# ORIGINAL ARTICLE



# Direct and plant community mediated effects of management intensity on annual nutrient leaching risk in temperate grasslands

Antonios Apostolakis<sup>®</sup> · Ingo Schöning<sup>®</sup> · Valentin H. Klaus<sup>®</sup> · Beate Michalzik<sup>®</sup> · Wolf-Anno Bischoff · Runa S. Boeddinghaus · Ralph Bolliger<sup>®</sup> · Markus Fischer · Norbert Hölzel · Ellen Kandeler · Till Kleinebecker<sup>®</sup> · Peter Manning<sup>®</sup> · Sven Marhan<sup>®</sup> · Margot Neyret · Yvonne Oelmann · Daniel Prati · Mark van Kleunen · Andreas Schwarz · Elisabeth Schurig · Marion Schrumpf<sup>®</sup>

Received: 27 September 2021 / Accepted: 21 May 2022 © The Author(s) 2022

Abstract Grassland management intensity influences nutrient cycling both directly, by changing nutrient inputs and outputs from the ecosystem, and indirectly, by altering the nutrient content, and the diversity and functional composition of plant and microbial communities. However, the relative importance of these direct and indirect processes for the leaching of multiple nutrients is poorly studied. We measured the annual leaching of nitrate, ammonium,

**Supplementary Information** The online version contains supplementary material available at https://doi.org/10.1007/s10705-022-10209-1.

A. Apostolakis (⊠) · I. Schöning · M. Schrumpf Max-Planck-Institute for Biogeochemistry, Hans-Knöll-Str. 10, 07745 Jena, Germany e-mail: apostolakis@bgc-jena.mpg.de

A. Apostolakis · B. Michalzik Department of Soil Science, Friedrich Schiller University Jena, Löbdergraben 32, 07743 Jena, Germany

V. H. Klaus Institute of Agricultural Sciences, ETH Zürich, Universitätstr. 2, 8092 Zurich, Switzerland

B. Michalzik German Centre for Integrative Biodiversity Research (iDiv) Halle -Jena-Leipzig, Deutscher Platz 5e, 04103 Leipzig, Germany

W.-A. Bischoff · A. Schwarz TerrAquat GmbH, Schellingstr. 43, 72622 Nürtingen, Germany phosphate and sulphate at a depth of 10 cm in 150 temperate managed grasslands using a resin method. Using Structural Equation Modeling, we distinguished between various direct and indirect effects of management intensity (i.e. grazing and fertilization) on nutrient leaching. We found that management intensity was positively associated with nitrate, ammonium and phosphate leaching risk both directly (i.e. via increased nutrient inputs) and indirectly, by changing the stoichiometry of soils, plants and microbes. In contrast, sulphate leaching risk was negatively associated with management intensity, presumably due to increased outputs with moving

R. S. Boeddinghaus · E. Kandeler · S. Marhan Institute of Soil Science and Land Evaluation, Soil Biology Department, University of Hohenheim, 70599 Stuttgart, Germany

R. Bolliger · M. Fischer · D. Prati Institute of Plant Sciences, University of Bern, Altenbergrain 21, 3013 Bern, Switzerland

N. Hölzel Institute of Landscape Ecology, University of Münster, Heisenbergstr. 2, 48149 Münster, Germany

T. Kleinebecker Institute of Landscape Ecology and Resource Management, Justus-Liebig-University Gießen, Heinrich-Buff-Ring 26-32, 35392 Gießen, Germany and grazing. In addition, management intensification shifted plant communities towards an exploitative functional composition (characterized by high tissue turnover rates) and, thus, further promoted the leaching risk of inorganic nitrogen. Plant species richness was associated with lower inorganic nitrogen leaching risk, but most of its effects were mediated by stoichiometry and plant community functional traits. Maintaining and restoring diverse plant communities may therefore mitigate the increased leaching risk that management intensity imposes upon grasslands.

**Keywords** Annual nutrient leaching · Inorganic nitrogen · Phosphate · Sulphate · Temperate grasslands · Grassland management

## Introduction

Grasslands are widespread and highly diverse ecosystems providing a multitude of ecosystem functions and services, including nutrient cycling and food production (Bengtsson et al. 2019). Management intensification of grassland ecosystems places them at considerable risk of nutrient losses via leaching, especially for nitrate (Klaus et al. 2018 and references therein), which represent the loss of important resources and, at the same time, potential threats to human health and aquatic ecosystems (EEA 2018; EU-Nitrate Directive 2018; WHO 2011). Compared to nitrogen (N), management effects on leaching of other elements, like phosphorus (P) and sulphur (S), are less studied in grasslands, despite their agricultural and environmental importance (Eriksen 2009;

P. Manning · M. Neyret

Senckenberg Biodiversity and Climate Research Centre (SBiK-F), Senckenberganlage 25, 60325 Frankfurt, Germany

Y. Oelmann · E. Schurig Geoecology, University of Tübingen, Rümelinstr. 19-23, 72070 Tübingen, Germany

M. van Kleunen Department of Biology, University of Konstanz, 87464 Constance, Germany

M. van Kleunen

Zhejiang Provincial Key Laboratory of Plant Evolutionary Ecology and Conservation, Taizhou University, Taizhou 318000, China Withers et al. 2014; Gallejones et al. 2012). Unravelling the relationships between ecosystem management, biotic drivers and nutrient leaching will provide important insights for sustainable grassland management and could help secure associated ecosystem services.

In unfertilized grasslands, inorganic nutrients that are available for leaching represent the net balance between nutrient inputs to the soil solution (e.g. by organic matter mineralization or dissolution) and outputs from the soil solution (e.g. due to uptake by plants, or immobilization in microbial biomass), as well as nutrient exchange with the soil matrix (i.e. via (de)sorption to minerals). Management, such as fertilization and grazing, changes these relations by increasing both inputs and outputs of nutrients from the ecosystem. In the past, grasslands were considered to have low N leaching fluxes, and introducing temporary (mowed) grasslands in arable crop rotations can even reduce nitrate leaching (Kunrath et al. 2015). However, studies investigating the effects of grassland management on soil nitrate show increased leaching risk in pastures compared to meadows (Di and Cameron 2002; Ryden et al. 1984), and when broader management gradients are included, both fertilization and grazing equally increase nitrate leaching risk (Klaus et al. 2018). Grassland intensification can affect nutrient cycling *directly* by changing inputs and outputs (i.e. fertilization, grazing and harvest, Rumpel et al. 2015) and *indirectly* by changing plant communities (e.g. loss of diversity or functional change) and altering interactions between soils, plants and soil microorganisms (de Vries et al. 2012; Klaus et al. 2018). However, at present, it is unclear if management intensity predominantly drives leaching risk directly and/or indirectly by changing plant and/or microbial communities.

Grassland management, mainly fertilization, mowing and grazing, influences soil, plant and microbial properties (Herold et al. 2014; Kleinebecker et al. 2014; Boeddinghaus et al. 2019) with potential effects on nutrient leaching. For instance, management intensification reduces the C:N ratio of grassland soils and increases enzymatic activities (Herold et al. 2014) and, potentially, the decomposition of organic matter. In addition, fertilization increases plant biomass productivity and its nutrient concentrations, reducing its C:N stoichiometry, and it reduces plant diversity and root litter decomposition (Gilhaus et al. 2017; Klaus et al. 2011; Solly et al. 2014). Many studies have reported that a high plant species diversity reduces N and P availability and leaching in grasslands (de Vries et al. 2016; Leimer et al. 2015, 2016; Oelmann et al. 2011; Tilman et al. 1996), especially when compared to monocultures (Scherer-Lorenzen et al. 2003), possibly due to resource complementarity (Kahmen et al. 2006). However, plant diversity effects on nitrate leaching are still debated and have not yet been comprehensively assessed up to now, as most of the respective studies were conducted in experimental grasslands or mesocosm experiments. Such studies are useful to gain a mechanistic understanding of the underlying processes controlling soil and plant interactions. However, they often do not consider realistic management and/or environmental gradients. Thus, they might fail to capture the complexity of real-world ecosystems (Klaus et al. 2020a) since, for instance, management is an important determinant of plant community diversity and composition (Busch et al. 2018; Midolo et al. 2019). To overcome these constraints, and gain knowledge transferable to sustainable grassland management, we need to investigate grassland ecosystems along broad gradients of management intensity and environmental conditions (Manning et al. 2019).

Plant community composition, e.g. the abundance of functional groups or specific traits, can also play an important role in grassland nutrient cycling (Boeddinghaus et al. 2019; de Vries and Bardgett 2016). Plant functional groups are often used as predictors of nitrogen leaching. The group of legumes is, for example, associated with higher nitrate leaching, while the group of grasses indicates lower nitrate leaching (de Vries and Bardgett., 2016; Leimer et al. 2015, 2016). In addition, plant traits can be used to categorize plant communities as fast-growing, and exploitative or slow-growing and conservative (Bruelheide et al. 2018; Reich 2014; Díaz et al. 2016). Exploitative plant communities have higher specific leaf area and leaf N (and P) and lower leaf dry matter content than conservative communities (Díaz et al. 2016; Wright et al. 2004). Regarding belowground traits, exploitative communities have higher specific root length and lower root tissue density and fine root diameter than the conservative communities (Bergmann et al. 2020). Plant, and especially root, traits are thus broadly indicative of nutrient acquisition strategies of plant species and these are relevant for nutrient cycling in soils (de Vries and Bardgett, 2016). For instance, a high specific leaf area indicates high photosynthetic capacity, which relates to fast growth rates, short tissue lifespan and high decomposability, and high needs for nutrients (de Vries and Bardgett, 2016). Moreover, a high specific root length indicates a better screening of soil for nutrients, and a low fine root diameter indicates high nutrient uptake efficiency (Bardgett et al. 2014; Freschet et al. 2021). In managed grasslands, plant community traits are influenced by management-induced shifts in plant community composition (Allan et al. 2015; Busch et al. 2018) and might explain the effects of vegetation change on nutrient leaching.

Recent evidence highlights the importance of soil microbial communities and their properties for N cycling (Grigulis et al. 2013; Pommier et al. 2018). Understanding, and even controlling, plant-microbe interactions could provide opportunities for enhanced N retention in grasslands, with desired effects both on biomass production and N leaching mitigation. Studies on element cycling that include soil microbial properties are, in general, scarce compared to those addressing abiotic and/or plant effects (Pommier et al. 2018), and so are studies that simultaneously study the cycling of multiple nutrients in managed grasslands. However, management strongly changes microbial properties since, for instance, fertilization shifts the microbial community structure towards bacteriadominated communities (de Vries et al. 2006; Donnison et al. 2000; Herold et al. 2014), which are characterized by lower microbial C:N ratios and higher mineralization rates (Six et al. 2006; Strickland and Rousk, 2010). To elucidate the drivers of ecosystem (dis)services like nutrient leaching, a holistic approach that includes management, but also soil, plant and microbial effects and their interactions is needed (Koncz et al. 2015; Pommier et al. 2018).

