, p < 0.05). N₂O production in the tetracycline group ranged from 2.07 to 13.97 nmol N g⁻¹, as compared to 0.32–1.37 nmol N₂O-N g⁻¹ h⁻¹ in the control group. N₂ production in the treated soil ranged from 0 to 24.2 nmol ³⁰N₂-N g⁻¹ h⁻¹, as compared to 1.52–108 nmol ³⁰N₂-N g⁻¹ h⁻¹ measured in the controls. Average N₂O production was 12 times higher in the tetracycline group, as compared to 1.52–108 nmol ³⁰N₂-N g⁻¹ h⁻¹ measured in the controls. Average N₂O production was 12 times higher in the tetracycline group.
while $N_2$ production was inhibited by up to 84%. The $N_2O$-N/$N_2$-N ratio was 42 times higher in the tetracycline group than the controls. These results suggest that microbial reduction of $N_2O$ to $N_2$ is strongly inhibited and $N_2O$ production is enhanced in soils exposed to tetracycline.

The stimulatory effect of antibiotics on $N_2O$ production has recently been observed in agricultural soils and estuarine sediments exposed to other antibiotics, such as sulfamethazine and narasin, for similar time periods and in lower concentrations than this study (DeVries et al., 2015; Hou et al., 2015; Yin et al., 2017). Observed increases in $N_2O$ production have been attributed to a shift in the end-products of denitrification from $N_2$ to $N_2O$ driven by the stronger inhibition of $N_2O$-reducing bacteria than $N_2O$-producing bacteria (DeVries et al., 2015; Hou et al., 2015; Yin et al., 2017). However, different antibiotics, such as norfloxacin or a mixture of sulfadiazine,
sulfamethoxazole, florfenicol, and chloramphenicol, were shown to reduce N₂O production across various time periods and low exposure concentrations (Sun et al., 2017; Yin et al., 2017). The antibiotic’s effect on net N₂O production seems to be dependent on the antibiotic type, dosage, and time of exposure, but no clear pattern has emerged in the literature. When considering tetracycline, we found only one study that measured how chronic exposure to tetracycline affected the production of different intermediates of denitrification in riverine sediments (Roose-Amsaleg et al., 2013). Roose-Amsaleg et al. (2013) reported no significant effect of tetracycline on NO₃⁻ reduction, or NO₂⁻ and N₂O production at variable tetracycline concentrations up to 10 mg L⁻¹. In our study, acute exposure to tetracycline in soil increased N₂O and inhibited N₂ production, indicating that the final denitrification step from N₂O to N₂ was susceptible to antibiotic exposure.

**Effects of Tetracycline on Bacterial and Fungal Abundance**

The abundance of bacterial 16S rRNA genes and fungal ITS is shown in Figure 2. The abundance levels observed in the control group ranged from 5.11 × 10¹⁰ to 5.79 × 10¹⁰ 16S rRNA gene copies per gram soil and from 1.27 × 10⁸ to 3.25 × 10⁸ ITS copies per gram soil. The tetracycline treatment had no significant effect (paired t-test, p > 0.05) on the abundance of either bacteria or fungi. However, the lowest levels of bacterial abundance and the highest levels of fungal abundance were observed in samples exposed to tetracycline, which resulted in a significant increase (paired t-test, p < 0.05) in the fungi:bacteria ratio in samples exposed to tetracycline compared to control (Figure 2).

The small subset of samples used for the molecular analysis (see section “Materials and Methods”) may have limited our capacity to detect smaller changes in bacterial or fungal absolute abundances. However, the subset used was enough to detect significant changes in the fungi:bacteria ratio, as hypothesized. Previous studies showed significant increases in the fungi:bacteria ratio in soils exposed to antibiotics, including tetracyclines (Thiele-Bruhn and Beck, 2005; Hammesfahr et al., 2008; Demoling et al., 2009; Ding and He, 2010; Gutiérrez et al., 2010). This is expected since tetracycline inhibits bacterial growth and may leave nutrients and habitat for fungal growth. The increase in the fungi:bacteria ratio suggests that the higher N₂O production observed could be associated with...
fungal denitrification (Shoun et al., 2012; Maeda et al., 2015; Mothapo et al., 2015). Various studies have shown that fungal denitrification is a major contributor to N2O production in different soils (Laughlin and Stevens, 2002; Long et al., 2013; Wei et al., 2014; Chen et al., 2015; Huang et al., 2017).

