Plant diversity effects on herbivory are related to soil biodiversity and plant chemistry

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Abstract

1. Insect herbivory is a key process in ecosystem functioning. While theory predicts that plant diversity modulates herbivory, the mechanistic links remain unclear. We postulated that the plant metabolome mechanistically links plant diversity and herbivory.

2. In late summer and in spring, we assessed individual plant above-ground herbivory rates and metabolomes of seven plant species in experimental plant communities varying in plant species diversity and resource acquisition strategies. In the same communities, we also measured plant individual biomass as well as soil microbial and nematode community composition.

3. Herbivory rates decreased with increasing plant species richness. Path modeling revealed that plant species richness and community resource acquisition strategy correlated with soil community composition. In particular, changes in nematode community composition were related to plant metabolome composition and thereby herbivory rates.

4. Synthesis. These results suggest that soil community composition plays an important role in reducing herbivory rates with increasing plant diversity by changing plant metabolomes.

KEYWORDS
above-ground–below-ground interactions, biodiversity-ecosystem function, chemical diversity, eco-metabolomics, herbivory, Jena experiment, metabolite profile
INTRODUCTION

Insect herbivory is an essential ecosystem process that can remove substantial amounts of biomass in grasslands within a single season (Meyer et al., 2017; Seabloom et al., 2017). Plant diversity can influence the abundance and diversity of insect herbivores (Haddad et al., 2001; Hertzog et al., 2016) as well as herbivory rates (Ebeling, Meyer, et al., 2014; Wan et al., 2020). Plant traits, in particular those associated with resource acquisition and competition, are considered to provide mechanistic links between plant diversity and herbivory (Loranger et al., 2013). These traits can reflect functional aspects of light interception, deep soil nutrient and water use, or resource use along a seasonal gradient (Ebeling, Pompe, et al., 2014; Marr et al., 2021). Whereas these traits may predict the performance of plant species in their niches, they also display plasticity in response to plant soil feedbacks and soil legacies (Delory et al., 2021; Xi et al., 2021). However, a recent study shows that the commonly used morphological and physiological traits only explain up to 12.7% of the variance in herbivory (van der Plas et al., 2020). Plant chemical composition may be a better predictor for individual herbivory, because many herbivores use plant metabolites to locate their host (Agrawal & Weber, 2015) while plants use metabolites to defend themselves (van Dam & van der Meijden, 2011). By using the plant’s metabolome, that is, the composition of all metabolites produced by an individual plant (Oliver et al., 1998), as an additional functional plant trait, we may gain deeper insights in the molecular mechanisms underlying differences in herbivory.

Several factors may explain differences in herbivory rates across plant diversity gradients. Higher levels of plant diversity may increase niche diversity by increasing spatial heterogeneity and the variety of food sources, thus, supporting more insect herbivores and increasing community-level herbivory rates (Ebeling, Meyer, et al., 2014). At individual plant level, however, increased plant diversity may lead to dilution effects which decrease herbivory, as it will be more difficult for specialized insect herbivores to localize their host plant (Castagneyrol et al., 2014; Finch & Collier, 2000). Indeed, in a previous study conducted in the Jena Experiment, individual herbivory decreased with increasing plant species richness (Scherber et al., 2006). Lastly, the abundance of predatory and parasitoid arthropods, which can reduce herbivore populations and thus plant community-level and individual herbivory via top-down control, is commonly higher in more diverse plant communities (Haddad et al., 2009; Hines et al., 2015; Schuldt et al., 2019; Wan et al., 2020).

In addition, differences in herbivory across plant species richness gradients may also be explained through changes in plant chemistry. Plant metabolomes change in response to abiotic variation. This metabolomic response to environmental conditions co-determines the defensive status of a plant (van Dam & van der Meijden, 2011). For instance, plants increase the synthesis of defensive metabolites following an attack by herbivores (Bezemer & van Dam, 2005; Karban & Baldwin, 1997). These induced responses can change defences both locally, that is, in the attacked tissue, and systemically, that is, throughout the plant (van Dam & Heil, 2011). In addition to herbivory-induced changes, plant diversity itself can affect plant metabolomes. Plant–plant interactions can alter the metabolome through competition, which may induce the production of volatile (Baldwin et al., 2006) and nonvolatile allelopathic compounds (Fernandez et al., 2016). Seen the broad biological activity spectrum of plant metabolites, these changes are likely to affect herbivory rates (Broz et al., 2010). Lastly, soil legacy effects, which may result from systemically induced changes triggered by soil biota, such as microbes and nematodes (van Dam & Heil, 2011; Wondafrash et al., 2013) can also affect plant metabolomes (Ristok et al., 2019). Taken together, the plant metabolome both affects and reflects above- and below-ground interactions with insect herbivores, other plants and soil biota, in a species-specific and context-dependent way (Bezemer & van Dam, 2005; Ristok et al., 2019). Hence, we argue that measuring plant metabolomes will provide novel insights into the relationship between plant diversity and above-ground herbivory.