Reliable assessments of the drivers of nutrient leaching need to cover long time spans, for example due to the large variability in weather conditions and the timing of land management activities (de Vries et al. 2011; Klaus et al. 2018). This is even more important since recent evidence suggests that plant diversity effects on N leaching follow seasonal patterns with more pronounced effects in fall or winter (Compton et al. 2021; Leimer et al. 2015), seasons that are often ignored as most studies focus on the growth period only. However, accounting for both high spatial and temporal variability imposes considerable logistical constraints. Resin methods for the evaluation of nutrient availability and leaching have received increasing attention in the literature (Grahmann et al. 2018; Grunwald et al. 2020; Predotova et al. 2011; Willich and Buerkert, 2016), and are often successfully used to integrate the temporal effects of different drivers on nutrient leaching (Klaus et al. 2018, 2020b).

In this study, we measured the annual downward flux of nitrate, ammonium, phosphate and sulphate with the percolating water, i.e. leaching, at a depth of 10 cm in 150 grassland sites from spring 2018 to spring 2019. The grassland sites were located in three regions in Germany as part of the Biodiversity Exploratories project (Fischer et al. 2010), and covered broad management intensity gradients within each region. We used a resin method, namely the Self-Integrating Accumulators (SIAs), to determine the cumulative annual leaching, and not just the availability of nutrients in the soil solution (Bischoff, 2009). Considering that grassland management, here fertilization and grazing, has direct (via increased inputs and outputs) and indirect (via soils, plants and microbial biomass) effects on nitrate, ammonium, phosphate and sulphate leaching, we addressed the following hypotheses:

1. Fertilization and grazing intensity increase nutrient leaching risk from topsoils due to increased nutrient inputs (direct effects).

- 2. Fertilization and grazing intensity increase nutrient leaching risk by lowering the C:N stoichiometry of soils, plants and soil microbial biomass (indirect effects).
- 3. Fertilization and grazing intensity increase nutrient leaching risk (i) by shifting the functional composition of plant communities from conservative (i.e. slow-growing) communities at extensively managed grasslands to exploitative (i.e. fast-growing) communities at intensively managed grasslands, and (ii) by reducing plant species richness (indirect effects).

# Methods

# Study regions

This study was conducted in the *Biodiversity Exploratories* project, which investigates land use effects on biodiversity and ecosystem functions and services in real-world ecosystems (Fisher et al. 2010). The project includes three study regions in Germany; the Schwäbische-Alb (ALB), the Hainich-Dün (HAI) and the Schorfheide-Chorin (SCH). The study regions differed in their geology, topology, dominant soil types and/or climate (Table 1). Soils in ALB developed mainly on Jurassic limestone and were clay-rich Leptosols or Cambisols (IUSS Working Group WRB, 2014). In HAI, soils had a loamy or clayey texture due to the dominant geological substrate of loess over limestone and the main soil types were Cambisols,

Table 1Geographical,geological and climaticcharacteristics of the threestudy regions includedin the <i>Biodiversity</i>	Parameter	Study region		
		ALB	HAI	SCH
	Location in Germany	Southwest	Central	Northeast
<i>Exploratories</i> project after Fischer et al. (2010)	Geology	Calcareous bedrock with loam layer and karstic phenomenon	Central Calcareous bedrock with loess layer 285–550 Cambisols, Stagnosols and Vertisols	Post glacial landscape with sandy ridges and fens
	Elevation (m a.s.l.)	460-860	285-550	3-140
<i>ALB</i> Schwäbische-Alb; <i>HAI</i> Hainich-Dün; <i>SCH</i> Schorfheide-Chorin; a.s.l. = above sea level. Soil	Dominant soil types	Leptosols and Cambisols	Cambisols, Stagnosols and Vertisols	Histosols, Gleysols, Cambisols, Luvisols and Albeluvisols
types are given according to IUSS Working group (2007)	Annual temperature (°C)	6.0-7.0	6.5-8.0	8.0-8.5
	Annual precipitation (mm)	700–1,000	500-800	500-600

Stagnosols and Vertisols. Soils in SCH were drained Histosols with a loamy texture, Gleysols, Cambisols, Luvisols and Albeluvisols.

In each study region, 50 plots of 50 m $\times$ 50 m were established within agriculturally managed permanent grasslands (Fisher et al. 2010). Information regarding the regional management practices as well as soil, plant and microbial properties is

given in Table 2 (and Tables S1 and S2). Management intensity ranged from extensively managed grasslands to intensively grazed and/or fertilized pastures and meadows, covering a wide and realistic management intensity gradient. Fertilization, expressed as N additions, was applied as organic or mineral (chemical) fertilizers. Grazing, expressed as equivalent-livestock unit per area multiplied

Parameter	Study region			
	ALB	HAI	SCH	
Management practices				
Fertilization (kg N ha <sup><math>-1</math></sup> y <sup><math>-1</math></sup> )	$58.2 \pm 87.8$	$39.1 \pm 52.0$	$3.0 \pm 13.0$	
Grazing (LU days ha <sup>-1</sup> y <sup>-1</sup> )	$77.8 \pm 117.4$	$101.5 \pm 138.9$	186.6±197.5	
Mowing (cuts $y^{-1}$ )	$1.3 \pm 1.1$	$1.0 \pm 0.7$	$0.8 \pm 0.6$	
Plant properties				
Aboveground biomass (g m <sup>-2</sup> )	$178.5 \pm 96.5$	$145.4 \pm 69.7$	$364.6 \pm 156.2$	
Aboveground N (g N kg <sup>-1</sup> )	$18.4 \pm 2.9$	$18.6 \pm 2.0$	17.4±3.4	
Aboveground P (g P kg <sup><math>-1</math></sup> )	$2.8 \pm 0.4$	$3.1 \pm 0.3$	$2.7 \pm 0.3$	
Species richness (per 16 m <sup>2</sup> )	$30.4 \pm 8.8$	$37.2 \pm 13.3$	$25.8 \pm 7.2$	
Cumulative legume species cover (%)	$8.2 \pm 9.0$	$13.6 \pm 10.9$	$9.3 \pm 11.7$	
Cumulative grass species cover (%)	$46.5 \pm 18.7$	$42.1 \pm 18.6$	$87.4 \pm 28.5$	
CWM plant traits				
Specific leaf area, SLA $(mm^2 mg^{-1})$	$23.2 \pm 3.2$	$23.7 \pm 1.8$	$23.8 \pm 2.4$	
Leaf dry matter content, LDMC (g $g^{-1}$ )	$0.26 \pm 0.03$	$0.25 \pm 0.02$	$0.26 \pm 0.02$	
Leaf N (mg $g^{-1}$ )	$24.2 \pm 4.0$	$25.7 \pm 2.5$	$25.3 \pm 2.7$	
Leaf P (mg $g^{-1}$ )	$2.3 \pm 0.5$	$2.4 \pm 0.3$	$2.4 \pm 0.4$	
Fine root diameter, fineRD (mm)	$0.19 \pm 0.01$	$0.20 \pm 0.02$	$0.18 \pm 0.01$	
Root tissue density, RTD (mg $cm^{-3}$ )	$53.1 \pm 12.7$	$51.1 \pm 9.7$	$50.2 \pm 11.3$	
Specific root length, SRL (m $g^{-1}$ )	$436 \pm 48$	$414 \pm 43$	$452 \pm 63$	
Microbial properties				
Biomass carbon ( $\mu g C g^{-1}$ )	$816 \pm 152$	$729 \pm 159$	$583 \pm 353$	
Biomass nitrogen ( $\mu$ g N g <sup>-1</sup> )	$137 \pm 39$	$140 \pm 41$	$136 \pm 97$	
Soil properties (at 0–10 cm depth)				
Bulk density (g $cm^{-3}$ )	$0.76 \pm 0.11$	$0.79 \pm 0.11$	$0.85 \pm 0.4$	
Clay $(g kg^{-1})$	$536.1 \pm 134.0$	$422.6 \pm 130.3$	173.8±87.0	
Silt $(g kg^{-1})$	$406.6 \pm 114.6$	$520.1 \pm 122.7$	372.1 ± 191.7	
Sand $(g kg^{-1})$	$57.4 \pm 45.0$	$57.6 \pm 23.1$	$454.2 \pm 221.0$	
pH	$6.3 \pm 0.6$	$6.9 \pm 0.5$	$6.5 \pm 0.9$	
Cation exch. capacity (cmol $kg^{-1}$ )	$23.0 \pm 8.54$	$21.9 \pm 9.67$	$21.0 \pm 17.4$	
Aluminum (oxalate extraction, $g kg^{-1}$ )	$0.56 \pm 0.73$	$0.36 \pm 0.22$	$0.17 \pm 0.21$	
Soil organic C (g C kg <sup><math>-1</math></sup> )	$69.8 \pm 14.3$	$49.9 \pm 12.1$	$95.5 \pm 90.7$	
Soil N (g N kg <sup><math>-1</math></sup> )	$6.8 \pm 1.5$	$4.9 \pm 1.2$	$10.4 \pm 9.0$	
Labile and moderately labile inorganic P	$19.3 \pm 16.6$	$11.7 \pm 8.1$	$29.8 \pm 17.1$	
$(P_{labile,inorg}) (mg P kg^{-1})$				
Total S (g S $kg^{-1}$ )	$0.9 \pm 0.2$	$0.6 \pm 0.1$	$1.4 \pm 1.3$	
Depth (cm)	$41.5 \pm 21.6$	$60.2 \pm 18.3$	$42.8 \pm 50.1$	

Table 2Mean valuesand standard deviation ofmanagement practices andplant, microbial and soilproperties of the three studyregions; Schwäbische-Alb(ALB), Hainich-Dün (HAI)and Schorfheide-Chorin(SCH)

LUlivestock units;

*CWM* community weighted mean. Fertilization, grazing and mowing intensity data (averaged for the period 2015–2018) were obtained from Ostrowski et al. (2020) with grazing days (i.e. the days the animals are on the plot; Blüthgen et al. 2012), included mainly cattle or sheep grazing.

## Soil properties

In May 2017, we collected 14 soil samples of the upper 10 cm along two intersecting 20 m transects in each of the 150 grasslands. Soil samples were used to prepare one composite sample per grassland. All soil samples were air-dried and sieved to < 2 mm, while a portion was ground for elemental analysis. Total carbon (C) and total N were determined by dry combustion at 1,100 °C with an elemental analyser VarioMax (Elemental, Hanau, Germany). Soil inorganic C was determined through the same process after removing organic carbon (OC) by exposing 250 mg of soil to 450 °C for 16 h. Soil OC was calculated as the difference between total and inorganic C. Soil pH was measured in the extraction solution of 10 g of soil with 25 mL of CaCl<sub>2</sub> (0.01 M) with a glass-electrode and a pH meter (WTW pH meter 538, Weilheim, Germany). Total soil S concentration was determined with **ICP-OES** (Acros, SPECTRO Analytical Instruments GmbH, Kleve, Germany) in the extracted solution of an aliquot of 1.5 g of soil with 3.5 mL HNO<sub>3</sub> and 10.5 mL HCl.

Soil bulk density was determined by drying a known volume of soil collected in May 2014 at 105 °C to constant weight. Soil texture analysis of composite samples collected in May 2011, was performed by destruction of the organic matter with hydrogen peroxide, dispersion of soil and clay aggregates into discrete units, and separation of particles of different sizes by sieving and sedimentation (DIN ISO 11277 2002; Solly et al. 2014). Soil oxalate extractable Al (in the composite samples collected in May 2011) was determined with ICP-OES (Optima 3300 DV, PerkinElmer) as a measure for non and poorly crystalline oxi-hydroxides. In soil samples collected in May 2014, we extracted NaHCO<sub>3</sub>-P and NaOH-P with subsequent extractions and we determined inorganic P with a continuous flow analyser (CFA, AA3, XY2, Seal-Analytic, Norderstedt, Germany). Labile and moderately labile inorganic P (Plabile.inorg) was calculated as their sum (Sorkau et al. 2018).

