


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Journal Article**Author(s):**

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Publication date:

2016-07

Permanent link:

<https://doi.org/10.3929/ethz-b-000114942>

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Originally published in:

Oecologia 181(3), <https://doi.org/10.1007/s00442-016-3611-8>

Plant species diversity affects soil–atmosphere fluxes of methane and nitrous oxide

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Received: 20 November 2015 / Accepted: 8 March 2016 / Published online: 2 April 2016
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Abstract Plant diversity effects on ecosystem functioning can potentially interact with global climate by altering fluxes of the radiatively active trace gases nitrous oxide (N₂O) and methane (CH₄). We studied the effects of grassland species richness (1–16) in combination with application of fertilizer (nitrogen:phosphorus:potassium = 100:43.6:83 kg ha⁻¹ a⁻¹) on N₂O and CH₄ fluxes in a long-term field experiment. Soil N₂O emissions, measured over 2 years using static chambers, decreased with species richness unless fertilizer was added. N₂O emissions increased with fertilization and the fraction of legumes in plant communities. Soil CH₄ uptake, a process driven by methanotrophic bacteria, decreased with plant species numbers,

irrespective of fertilization. Using structural equation models, we related trace gas fluxes to soil moisture, soil inorganic N concentrations, nitrifying and denitrifying enzyme activity, and the abundance of ammonia oxidizers, nitrite oxidizers, and denitrifiers (quantified by real-time PCR of gene fragments amplified from microbial DNA in soil). These analyses indicated that plant species richness increased soil moisture, which in turn increased N cycling-related activities. Enhanced N cycling increased N₂O emission and soil CH₄ uptake, with the latter possibly caused by removal of inhibitory ammonium by nitrification. The moisture-related indirect effects were surpassed by direct, moisture-independent effects opposite in direction. Microbial gene abundances responded positively to fertilizer but not to plant species richness. The response patterns we found were statistically robust and highlight the potential of plant biodiversity to interact with climatic change through mechanisms unrelated to carbon storage and associated carbon dioxide removal.

Communicated by Jennifer Funk.

Electronic supplementary material The online version of this article (doi:10.1007/s00442-016-3611-8) contains supplementary material, which is available to authorized users.

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Keywords Functional genes · Jena experiment · Microbial activities · Nitrification and denitrification · Structural equation modeling

Introduction

Recent syntheses of experimental research have shown that effects of plant diversity loss on ecosystem functioning can be of similar magnitude as those of other global changes, including warming and drought (Hooper et al. 2012). However, the effects of plant diversity interact with other global changes, with mechanisms operating at spatial scales ranging from the plot scale to the globe. For example, drivers of global change may affect ecosystem functions directly

and indirectly via alterations of plant diversity (Balvanera et al. 2006). Changes in plant diversity can alter ecosystem carbon and nutrient cycling, thereby changing the ecosystem–atmosphere exchange rates of greenhouse gases and thus affecting climatic drivers at the global scale.

Carbon dioxide (CO₂) is the dominant anthropogenic greenhouse gas, but nitrous oxide (N₂O) and methane (CH₄) also are important drivers of anthropogenic climate change. Despite their low atmospheric volume-mixing ratios, these gases account for over 25 % of the radiative forcing increase since pre-industrial times because of their high warming potential relative to CO₂ (IPCC 2007). To date, most experimental plant diversity studies have focused on ecosystem functions related to plant primary productivity and the ecosystem's carbon cycle (e.g., biomass, photosynthesis, leaf area, litter decomposition), which are processes directly relevant to the land–atmosphere exchange of CO₂. However, the effects of plant diversity on land–atmosphere fluxes of N₂O and CH₄ have only been studied in few mesocosm studies (N₂O: Abalos et al. 2014; Chang et al. 2014; Sun et al. 2013; N₂O and CH₄: Niklaus et al. 2006), and relatively little is yet known on responses of the underlying nitrogen (N) cycling processes to plant diversity (Le Roux et al. 2013).

N₂O is released from soils as an intermediate or by-product of microbial N cycling (see Cabello et al. 2009 for a comprehensive overview of soil N transformations). Under oxic conditions, some N₂O produced as a by-product of nitrification will escape to the atmosphere. Under more anoxic conditions, nitrate (NO₃[−]) is sequentially reduced to N₂O, which generally is lost from soils if it is not further reduced to molecular nitrogen (Firestone and Davidson 1989). Soil N₂O emissions are controlled by many factors (Robertson and Tiedje 1987) including the availability of inorganic N, in particular ammonium (NH₄⁺) and NO₃[−]. Soil moisture is also an important factor in the release of N₂O from the soil because it affects gas diffusivity and oxygen supply. At a low water-filled pore space, oxygen is readily available and soil organic matter is mineralized, yielding NH₄⁺ and promoting nitrification and associated N₂O emissions. At a high water-filled pore space, nitrification ceases and NO₃[−] is consumed by denitrification, which is the dominant source of N₂O emissions from soils. At an even higher water-filled pore space, N₂O emissions drop because most of the N₂O produced is reduced to molecular N.

Soils can be both sources and sinks of atmospheric CH₄. On an annual basis, most aerobic soils are a net sink for atmospheric CH₄, although periodic emissions can occur when soils are wet. This soil CH₄ sink is essentially driven by the consumption of CH₄ by methanotrophic bacteria, which use CH₄ as a source of carbon and energy (Dunfield 2007). At a very low redox potential (typically found in

wetlands), however, methanogenic archaea produce CH₄ through several pathways which involve the consumption of organic substrates or CO₂ (Conrad 1996). Non-microbial soil CH₄ emissions have recently been described (Wang et al. 2013), but the quantitative importance of this process in natural ecosystems remains to be evaluated. Methanogens and methanotrophs often co-occur in soils, with methanotrophs acting as a biofilter that oxidizes the CH₄ produced by methanogens in anoxic soil domains. On the other hand, occasional methanogenesis can prime methanotrophic activity, leading to increased uptake of atmospheric CH₄ (West and Schmidt 2002). Several links have been established between CH₄ oxidation and N transformations, with complex positive and negative correlations demonstrated between the two (Bodelier and Laanbroek 2004). Ammonia (NH₃) has been shown to inhibit CH₄ oxidation, at least in laboratory cultures (Hanson and Hanson 1996), but on the other hand inorganic N is an essential nutrient for most methanotrophs. In addition, differential stratification of these processes may occur in the soil, with effects that may be restricted to some soil layers and therefore not necessarily generate effects on soil–atmosphere CH₄ fluxes (Stiehl-Braun et al. 2011a).