Effects of Tetracycline on Prokaryotic Community Composition

A total of 201,980 16S rRNA gene sequences were obtained for taxonomic analysis following screening and filtering of the prokaryotic sequences. Bacteria was clearly the most abundant domain with relative abundances ranging from 89 to 96%, while archaeal sequences represented less than 12% of the communities. The tetracycline treatment shifted the relative abundance away from Bacteria (paired t-test, $p = 0.00426$) and toward Archaea (paired t-test, $p = 0.00440$) by 4%. This was expected since tetracycline is an antibacterial compound that would mostly inhibit bacterial growth. The relative abundances of the most represented prokaryotic phyla (16S rRNA > 1% of total reads) in control and tetracycline groups are shown in Figure 3. Eleven abundant phyla were identified. All samples except one were dominated by Acidobacteria, representing 19–30% of the sequences, followed by Proteobacteria (17–26%), Firmicutes (0.5–17%), Actinobacteria (8.3–16%), and Thaumarchaeota (3.7–11%). The prokaryotic phyla that were significantly different between control and tetracycline groups were Actinobacteria (paired t-test, $p = 0.00203$), Chloroflexi (paired t-test, $p = 0.00287$), Firmicutes (paired t-test, $p = 0.00210$), and Thaumarchaeota (paired t-test, $p = 0.00430$). The most drastic effect was the decline in the relative abundance of Firmicutes, from an average of 14% in controls to 0.9% in the tetracycline samples. The Firmicutes classes most affected by the tetracycline treatment were Bacilli and Clostridia (Supplementary Figure S1). These results show that specific bacterial taxa are more susceptible to tetracycline exposure despite its broad-spectrum antibacterial properties.

Firmicutes is likely to be less resistant to tetracycline according to previous studies, which report a decrease in the ratio of Gram-positive:Gram-negative bacteria when exposed to tetracycline and other antimicrobials (Hund-Rinke et al., 2004; Ding and He, 2010; Gutiérrez et al., 2010). Indeed, the vast majority of tetracycline-resistant bacteria are Gram-negative (Schnabel and Jones, 1999). The observed decrease in the relative abundance of Firmicutes may have important consequences for N2O reduction since bacteria harboring nosZ genes without possessing any nir
or nor gene are mainly found amongst the Bacteroidetes and Firmicutes phyla (Graf et al., 2014).

Effects of Tetracycline on Fungal Community Composition

A total of 369,805 (ITS) sequences were obtained for taxonomic analysis following screening and filtering of the fungal sequences. The relative abundances of the most represented fungal phyla (ITS > 1% of total reads) in control and tetracycline groups are shown in Figure 4. Five abundant phyla were identified. All samples were dominated by Zygomycota, representing 47–80% of the sequences, followed by Ascomycota (18–49%).

The tetracycline treatment did not have significant effects (paired t-test, p > 0.05) on the relative abundance of any fungal phyla, including the N₂O producing Ascomycota, Basidiomycota, and Zygomycota (Maeda et al., 2015; Mothapo et al., 2015). Most fungal denitrifiers belong to the Sordariomycetes, Eurotiomycetes, and Saccharomycetes classes (Mothapo et al., 2015). No significant differences between treatments were detected in the relative abundance of any of these classes (Supplementary Figure S2). Although the present study shows a tetracycline impact on N₂O production and the fungi:bacteria abundance ratio, no individual impacts appear to occur in fungal denitrifier taxa.