The aim of our study was to jointly analyse the relationships between plant diversity, soil biota communities, plant metabolomes, and above-ground herbivores to provide a mechanistic framework for above-below-ground multitrophic interactions in grasslands. Here, we analysed the metabolomes and individual plant herbivory of three grass and four forb species in experimental plant communities manipulated to vary in spatial or temporal resource acquisition traits (Ebeling, Pompe, et al., 2014). Our species selection covered a range of functional traits related to resource acquisition (Ebeling, Pompe, et al., 2014) and included both grasses and forbs, because their metabolomes and response to the abiotic and biotic environment may differ (Dietz et al., 2019, 2020; Hubert et al., 2020).

All plants were grown in 34 experimental plant communities that varied in plant diversity, that is, species richness and functional trait diversity (Ebeling, Pompe, et al., 2014). We tested if and how plant diversity alters the secondary metabolome and how this relates to herbivory. We hypothesized that (1) plant species richness and the resource acquisition strategy of the plant community affect individual plant herbivory. Moreover, we calculated partial-least-squares path models to explore if (2) plant species richness and plant community resource acquisition traits directly or indirectly, via the soil biota, relate to the plant’s metabolome and thereby may explain variation in herbivory. Our hypotheses are based on observations that plant species richness affects soil community composition (Eisenhauer et al., 2010) and that differences in soil biota can affect the plant’s metabolome and thereby herbivory (Hubert et al., 2020; Ristok et al., 2019). Additional paths in our models accounted for relationships between the soil microbial community and soil nematode community (Dong & Zhang, 2006) as well as for potential direct relationships between plant species richness and individual plant herbivory rates (e.g. due to dilution effects; Castagneyrol et al., 2014; Scherber et al., 2006). We inferred similar paths for functional trait composition, accounting for observations in which plant communities containing tall-statured species with large leaves and deep roots increased individual plant herbivory, as these species may have
provided more niches for insect herbivores (Loranger et al., 2012). We also modelled the relationship of growth and flowering time with herbivory as plant chemistry is known to change with ontogeny (Barton & Koricheva, 2010; Boege, 2005). Lastly, we included the relationship between plant biomass and herbivory in our path models, because soil community composition can affect plant biomass, which in turn may affect herbivory, whereby larger plants may incur more herbivory (Windig, 1993).

We show that increasing plant species richness reduces individual plant herbivory. Furthermore, our path models suggest that soil community composition, especially the composition of the nematode community, and plant individual metabolomes are key players in the relationship between plant diversity and above-ground herbivory.

2 | METHODS

2.1 | Experimental design

The trait-based experiment (Ebeling, Pompe, et al., 2014) was established in 2010 within the ‘Jena Experiment’ (www.the-jena-experiment.de) field site, Thuringia, Germany; 50°55′N, 11°35′E, 130m a.s.l. (Roscher et al., 2004; see Appendix S1 for details). We sampled 34 plots that differed in plant species richness (1, 2, 4, and 8 species) and plant functional trait dissimilarity (see Table S1 for information on the plant community of each plot). The functional trait dissimilarity was based on traits that reflect spatial and temporal resource acquisition strategies. We chose plant height, leaf area, rooting depth and root length density to reflect spatial resource acquisition. To reflect temporal resource acquisition, we chose growth starting date and flowering onset. All plots were arranged in three blocks, mown in June and September, and weeded three times per year.