While the fundamental processes involved in soil N₂O and CH₄ transformations are reasonably well understood, the ecology of the organisms involved in complex field ecosystems is far from being as clear (see Bodelier and Laanbroek 2004; Robertson 1989). Predicting N₂O and CH₄ fluxes is difficult given the large number of factors and organisms involved, which interact in ways difficult to predict and change in space and time. Nevertheless, we argue that plant diversity has the potential to alter trace gas fluxes via a range of mechanisms that can reasonably be predicted and tested. First, a more species-rich plant community often leads to tighter N cycling, with a relatively lower accumulation of inorganic N (Ewel et al. 1991; Niklaus et al. 2001; Tilman et al. 1996), which in turn could affect nitrification and denitrification and their associated N₂O emissions and, possibly, also CH₄ dynamics. Second, increased productivity at higher plant diversity may also translate into an improved supply of organic substrates for soil microbial communities, which could increase heterotrophic activity, lead to higher oxygen consumption, and increase the amount of soil volume with a redox potential sufficiently low for denitrification (Sexstone et al. 1985). Third, water use may increase with plant diversity, resulting in reduced soil moisture and increased soil diffusivity (Caldeira et al. 2001; Leimer et al. 2014a; Spehn et al. 2000), which may affect both N transformations and soil CH₄ uptake (Ball et al. 1997). Fourth, altered root densities and root architectures may lead to changes in soil aeration, spatio-temporal organic C deposition, or N use patterns. Fifth, atmospheric nitrogen (N₂) fixation by legumes will have particularly

strong impacts on N cycling and has been identified as strong determinant of ecosystem responses in many plant diversity experiments (Oelmann et al. 2007; Spehn et al. 2002). With respect to CH₄, the study by Niklaus et al. (2006) hints at the complexity of interactions at play, with plant diversity effects on CH₄ consumption changing from positive to negative when a soil disturbance treatment was applied in the field.

Here, we present the first study investigating the effects of plant species diversity and fertilization, together with their interaction, on fluxes of N₂O and CH₄ under field conditions. We investigated the dynamics of these trace gases and the underlying mechanisms at several levels. First, we measured in situ flux rates of N₂O and CH₄; second, we assessed the enzymatic potential of key N transformations functionally linked to these fluxes; third, we quantified the abundance of bacterial nitrifiers and denitrifiers using quantitative PCR of selected functional and ribosomal genes. We further assessed soil environmental conditions (temperature, moisture) and concentrations of inorganic N species. We were particularly interested in determining whether higher plant species richness would lead to more complete N capture and therefore reduced nitrification and denitrification rates and associated N₂O emissions. An additional aim was to test whether the plant diversity effects in our experimental system operated via changes in soil moisture and whether there was evidence that nitrification promotes soil CH₄ uptake by the removal of potentially inhibiting NH₄⁺.

Materials and methods

Study site and experimental design

We studied soil–atmosphere trace gas fluxes in a large grassland biodiversity experiment near Jena, Germany (50°55′N, 11°35′W; 130 m a.s.l.). In 2002, experimental grassland communities varying in diversity were established on an Eutric Fluvisol with a texture progressing from sandy loam to silty clay with increasing distance from the

adjacent Saale river. In the present study we focused on 78 plots (20 × 20 m) sown with one, two, four, eight, or 16 herbaceous species (see Roscher et al. 2004a for a complete description of the experimental design). Plot community composition was determined by random selection of species from a 60-species pool, with the constraint that species richness and plant functional type richness were as orthogonal as possible. Plant species had previously been assigned to functional types (grasses, small non-legume herbs, tall non-legume herbs, and legumes) based on a cluster analysis combining a large number of morphological and functional traits (Roscher et al. 2004a). Starting in 2005, factorial management treatments were established (Weigelt et al. 2009) in subplots (1.6 × 4 m) located along one side of the large plots. In the present study, we analyzed data from the main plot (“control” subplots) and the subplot that was managed identically except that fertilizer pellets were added after each mowing (“fertilized” subplots), at a rate of 100 kg N ha⁻¹ a⁻¹, 43.6 kg phosphorus (P) ha⁻¹ year⁻¹ and 83 kg potassium (K) ha⁻¹ year⁻¹. In total, the study thus presents a split plot design with 70 plots (unit of replication for plant species composition and thus plant species richness) and 2 × 70 = 140 subplots (unit of replication for fertilizer application).

Soil–atmosphere trace gas fluxes

We measured soil–atmosphere fluxes of N₂O and CH₄ on 20 July and 17 October 2007 and on 11 March, 29 April, and twice on 6 June 2008 (one daytime and one nighttime measurement; Table 1), using static chambers that had been installed in all control and fertilized subplots that were part of this study. The static chambers were 32 cm in diameter and lowered 14 cm into the ground, leaving 11 cm extending above ground. During flux measurements, air-tight lids were fitted over the static chambers, and headspace samples were collected 5, 20 and 35 min after chamber closure. These samples were injected into pre-evacuated exetainers and analyzed for N₂O and CH₄ concentrations in the laboratory (Agilent 6890 gas chromatograph equipped with a flame ionization and an electron capture detector;

Table 1 Environmental and meteorological conditions during the nitrous oxide and methane flux measurements

Date	Soil temperature (°C)	Soil moisture (m ³ H ₂ O m ⁻³)	Precipitation	
			Past 24 h (mm)	Past week (mm)
20 June 2007	21.6	0.28	0.0	1.0
18 Oct 2007	10.1	0.31	0.0	0.0
11 March 2008	6.8	0.37	0.0	0.1
21 April 2008	8.9	0.36	0.0	0.1
6 June 2008	20.6	0.23	0.0	0.7

Soil temperature and moisture refer to a soil depth of 8 cm and are daily averages. Note that the data for 6 June 2008 refer to 1 daytime and 1 nighttime measurement

Agilent Technologies Inc., Santa Clara, CA). Concentrations were determined based on calibration against four standard gases with four different concentrations of CH₄ and N₂O (see Hartmann et al. 2011 for details of the experimental setup). Gas exchange rates were calculated by linear regression of headspace concentrations against time. Concentration changes were linear over time, and regressions explained >95 % of the variation in gas concentration ($R^2 > 0.95$), unless flux rates were very low.