Effects of Tetracycline in the Community Richness and Diversity

Principle coordinate analysis represents β-diversity of microbial communities based on the calculated dissimilarities among samples (Figure 5). The first two principal coordinates represented 58.6% and 80.3% of the variation in bacterial and archaeal communities (Figures 5A,B), respectively. The archaeal β-diversity was not significantly affected by the tetracycline treatment (PERMANOVA, p > 0.05), while the bacterial communities had a distinct clustering of control samples separated from the tetracycline samples. This result shows a significant difference (PERMANOVA, p < 0.05) between the compositions of bacterial communities in control and tetracycline groups. Previous studies also reported significant effects of different antibiotics (sulfadiazine, tylosin, amoxicillin, and ciprofloxacin) on bacterial community structure in soils and marine sediments (Westergaard et al., 2001; Zielezny et al., 2006; Binh et al., 2007; Hammesfahr et al., 2008; Näslund et al., 2008). When considering tetracyclines, Zielezny et al. (2006) did not find significant effects in the bacterial community structure after exposure to 1–50 mg kg⁻¹ of chlortetracycline for 48 h. This result contrasts with our study, where the effects of tetracycline on bacterial community structure were significant, probably due to the higher dosage and longer duration of exposure tested (1 week). Since tetracyclines are protein synthesis inhibitors with bacteriostatic action (i.e., limits bacterial growth) rather than bactericidal it is possible that an incubation period longer than 48 h is necessary to detect changes in community structure.

Contrasting to the bacterial communities, no clear patterns were observed in fungal β-diversity (Figure 5C). The first...
The effects of tetracycline on species richness and microbial \( \alpha \)-diversity are displayed in Table 2. Interestingly, bacterial \( \alpha \)-diversity, estimated through calculation of the Shannon Index \( H' \), was significantly higher (paired \( t \)-test, \( p < 0.05 \)) in the tetracycline group (average \( H' = 6.36 \)) when compared to controls (average \( H' = 5.98 \)). An increase in bacterial diversity after antibiotic exposure was not expected since previous studies have shown that antibiotic exposure decreases bacterial diversity in the soil ecosystem (Westergaard et al., 2001; Zielezny et al., 2006; Cycoñ et al., 2016; Ding and He, 2010). However, temporary increases (after 4 days of exposure) in bacterial diversity of agricultural soils exposed to antibiotics have also been reported (Hammesfahr et al., 2008). The increase in bacterial diversity was probably caused by the inhibition of the most dominant taxa and a slight increase in the numbers of rare taxa. This explanation is supported by the significant increase in community evenness (paired \( t \)-test, \( p < 0.05 \)), estimated through calculation of the Shannon index-based evenness \( E_H \), and the non-significant difference observed in both indexes of species richness (Chao and Ace; paired \( t \)-test, \( p > 0.05 \)). We found that higher community diversity in the tetracycline group was associated with higher \( N_2O \) emission, while previous studies showed that these two variables are negatively correlated (Wagg et al., 2014; Samad et al., 2016). These studies, however, tested a broader range of bacterial diversity and statistically tested the direct relationships between microbial diversity and \( N_2O \) emissions. In our study, the observed higher bacterial diversity in the tetracycline group with higher \( N_2O \) production could be sporadic and the relationship between microbial diversity and \( N_2O \) emissions is outside of the scope of this work.

Similarly to the \( \beta \)-diversity patterns, no significant changes (paired \( t \)-test, \( p > 0.05 \)) were detected in fungal species richness, \( \alpha \)-diversity, or evenness with tetracycline exposure (Table 2). As expected, tetracycline had no impact on the composition of the fungal communities at any taxonomical level.