#### Microbial soil properties

Aliquots of sieved soil (from the composite samples collected in May 2017) were used for microbial analysis. We determined the microbial C and N concentration with the chloroform-fumigation-extraction method (CFE), according to Vance et al. (1987) and Keil et al. (2011). Fumigated sample replicates were incubated with CHCl<sub>3</sub> for 24 h. C and N were extracted from each fumigated and non-fumigated replicate (5 g) with 40 ml 0.5 M K<sub>2</sub>SO<sub>4</sub>. The suspension was horizontally shaken (30 min, 150 rpm) and centrifuged (30 min, 4400×g). C and N concentrations in diluted extracts (1:4, extract: deionized H<sub>2</sub>O) were measured with a TOC/TN analyser (Multi N/C 2100S, Analytik Jena AG, Jena, Germany). No correction factors accounting for the extractable fraction of microbial C and N were used to calculate microbial biomass.

### Plant properties

Plant communities surveyed May were in 2018. Standing biomass was measured in eight  $0.5 \text{ m} \times 0.5 \text{ m}$  subplots in each grassland, which were fenced to prevent livestock grazing or mowing prior to harvest. Aboveground vegetation was clipped at 2 cm height and dried at 80 °C for 48 h. Dry mass was then weighed at one decimal of the gram, and ground for chemical analysis. In a nearby 4 m×4 m subplot, we recorded all vascular plant species and estimated their percentage cover. Afterwards, we calculated the plant species richness, and the cover of two main functional groups, legumes, which typically have nitrogen rich tissues, and grasses. We determined the concentrations of C, N and P in aboveground plant biomass with a Near-Infrared Reflectance spectrometer (SpectraStar 2400, Unity Scientific, Columbia, MD, USA) (Busch et al. 2018; Kleinebecker et al. 2011). The reflectance spectrum of each sample, averaged over 24 scans, was recorded from 1250 to 2350 nm with 1 nm intervals.

We used plant species identity, and their abundances in our grasslands, to calculate the Community Weighted Mean (CWM) traits, for four above-(specific leaf area, SLA; the leaf dry matter content, LDMC; leaf N and P) and three belowground (fine root diameter, fineRD; root tissue density, RTD; specific root length, SRL) traits (Table 2). Aboveground traits were extracted from the TRY database (Kattge et al. 2020), and belowground traits were measured in a greenhouse experiment on plant species from the Biodiversity Exploratories grasslands (Lachaise et al. 2021). First, we matched the species identities found in our grassland sites to the TRY database (including synonyms) and excluded (i) duplicates, (ii) non-mature plants, (iii) non-healthy or dead plants, (iv) measurement contributions from experimental manipulations, and (v) measurement contributions where mean data were not reported. Then, for each species-trait combination, we averaged the trait value for each contributing dataset and excluded outlying datasets. We calculated average trait values for each species over all contributing datasets, weighted by the number of replicates in each dataset. Finally, the community weighted mean was calculated as the average trait values of all species, weighted by their relative abundance.

In our grasslands, the cover of legume species was negatively correlated with the CWM leaf dry matter content and specific root length, and, to some extent, it was positively correlated with leaf N and fine root diameter (Fig. 1). Moreover, the cover of grass species was negatively correlated with fine root diameter and root tissue density, and positively correlated with specific root length. Specific leaf area was not related to the plant functional group of legumes or grasses. Grazing intensity was negatively correlated with specific root length and leaf dry matter content, and was positively related to leaf N (and P). Fertilization was positively correlated with specific leaf area, and, to some extent, negatively related to leaf dry matter content and root tissue density.

#### Annual nutrient leaching

Annual, downward fluxes of nitrate, ammonium, phosphate and sulphate with the percolating water, i.e. leaching, were measured with the Self-Integrating Accumulators (SIAs) provided by the TerrAquat GmbH (Nürtingen, Germany, www.terraquat-gmbh. com; Bischoff, 2009). SIAs consisted of a PVC cylindrical cartridge (diameter and height: 10 cm) with a net at the lower end and were filled with a mixture of cation and anion exchange resins with a silica sand of specified grain size distribution.

SIAs were installed below the upper 10 cm of soil, and we measured the leaching from the upper



Fig. 1 Principal component analysis among seven plant (community weighted mean) traits (listed in Table 2), two plant functional groups (i.e. cumulative cover of Legumes and Grasses) and management intensities (Grazing and Fertilization) for the three study regions; Schwäbische-Alb (ALB), Hainich-Dün (HAI) and Schorfheide-Chorin (SCH). Abbreviations: fineRD=fine root diameter, LDMC=leaf dry matter content, Leaf N (or P)=leaf nitrogen (or phosphorus) concentration, RTD=root tissue density and SLA=specific leaf area, SRL=specific root length

10 cm of soil for three reasons. Firstly, we wanted our leaching data to match the soil and microbial properties data, which were also measured at the upper 10 cm of the soil (Table 2). Secondly, we wanted to follow a standardized methodology in every grassland, because the total soil depth varied among and within the three study regions (Table 2), and it was often  $\leq 20$  cm deep (in 58 out of the 150 grasslands). Finally, we were interested in biotic drivers of nutrient leaching, which are probably easier identified below the biologically most active topsoil layer than in the deeper subsoil, where geochemical drivers might become more important. Since we measured nutrient leaching from the upper 10 cm of soil, and not below the rooting zone, we cannot conclude on nutrient losses from the ecosystem, but our data indicate their loss risk.

We installed four SIAs in each of the 150 grasslands (total of 600 SIAs) in spring 2018, a year in which a summer drought occurred in central Europe (Apostolakis 2022). We selected four 1 m×1 m subplots at a 2 m extension of the cardinal transects used for soil sampling. In each subplot, we opened a 40 cm×40 cm pit down to 30 cm. We selected the wall that was against the slope and, on this wall, we opened a lateral cubic tunnel (side of 15 cm) at a depth of 10 to 25 cm. We lined the floor of the tunnel with silica sand. SIAs were saturated with deionized water from below and then placed in the lateral tunnels, so that SIAs were installed under undisturbed soil. The empty space around and above the SIAs was filled with silica sand, and the pit was re-filled with the removed soil. The silica sand ensured that SIAs were not in direct contact with soil impeding, thus, direct exchange reactions between soils and resins and, at the same time, it allow the undisturbed penetration of water.

SIAs stayed in the soil for one year, until spring 2019. For the installation and removal, we visited the study regions in the same order. Installation and removal dates were recorded together with field observations (e.g. presence of roots and soil fauna) for a quality control of the samples. After removal, SIAs were stored at 4 °C, and each SIA was separated to three resin samples by separating the resin column into three resin layers. The first resin layer accounted for nutrient leaching, and had a depth of 5 cm. The second resin layer was a control, and the third layer served as a buffer for upward nutrient movement, and they had a length of 1 and 4 cm, respectively. Each resin sample was mixed to ensure homogeneity, weighed, and stored at 4 °C. To calculate leaching, we analysed the first two layers for their nutrient concentrations. If the second layer was contaminated with NO<sub>3</sub> (>10% of first layer NO<sub>3</sub>-N concentration), then the SIA was excluded from our dataset. Out of the 600 SIAs, five SIAs were excluded due to disturbances by roots and/or soil fauna, and 34 SIAs due to a contaminated second layer.

Resin samples of 15 g were weighed. After adding 100 mL of 1 M NaCl (Natriumchlorid zur Analyse Acs, Merck KGaA, Darmstadt, Germany) the resin was shaken for 45 min. The extracted solution was filtered by gravity using N-free filters (MN 619 DE 1/4, Macherey–Nagel Gmbh & Co, Germany), and the filtrate was transferred to a 15-mL falcon tube, which was stored at -20 °C. Ammonium (NH<sub>4</sub>-N), nitrite (NO<sub>2</sub>-N) and nitrite and nitrate (NO<sub>2</sub>-N+NO<sub>3</sub>-N) and phosphate (PO<sub>4</sub>-P) concentrations in the solution were determined by a flow-injection analyser (Quickchem QC85S5, Lachat Instruments, Hach Company, Dreieich, Germany) after applying appropriate dilutions. Nitrate concentration was calculated as the difference of [NO<sub>2</sub>-N+NO<sub>3</sub>-N] from NO<sub>2</sub>-N

concentration. Sulphate  $(SO_4-S)$  concentration was determined by an ion chromatography system (Dionex-DX 500, Thermo Fisher Scientific GmbH, Dreieich, Germany), after applying appropriate dilutions. Nutrient concentrations of the first and second layer were used to calculate leaching fluxes.

SIAs might underestimate the effects of grazing intensity on nutrient leaching compared to the effects of fertilization intensity due to the higher spatial heterogeneity of nutrient inputs through dung and urine patches compared to the more homogeneous nutrient inputs through fertilization. However, i) the high number of grazed sites included in our study (118 out of 150 permanent grasslands), ii) the use of average management history (fertilization and grazing intensities averaged over 2015–2018) instead of management applied only in 2018, and iii) the determination of leaching as an integrating measurement over a year, support the overall representation of management effects on nutrient leaching risk.

# Data analysis

We tested for differences of annual nutrient leaching in the three study regions using analysis of variance (ANOVA) models and the Tukey's (honest significant difference) test. We use a backward stepwise function to assess the impact of (i) study regions, (ii) plant properties (aboveground biomass, species richness, legume cover and grass cover), (iii) soil OC, (iv) microbial biomass C and (v) grassland management (fertilization and grazing intensities) on nutrient leaching based on the Akaike Information Criterion. Using the variables selected by the stepwise function, we modelled nutrient leaching with analysis of covariance (ANCOVA). ANCOVA models were performed using a type II sum of squares, which is not influenced by the order in which the explanatory variables are introduced in the model (Zuur et al. 2009). For every model, we used diagnostic plots to test the assumptions of linearity, normality and homoscedacity of variance in the residuals, and to check for outliers. Heteroscedasticity was tackled with logarithmic or square-root transformations. Collinearities between explanatory variables were assessed with the variance inflation factor and values < 3.0 were accepted. We followed the same approach to separately investigate the importance of several physico-chemical soil properties (Table S3) and CWM traits on nutrient leaching (Table S4) listed in Table 2.

We used Structural Equation Modeling (Shipley, 2016) to investigate how fertilization and grazing, and their impacts on soils, plants and soil microorganisms, affect nutrient leaching. Grazing and fertilization intensities were included as exogenous variables. Plant species richness is an important predictor of soil functions (Chen et al. 2019; Leimer et al. 2016) and sensitive to management intensification (Midolo et al. 2019), and it was included as an endogenous variable. Substrate stoichiometry controls the cycling of C and nutrients in soils (Manzoni et al. 2008; Schleuss et al. 2021), and so we used stoichiometric ratios (C:N, C:P or C:S) of soils, plants and soil microbial biomass as endogenous variables in our models. We removed the effects of study regions (with one-way ANOVA models) from all endogenous variables, and we used their residuals in our Structural Equation Models. Proposed causal structures with p-values  $\geq 0.05$  were accepted. In addition, we calculated the root mean square errors of approximation (RMSEAs) and the standardized root mean square residuals (SRMRs), which both should be  $\leq 0.05$ . The high number of observations compared with the number of estimated parameters in our models provided safety against multivariate non-normality issues. As an additional measure, we re-ran our models using bootstrapping. The p-values of bootstrapped models (not shown) were always similar to those of the classic models. We followed the same approach to separately investigate the importance element pools (C, N, P or S, Fig. S1) and plant CWM traits (selected based on Table S4) on nutrient leaching.