Soil sampling

In October 2007 and again in June 2008, we collected eight soil cores (diameter 1.5 cm; depth 15 cm) per subplot within 50 cm of the respective static chamber used for the N₂O and CH₄ flux measurements. Samples were pooled by subplot and sieved (mesh size 2 mm). A sub-sample was stored at −18 °C for molecular analyses, while a second subsample was stored at 4 °C for potential nitrifying and denitrifying activity analysis.

Nitrifying and denitrifying enzyme activity

Nitrifying (NEA) and denitrifying (DEA) enzyme activities were determined in laboratory assays in which the transformation rates of NH₄⁺ and NO₃[−] were quantified under standardized conditions, with no limitations in water availability nor substrate concentration. The goal of this analysis was to quantify potential activity rates in the absence of constraint by substrate availability or environmental conditions that may have existed in the field. NEA was measured at soil pH using the method described in Patra et al. (2005) and was determined as the difference between soil NO₃[−] concentration before and after a 7-h incubation of 10 g of soil under conditions favoring nitrification (28 °C, addition of 0.2 mg (NH₄)₂SO₄-N g^{−1} dry soil, soil moisture equivalent to 70 % water-holding capacity). DEA was assessed over a short time by making all the factors affecting the denitrification rate non-limiting (Smith and Tiedje 1979; Tiedje et al. 1989). DEA was determined as the linear production rate of N₂O, as measured by gas chromatography (Agilent P200; Agilent Technologies Inc.), during an 8-h incubation of 10 g soil under conditions favoring denitrification (28 °C; 90:10 He-C₂H₂ atmosphere providing anaerobic conditions and inhibition of N₂O-reductase activity; addition of 1 mg glucose-C, 1 mg glutamic acid-C, and 0.1 mg NO₃[−]-N g^{−1} dry soil; soil moisture content equivalent to 100 % water-holding capacity).

Nitrifier and denitrifier abundances

Microbial DNA was extracted from the soil samples using the PowerSoil DNA isolation kit (MO BIO Laboratories, Carlsbad, CA). The abundance of beta-proteobacterial ammonia oxidizers, i.e., known ammonia-oxidizing bacteria (AOB) in soil, was measured by quantitative PCR (qPCR) according to Le Roux et al. (2008), targeting 16S rRNA gene sequences that are specific for this group (Hermansson and Lindgren 2001). As reference, we used 5.72·10⁸–5.72·10² copies of standard DNA (purified AOB 16S rRNA gene PCR product of ATCC19718 *Nitrosomonas europaea*).

The abundance of copies of the nitrite (NO₂[−]) oxidoreductase gene (*nxrA*) from *Nitrobacter*-like nitrite-oxidizing bacteria (NOB) was quantified according to Attard et al. (2010). Tenfold standard serial dilutions ranging from 10⁷ to 10¹ *nxrA* copies of genomic DNA from *Nitrobacter hamburgensis* X14 (DSMZ 10229) were used.

Denitrifier abundance was estimated by qPCR targeting fragments of the *nirK* and *nirS* genes encoding the copper and cdI nitrite reductases, respectively. Quantitative PCR assays were carried out from soil DNA with Lightcycler 480 (Roche Diagnostics, Meylan, France). For *nirK*, thermal cycling was performed and a standard curve constructed according to Henry et al. (2004), except that the last step of the PCR cycling profile was performed for 45 cycles at an annealing temperature of 58 °C. *nirS* fragments were amplified as described by Baudoin et al. (2009), using tenfold serial dilutions of genomic DNA of *Pseudomonas aeruginosa* PA14 as standards. We checked melting curves and the lengths of the amplified *nirK* or *nirS* fragments after running an agarose gel of randomly chosen final PCR products.

For all of the qPCR assays, measurements were performed in triplicate. Melting curve analysis confirmed the specificity of the amplifications, and possible inhibitory effects of co-extracted humic compounds in soil extracts were checked by dilution series, but no inhibition was observed.

Soil moisture and inorganic N concentrations

Soil moisture was determined gravimetrically in all fresh samples collected for molecular analysis. Soil NH₄⁺ and NO₃[−] concentrations were determined colorimetrically (SAN+; Skalar Analytical B.V., the Netherlands) in fresh soil samples (20 g dry mass equivalent) following extraction with 60 mL of 2 M KCl (30 min) and filtration.

Data analysis

All data were analyzed by analysis of variance based on mixed-effects models reflecting the experimental design (ASReml 3.0; VSN International, Hemel Hempstead, UK). The models included the main design variables plant species richness (as log-linear contrast) and fertilizer application, and their interaction. The specific composition of plant species communities was included as a random term to ensure adequate testing of the species richness effect (note that this random term is equivalent to “plot” since specific plant community compositions are not replicated at the plot level). The Jena Experiment is located on a flood plain adjacent to the Saale river, and spatial, treatment-independent gradients in soil conditions across the site affected the processes we investigated. We therefore fitted the Cartesian coordinates X and Y , which indicate the plot’s position, plus their second-order terms ($X \cdot X$, $Y \cdot Y$, $X \cdot Y$) to account for non-linearity in spatial gradients and for interactions between the two cardinal directions (i.e., allowing for a response surface that may be curved in a direction that does not align with the main coordinate axes). This spatial model explained more variance than fitting the design’s standard blocks which were placed perpendicular to the river.