2.2 | Secondary metabolome sampling and sample processing

We sampled twice under different environmental conditions to account for seasonal variation in the plants’ metabolomes. Initially, we sampled above-ground biomass of eight common central European grassland species (grasses: Anthoxanthum odoratum L., Dactylis glomerata L., Holcus lanatus L., Phleum pratense L., forbs: Geranium pratense L., Leucanthemum vulgare (Vail.) Lam., Plantago lanceolata L., and Ranunculus acris L.) in 34 plots on 24–25 August 2015 and 31 May–1 June 2016, each time just before mowing. We sampled three individuals per species and plot during both sampling campaigns (theoretical number of samples = 504). Due to an unforeseen low abundance of A. odoratum individuals in August 2015, we failed to sample three individuals per plot. Therefore, we decided to exclude A. odoratum from our analyses (i.e. 54 samples). Furthermore, we excluded seven samples (4 samples of P. pratense, 2 samples of R. acris, 1 sample of G. pratense) due to contamination during sample processing (i.e. the final number of analysed samples = 443). We harvested the shoot biomass by cutting the plants c. 1 cm above ground and removed all inflorescences. All samples were taken between 15.00 and 19.00 h each sampling day to minimize diurnal variation. All samples were processed, extracted and analysed according to Ristok et al. (2019) with slight changes (see Appendix S1). In short, we extracted 20 mg dried ground plant tissue of each sample in 1 ml of extraction buffer (methanol/50 mM acetate buffer, pH 4.8; 50/50 [v/v]). The samples were homogenized for 5 min at 30 Hz using a ball mill (Retsch mixer mill MM 400), and subsequently centrifuged (20,000 g, 10 min, 4°C). The supernatant was collected in a 2 ml Eppendorf tube. We repeated the extraction procedure with the remaining pellet and combined the supernatant with the first one. We centrifuged (20,000 g, 5 min, 4°C) all extracts, transferred 200 μl to an HPLC vial and added 800 μl extraction buffer, resulting in a 1:5 dilution.

We performed chromatographic separation of all diluted extracts by injecting 2 μl on a Thermo Scientific Dionex UltiMate 3000 (Thermo Scientific Dionex) UPLC unit, equipped with a C18 column (Acclaim RSLC 120 C18, 2.2 μm, 120 Å, 2.1 × 150 mm, Thermo Fisher Scientific). We applied the following binary elution gradient at a flow rate of 0.4 ml min⁻¹ and a column temperature of 40°C: 0–2 min, 95% A (water and 0.05% formic acid), 5% B (acetonitrile and 0.05% formic acid); 2–12 min, 5 to 50% B; 12–13 min, 50 to 95% B; 13–15 min, 95% B; 15–16 min, 95 to 5% B; 16–20 min, 5% B.

Metabolites were analysed on a liquid chromatography quadrupole time-of-flight mass spectrometer (LC-qToF-MS; Bruker maXis impact HD; Bruker Daltonik) with an electrospray ionization source operated in negative mode (Appendix S1).

2.3 | LC-MS data processing and metabolite prediction

The LC-MS data are presented as a list of features described by mass-to-charge ratios, retention times, and intensities. We processed LC-MS data as in Ristok et al. (2019) with minor changes (see Appendix S1). We predicted metabolite structures through the comparison of LC-MS data with literature references. We submitted high-resolution mass-to-charge values to the MassBank of North America (MoNA, http://mona.fiehnlab.ucdavis.edu/) spectral database. We used a mass tolerance of 0.5 D for comparison. Furthermore, we calculated high-resolution molecular weights, molecular formulae for putative molecular ions in neutral form, and particle weights for mass spectrometry generated fragments using ChemDraw Ultra 8.0 (www.cambridgesoft.com).

2.4 | Leaf herbivory rate assessment

For each plant, we counted the total number of leaves and the number of leaves with herbivore damage 1 day before we sampled...
above-ground biomass. We categorized herbivore damage for each damaged leaf that had signs of sucking, chewing, and mining. The damage categories were 1%–10%, 10%–20%, 20%–30%, 30%–40%, 40%–50%, 50%–60%, 60%–70%, 70%–80%, 80%–90%, and 90%–100%. We multiplied the number of leaves in each category with their damage level (0.1 for 1%–10%, 0.2 for 10%–20%, etc.), and summed across all categories. Finally, we calculated relative herbivory rate for each sample by dividing the summed herbivory by the total number of leaves.

2.5 | Soil sampling

In each plot, we took soil samples to a depth of 10 cm using a metal corer (diameter 2 cm) on 27 August 2015 and 6 June 2016. We pooled and homogenized five subsamples per plot to account for spatial heterogeneity. We sieved soil samples to 2 mm. We stored one part at −20°C for phospholipid fatty acid analysis and the other part at 4°C for nematode extraction.