Structural Equation Modeling is among the few (semi)statistical methods claiming to prove, or disprove, causal relationships in a proposed structure. Natural ecosystems are characterized by complex interactions between processes involved in element cycling. This makes the development of a unique causal structure challenging. We recognise that results from our Structural Equation Models represent associations between variables, and not necessarily causal relationships. However, we interpret these associations as evidence of management effects and, for simplicity, we often use terms such as 'effects' and 'drivers' hereafter.

Statistical analysis was performed with the R statistical software (R Core Team 2019). Figures were designed with the *ggplot2* package. ANOVA and ANCOVA models were performed with the *lm* function from *stats* package, and the backward stepwise elimination with the *step* function. The variance inflation factor was calculated with the *vif* function of the *faraway* package (Faraway, 2016). Tukey's test and PCA were performed with the *TukeyHSD* and the *prcomp* functions, respectively, of the *stats* package. The Structural Equation Modeling was performed with the *sem* function from the *lavaan* package (Rosseel, 2012).

# Results

Annual nutrient leaching from the topsoil in grassland

The median nitrate (annual) leaching in a depth of 10 cm was 7.4 kg NO<sub>3</sub>-N ha<sup>-1</sup> (interquartile range: 2.2–22.2 kg NO<sub>3</sub>-N ha<sup>-1</sup>, Table S1) over the three study regions and increased in the order ALB < SCH = HAI (Fig. 2a). The median leaching of the less mobile ammonium was 4.4 kg NH<sub>4</sub>-N ha<sup>-1</sup> (interquartile range: 3.8-6.0 kg NH<sub>4</sub>-N ha<sup>-1</sup>) and increased in the order HAI < ALB = SCH (Fig. 2b). The cumulative inorganic N (NH<sub>4</sub>-N+NO<sub>3</sub>-N) leaching followed the patterns of nitrate leaching and increased in the order ALB < SCH = HAI, underlining the importance of nitrate for total N leaching losses. Interestingly, nitrate and ammonium leaching show comparable fluxes despite that nitrate is more mobile than ammonium. This might be due to the relatively shallow depth of our measurements since, within the rooting zone, nutrient inputs (e.g. from fertilization or decomposition) might be more important for leaching than nutrient mobility. The three study sites differ in atmospheric N inputs, which can be relevant for inorganic N leaching since, for instance, the annual bulk deposition of total N in SCH ( $11.8 \pm 1.7$  kg N ha<sup>-1</sup>, Schwarz et al. 2014) is nearly four times higher than the annual fertilization inputs (Table 2).

The median phosphate leaching flux was 0.8 kg  $PO_4$ -P ha<sup>-1</sup> (interquartile range: 0.6–1.7 kg  $PO_4$ -P ha<sup>-1</sup>, Table S1) over the three study regions and there were no detectable differences between the regions (Fig. 2c). Lastly, the median sulphate leaching flux was 21.8 kg  $SO_4$ -S ha<sup>-1</sup> (interquartile range: Fig. 2 Annual leaching of (a) nitrate (NO<sub>3</sub>-N), (b) ammonium ( $NH_4$ -N), (c) phosphate ( $PO_4$ -P) and (d) sulphate  $(SO_4-S)$  in a depth of 10 cm for the three study regions; Schwäbische-Alb (ALB), Hainich-Dün (HAI) and Schorfheide-Chorin (SCH). Bars show average values and error bars represent standard deviations. Numbers at the base of the bars show the number of observations. Lower-case letters indicate significant differences (p-value < 0.05) among the study regions based on ANOVA and Tukey's HSD models



13.7–29.5 kg SO<sub>4</sub>-S ha<sup>-1</sup>), and leaching in SCH was significantly higher than ALB (Fig. 2d), but not than HAI. The regional patterns of nutrient leaching differed from the patterns of soil N,  $P_{labile,inorg}$  and total S concentrations for nitrate, phosphate and sulphate leaching, while ammonium leaching differences between the study regions matched those of soil N concentrations (Table 2).

Management effects on nutrient leaching

Including management intensities and selected soil, plant and microbial properties in our ANCOVA models for nitrate, ammonium and phosphate annual leaching (Table 3) considerably increased the explained variance compared to models including only soil properties (Table S3). Our models explained

Annual leaching	NO <sub>3</sub> -N		NH <sub>4</sub> -N	
	df = 138	Model p	df = 135	Model p
	$R^2 = 0.39$	< 0.001	$R^2 = 0.27$	< 0.001
Parameter	<i>t</i> -value	<i>p</i> -value	<i>t</i> -value	<i>p</i> -value
Intercept	3.51	< 0.001	2.96	0.004
Region HAI	6.64	< 0.001	- 3.14	0.002
Region SCH	3.51	< 0.001	- 1.11	0.270
Fertilization	2.54	0.012	2.74	0.007
Grazing	2.21	0.029	-	_
Plant abov. biomass	_	-	3.19	0.002
Plant species richness	- 2.89	0.004	-	_
Legume cover	1.83	0.069	-	_
Grass cover	- 3.55	< 0.001	-	_
Microbial C	1.38	0.170	-2.49	0.014
Soil OC	1.50	0.137	1.99	0.048
Annual leaching	PO <sub>4</sub> -P		SO <sub>4</sub> -S	
	df = 142	Model p	df = 142	Model p
	$R^2 = 0.23$	< 0.001	$R^2 = 0.38$	< 0.001
Parameter	<i>t</i> -value	<i>p</i> -value	<i>t</i> -value	<i>p</i> -value
Intercept	0.89	0.374	6.29	< 0.001
Region HAI	- 1.30	0.195	5.16	< 0.001
Region SCH	- 3.60	< 0.001	3.46	< 0.001
Fertilization	2.31	0.022	-	_
Grazing	2.57	0.011	-	_
Plant abov. biomass	3.06	0.003	- 2.29	0.024
Plant species richness	-	-	-	-
Legume cover	-	-	- 1.63	0.105
Grass cover	-	-	1.69	0.094
Microbial C	- 2.20	0.030	6.46	< 0.001
Soil OC	_	_	_	_

Table 3 ANCOVA models (SS type II) of nitrate, ammonium, phosphate and sulphate annual leaching with management intensities and plant, microbial and soil properties. Full models were stepwise reduced.

Study region effects are given relative to the Schwäbische-Alb region. Significant predictors are given in bold

HAI Hainich-Dün; SCH Schorfheide-chorin; Microbial C microbial carbon concentration; Soil OC soil organic carbon concentration

39, 27, 23 and 38% of the variance in nitrate, ammonium, phosphate and sulphate leaching, respectively (Table 3). Study regions were important predictors of leaching for every nutrient. Fertilization intensity increased nitrate, ammonium and phosphate annual leaching, while grazing had an increasing effect only on nitrate and phosphate leaching. Management was not important for sulphate leaching. Plant aboveground biomass was positively related to ammonium and phosphate leaching, but it was negatively linked to sulphate leaching. Plant species richness was negatively associated with nitrate leaching, and it was not important for the other nutrients. The relative cover of grasses was negatively related to nitrate leaching, and that of legumes was marginally (and positively) related to nitrate leaching. Soil OC was positively related to ammonium leaching. Finally, microbial biomass C was negatively associated with ammonium and phosphate leaching, but it was positively associated with sulphate leaching. According to our structural equation models, fertilization intensity had (direct) positive effects on nitrate, ammonium and phosphate leaching risk, but not on sulphate, and these effects were not explained by plant species richness or soil, plant and microbial stoichiometry (Fig. 3). Similarly, grazing intensity had direct positive effects on nitrate and phosphate leaching, but not on ammonium or sulphate. Plant species richness was directly associated with ammonium and sulphate leaching, and these relationships were negative and positive, respectively. In contrast to our ANCOVA model (Table 3), plant species richness was not directly related to nitrate leaching (Fig. 3a).

Interestingly, soil, plant and microbial C:N ratios were not associated with fertilization intensity directly, but rather indirectly through plant species losses caused by fertilization. In contrast, grazing intensity was not related to C:N ratios directly or indirectly. A higher plant species richness was associated with higher C:N ratios of soils, plants and microbial biomass. Both plant and microbial C:N ratios had a negative relationship with nitrate leaching, while soil and plant C:N ratios had a positive relationship with ammonium leaching. In a similar model with N pools instead of C:N ratios, microbial and plant N were



**Fig. 3** Structural Equation Models for (a) nitrate ( $NO_3$ -N), (b) ammonium ( $NH_4$ -N), (c) phosphate ( $PO_4$ -P) and (d) sulphate ( $SO_4$ -S) annual leaching explained by fertilization and grazing intensities, plant species richness and stoichiometry of soils, plants and microbial biomass. Single-headed arrows indicate direct causal paths and double-headed arrows indicate covari-

ances. Solid, blue arrows indicate positive effects and dashed, red arrows indicate negative effects. Grey arrows indicate paths of marginal significance. Standardized path coefficients are given together with their significance level (\*p<0.050; \*\*p<0.010; \*\*\*p<0.001). Coefficients of determination are given for the response variables

positively linked to nitrate leaching, and plant N was negatively linked to ammonium (Fig. S1a, b).

Grazing and fertilization intensities were negatively associated with C:P ratios of plants and soils and with microbial biomass C (no microbial P data available, Fig. 3c). Apart from management, soil C:P ratio was the only parameter related to phosphate leaching, and this relationship was negative. In a similar model with P pools instead of C:P ratios, soil  $P_{labile,inorg}$  was positively related to phosphate leaching, while plant P and microbial C were not important (Fig. S1c).

For S, no information on plants or microbes was available. Microbial C and plant species richness were (directly) positively related to sulphate leaching (Fig. 3d). However, plant species richness was negatively linked to microbial C, and, therefore, its effect on sulphate leaching was offset. Soil C:S ratio was not associated with fertilization or grazing intensities, and it was only marginally associated with sulphate leaching. In a similar model with soil S concentration instead of C:S ratio, soil S was positively related to sulphate leaching, and there were no other effects from management, plants or microbial biomass on sulphate leaching (Fig. S1d). The net effects of fertilization and grazing on sulphate leaching were negative and rather small (Table S5).

Effects of plant community traits on nutrient leaching

Based on backwards stepwise analyses and ANCOVA models, we identified CWM plant traits that were important for nutrient leaching (Table S4). Each nutrient had a different set of important CWM plant traits, while for sulphate leaching none of the traits was important. Our Structural Equation Models show that fertilization and grazing intensities exhibited multiple indirect effects on nutrient leaching via CWM plant traits, in addition to their direct effects (Fig. 4). Interestingly, while plant species richness was only affected by fertilization, CWM plant traits were, typically, influenced by both grazing and fertilization, though grazing acted more strongly on aboveground traits (with the exception of specific root length) than fertilization. Fertilization and grazing intensification seem to have shifted plant communities towards more resource acquisitive, fast-growing species with higher leaf N and P, and lower leaf dry matter content and root tissue density (Fig. 1). However, grazing was also negatively associated with specific root length.