The different variables necessitated a range of transformations due to their specific distributions. In part, these distributions result from biological processes; however, they also are the result of specific measurement procedures, such as the PCR reaction, which is an exponential process and results in a corresponding distribution of measurement errors. All gene abundances were therefore log-transformed. Soil CH_4 uptake rates as well as nitrification and denitrification enzyme activities were analyzed as untransformed values. Finally, soil N_2O emissions resembled a log-normal distribution at large values but included very small and negative values due to measurement error, precluding log-transformation (which further would have over-emphasized differences between very small fluxes). We found that taking the fourth square root (with sign reconstruction for negative values) and shifting the abscissa of the first and third quadrant so that the two legs of the function merged with unity slope resulted in a near-perfect normal distribution of residuals. However, the results of the analysis did not change fundamentally when other transformations were chosen.

After testing for the significance of effects of species richness and fertilization, respectively, we used structural equation modeling to explore possible causal relationships between trace gas fluxes, soil environmental variables (water content, NH_4^+ and NO_3^- concentrations), potential nitrifying and denitrifying activities, and gene abundances. The goal was to develop an ecologically sensible,

parsimonious model for which the model-implied covariance between variables matched the observed covariance structure of the data. These covariance matrices were then compared using χ^2 tests, with significant results indicating a poor fit. Models were fitted by maximum likelihood using the lavaan software (<http://lavaan.ugent.be>). As with the mixed model analyses, data were first corrected for spatial gradients across the experimental site. Since not all data were present on all dates, these analyses were carried out on temporal averages of all available measurements, the underlying assumption being that effects and correlations were reasonably stable over time. Structural models were constructed under the premises that (1) processes upstream in terms of N transformation influence downstream processes (e.g., nitrification can influence denitrification), but not vice versa; (2) gene abundance can influence microbial community activity, but not vice versa; (3) trace gas fluxes can be influenced by all variables; (4) the experimental treatments can directly influence all variables. In an educated stepwise process, we then searched for the most parsimonious model that explained the observed covariance structure adequately (insignificant χ^2 test; P was >0.5 for all models, indicating a very good fit). This was achieved by omitting paths with small, non-significant coefficients.

Results

Soil N_2O emissions

Averaged over all sampling dates, soil N_2O emissions were left unaffected by plant species richness but they increased slightly with fertilizer application ($P = 0.003$). When we analyzed sampling dates separately, we found a significant decrease of N_2O emissions with diversity on 21 April 2008 ($P = 0.02$) and significant species richness by fertilizer application interactions on 20 June 2007 ($P = 0.01$) and 11 March 2008 ($P = 0.05$), with the same effect being marginally significant on 21 April 2008 ($P = 0.07$). These interactions resulted from the difference in N_2O emissions between fertilized and control subplots being larger at high plant species richness than at low plant species richness.

Emissions of N_2O were strongly controlled by the presence of legumes. We therefore separately analyzed plots that were sown with and without legumes (Fig. 1). In the absence of legumes, species richness slightly reduced N_2O emissions, while N application slightly increased emissions (both effects being marginally significant at $P = 0.06$). In plots with legumes, N_2O emissions were significantly correlated with the fraction of legume biomass recovered in the summer harvest ($P = 0.001$) and increased with fertilizer application ($P = 0.02$), but they were unrelated to plant species richness.

Fig. 1 Soil nitrous oxide (N_2O) emission rates for plots without (left) and with (right) legumes planted. Individual symbols Values for each plot (gray and white circles values for control and fertilized plots, respectively), black pie shape inside symbol fraction of legumes in aboveground plant biomass harvested in June 2007 and 2008. Square symbols and lines Means predicted by the model for both fertilizer treatments, shaded areas corresponding standard errors (SE)

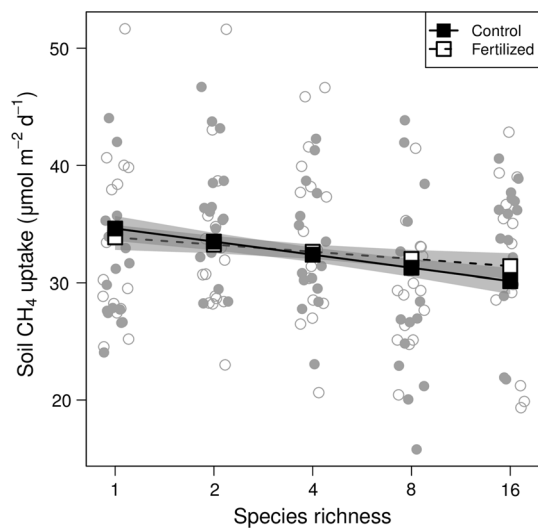
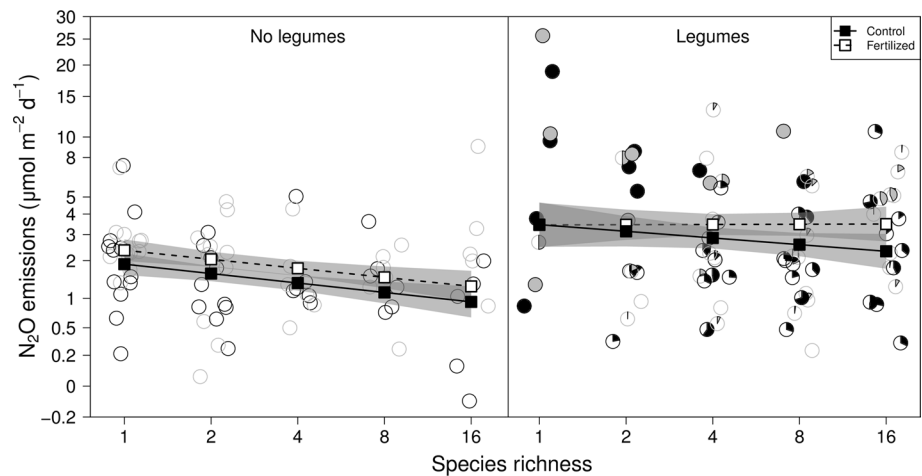