2.6 | Phospholipid fatty acid analysis

We measured phospholipid fatty acids (PLFA) as a proxy for the soil microbial community composition, based on the abundance of functional groups (Wixon & Balser, 2013). We extracted PLFAs following the Frostegård et al. (1991) protocol as described in Wagner et al. (2015). We analysed all samples on a gas chromatograph (see Appendix S1). We used the following PLFA-markers (ng g−1 dry weight soil) as bacterial markers: (a) gram-negative bacteria: cy17:0 and cy19:0; (b) gram-positive bacteria: i15:0, a15:0, i16:0, and i17:0; and (c) widespread in bacteria: 16:1ω7c, 16:1ω6c, 18:1ω7c, and 18:2ω6c. As fungal markers we used: (a) saprophytic fungi: 18:1ω9t, 18:2ω6t, and 18:2ω6c; and (b) arbuscular mycorrhizal fungi: 20:1 (Ruess & Chamberlain, 2010; Wagner et al., 2015). We summed up all markers within each group of bacteria and fungi to receive a representative value.

2.7 | Nematode extraction and identification

We extracted nematodes from 25 g fresh soil using a modified Baermann method (Ruess, 1995; Wagner et al., 2015). We counted all nematodes at 100x magnification and identified at least 100 randomly chosen nematodes (if available) at 400x magnification using a Leica DMI 4000B light microscope. Nematodes were identified to genus or family level following Bongers (1994). We classified all nematodes into plant feeders, fungal feeders, bacterial feeders, predators and omnivores. Moreover, we assigned all nematodes a c-p score (colonization-persistence gradient) that ranged from 1 to 5 (Bongers & Bongers, 1998). Finally, we combined the trophic group and c-p score to create functional nematode guilds as a proxy for nematode community structure (Cesarz et al., 2015; Ferris et al., 2001).

2.8 | Statistical analysis

We analysed our data in R v3.5 (R Core Team, 2017: http://www.r-project.org) using the packages vegan (Oksanen et al., 2017), pairwiseAdonis (Arbizu, 2017), lme4 (Bates et al., 2014), lmerTest (Kuznetsova et al., 2016), effects (Fox & Weisberg, 2019) and plsPM (Sanchez et al., 2017).

Based on earlier studies in the same experiment (Beugnon et al., 2019; Steinauer et al., 2017), we calculated community mean scores (CMS) to represent resource acquisition strategy (spatial and temporal). We based our CMS calculations on the original PCA species scores calculated when the experiment was designed (Ebeling, Pompe, et al., 2014; Fischer et al., 2016), and on the relative species-specific cover for each plant community recorded in August 2015 and May 2016, respectively. In short, the six functional traits plant height, leaf area, rooting depth, root length density, growth starting date, and flowering onset were analysed in a standardized PCA. The first PCA axis arranged species according to their spatial resource acquisition strategy. The second PCA axis arranged species according to their temporal resource acquisition strategy (Ebeling, Pompe, et al., 2014). Plots with high community mean scores on the first PCA axis (CMS_PCA1) were mostly dominated by tall-statured species with deep roots and large leaves. In contrast, plots with negative community mean scores on the first PCA axis contained a high proportion of small-statured species with dense shallow roots and small leaves. Plots with high community mean scores on the second PCA axis (CMS_PCA2) contained mostly late growing and late flowering species (Fischer et al., 2016).

We tested our first hypothesis by calculating linear mixed effects models. We fitted herbivory rate (log-transformed) as the response variable. As predictor variables, we fitted sampling campaign (categorical; August 2015 or May 2016), plant functional group identity (categorical; grass or forb), and either plant species richness (metric; 1, 2, 4 or 8) or either CMS_PCA1 or CMS_PCA2 (metric), as well as the two-way and three-way interactions.

We tested for the overall and pairwise differences in shoot metabolome composition among the different sown plant species richness levels by calculating permutational multivariate analyses of variance using distance matrices. We log +1 transformed the metabolite intensity data to achieve multivariate normality, and used Bray–Curtis dissimilarity to calculate the distance matrices. All analyses were permuted 9999 times. Each analysis was species-specific and sampling campaign-specific. We were not able to calculate pairwise comparisons of the metabolome composition between plants grown in monoculture (lowest plant species richness level) and in the highest diversity plot (8 species mixture). This is due to the experimental design (see Table S1). For each species, only one monoculture plot
was present. In addition, there was only one 8-species plot. This meant there were not enough replicates to run permutational multivariate analyses of variance and, as such, the pairwise comparisons between monoculture and the 8-species plot were excluded from the analyses.