Fine root diameter directly increased, and leaf dry matter content directly decreased, nitrate leaching (Fig. 4a). Microbial N mediated positive indirect effects from fine root diameter and leaf dry matter content on nitrate leaching. Interestingly, plant species richness directly decreased nitrate leaching, even after including selected CWM plant traits and microbial N in our model. In contrast, plant and microbial C:N ratios (from field measurements) fully mediated the effects of plant species richness on nitrate leaching (Fig. 3a), but its net effect was negative in both models (Table S5).

Root tissue density was marginally (and positively) linked to ammonium leaching, and leaf dry matter content was marginally (and negatively) linked to it (Fig. 4b, Table S4). Plant species richness was again negatively associated with ammonium leaching (as in Fig. 3b). Leaf P increased phosphate leaching, and it mediated indirect positive effects from both fertilization and grazing intensities on phosphate leaching (Fig. 4c). Specific root length was positively linked to sulphate leaching, and it mediated an indirect negative effect from grazing intensity on sulphate leaching (Fig. 4d). Finally, leaf N marginally (and negatively) related to sulphate leaching, and it mediated indirect negative effects by both fertilization and grazing intensities.

## Discussion

Intensively managed grasslands have an elevated nutrient leaching risk

Previous studies stressed the importance of grazing, and the associated dung and urine patches, for nitrate leaching (Ledgard et al. 2009), and they associated (grazed) pastures with higher nitrate leaching risk than (mowed) meadows (Cameron et al. 2013; Ryden et al. 1984). In this study, nitrate leaching risk significantly increased with both fertilization and grazing intensity (Table 3, Fig. 3). We observed that fertilized grasslands also have an elevated nitrate leaching risk, and we extended previous findings from mostly short-term observations (Klaus et al. 2018) to an annual timescale. In addition, ammonium annual leaching



**Fig. 4** Structural Equation Models for (a) nitrate (NO<sub>3</sub>-N), (b) ammonium (NH<sub>4</sub>-N), (c) phosphate (PO<sub>4</sub>-P) and (d) sulphate (SO<sub>4</sub>-S) annual leaching explained by fertilization and grazing intensities, plant species richness and selected CWM plant traits (LDMC: leaf dry matter content; Fine root diam.: fine root diameter; Root tis. dens.: root tissue density; Leaf P or N: leaf phosphorus or nitrogen; Specific root length). Single-

risk increased with fertilization intensity, but not with grazing (Table 3). This is possibly because grazing increases N inputs as urea, which is quickly oxidized to nitrate (in various soil types, Soulides and Clark, 1958; Tzanakakis et al. 2018), especially since management intensification increases nitrification rates (Stempfhuber et al. 2014). Considering both nitrate and ammonium, we conclude that total inorganic N leaching risk increases both with grazing and fertilization intensification across an annual timescale.

Agricultural activities are responsible for excess P losses via leaching, which contribute to the degradation of (surface) aquatic ecosystems

headed arrows indicate direct causal paths and double-headed arrows indicate covariances. Solid, blue arrows indicate positive effects and dashed, red arrows indicate negative effects. Grey arrows indicate paths of marginal significance. Standardized path coefficients are given together with their significance level (\*p < 0.050; \*\*p < 0.010; \*\*\*p < 0.001). Coefficients of determination are given for the response variables

due to eutrophication (SOER 2020). We found that managed grasslands can also have a high P leaching risk, since annual phosphate leaching at 10 cm soil depth were frequently greater than 0.8 kg  $PO_4$ -P ha<sup>-1</sup> (overall median) (Table S1, Fig. 2). Such fluxes are higher than leaching losses from the subsoil of other agricultural land-uses, like croplands (Leinweber et al. 1999). However, P leached from the topsoil, as measured in our study, can still be fixed in the subsoil, or be taken up by plants (Kuhlmann and Baumgärtel, 1991), and so it may not reach aquatic ecosystems. Therefore, our study can only identify an increased nutrient loss risk due to management intensification. However, the P sorption capacity in subsoils depends on soil texture and pH, and it could, at some point, be exceeded with long term excess P supply (Gocke et al. 2021; Werner et al. 2006), especially in sandy soils, such as those in SCH (Table 2). Overall, we found that phosphate leaching significantly increased with management intensity (Table 3), and that fertilization and grazing equally increased phosphate loss risk.

Compared to N and P, little is known for S cycling in temperate managed grasslands and, usually, S leaching is not measured as an actual flux but it is, instead, estimated based on soil solution S concentrations and water balance modeling. In this study, we measure sulphate leaching at 10 cm soil depth and the overall median was 21.8 kg  $SO_4$ -S ha<sup>-1</sup> (interquartile range: 13.7–29.5 kg  $SO_4$ -S ha<sup>-1</sup>, Table S1), which agrees with reported values from grassland and cropland subsoils (4–45 kg SO<sub>4</sub>-S ha<sup>-1</sup>; Erisken et al. 2000; Gallejones et al. 2012). Due to ample atmospheric S depositions in the past, temperate soils are typically considered to be S-rich. However, concerns that S might be a limiting resource for plant production and quality have been raised (Eriksen, 2009; Mathot et al. 2008). In this study, sulphate was the only nutrient for which management intensity reduced its leaching, and this operated indirectly (Fig. S1d). Both grazing and fertilization reduced soil S concentrations and, thus, sulphate leaching. A possible explanation is that management intensification increases plant productivity, which creates higher S demands and plant uptake (Eriksen 2009), eventually leading to increased export of S from the ecosystem with harvest or grazing. This is supported by a negative relationship between aboveground plant biomass and sulphate leaching in our grasslands (Table 3). Another explanation for management-dependent reductions in soil S is the desorption of sulphate from soil minerals due to phosphate additions, since phosphate is adsorbed more strongly than sulphate (Bolan et al. 1988). However, this process should have led to increased sulphate leaching risk with increased fertilization, which we did not observe. The lack of information on P and S fertilization rates might obscure additional effects from fertilization on sulphate leaching.

Fertilization, but not grazing, alters stoichiometry towards increased nutrient leaching risk

The positive effect of plant N on nitrate leaching (Fig. S1a) indicates faster mineralization of N-rich litter under high management intensity. Meanwhile, the negative effect of plant N on ammonium leaching (Fig. S1b) could be similarly explained by higher nitrification rates in intensively managed grasslands (Stempfhuber et al. 2014) or, alternatively, by a preference for ammonium uptake by plants. In contrast to plant N, plant biomass, which increases with management intensification (Klaus et al. 2018), was positively related to ammonium leaching (Table 3). This could be explained by increased above- and below-ground litter inputs and, thus, increased soil organic N (Leimer et al. 2016), which is then mineralized to ammonium.

In contrast to ammonium, nitrate leaching risk was also controlled by microbial properties. Microbial nitrification, following soil organic N mineralization, could explain the positive relationship between microbial biomass N and nitrate leaching. Management intensity, and especially fertilization, was shown to increase the nitrification potential of the studied soils (Meyer et al. 2013; Stempfhuber et al. 2014). Therefore, fast nitrification of ammonium at fertilized sites with low C:N ratios could be the reason for the observed reduction of ammonium leaching with increasing microbial biomass C (Table 3) and for the observed increase of ammonium leaching risk with increasing soil C:N ratio (Fig. 3b).

Interestingly, microbial C:N ratios were negatively linked with nitrate leaching risk (Fig. 3a), indicating that leaching was not only affected by changes in plant communities, but also microbial communities (Fig. S2). This may be because the less nutrientdemanding, more fungal-dominated communities at extensively managed, botanically diverse grasslands capture N more effectively and prevent leaching (de Vries et al. 2006). Alternatively, it may be that the turnover and decomposition of fungal-derived, N-poor necromass is low. In contrast to the latter, evidence from short-term observations suggests that microbial communities which are not N-limited, increase N retention in the plant-soil system (de Vries and Bardgett 2016). To our knowledge, our work is among the few studies showing such a clear relationship between microbial biomass and annual nitrate leaching in managed grasslands under field conditions.

Microbial biomass N had a positive effect on nitrate leaching risk (Fig. S1a). In experimental grasslands, the negative effect of microbial biomass on the leaching of dissolved organic N has been attributed to enhanced mineralization (Leimer et al. 2016), which could explain higher soil nitrate concentrations and, thus, leaching. Previous work has shown that N additions can increase or decrease microbial biomass (Dietrich et al. 2017; Lu et al. 2011). Soil microbial communities are also influenced by plant communities, mainly through changes in N availability (Moreau et al. 2015), and perhaps to a greater extent than by management intensity in our study sites (Boeddinghaus et al. 2019). In our study, management intensity directly reduced microbial biomass N (and C), but the net effect was still close to neutral, because the fertilization-induced reduction in species richness increased it again (Fig. S1a). This suggests that relations between long-term fertilizer additions and microbial biomass in managed grasslands are modified by plant communities and that the observed responses of microbial activity to short-term fertilizer additions to soils in the laboratory, where plant communities are absent, might not be representative of field conditions.

Phosphate leaching risk was mainly associated with soil  $P_{labile,inorg}$  and C:P ratio, which were both positively related to plant P and C:P ratios (Fig. 3c, S1c). Accordingly, fertilization-induced reductions in plant species richness were positively associated with soil  $P_{labile,inorg}$  and, thus, with phosphate leaching risk. In contrast to inorganic N, phosphate leaching was negatively related to soil pH (Table S3), indicating high P immobilization as calcium phosphate in alkaline soils.

Exploitative plant communities increase nutrient leaching risk in intensively managed grasslands

We observed that the functional group of grasses was negatively associated with annual nitrate leaching risk, while the group of legumes was positively (but marginally) associated with it (Table 3). Neither of them was as important for ammonium, phosphate and sulphate leaching risk. A positive relationship between legumes and nitrate leaching can be explained by increased N availability caused by their N-fixing ability and N-rich tissues (Leimer et al. 2015). Grasses decrease nitrate leaching due to increased N acquisition and/or more efficient use of N and water (de Vries and Bardgett 2016; Leimer et al. 2016) and due to their, typically, lower tissue N content. Indeed, we observed that legume cover was positively correlated with CWM leaf N, and that grass cover was negatively correlated with CWM fine root diameter (Fig. 1), which could, at least partly, explain the functional group effects on nutrient leaching.

Plant traits give hints on the importance of plant nutrient acquisition for nutrient leaching. In accordance with the observation that phosphate and sulphate leaching risk were more related to soil properties than inorganic N (Table S3), plant traits were less important for them, too (Table S4). We identified the CWM leaf dry matter content as an important predictor of inorganic N leaching (Fig. 4), reducing both, nitrate and ammonium leaching risk. This effect could be explained by a slower and more efficient N cycling in conservative than exploitative plant communities (Kleinebecker et al. 2014), and/or by lower N inputs in the soil system since leaf dry matter content was negatively related to i) fertilization, ii) legume cover and iii) CWM leaf N (Fig. 1). The positive relationship between CWM fine root diameter and nitrate leaching risk could be explained by a reduced nutrient uptake efficiency with coarser roots. For ammonium, leaching risk was related to root tissue density (Fig. 4b), where a higher root tissue density would indicate coarser fine roots (Fig. 1) and a reduced nutrient uptake potential.