Fig. 2 Net soil methane (CH_4) uptake rate as a function of plant species richness and fertilizer application. Gray and white circles Values for each control and fertilized subplot, respectively. Squares and lines model-predicted means for both fertilizer treatments, shaded areas corresponding SE

Soil CH_4 uptake

Averaged over all sampling dates, soil CH_4 uptake rates decreased with plant species richness ($P = 0.02$), but they remained unaffected by fertilizer application (Fig. 2) and by the abundance of legumes. This effect was remarkably stable over time, with a decrease in CH_4 uptake with species richness detected on 18 October 2007 ($P = 0.05$) and 6 June 2008 ($P = 0.002$), a marginally significant decrease found on 20 June 2007 ($P = 0.08$) and non-significant trends in the same direction found on the other sampling dates.

Nitrifying and denitrifying enzyme activities and functional gene abundances

Nitrifying enzyme activity increased with plant species richness ($P < 0.05$), irrespective of fertilizer application (Table 2). In contrast, DEA increased with fertilizer application ($P < 0.001$), irrespective of plant species richness (Table 2).

Plant species richness did not affect functional gene abundances, although *nxrA* showed a marginally significant ($P = 0.06$) and weak decrease with plant diversity. In contrast, the abundances of AOB, *Nitrobacter*-like NOB and *nirK*- and *nirS*-like denitrifiers increased with fertilizer application (Table 2).

Structural equation models explaining N_2O and CH_4 emissions

In the absence of sown legumes, N_2O emissions were affected negatively by plant species richness, as indicated by a strong direct negative path coefficient (Fig. 3, top). Plant species richness increased soil moisture, resulting in increased potential nitrification (NEA), which in turn stimulated DEA. However, DEA was not significantly linked to N_2O emissions. When plant communities contained legumes (Fig. 3, bottom), the percentage of legumes (measured as aboveground biomass fraction harvested in June) dominated the effects on N_2O emissions. Since the fraction of legumes in the plant community was negatively correlated with species richness and fertilizer application, these treatments thus reduced emissions by this pathway. However, plant species richness stimulated N_2O emissions via increases in soil moisture, which in turn increased NEA (and DEA, but as in the legume-free plots no significant link between DEA and N_2O fluxes was detected).

Table 2 Effects of plant species richness (logSR) and fertilizer application on nitrifying and denitrifying enzyme activities and the abundance of ammonia-oxidizing bacteria, *Nitrobacter*-like nitrite oxidizing bacteria, and *nirK*- and *nirS*-like denitrifiers

Parameters	Significance of effects			Effect sizes	
	logSR	Fertilizer application	logSR × fertilizer application	logSR (%)	Fertilizer application (%)
NEA	*	n.s.	n.s.	+10	
DEA	n.s.	***	n.s.		+7
AOB	n.s.	***	n.s.		+51
NOB	(*)	**	n.s.	(−27)	+47
<i>nirK</i>	n.s.	*	n.s.		+11
<i>nirS</i>	n.s.	***	n.s.		+138

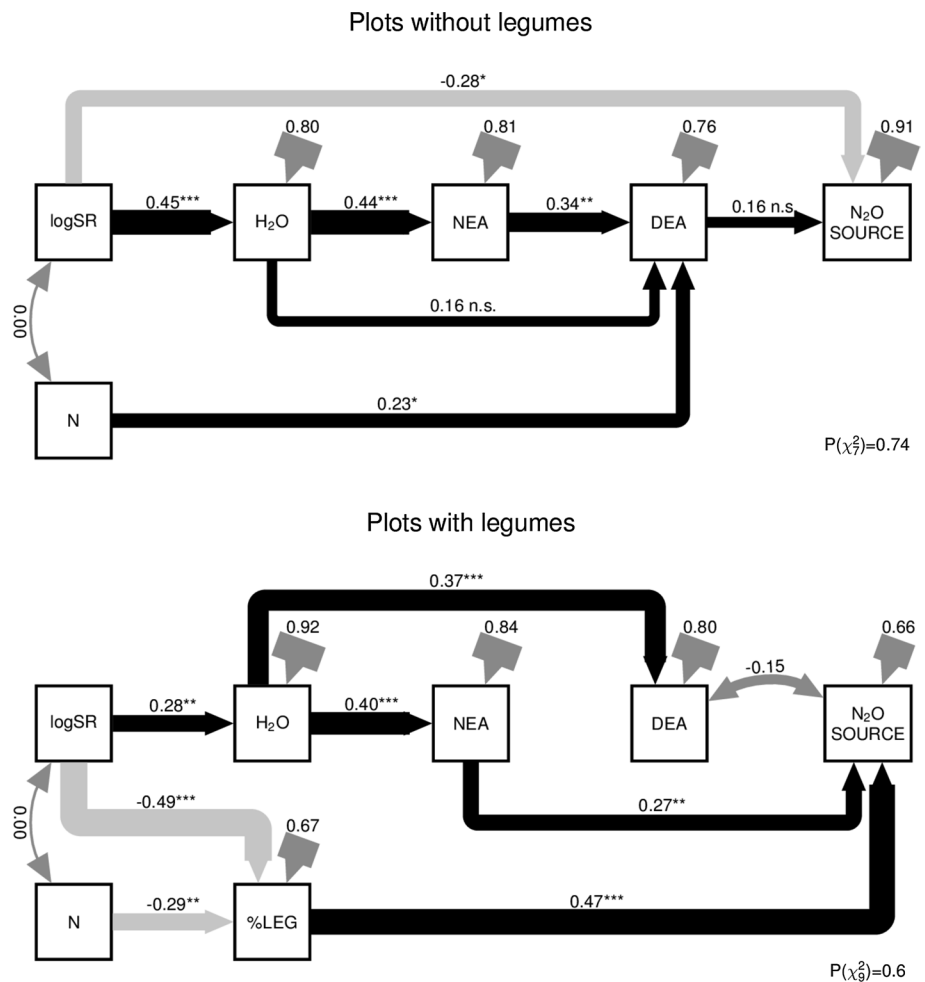
Plant species richness effect is expressed as the logSR (log-linear effect)