**Effects of Tetracycline on NosZ-Carrying Community Composition**

By employing metabolic inference using PAPRICA (Bowman and Ducklow, 2015), we were able to identify the 16S rRNA
FIGURE 6 | Difference in the nosZ-carrying microbial community composition between tetracycline and control samples for nosZ-I (A) and nosZ-II (B) bacteria. Each dot denotes a taxon that is either higher (log₂ fold change > 0) or lower (log₂ fold change < 0) in the tetracycline treated samples when compared to the control samples. Taxa are organized by family in decreasing order and colored by genus. Normalized abundances were obtained by metabolic inference of 16S rRNA gene sequences using PAPRICA and tested for significant effects (p < 0.05) of treatment by fitting the data into a generalized linear model (GLM) based on the negative binomial distribution. Taxa with significant difference between tetracycline and control samples are marked with * in the legend.
sequences belonging to the bacterial taxa carrying either nosZ-I or nosZ-II genes. Within the nosZ-I community, Gammaproteobacteria was the most abundant class, followed by Alphaproteobacteria (Proteobacteria) (Supplementary Figure S3). A few sequences belonging to the Caldilineae class (Chloroflexi) and Betaproteobacteria (Proteobacteria) were also observed in this study. The nosZ-II communities were more taxonomically diverse at the phylum and class levels (Supplementary Figure S4). Sixteen classes of nosZ-II-carrying taxa were identified and virtually all samples were dominated by Betaproteobacteria (Bacteroidetes), Deltaproteobacteria (Proteobacteria), and Opitutae (Verrucomicrobia).

The effects of tetracycline on nosZ-carrying taxa were evaluated based on the changes in families and genera of bacteria.
carrying \textit{nosZ-I} or \textit{nosZ-II} genes (Figure 6). The tetracycline group had a significantly higher abundance of the \textit{nosZ-I} carrying \textit{Rhodanobacter} spp. (Gammaproteobacteria) than the control group (negative binomial GLM, \(p = 0.0038\)) (Figure 6A). On the other hand, the tetracycline treatment significantly decreased the abundance of two \textit{nosZ-II} carrying taxa, \textit{Bacillus} spp. (Firmicutes; negative binomial GLM, \(p = 0.0008\)) and \textit{Anaeromyxobacter} spp. (Deltaproteobacteria; negative binomial GLM, \(p = 0.0375\)) (Figure 6B). Abundances of both taxa were approximately eight times lower in the tetracycline group.

The observed decrease in \textit{nosZ-II} carrying \textit{Bacillus} spp. is consistent with the decrease in relative abundance of Firmicutes and provides evidence that some of the affected Firmicutes bacteria were potential \textit{nosZ-II} carriers. The microbes harboring \textit{nosZ} genes without possessing any \textit{nir} or \textit{nor} gene are frequently found amongst the Firmicutes phylum (Graf et al., 2014). This underscores the importance of some \textit{nosZ} carrying Firmicutes as \(\text{N}_2\text{O}\) consumers, rather than \(\text{N}_2\text{O}\) producers, in the soil ecosystem. The other genus that had lower abundances in the tetracycline group, \textit{Anaeromyxobacter} spp. is also known as a non-denitrifying \(\text{N}_2\text{O}\) reducer, i.e., these bacteria do not possess all other denitrification genes besides \textit{nosZ} (Sanford et al., 2012; Hallin et al., 2017). This finding suggests that non-denitrifying \(\text{N}_2\text{O}\) reducers are more susceptible to antibiotic disturbance than denitrifying \(\text{N}_2\text{O}\) reducers. Both \textit{nosZ-II} taxa significantly affected by the tetracycline exposure are likely important \(\text{N}_2\text{O}\) consumers rather than \(\text{N}_2\text{O}\) producers. The observed decreases in abundances of \textit{Bacillus} (Firmicutes) and \textit{Anaeromyxobacter} (Deltaproteobacteria) may thus explain the observed increase in \(\text{N}_2\text{O}/\text{N}_2\) ratio in the same treatment. These decreases also indicate that, despite tetracycline being a broad-spectrum antibiotic, there are functional bacterial taxa more susceptible than others.