To date, it is not clear whether conservative, slow-growing plant communities, or exploitative, fast-growing plant communities lead to a greater N retention in the soil, plant and soil microorganisms' system. For instance, conservative plant communities can lead to decreased N retention due to low N acquisition rates (de Vries and Bardgett, 2016), while exploitative plant communities can lead to decreased N retention due to enhanced N cycling and high N availability (de Vries et al. 2012; Grigulus et al. 2013; Robson et al. 2010). We observed a differentiation between the effects of above- and belowground traits on inorganic N leaching risk, similar to findings on N-related ecosystem functions by de Vries and Bardgett (2016). Conservative aboveground plant traits reduced inorganic N leaching risk, while conservative belowground traits increased it (Fig. 4). Regarding our third hypothesis, we suggest that management intensity, and especially fertilization, shifted mostly the aboveground plant traits towards values more representative of exploitative communities (Fig. 1; Allan et al. 2015; Busch et al. 2018; Zobel et al. 2007), which are characterized by a fast tissue turnover and decomposition (Lavorel et al. 2011) and, thus, influenced their association with annual N leaching risk.

## Plant diversity reduces inorganic N leaching risk

Previous studies have reported a negative relationship between plant diversity and nitrate availability or leaching in managed and experimental grasslands (Leimer et al. 2016; Niklaus et al. 2001), but effects on ammonium leaching are less often described (Leimer et al. 2015). Here, plant species richness was negatively associated both with nitrate and ammonium leaching risk, even after considering fertilization intensity (Table 3, Fig. 3b). Interestingly, the negative effect of plant species richness on nitrate leaching was completely mediated by higher plant and microbial C:N ratios (and lower N concentration) in more botanically diverse grasslands (Fig. 3a, S1a), while aboveground plant biomass was not selected as a predictor for nitrate leaching risk. In contrast to nitrate, the negative effect of plant species richness on ammonium leaching was not completely mediated by stoichiometry, element pools or CWM plant traits (Fig. 3b, S1b, 4b). However, it could be explained by the negative relationship between plant (aboveground) biomass and species richness in managed grasslands (Klaus et al. 2018) since plant biomass was positively associated with ammonium leaching (Table 3). Alternatively, it could be explained by niche complementarity and ammonium uptake by plants in species-rich grasslands, as shown for plant species identity and functional groups (Kahmen et al. 2006). In addition, the negative relationship between plant species richness and plant N led to a positive indirect effect on ammonium leaching risk (Fig. S1b). The negative relationship between plant species richness and plant N could be explained by fertilization that simultaneously causes plant biomass N enrichment and plant species losses, and/or by an increased plant N use efficiency with increasing plant diversity in grasslands (Dias et al. 2010; Fargione et al. 2007). To conclude, in terms of net effects, plant species richness was negatively associated both with nitrate and ammonium leaching risk (Table S5) and most of its effects on inorganic N leaching were mediated by the stoichiometry of soils, plants and soil microorganisms, as well as several CWM plant traits.

# Conclusions

By applying a systems perspective, we identified that management intensity, and especially fertilization, not only promote nutrient leaching risk directly, but also indirectly by changing the properties of soils, plants and microbes. Most of the indirect effects of management on nutrient leaching were mediated by fertilization-induced reductions in plant species richness. While ceasing fertilization will reduce its direct effect on N and P leaching over a relatively short period in time, some of these changes, such as a loss in plant diversity and functional changes in plant communities, are not immediately reversible. Thus, management intensification changes the risk of nutrient leaching on the long-term and active (restoration) measures to maintain diverse plant communities are needed, which might, in return, mitigate the increased leaching risk that management intensification imposes on grasslands. Finally, we showed that, when considering direct and indirect effects, both fertilization and grazing lead to higher nitrate and phosphate leaching risk in the topsoil, while only fertilization intensity associates with higher ammonium leaching risk. As such, shifting grassland management towards (mowed) meadows instead of (grazed) pastures should not be considered as a management strategy for N leaching mitigation, as long as meadows are fertilized.

Acknowledgements The authors thank Theresa Kloetzing, Steffen Ferber, Marco Poehlmann, for organizing and conducting lab and field work relating to soil properties. The authors thank Ines Hilke for conducting the analysis of nitrate, ammonium, phosphate and sulphate. The authors thank Martin Freitag and Verena Busch for organizing and conducting lab and field work relating to plant properties and Tom Lachaise for belowground plant traits. We thank the managers of the three Exploratories, Kirsten Reichel-Jung, Florian Staub, Juliane Vogt, Miriam Teuscher and all former managers for their work in maintaining the plot and project infrastructure; Christiane Fischer for giving support through the central office, Andreas Ostowski for managing the central database, and, Eduard Linsenmair, Dominik Hessenmöller, François Buscot, Ernst-Detlef Schulze, Wolfgang W. Weisser and the late Elisabeth Kalko for their role in setting up the Biodiversity Exploratories project.

We thank the administration of the Hainich national park, the UNESCO Biosphere Reserve Swabian Alb and the UNESCO Biosphere Reserve Schorfheide-Chorin as well as all land owners for the excellent collaboration. Field work permits were issued by the responsible state environmental offices of Baden-Württemberg, Thüringen, and Brandenburg. This work has been funded by the DFG Priority Program 1374 "Biodiversity-Exploratories'' (SCHR 1181/2-3) and the Max Planck Society.

Authors' contribution Antonios Apostolakis designed and carried out the field and lab work related to nutrient leaching, analysed and discussed the data, and was responsible for writing this manuscript. Ingo Schöning, Beate Michalzik and Marion Schrumpf designed the experiment, discussed the data, and reviewed drafts of the manuscript and IS carried out also part of the field work. Valentin H. Klaus and Peter Manning discussed the data and reviewed drafts of the manuscript. Wolf-Anno Bischoff and Andreas Schwarz provided expertise on the resin method used for the nutrient leaching measurements. Yvonne Oelmann and Elisabeth Schurig contributed soil (Olsen) P data. Runa S. Boeddinghaus, Ellen Kandeler and Sven Marhan contributed data related to microbial properties. Valentin H. Klaus, Ralph Bolliger, Norbert Hölzel, Till Kleinebecker, Daniel Prati and Markus Fischer contributed data related to plant properties. Margot Neyret, Peter Manning, and Mark van Kleunen contributed above- and belowground plant (community weighted mean) trait data. All authors contributed to the manuscript and approved the final version.

**Funding** Open Access funding enabled and organized by Projekt DEAL. This work has been funded by the DFG Priority Program 1374 "Biodiversity- Exploratories'' (SCHR 1181/2–3) and the Max Planck Society.

**Availability of data:** This work is based on data elaborated by several projects of the *Biodiversity Exploratories* project. Part of the data used are available at database of the project (https://www.bexis.uni-jena.de/ddm/publicsearch/index). However, to respect the rights of PhD students of individual projects, the remaining data will be made publicly available at the same website after an embargo period of 3 years, according to the data and publication policy of the *Biodiversity Exploratories* project.

#### Declarations

**Conflict of interest** The authors declare that they have no conflict of interest.

**Open Access** This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds

the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/.

## References

- Allan E, Manning P, Alt F, Binkenstein J, Blaser S, Blüthgen N, Böhm S, Grassein F, Hölzel N, Klaus VH, Kleinebecker T, Morris EK, Oelmann Y, Prati D, Renner SC, Rillig MC, Schaefer M, Schloter M, Schmitt B, Schöning I, Schrumpf M, Solly E, Sorkau E, Steckel J, Steffen-Dewenter I, Stempfhuber B, Tschapka M, Weiner CN, Weisser WW, Werner M, Westphal C, Wilcke W, Fischer M (2015) Land use intensification alters ecosystem multifunctionality via loss of biodiversity and changes to functional composition. Ecol Lett 18:834–843. https://doi. org/10.1111/ele.12469
- Apostolakis A (2022) Land use effects on carbon and nutrient fluxes in soils. PhD thesis. Friedrich-Schiller-Universität Jena
- Bardgett RD, Mommer L, De Vries FT (2014) Going underground: root traits as drivers of ecosystem processes. Trends Ecol Evol 29:692–699. https://doi.org/10.1016/j. tree.2014.10.006
- Bengtsson J, Bullock JM, Egoh B, Everson C, Everson T, O'Connor T, O'Farrell PJ, Smith HG, Lindborg R (2019) Grasslands—more important for ecosystem services than you might think. Ecosphere. https://doi.org/10.1002/ecs2. 2582
- Bergmann J, Weigelt A, Van Der Plas F, Laughlin DC, Kuyper TW, Guerrero-Ramirez N, Valverde-Barrantes OJ, Bruelheide H, Fresche GT, Iversen CM, Kattge J, McCormack ML, Meier IC, Rillig MC, Roumet C, Semchenko M, Sweeney CJ, Van Ruijven J, York LM, Mommer L (2020) The fungal collaboration gradient dominates the root economics space in plants. Sci Adv 6:1–10. https://doi.org/ 10.1126/sciadv.aba3756
- Bischoff W-A (2009) Development and applications of the self-integrating accumulators: a method to quantify the leaching losses of environmentally relevant substances. Hohenheimer Bodenkundliche Hefte, Heft. Kandeler E, Kuzyakov Y, Stahr K, Streck T, Kaupenjohann M (eds) University of Hohenheim, Stuttgart
- Blüthgen N, Dormann CF, Prati D, Klaus VH et al (2012) A quantitative index of land-use intensity in grasslands: integrating mowing, grazing and fertilization. Basic Appl Ecol 13:207–220. https://doi.org/10.1016/j.baae.2012.04. 001
- Boeddinghaus RS, Marhan S, Berner D, Boch S, Fischer M, Hölzel N, Kattge J, Klaus VH, Kleinebecker T, Oelmann Y, Prati D, Schäfer D, Schöning I, Schrumpf M, Sorkau E, Kandeler E, Manning P (2019) Plant functional trait shifts explain concurrent changes in the structure and function of grassland soil microbial communities. J Ecol 107:2197–2210. https://doi.org/10.1111/1365-2745.13182
- Bolan NS, Syers JK, Tillman RW, Scotter DR (1988) Effect of liming and phosphate additions on sulphate leaching in soils. J Soil Sci 39:493–504. https://doi.org/10.1111/j. 1365-2389.1988.tb01234.x