In addition, we calculated two metrics of metabolite diversity: (a) the richness of secondary metabolites, that is, the number of metabolites within a plant individual; and (b) the Shannon diversity of secondary metabolites, that is the abundance-weighted diversity of metabolites expressed as the exponential of the Shannon-Weaver index (Hill, 1973) based on plant individual-level metabolite intensities. We also calculated community-weighted mean (CWM) trait values for each trait considered in the design of the Trait-Based-Experiment (Roscher et al., 2012). Here, we based the calculations on the relative species-specific cover for each plant community. We calculated linear mixed effects models to test for the effect of sown plant species richness or CWM trait values on the richness or Shannon diversity of secondary metabolites. We fitted either the richness or the Shannon diversity of secondary metabolites as response variables. As predictor variables, we fitted sampling campaign, plant functional group identity, and either sown plant species richness or each of the CWM traits separately (metric, scaled), as well as the two-way and three-way interactions.

Finally, we tested for the relationship between richness or Shannon diversity of secondary metabolites and plant-individual herbivory using linear mixed effects models. We fitted herbivory rate (log-transformed) as the response variable. As predictor variables, we fitted sampling campaign, plant functional group identity, and either richness or the Shannon diversity of secondary metabolites, as well as the two-way and three-way interactions.

For all linear mixed effects models calculated in our study, we fitted plot nested in block and species identity as independent random effects. We performed backwards model simplification, first removing nonsignificant interactions and then nonsignificant predictors, until the change in Akaike Information Criterion (AIC) was <2. Finally, the most parsimonious model with the lowest AIC was chosen. All linear mixed effects models were based on restricted-maximum likelihood estimation and Type I analysis of variance with Satterthwaite approximation for degrees of freedom.

To test our second hypothesis, we calculated partial-least-squares path models (PLS-PM; see Appendix S1; Sanchez, 2013). We hypothesized direct links from the experimental design variables plant species richness and resource acquisition traits to microbial and nematode community composition (De Deyn & Van der Putten, 2005; Strecker et al., 2016), as well as to plant individual biomass (Tilman et al., 2001), plant metabolome (Schering et al., 2010), and individual plant herbivory (Scherber et al., 2006; for details on latent variables see Table 1). In addition, we hypothesized links from the microbial to the nematode community composition (Dong & Zhang, 2006) as well as from either soil biota community to plant biomass (van der Putten et al., 2013) and to the composition of the plant metabolome (Huberty et al., 2020; Ristok et al., 2019). Furthermore, we hypothesized links from plant biomass to metabolome (de Jong, 1995; Fernandez et al., 2016) and herbivory (Barnes et al., 2020). Finally, we hypothesized a link from plant metabolome to herbivory (van Dam & van der Putten, 2001; Strecker et al., 2016). As such, we fitted sampling campaign, plant functional group identity, and either richness or the Shannon diversity of secondary metabolites as deterministic variables that determined the metabolome and, in turn, the biomass and herbivory, as well as the two-way and three-way interactions.

Table 1: Description of the latent and observed variables used in the PLS-PM models

| Latent variable  | Description                                                                                                                                 |
|------------------|-----------------------------------------------------------------------------------------------------------------------------------------------|
| Resource acquisition traits | The latent variable represents the community-weighted mean (CWM) trait values of maximum plant height, leaf area, root length density, growth start, and flowering start. We based the calculations of the CWM trait values on the relative species-specific cover for each plant community. For instance, a positive relationship of resource acquisition traits with the latent variable biomass means higher values in the CWM traits correlated with a higher plant biomass |
| Microbial community | The latent variable represents the microbial biomass for gram-negative bacteria, gram-positive bacteria, undefined bacteria, saprophytic fungi, and arbuscular mycorrhizal fungi. For instance, a negative relationship of microbial community with the latent variable biomass indicates that a higher biomass of soil bacteria and fungi correlated with a lower individual plant biomass |
| Nematode community | The latent variable represents the relative abundance of functional nematode guilds. A functional nematode guild is the combined information of trophic guild and colonizer-persistence score. For instance, a positive relationship of nematode community with the latent variable metabolome means a greater relative abundance of certain functional guilds correlated with a higher concentration of metabolites |
| Metabolome | The latent variable represents the abundance of secondary plant metabolites. For instance, a negative relationship of metabolome with the latent variable herbivory means that a higher abundance of metabolites is correlated with a lower herbivore damage on individual plants |