Effect sizes are given for model-predicted changes along the full plant species gradient of 1–16 species. Gene abundances are modeled on a log scale, and effect sizes thus are given from the back-transformed model prediction

NEA nitrifying enzyme activity, DEA denitrifying enzyme activity, AOB ammonia-oxidizing bacteria, NOB *Nitrobacter*-like nitrite oxidizing bacteria, *nirK*, *nirS* *nirK*- and *nirS*-like denitrifiers, respectively

(*), *, **, *** Significant difference at (*) $P < 0.1$, * $P < 0.05$, ** $P < 0.01$, *** $P < 0.001$. n.s. Not significant

Fig. 3 Path diagram with standardized path coefficients showing effects of plant species richness (log-transformed, logSR) and fertilizer (N) on soil moisture (H_2O), soil microbial nitrification and denitrification enzyme activities (NEA and DEA, respectively), and soil N_2O emissions (N_2O SOURCE), separately for plots without (top) and with (bottom) legumes. Arrows Standardized path coefficients (gray negative; black positive) together with their significance (* $P < 0.05$, ** $P < 0.01$, *** $P < 0.001$, n.s. not significant). %LEG Mean fraction of legume aboveground biomass harvested in June 2007 and 2008. χ^2 statistics indicate no significant deviation of measured and model-implied covariance structure



Soil NH_4^+ or NO_3^- concentrations had no explanatory power in the analysis and were therefore dropped from all models.

While the net effect of plant species richness on soil CH_4 oxidation was negative, our path analysis suggested that a positive indirect effect of richness via increased soil

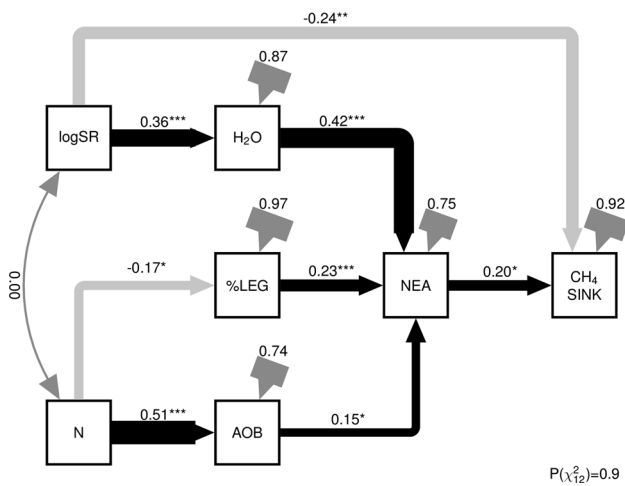


Fig. 4 Path diagram showing effects of plant species richness and fertilizer application on soil moisture, nitrification enzyme activity, the abundance of ammonia-oxidizing bacteria (AOB) and soil CH₄ uptake (*CH₄ SINK*). See caption to Fig. 3 for all other abbreviations. *Arrows* Standardized path coefficients (gray negative, black positive) together with their significance (**P* < 0.05, ***P* < 0.01, ****P* < 0.001). χ^2 statistics indicate no significant deviation of measured and model-implied covariance structure

moisture and increased NEA was also at play. However, this positive effect was masked by a larger direct negative effect on CH₄ uptake (Fig. 4). Fertilizer application stimulated AOB abundance, which positively correlated with

nitrification activity. Legume abundance also boosted NEA. Ultimately, NEA had a positive effect on CH₄ uptake. Soil NH₄⁺ concentration had no explanatory power in this analysis.

Structural equation models linking functional gene abundances with NEA and DEA

Path analysis did not reveal any significant links between potential nitrification and denitrification activities on one hand, and the abundances of AOB, *Nitrobacter*-like NOB or *nirK*-like denitrifiers on the other hand (Fig. 5). Due to limited amounts of extracted soil DNA available, *nirS* abundances could only be determined for both sampling dates in 83 plots, for one sampling date in another 63 plots, and not at all in 10 plots. Analysis of the subset for which both *nirS* and *nirK* abundances were available revealed a similar pattern, i.e., no link between *nirS* and enzyme activities (Online Resource 1).

At the level of enzyme activities, the effects of plant species richness were mediated by soil moisture, while the effects of fertilizer application were mediated by a reduced legume fraction in the plant communities. At the level of microbial functional group abundances, N fertilization had a clear positive effect on the abundance of AOB, with cascading effects on the abundance of *Nitrobacter*-like NOB and *nirK*-like denitrifiers (Fig. 5). *nirS*-like denitrifier abundances were affected positively by N fertilization, both