- Bruelheide H, Dengler J, Purschke O et al (2018) Global trait–environment relationships of plant communities. Nat Ecol Evol 2:1906–1917. https://doi.org/10.1038/ s41559-018-0699-8
- Busch V, Klaus VH, Penone C, Schäfer D, Boch S, Prati D, Müller J, Socher SA, Niinemets Ü, Peñuelas J, Hölzel N, Fischer M, Kleinebecker T (2018) Nutrient stoichiometry and land use rather than species richness determine plant functional diversity. Ecol Evol 8:601–616. https://doi.org/ 10.1002/ece3.3609
- Cameron KC, Di HJ, Moir JL (2013) Nitrogen losses from the soil/plant system: a review. Ann Appl Biol 162:145–173. https://doi.org/10.1111/aab.12014
- Compton JE, Pearlstein SL, Erban L, Coulombe RA, Hatteberg B, Henning A, Brooks JR, Selker JE (2021) Nitrogen inputs best predict farm field nitrate leaching in the Willamette Valley. Oregon Nutr Cycl Agroecosyst 120:223– 242. https://doi.org/10.1007/s10705-021-10145-6
- de Vries FT, Bardgett RD (2016) Plant community controls on short-term ecosystem nitrogen retention. New Phytol 210:861–874. https://doi.org/10.1111/nph.13832
- de Vries FT, Hoffland E, van Eekeren N, Brussaard L, Bloem J (2006) Fungal/bacterial ratios in grasslands with contrasting nitrogen management. Soil Biol Biochem 38:2092– 2103. https://doi.org/10.1016/j.soilbio.2006.01.008
- de Vries FT, van Groenigen JW, Hoffland E, Bloem J (2011) Nitrogen losses from two grassland soils with different fungal biomass. Soil Biol Biochem 43:997–1005. https:// doi.org/10.1016/j.soilbio.2011.01.016
- de Vries FT, Bloem J, Quirk H, Stevens CJ, Bol R, Bardgett RD (2012) Extensive management promotes plant and microbial nitrogen retention in temperate grassland. PLoS ONE 7:1–12. https://doi.org/10.1371/journal.pone.00512 01
- Di HJ, Cameron KC (2002) Nitrate leaching in temperate agroecosystems: sources, factors and mitigating strategies. Nutr Cycl Agroecosyst 64:237–256. https://doi.org/10. 1023/A:1021471531188
- Dias ATC, van Ruijven J, Berendse F (2010) Plant species richness regulates soil respiration through changes in productivity. Oecologia 163:805–813. https://doi.org/10.1007/ s00442-010-1569-5
- Díaz S, Kattge J, Cornelissen JHC, Wright IJ, Lavorel S, Dray S, Reu B, Kleyer M, Wirth C, Colin Prentice I, Garnier E, Bönisch G, Westoby M, Poorter H, Reich PB, Moles AT, Dickie J, Gillison AN, Zanne AE, Chave J, Joseph Wright S, Sheremet Ev SN, Jactel H, Baraloto C, Cerabolini B, Pierce S, Shipley B, Kirkup D, Casanoves F, Joswig JS, Günther A, Falczuk V, Rüger N, Mahecha MD, Gorné LD (2016) The global spectrum of plant form and function. Nature 529:167–171. https://doi.org/10.1038/nature16489
- Dietrich P, Buchmann T, Cesarz S, Eisenhauer N, Roscher C (2017) Fertilization, soil and plant community characteristics determine soil microbial activity in managed temperate grasslands. Plant Soil 419:189–199. https://doi.org/ 10.1007/s11104-017-3328-4
- DIN ISO 11277 (2002) Soil quality: determination of particle size distribution in mineral soil material – method by sieving and sedimentation. General: ISO
- Donnison LM, Griffith GS, Hedger J, Hobbs PJ, Bardgett RD (2000) Management influences on soil microbial

communities and their function in botanically diverse haymeadows of northern England and Wales. Soil Biol Biochem 32:253–263. https://doi.org/10.1016/S0038-0717(99)00159-5

- Eriksen J (2009) Chapter 2 soil sulfur cycling in temperate agricultural systems. Adv Agron 102:55–89. https://doi. org/10.1016/S0065-2113(09)01002-5
- EU-Nitrate Directive (2018) Report from the commission to the council and the European Parliament.
- European Environment Agency (EEA) (2018) Environmental indicator report 2018. In support to the monitoring of the Seventh Environment Action Programme
- Faraway J (2016) faraway: Functions and datasets for books by Julian Faraway. R package version 1.0.7. https://CRAN.R-project.org/package=faraway
- Fargione J, Tilman D, Dybzinski R, Lambers JHR, Clark C, Harpole WS, Knops JMH, Reich PB, Loreau M (2007) From selection to complementarity: shifts in the causes of biodiversity-productivity relationships in a long-term biodiversity experiment. Proc r Soc B Biol Sci 274:871–876. https://doi.org/10.1098/rspb.2006.0351
- Fischer M, Bossdorf O, Gockel S, Hänsel F et al (2010) Implementing large-scale and long-term functional biodiversity research: the biodiversity exploratories. Basic Appl Ecol 11:473–485. https://doi.org/10.1016/j.baae.2010.07.009
- Freschet GT, Roumet C, Comas LH et al (2021) Root traits as drivers of plant and ecosystem functioning: current understanding, pitfalls and future research needs. New Phytol. https://doi.org/10.1111/nph.17072
- Gallejones P, Castellón A, del Prado A, Unamunzaga O, Aizpurua A (2012) Nitrogen and sulphur fertilization effect on leaching losses, nutrient balance and plant quality in a wheat-rapeseed rotation under a humid Mediterranean climate. Nutr Cycl Agroecosystems 93:337–355. https://doi.org/10.1007/s10705-012-9520-2
- Gilhaus K, Boch S, Fischer M, Hölzel N, Kleinebecker T, Prati D, Rupprecht D, Schmitt B, Klaus VH (2017) Grassland management in Germany: effects on plant diversity and vegetation composition. Tuexenia; 37: 379–397. https:// doi.org/10.14471/2017.37.010
- Gocke MI, Don A, Heidkamp A, Schneider F, Amelung W (2021) The phosphorus status of German cropland an inventory of top- and subsoils. J Plant Nutr Soil Sci 184:51–64. https://doi.org/10.1002/jpln.202000127
- Grahmann K, Verhulst N, Palomino LM, Bischoff WA, Govaerts B, Buerkert A (2018) Ion exchange resin samplers to estimate nitrate leaching from a furrow irrigated wheat-maize cropping system under different tillage-straw systems. Soil Tillage Res 175:91–100. https://doi.org/10. 1016/j.still.2017.08.013
- Grigulis K, Lavorel S, Krainer U, Legay N, Baxendale C, Dumont M, Kastl E, Arnoldi C, Bardgett RD, Poly F, Pommier T, Schloter M, Tappeiner U, Bahn M, Clément JC (2013) Relative contributions of plant traits and soil microbial properties to mountain grassland ecosystem services. J Ecol 101:47–57. https://doi.org/10.1111/1365-2745.12014
- Grunwald D, Panten K, Schwarz A, Bischoff W-A, Schittenhelm S (2020) Comparison of maize, permanent cup plant and a perennial grass mixture with regard to soil and water

protection. GCB Bioenergy 12:694–705. https://doi.org/ 10.1111/gcbb.12719

- Herold N, Schöning I, Gutknecht J, Alt F, Boch S, Müller J, Oelmann Y, Socher SA, Wilcke W, Wubet T, Schrumpf M (2014) Soil property and management effects on grassland microbial communities across a latitudinal gradient in Germany. Appl Soil Ecol 73:41–50. https://doi.org/10. 1016/j.apsoil.2013.07.009
- IUSS Working Group WRB, 2014. World Reference base for soil resources 2014, Update 2015. international soil classification system for naming soils and creating legends for soil maps. World Soil Resources Reports No. 106, Rome: FAO
- Kahmen A, Renker C, Unsicker SB, Buchmann N (2006) Niche complementarity for nitrogen: an explanation for the biodiversity and ecosystem functioning relationship? Ecology 87(5), 1244–1255. http://www.jstor.org/stable/ 20069065
- Kattge J, Bönisch G et al (2020) TRY plant trait database: enhanced coverage and open access. Glob Chang Biol 26:119–188. https://doi.org/10.1111/gcb.14904
- Keil D, Meyer A, Berner D, Poll C, Schützenmeister A, Piepho HP, Vlasenko A, Philippot L, Schloter M, Kandeler E, Marhan S (2011) Influence of land-use intensity on the spatial distribution of N-cycling microorganisms in grassland soils. FEMS Microbiol Ecol 77:95–106. https://doi. org/10.1111/j.1574-6941.2011.01091.x
- Klaus VH, Kleinebecker T, Hölzel N, Blüthgen N, Boch S, Müller J, Socher SA, Prati D, Fischer M (2011) Nutrient concentrations and fibre contents of plant community biomass reflect species richness patterns along a broad range of land-use intensities among agricultural grasslands. Perspect Plant Ecol Evol Syst 13:287–295. https://doi.org/10. 1016/j.ppees.2011.07.001
- Klaus VH, Kleinebecker T, Busch V, Fischer M, Hölzel N, Nowak S, Prati D, Schäfer D, Schöning I, Schrumpf M, Hamer U (2018) Land use intensity, rather than plant species richness, affects the leaching risk of multiple nutrients from permanent grasslands. Glob Chang Biol 24:2828–2840. https://doi.org/10.1111/gcb.14123
- Klaus VH, Friedritz L, Hamer U, Kleinebecker T (2020a) Drought boosts risk of nitrate leaching from grassland fertilisation. Sci Total Environ 726:137877. https://doi.org/ 10.1016/j.scitotenv.2020.137877
- Klaus VH, Whittingham MJ, Báldi A, Eggers S, Francksen RM, Hiron M, Lellei-Kovács E, Rhymer CM, Buchmann N (2020b) Do biodiversity-ecosystem functioning experiments inform stakeholders how to simultaneously conserve biodiversity and increase ecosystem service provisioning in grasslands? Biol Conserv 245:108552. https:// doi.org/10.1016/j.biocon.2020.108552
- Kleinebecker T, Klaus VH, Hölzel N (2011) Reducing sample quantity and maintaining high prediction quality of grassland biomass properties with near infrared reflectance spectroscopy. J near Infrared Spectrosc 19:495–505. https://doi.org/10.1255/jnirs.957
- Kleinebecker T, Hölzel N, Prati D, Schmitt B, Fischer M, Klaus VH (2014) Evidence from the real world: 15N natural abundances reveal enhanced nitrogen use at high plant diversity in Central European grasslands. J Ecol 102:456– 465. https://doi.org/10.1111/1365-2745.12202