| Observed variable  | Description |
|-------------------|-------------|
| Species richness | The observed variable represents the plot level sown plant species richness from 1 to 8 |
| Biomass           | The observed variable represents the above-ground dry biomass of individual plants |
| Herbivory         | The observed variable represents the herbivory rate on individual plants |
Meijden, 2011; Figure 4a). We calculated three separate PLS-PMs: (a) the full model using all data of the seven plant species; (b) the grasses-only model using only data of the three grass species; and (c) the forbs-only model using only data of the four forb species. All data were scaled, and we used bootstrapping (n = 200) to calculate confidence intervals for the effect sizes within the path models. We simplified all path models by reducing the number of outer model observable variables (so-called indicators), until the most parsimonious solution was achieved (Sanchez, 2013). Indicators are always positively correlated with their latent variable; a low value of the latent variable relates to a low value in all respective indicators, and vice versa. Latent variables are estimated as a weighted linear combination of their indicators (Sanchez, 2013).

Across all three models, the latent variable resource acquisition traits were characterized by the community-weighted means of plant height, leaf area, rooting depth, root length density, growth starting date and flowering start. The latent variable soil microbial community was characterized by gram-negative, gram-positive, general bacteria, saprophytic fungi, and arbuscular mycorrhizal fungi. The effects of the latent variable nematode community were mostly driven by bacterial feeders with a colonization-persistence score (c-p) of 1 and 3, predators (c-p 4 and 5), fungivores (c-p 4), omnivores (c-p 2 and 3) and plant feeders (c-p 2 and 3).

3 | RESULTS

3.1 | The effects of plant species richness and resource acquisition strategy

Herbivory rates decreased significantly with plant species richness (Table S2; Figure 1a) and were lower in May 2016 than in August 2015 (Table S2; Figure 1). Together, plant species richness and sampling campaign explained 26% (marginal $R^2$ value, hereafter $R^2_{\text{marg}}$) of the total variation in herbivory rates.

We found no significant effect of the community's spatial resource acquisition strategy on herbivory rates (Table S2). In other words, the relative abundance of tall-statured species with large leaves and deep roots did not significantly affect the plant-individual herbivory rates (Figure 1b). However, when we tested for the effect of community spatial resource acquisition strategy within plant functional groups, we found a significant effect for grasses (Table S3). Grasses growing in communities that predominantly contained tall-statured plants with large leaves suffered lower herbivory rates (Figure S1a; $R^2_{\text{marg}} = 0.35$). In contrast, we observed no significant effect of spatial resource acquisition strategy on individual herbivory in forbs (Table S3; Figure S2a). When we tested for the effect of the temporal resource acquisition strategy of the plant communities on herbivory rates, we only found a marginally significant relationship (Table S2). More specifically, plant-individual herbivory tended to be greater in communities containing mostly later growing and flowering species (Figure 1c). Separate analyses for each plant functional group showed that this effect was significant for grasses ($R^2_{\text{marg}} = 0.36$; Table S3; Figure S1b), but not for forbs (Table S3; Figure S2b).

We also tested for the effects of plant species richness on plant metabolome composition and on metabolite diversity, that is, richness or Shannon diversity of secondary metabolites. We found a significant effect of plant species richness on the metabolome composition across most plant species in at least one sampling campaign, except for Holcus lanatus (Table 2). Consecutive pairwise comparisons revealed that the metabolome of plants grown in monocultures most frequently differed from the metabolome of plants grown in more diverse communities (Table S4).

We did not find an effect of plant species richness on the richness of secondary metabolites (Figure 2a). Rather, we observed an effect of plant functional group identity on the richness of secondary metabolites ($F_{1,5} = 6.69; p = 0.049$; Table S5a). Forb species had significantly more secondary metabolites ($396 \pm 8.1; \text{mean} \pm \text{SE}$) than grass species ($311 \pm 8.5; \text{mean} \pm \text{SE}$). Together with sampling campaign, plant functional group identity could explain 68% ($R^2_{\text{marg}}$) of the total variation in secondary metabolite richness. We also discovered that the effect of plant species richness on the Shannon diversity of secondary metabolites depended on the functional group identity ($F_{1,138} = 5.35; p = 0.022$; Table S5a). Increasing plant species richness increased the Shannon diversity of secondary metabolites in grasses, while it reduced the Shannon diversity of secondary metabolites in forbs (Figure 2b). Sampling campaign, plant functional group identity, and plant species richness together explained 49% ($R^2_{\text{marg}}$) of the total variation in the Shannon diversity of secondary metabolites.