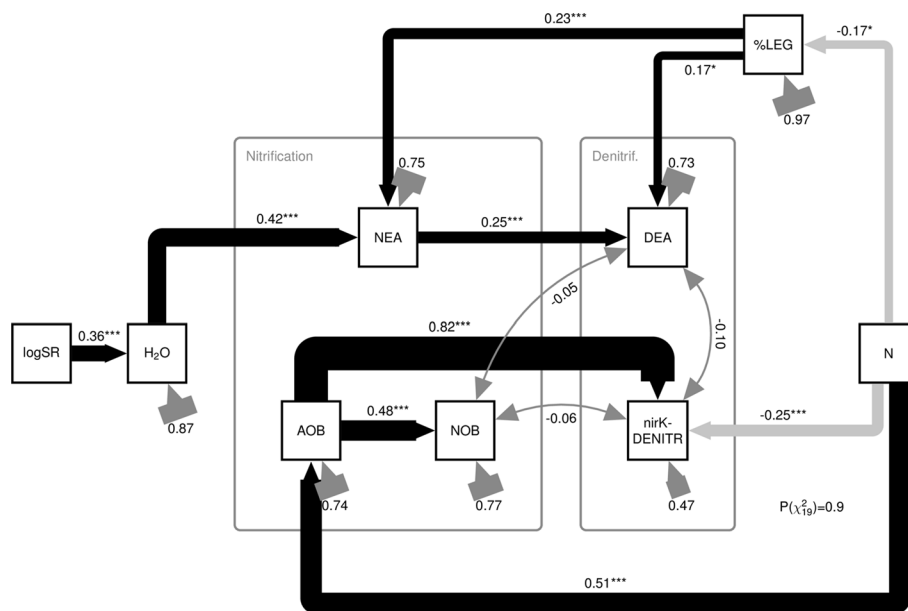


Fig. 5 Path diagram showing relationships between species richness, fertilizer application, nitrifying and denitrifying enzyme activities, and abundances of key microbial functional groups (*NOB Nitrobacter*-like nitrite oxidizers, *nirK-DENITR nirK*-like denitrifiers). See captions to Figs. 3 and 4 for all other abbreviations. *Arrows* Stand-

ardized path coefficients (gray negative, black positive) together with their significance (**P* < 0.05, ***P* < 0.01, ****P* < 0.001). The correlation between the exogenous variables logSR and N is zero and omitted for clarity. χ^2 statistics indicate no significant deviation of measured and model-implied covariance structure

directly and indirectly via AOB and *Nitrobacter*-like NOB (Electronic Supplementary Material 1).

Discussion

Plant species richness decreased soil N₂O emissions—at least in the absence of legumes—and decreased soil CH₄ uptake. We also detected plant diversity effects on underlying soil microbial processes and the abundances of soil microbial groups related to these processes. However, structural equation modeling suggested that these different effects were linked in a complex way, with net effects on trace gas fluxes emerging from a combination of direct and indirect species richness effects; in this context, “direct” denotes effects not likely related to the dependent variables observed.

Structural equation modeling suggests that the effects on N₂O fluxes were at least in part mediated by positive effects of plant species numbers on soil moisture, which in turn led to accelerated inorganic N cycling, as evidenced by increased NEA and DEA. Positive path coefficients to N₂O fluxes (Fig. 3) suggest that this increase in N transformation capacity indeed stimulated N₂O emissions. At the same time, the structural equation models support a direct negative effect of plant species richness on N₂O emissions, which counteracted the positive effects of increased enzyme activities. A possible mechanism underlying this link may stem from disproportionate effects of species richness on soil inorganic N concentrations and the enzymatic potential (in our study: NEA and DEA) to transform this inorganic N. Many studies have shown more efficient soil inorganic N capture and thus lower soil NO₃[−] concentrations in more species-rich plant communities (Ewel et al. 1991; Niklaus et al. 2001; Tilman et al. 1996). In the Jena experiment, Leimer et al. (2014b) found that soil NO₃[−] concentrations were reduced at high diversity if no or only few legumes were present, which is compatible with the general trend found in other studies. In our analysis, however, soil inorganic N concentrations were unrelated to the explanatory variables analyzed. One reason for this difference may be that our measurements were not frequent enough to estimate robust average concentrations of these very dynamic parameters. When legumes were present in experimental communities, their fractional contribution to plant community biomass was positively related to nitrification and soil N₂O emissions. However, the negative direct effect from species richness to N₂O emissions that we identified in the absence of legumes vanished in the structural equation models (Fig. 3). This observation is compatible with the finding by Leimer et al. (2014b) that soil NO₃[−] concentrations increased with diversity when a large fraction of species were legumes, indicating that a high legume

diversity also led to high NO₃[−] concentrations and that this effect exceeded the higher inorganic N capture efficiency in more diverse plant communities that was otherwise found.

Net soil CH₄ uptake decreased persistently with plant species richness. The soil’s CH₄ balance is determined by the difference between methanogenesis and CH₄ oxidation, with the latter dominated by methanotrophic bacteria. We did not observe net CH₄ emissions during our study, indicating that CH₄ oxidation consistently outweighed methanogenesis; in fact, we think that methanogenesis did not play an important role in the present system and that the observed patterns were thus solely caused by variation in methanotrophic activity. Soil CH₄ oxidation is often limited by gas transport rates, which in turn depend on water-filled pore space (Ball et al. 1997). Increased soil moisture at high plant species richness could therefore explain the observed pattern. However, structural equation modeling suggested indirect positive effects of moisture on CH₄ oxidation via increased nitrification (Fig. 4). This mechanism is plausible, given that methanotrophy can be inhibited by high NH₄⁺ concentrations; under these conditions, all processes consuming soil NH₄⁺ will indirectly promote CH₄ oxidation. Apart from nitrification, NH₄⁺ assimilation by plants and immobilization in microbial biomass would exert such an effect. Path coefficients further suggested a soil moisture-independent direct negative effect of species richness on soil CH₄ uptake. This effect persisted even if a (non-significant) link from soil moisture to CH₄ uptake was added to the model, suggesting that this effect was unrelated to soil moisture. The potential mechanisms supporting this link remain elusive; however, it also should be noted that path models do not provide strict tests of causalities but rather indicate possible mechanistic links based on correlations. The ecology of soil methanotrophic bacteria consuming atmospheric CH₄ is only poorly understood to date, in part due to the lack of success in isolating such organisms from soils (Dunfield 2007). However, it is known from experiments with intact soil cores and with available laboratory cultures that CH₄ oxidation is negatively affected by many chemical compounds, including ethylene (Jäckel et al. 2004), some organic acids (Wieczorek et al. 2011), and terpenes (Amaral et al. 1998). One possibility is that the detrimental effect of these substances results from the lack of specificity of methane-monooxygenase, which leads to co-metabolic activity harmful to the bacteria, such as by suicide activation (Mahendra and Alvarez-Cohen 2006; Prior and Dalton 1985). In this light, one possible explanation for the soil moisture-independent reduction of CH₄ uptake in plant species-rich communities could be an increased production of such compounds inhibiting CH₄ oxidation.