### Effects of Tetracycline on Bacterial \textit{nosZ} Abundance

The abundances of \textit{nosZ-I} and \textit{nosZ-II} genes measured by qPCR are shown in Figure 7. Abundances of \textit{nosZ-I} and \textit{nosZ-II} genes in the control group ranged from 1.04 to 2.13 \times 10^9 copies per gram soil and from 0.45 to 1.03 \times 10^9 copies per gram soil, respectively. The \textit{nosZ-I} abundance was not significantly affected by tetracycline (paired \(t\)-test, \(p > 0.05\)), while the \textit{nosZ-II} abundance was significantly lower (paired \(t\)-test, \(p < 0.05\)) in the tetracycline group (Figure 7).

Together, \textit{nosZ-I} and \textit{nosZ-II} comprised 2.9–5.5 % of the total number of 16S rRNA gene copies, which are in accordance with reports in the literature for \textit{nosZ} relative abundances (Jones et al., 2013, 2014; Domeignoz-Horta et al., 2015). Taking the effect of tetracycline into account, the \textit{nosZ-II} community appears to be more sensitive to the antibiotic than the \textit{nosZ-I} community. This result supports the observed taxonomical changes described above for the \textit{nosZ} carrying taxa. The \textit{nosZ-II} community was previously reported to be more sensitive than \textit{nosZ-I} to other environmental factors, such as pH, calcium concentration, soil moisture, total N, and crop rotation systems (Jones et al., 2014; Domeignoz-Horta et al., 2015; Samad et al., 2016). The significant decrease of \textit{nosZ-II} abundance reported here for samples exposed to tetracycline corresponds to the observed increase in \(\text{N}_2\text{O}/\text{N}_2\) ratio in the same treatment, which underscores the importance of this clade as \(\text{N}_2\text{O}\) consumers.

Abundance of \textit{nosZ-I} and \textit{nosZ-II} genes measured by qPCR were compared with the inferred abundance of both genes by PAPRICA (Figure 8). A significant, positive linear correlation was observed between the two estimates of the \textit{nosZ} clades. The inferred abundances of \textit{nosZ-I} were on average 17% lower than those determined by qPCR, while inferred \textit{nosZ-II} abundances were on average two times higher than qPCR results.

A similar significant relationship between inferred and qPCR abundances was previously reported for the \textit{nosZ-I} gene in the oyster microbiome (Arñken et al., 2017). These findings suggest that 16S rRNA gene-based metabolic inference is a promising approach to evaluate functional gene abundances and community composition in environmental samples. Despite the small sampling size of our study and the intrinsic limitations of a reference database, this approach could be of great relevance to the microbial ecology field as 16S rRNA gene high throughput sequencing becomes increasingly available.

### CONCLUSION

We present several lines of evidence that unveil the microbial community changes associated with the antibiotic impacts on soil denitrification. The increase in \(\text{N}_2\text{O}/\text{N}_2\) ratio from denitrification is paralleled by a greater fungi:bacteria ratio and lower abundance of \textit{nosZ-II} carrying bacteria. Non-denitrifying \(\text{N}_2\text{O}\) reducers belonging to the Firmicutes and Deltaproteobacteria phyla appear to be particularly susceptible to an acute exposure to tetracycline and may be crucial for soil \(\text{N}_2\text{O}\) sink capacity. Future research aiming to predict \(\text{N}_2\text{O}\) emissions based on microbial community structure under antibiotic-disturbed environments would potentially benefit from targeted approaches to these taxa.

Overall, the findings of this study emphasize the importance of microbial community dynamics to fundamental ecosystem processes that can have major impacts on the emissions of a potent greenhouse gas such as \(\text{N}_2\text{O}\).

### AUTHOR CONTRIBUTIONS

BS and RP conceived and designed the experiments. MS, BS, and TS performed the experiments and acquired the data. MS analyzed the data. MS, BS, and RP wrote the paper. All authors approved the final submitted manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2018.02121/full#supplementary-material
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