Moreover, we analysed the extent to which resource acquisition strategy affected metabolite diversity. We found that some traits associated with spatial resource acquisition strategy can increase or decrease the richness of metabolites dependent on sampling campaign and functional group identity (Table S5). The community-weighted mean of rooting depth reduced the richness of metabolites in grasses in May 2016, but increased their richness in August 2015 (Table S6; Figure 2d), while plant height (Figure 2c) and root length density (Figure 2e) only tended to have a similar effect (Table S6). We observed a similar significant effect of plant height on the richness of metabolites in forbs (Table S6; Figure 2f). In contrast, rooting depth (Figure 2g) and root length density (Figure 2h) increased the richness of metabolites in forbs in May 2016, but reduced their richness in August 2015 (Table S6). In addition, we discovered that the community-weighted mean of leaf area, a trait associated with spatial resource acquisition, and the community-weighted mean of flowering onset, a trait associated with temporal resource acquisition, had similar effects on the Shannon diversity of metabolites (Table S5). Leaf area increased Shannon diversity of metabolites in grasses (Table S6; Figure 2i). In forbs, leaf area reduced the Shannon diversity in May 2016, while it increased the Shannon diversity of metabolites in August 2015 (Table S6; Figure 2k). The community-weighted mean of flowering onset tended to have a similar effect in grasses (Table S6; Figure 2j) and forbs (Table S6; Figure 2l) as leaf area had.
Finally, we analysed the relationship between the richness or Shannon diversity of secondary metabolites and plant individual herbivory rates. We observed an interactive effect of sampling campaign and plant functional group identity on the relationship between richness of secondary metabolites and plant-individual herbivory (Table S2; Figure 3a,b). In grasses, herbivory slightly increased with increasing richness of secondary metabolites in August 2015, while herbivory decreased with metabolite richness in May 2016. In forbs, we found a strong positive relationship between the richness of secondary metabolites and plant-individual herbivory in May 2016 and a weak positive relationship in August 2015. Regarding the Shannon diversity of secondary metabolites, we discovered a significant relationship with plant-individual herbivory that differed between sampling campaigns but not between plant functional groups (Table S2; Figure 3c). Herbivory increased with increasing Shannon diversity of secondary metabolites in August 2015, while herbivory decreased with Shannon diversity in May 2016.

3.2 | The relationships between plant diversity, soil community composition, plant metabolomes, and above-ground herbivory

By analysing the significant direct paths in our full-model PLS-PM (Figure 4b; Goodness-of-Fit (GoF) = 0.15), we found a negative relationship between plant species richness and nematode community composition, and a positive relationship between microbial community composition and nematode community composition. Nematode
community composition was positively related to plant metabolomes, which itself was negatively related to plant individual herbivory (for details of each latent variable see Table 1; for all direct, indirect, and total path coefficients see Table S7). Our most parsimonious model predicted 34% of the total variation in the secondary metabolome, and 22% of the total variation in individual herbivory. Plant species richness was negatively correlated with the relative abundance of predatory, omnivorous and plant feeding nematodes (Figure S3). The spatial resource acquisition trait plant height was positively correlated with the relative abundance of predators, fungivores, omnivores, and plant feeders. In contrast, leaf area was negatively correlated with the relative abundance of bacterial feeders, fungivores, omnivores and plant feeders. Rooting depth and root length density negatively correlated with bacterial feeders, predatory nematodes and omnivores, but positively correlated with plant feeders. Conversely, the temporal resource acquisition traits growth starting date and flowering start were negatively correlated with the relative abundance of plant feeders (Figure S3). In addition, we extracted the 100 most important metabolite mass spectra that characterized the metabolome, that is, the metabolites with the strongest positive correlation with the latent variable ’metabolome’. We could assign molecular formulas and structures to 13 mass spectra (Table S8; Figures S4–S16). These metabolites were mainly phenolics, their precursors or their derivatives, which are all products of the shikimic acid pathway. Moreover, these compounds are known to respond to phytopathogenic nematode infection (Ohri & Pannu, 2010) and play a role in plant-herbivore interactions (Whitehead et al., 2021). As part of the nematode community composition, the relative abundance of bacterial feeders, predators, omnivores, and plant feeders showed the strongest positive correlations with the concentration of the assigned metabolites. Especially sinapic acid, a flavonol, the chlorogenic acid dimers, and quinic acid, were negatively correlated with plant herbivory (Figure S17).