- Koncz P, Balogh J, Papp M, Hidy D, Pintér K, Fóti S, Klumpp K, Nagy Z (2015) Higher soil respiration under mowing than under grazing explained by biomass differences. Nutr Cycl Agroecosystems 103:201–215. https://doi.org/10. 1007/s10705-015-9732-3
- Kuhlmann H, Baumgärtel G (1991) Potential importance of the subsoil for the P and Mg nutrition of wheat. Plant Soil 137:259–266. https://doi.org/10.1007/BF00011204
- Kunrath TR, de Berranger C, Charrier X, Gastal F, de Faccio Carvalho PC, Lemaire G, Emile JC, Durand JL (2015) How much do sod-based rotations reduce nitrate leaching in a cereal cropping system? Agric Water Manag 150:46– 56. https://doi.org/10.1016/j.agwat.2014.11.015
- Lachaise T, Bergmann J, Rillig MC, van Kleunen M (2021) Below- and aboveground traits explain local abundance, and regional, continental and global occurrence frequencies of grassland plants. Oikos 130:110–120. https://doi. org/10.1111/oik.07874
- Lavorel S, Grigulis K, Lamarque P, Colace MP, Garden D, Girel J, Pellet G, Douzet R (2011) Using plant functional traits to understand the landscape distribution of multiple ecosystem services. J Ecol 99:135–147. https://doi.org/10. 1111/j.1365-2745.2010.01753.x
- Ledgard S, Schils R, Eriksen J, Luo J (2009) Environmental impacts of grazed clover/grass pastures. Irish J Agric Food Res 48:209–226
- Leimer S, Oelmann Y, Wirth C, Wilcke W (2015) Time matters for plant diversity effects on nitrate leaching from temperate grassland. Agric Ecosyst Environ 211:155–163
- Leimer S, Oelmann Y, Eisenhauer N, Milcu A, Roscher C, Scheu S, Weigelt A, Wirth C, Wilcke W (2016) Mechanisms behind plant diversity effects on inorganic and organic N leaching from temperate grassland. Biogeochemistry 131:339–353. https://doi.org/10.1007/ s10533-016-0283-8
- Leinweber P, Meissner R, Eckhardt KU, Seeger J (1999) Management effects on forms of phosphorus in soil and leaching losses. Eur J Soil Sci 50:413–424. https://doi.org/10. 1046/j.1365-2389.1999.00249.x
- Lu M, Zhou X, Luo Y, Yang Y, Fang C, Chen J, Li B (2011) Minor stimulation of soil carbon storage by nitrogen addition: a meta-analysis. Agric Ecosyst Environ 140:234– 244. https://doi.org/10.1016/j.agee.2010.12.010
- Manning P, Loos J, Barnes AD, Batáry P, Bianchi FJJA, Buchmann N, De Deyn GB, Ebeling A, Eisenhauer N, Fischer M, Fründ J, Grass I, Isselstein J, Jochum M, Klein AM, Klingenberg EOF, Landis DA, Lepš J, Lindborg R, Meyer ST, Temperton VM, Westphal C, Tscharntke T (2019) Chapter ten: transferring biodiversity-ecosystem function research to the management of 'real-world' ecosystems. Adv Ecol Res 61:323–356. https://doi.org/10.1016/bs.aecr.2019.06.009
- Manzoni S, Jackson RB, Trofymow JA, Porporato A (2008) The global stoichiometry of litter nitrogen mineralization. Science (80-.). 321:684–686. https://doi.org/10.1126/scien ce.1159792
- Mathot M, Mertens J, Verlinden G, Lambert R (2008) Positive effects of sulphur fertilisation on grasslands yields and quality in Belgium. Eur J Agron 28:655–658. https://doi. org/10.1016/j.eja.2007.12.004

- Meyer A, Focks A, Radl V, Keil D, Welzl G, Schöning I, Boch S, Marhan S, Kandeler E, Schloter M (2013) Different land use intensities in grassland ecosystems drive ecology of microbial communities involved in nitrogen turnover in soil. PLoS ONE. https://doi.org/10.1371/journal.pone. 0073536
- Midolo G, Alkemade R, Schipper AM, Benítez-López A, Perring MP, de Vries W (2019) Impacts of nitrogen addition on plant species richness and abundance: a global metaanalysis. Glob Ecol Biogeogr 28:398–413. https://doi.org/ 10.1111/geb.12856
- Moreau D, Pivato B, Bru D, Busset H, Deau F, Faivre C, Matejicek A, Strbik F, Philippot L, Mougel C (2015) Plant traits related to nitrogen uptake influence plantmicrobe competition. Ecology 96:2300–2310. https://doi.org/10. 1890/14-1761.1
- Niklaus PA, Kandeler E, Leadley PW, Schmid B, Tscherko D, Korner C (2001) A link between plant diversity, elevated CO2 and soil nitrate. Oecologia 127:540–548. https://doi. org/10.1007/s004420000612
- Oelmann Y, Richter AK, Roscher C, Rosenkranz S, Temperton VM, Weisser WW, Wilcke W (2011) Does plant diversity influence phosphorus cycling in experimental grasslands? Geoderma 167–168:178–187. https://doi.org/10.1016/j. geoderma.2011.09.012
- Ostrowski A, Lorenzen K, Petzold E, Schindler S (2020) Land use intensity index (LUI) calculation tool of the Biodiversity Exploratories project for grassland survey data from three different regions in German since 2006. Zenodo. 10.25504/FAIRsharing.6AmTXC
- Pommier T, Cantarel AAM, Grigulis K, Lavorel S, Legay N, Baxendale C, Bardgett RD, Bahn M, Poly F, Clément JC (2018) The added value of including key microbial traits to determine nitrogen-related ecosystem services in managed grasslands. J Appl Ecol 55:49–58. https://doi.org/10. 1111/1365-2664.13010
- Predotova M, Bischoff WA, Buerkert A (2011) Mineral-nitrogen and phosphorus leaching from vegetable gardens in Niamey. Niger J Plant Nutr Soil Sci 174:47–55. https:// doi.org/10.1002/jpln.200900255
- R Core Team (2019) R: a language and environment for statistical computing. R Found. Stat. Comput, Vienna
- Reich PB (2014) The world-wide "fast-slow" plant economics spectrum: a traits manifesto. J Ecol 102:275–301. https:// doi.org/10.1111/1365-2745.12211
- Robson TM, Baptist F, Clément JC, Lavorel S (2010) Land use in subalpine grasslands affects nitrogen cycling via changes in plant community and soil microbial uptake dynamics. J Ecol 98:62–73. https://doi.org/10.1111/j. 1365-2745.2009.01609.x
- Rosseel Y (2012) lavaan: an R package for structural equation modeling. J Stat Softw. https://doi.org/10.18637/jss.v048. i02
- Rumpel C, Crème A, Ngo P, Velásquez G, Mora M, Chabbi A (2015) The impact of grassland management on biogeochemical cycles involving carbon, nitrogen and phosphorus. J Soil Sci Plant Nutr. https://doi.org/10.4067/S0718-95162015005000034
- Ryden JC, Ball PR, Garwood EA (1984) Nitrate leaching from grassland. Nature 311:10. https://doi.org/10.1038/31101 0a0

- Scherer-Lorenzen M, Palmborg C, Prinz A, Schulze E (2003) The role of plant diversity and composition for nitrate leaching in grasslands. Ecology 84:1539–1552. https:// doi.org/10.1890/0012-9658(2003)084[1539:TROPDA] 2.0.CO;2
- Schleuss PM, Widdig M, Biederman LA, Borer ET, Crawley MJ, Kirkman KP, Seabloom EW, Wragg PD, Spohn M (2021) Microbial substrate stoichiometry governs nutrient effects on nitrogen cycling in grassland soils. Soil Biol Biochem 155:108168. https://doi.org/10.1016/j.soilbio. 2021.108168
- Schwarz MT, Bischoff S, Blaser S, Boch S, Schmitt B, Thieme L, Fischer M, Michalzik B, Schulze ED, Siemens J, Wilcke W (2014) More efficient aboveground nitrogen use in more diverse Central European forest canopies. For Ecol Manage 313:274–282. https://doi.org/10.1016/j.foreco. 2013.11.021
- Shipley B (2016) Cause and correlation in biology. Cambridge. https://doi.org/10.1017/CBO9781139979573
- Six J, Frey SD, Thiet RK, Batten KM (2006) Bacterial and fungal contributions to carbon sequestration in agroecosystems. Soil Sci Soc Am J 70:555–569. https://doi.org/ 10.2136/sssaj2004.0347
- Solly EF, Schöning I, Boch S, Kandeler E, Marhan S, Michalzik B, Müller J, Zscheischler J, Trumbore SE, Schrumpf M (2014) Factors controlling decomposition rates of fine root litter in temperate forests and grasslands. Plant Soil 382:203–218. https://doi.org/10.1007/ s11104-014-2151-4
- Sorkau E, Boch S, Boeddinghaus RS, Bonkowski M, Fischer M, Kandeler E, Klaus VH, Kleinebecker T, Marhan S, Müller J, Prati D, Schöning I, Schrumpf M, Weinert J, Oelmann Y (2018) The role of soil chemical properties, land use and plant diversity for microbial phosphorus in forest and grassland soils. J Plant Nutr Soil Sci 181:185–197. https://doi.org/10.1002/jpln.201700082
- Soulides DA, Clark FA (1958) Nitrification in grassland soils. Soil Sci Soc Am J. https://doi.org/10.2136/sssaj 1958.03615995002200040011x
- State and Outlook Environmental Report (2020) The European environment. European Environment Agency, Copenhagen
- Stempfhuber B, Welzl G, Wubet T, Schöning I, Marhan S, Buscot F, Kandeler E, Schloter M (2014) Drivers for ammonia-oxidation along a land-use gradient in grassland soils. Soil Biol Biochem 69:179–186. https://doi.org/10. 1016/j.soilbio.2013.11.007
- Strickland MS, Rousk J (2010) Considering fungal: bacterial dominance in soils - methods, controls, and ecosystem implications. Soil Biol Biochem 42:1385–1395. https:// doi.org/10.1016/j.soilbio.2010.05.007
- Tilman D, Wedin D, Knops J (1996) Productivity and sustainability influenced by biodiversity in grassland ecosystems. Nature 376:718–720. https://doi.org/10.1038/379718a0
- Tzanakakis VA, Apostolakis A, Nikolaidis NP, Paranychianakis NV (2018) Ammonia oxidizing archea do not respond to ammonium or urea supply in an alkaline soil. Appl Soil Ecol 132:194–198. https://doi.org/10.1016/j.apsoil.2018. 08.002
- Vance ED, Brookes PC, Jenkinson DS (1987) An extraction method for measuring soil microbial biomass C. Soil

Biol Biochem 19:703–707. https://doi.org/10.1016/0038-0717(87)90052-6

- Werner W, Trimborn M, Pihl U (2006) Prediction of the P-leaching potential of arable soils in areas with high livestock densities. J Zhejiang Univ Sci B 7:515–520. https:// doi.org/10.1631/jzus.2006.B0515
- Willich M, Buerkert A (2016) Leaching of carbon and nitrogen from a sandy soil substrate: a comparison between suction plates and ion exchange resins. J Plant Nutr Soil Sci 179:609–614. https://doi.org/10.1002/jpln.201600036
- Withers PJA, Neal C, Jarvie HP, Doody DG (2014) Agriculture and eutrophication: Where do we go from here? Sustain 6:5853–5875. https://doi.org/10.3390/su6095853
- World Health Organization (WHO) (2011) Nitrate and nitrite in drinking-water background documents for development of WHO guidelines for drinking-water quality. WHO, Geneva
- Wright I, Reich PB, Westoby M, Ackerly DD, Baruch Z, Bongers F, Cavender-Bares J, Chapin T, Cornelissen JH, Diemer M, Flexas J, Garnier E, Groom PK, Gulias J,

Hikosaka K, Lamont BB, Lee T, Lee W, Lusk C, Midgley JJ, Navas M-L, Nilnemets Ü, Oleksyn J, Osada N, Poorter H, Poot P, Prior L, Pyankov V, Roumet C, Thomas SC, Tjoelker MG, Veneklaas EJ, Villar R (2004) The worldwide leaf economics spectrum. Nature 428:821–827

- Zobel RW, Kinraide TB, Baligar VC (2007) Fine root diameters can change in response to changes in nutrient concentrations. Plant Soil 297:243–254. https://doi.org/10.1007/ s11104-007-9341-2
- Zuur AF, Ieno EN, Walker NJ, Saveliev AA, Smith GM (2009) Mixed effects models and extensions in ecology with R. Springer, Book Ser Stat Biol Health. https://doi.org/10. 1007/978-0-387-87458-6

**Publisher's Note** Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.