It is surprising that soil inorganic N concentrations were not significantly related to any other variable measured,

whereas species richness, fertilization, and legume presence had significant effects on these parameters. This could be due to the fact that soils are spatially heterogeneous, as are the biogeochemical transformations we investigated. For example, denitrification often occurs in the micro-anaerobic parts of soil aggregates (Sexstone et al. 1985), and recent micro-autoradiographic analyses indicate that CH_4 assimilation preferentially takes place on aggregate surfaces to which CH_4 can readily diffuse (Stiehl-Braun et al. 2011a, b). Directly relating system-level processes, such as land–atmosphere trace gas fluxes, to bulk soil measures of putative drivers, such as to soil inorganic N concentrations or enzymatic activities, disregards the micro-scale character of the interactions involved. When processes respond in a non-linear fashion to substrate availability, different effects will be found when a resource such as NO_3^- is homogeneously or heterogeneously distributed. Our measurements, as is typical for ecosystem-level field studies, integrate the small-scale heterogeneity and thus are unable to account for small-scale interactions in hot spots and at hot moments, which show episodically high biogeochemical transformation rates (McClain et al. 2003). We consider it likely that the marked effect of legumes is related to such heterogeneity. For example, high local concentrations of NO_3^- can occur in the vicinity of legume nodules, possibly creating hot spots of denitrification and N_2O emissions. Efficient capture of inorganic N by neighboring plants will not lower inorganic N concentrations at these locations, although it will reduce NO_3^- leaching losses. Such localized high concentrations will be difficult to detect in bulk soil measurements. This reasoning is consistent with a previous investigation in which we found nitrification to be only weakly related to NH_4^+ and denitrification only weakly related to NO_3^- , whereas legume presence had comparably large effects (Le Roux et al. 2013). With respect to CH_4 fluxes, high bulk soil NH_4^+ concentrations may not affect CH_4 oxidation if they occur in soil space that is distinct from the spatial niche of the active methanotrophs (Hartmann et al. 2011; Stiehl-Braun et al. 2011a).

The quantification of microbial guilds by functional gene abundance promised some insight into the community dynamics underlying N transformations. In our study, different experimental treatments were reflected differently in terms of functional gene abundances and enzyme activities, and no links between the two could be detected in the structural equation models. Soil moisture-mediated effects of plant species could be detected in enzyme activities but not in the community size of the respective microbial guilds. However, fertilizer effects did exert a cascading effect via increased NH_4^+ oxidizer communities to nitrite oxidizers and denitrifiers. Interestingly, no link from nitrite oxidizer gene abundances to *nirK* gene abundances was

detected (Fig. 5), although the latter is downstream of the first in terms of N transformation processes. This relationship can be understood in the light of NH_4^+ (and not nitrite) oxidation being the rate-limiting step of nitrification. Probably reflecting differences in ecology of denitrifier groups (Xie et al. 2014), *nirS* gene abundances were dependent on NH_4^+ oxidizer 16S genes abundances indirectly via nitrite oxidizer genes. Overall, however, NH_4^+ oxidizer community size appeared to control denitrifier community size, mostly directly but to some extent also indirectly via intermediate (in terms of N processing) nitrite oxidizer communities. Environmental controls rather than community size have also been detected in contexts other than species richness (e.g., fertilizer and drought effects: Hartmann et al. 2013; land use change: Attard et al. 2011), underlining that gene abundance data reflect biological activities only under specific conditions.

Our study revealed that many effects of plant species richness on microbial processes were mediated by effects on soil moisture. Reports on species richness effects on soil moisture are equivocal (Caldeira et al. 2001; Rosenkranz et al. 2012; Spehn et al. 2000); Caldeira et al. (2001) reported positive effects of species richness on grassland top soil moisture but negative effects at soil depths of >15 cm. In the Jena experiment, Rosenkranz et al. (2012) found similar positive effects on top soil moisture as we did, while Leimer et al. (2014a) detected no such effects. In general, patterns in soil moisture seem to have been caused by decreased evaporative losses under higher species richness due to soil surface shading, with a concomitant increase in water capture from deeper soil layers due to better soil exploration by roots. If processes are soil moisture driven, the depth at which the respective microbial transformations take place is therefore critical. N_2O production due to nitrification and denitrification often occurs in the top soil; soil CH_4 oxidation generally also dominates in the top soil, whereas methanogenesis often occurs deeper in the soil (Conrad 1996).

The patterns that emerged from our study, in particular from structural equation modeling, were robust and coherent, integrating well into our current understanding of the fundamental ecological processes. With respect to the fluxes of N_2O and CH_4 per se, one must note that the effects of species richness and fertilization were small compared to the total variation in our data, which commands a cautious interpretation (Low-Décarie et al. 2014). One reason for the large fraction of unexplained variance may be that the between-plant community variation is typically very high in experimental plant diversity experiments, in particular at low species richness. The observed diversity effects thus result from averaging over the large inter-community variation within plant species richness levels. In the Jena experiment, as in many field biodiversity studies, specific species

combinations are not replicated in independent plots, so that the diversity-independent composition effect cannot be separated from unrelated (random) variation, with the exception of simple contrasts, such as the presence or absence of specific plant functional types. When studying trace gas dynamics and the underlying microbial processes, the researcher is further confronted with large spatial and temporal variation inherent to the processes. Despite this challenging variability, the present field study is the first to present clear and consistent responses of soil trace gas exchange to plant species richness and fertilization. Controlled experimental studies as the present one complement observational research carried out under more natural conditions but in which it is more difficult to isolate effects of plant diversity from potentially confounding factors (e.g., Keil et al. 2015; Sutton-Grier et al. 2011).

Author contribution statement PAN and RLB conceived and designed the study with input from MSL and NB. Trace gas flux measurements and soil analyses were carried out by PAN, MSL, and RLB. XLR and FP analyzed functional gene abundances. PAN analyzed the data with input from XLR and RLB. PAN wrote the manuscript with input from all co-authors.

Acknowledgments We gratefully acknowledge Ingeborg Schininger for help with field sampling and laboratory sample preparation, and Nadine Guillaud (AME platform of UMR5557) for help with enzyme activity measurements.

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