Our full-model PLS-PM also indicated that plant herbivory was positively related to community-weighted resource acquisition traits and plant individual biomass (Figure 4b). Plant height, growth starting date, and flowering start were most strongly positively correlated with the latent variable ’resource acquisition traits’. Neither trait was individually correlated with plant herbivory (Figure S3). However, our path model suggests a synergistic effect on plant herbivory, that is, plant communities of tall-growing species with late growth and flowering start may increase plant individual herbivory. Plant biomass was negatively related to resource acquisition traits, plant species richness, nematode community composition, and plant metabolome. Lastly, the microbial community composition was positively related to plant species richness and negatively related to resource acquisition traits. We performed sensitivity analyses and calculated two alternative full-model PLS-PMs: (a) a path model...
FIGURE 4 Hypothesis-based conceptual partial-least-squares path model (a) as well as path model including data across both sampling campaigns and all plant species (b), only across all grasses (c), and only across all forbs (d). Species richness represents the plot-level sown plant species richness. Resource acquisition traits represent the community-weighted mean traits maximum plant height, leaf area, rooting depth, root length density, growth starting date, and flowering start. Microbial community represents PLFA-based estimates on plot-level gram-negative, gram-positive, and undefined bacteria, as well as arbuscular mycorrhizal fungi and all other fungi abundance. Nematode community represents plot-level summed relative abundance of functional nematode guilds, that is, bacterial-feeding, carnivorous, fungal-feeding, omnivorous and plant-feeding. Biomass represents plant-individual above-ground dry biomass. Metabolome represents plant-individual secondary metabolite composition. Herbivory represents plant-individual herbivory rate expressed as the proportion of damaged leaves to the total number of leaves. All data is scaled. Variables taken at plot level are highlighted by a grey-shaded background. Variables taken at the plant-individual level are highlighted by a white-shaded background. Black arrows display significantly positive relationships. Red arrows display significantly negative relationships. Numbers on arrows are path coefficients. Numbers in the round boxes display the explained variation ($R^2$).
that links herbivory to plant metabolome, which would account for herbivore-induced responses (GoF = 0.14; Figure S18a), and (b) a path model that directly links microbial community composition and nematode community composition to plant individual herbivory (GoF = 0.14; Figure S18b). The Goodness-of-Fit of both models was similar to our original full-model PLS-PM. However, the strength of the direct path between metabolome and herbivory was stronger in the original full-model PLS-PM (path coefficient of −0.34) than in the first alternative model (path coefficient of −0.23). Moreover, the first alternative model predicted only 13% of the total variation in individual plant herbivory, whereas the original model predicted 22%. In the second alternative model, both direct paths from microbial community composition and nematode community composition to plant individual herbivory were nonsignificant, thus this model was not improving upon our original full-model PLS-PM.

Based on our observations that functional group identity affects plant metabolomes and herbivory, we calculated two additional PLS-PMs: a grasses-only model (Figure 4c; GoF = 0.21) and a forbs-only model (Figure 4d; GoF = 0.18). In the grasses-only model both the microbial and the nematode community composition were positively related to plant metabolome. In contrast to the full-model, the resource acquisition traits were not related to microbial community composition, but microbial community composition was positively related to the plant metabolome (see Table S7). In the forbs-only model, the plant metabolome was negatively related to nematode community composition, but positively related to biomass and herbivory. Moreover, the resource acquisition traits were not related to any other latent variable, plant individual biomass was not related to herbivory, but plant species richness was negatively related to herbivory (see Table S7). Both functional group models, however, explained more variation in individual plant herbivory (grasses-only model 32%, forbs-only model 49%) than the full-model, suggesting that in grasses and forbs different mechanisms may link plant diversity and soil community composition with plant metabolomes and herbivory.

**4 | DISCUSSION**

Our study highlights how relationships between different facets of biodiversity in plant communities can shape the plant’s metabolome, thereby providing a mechanistic explanation for reduced above-ground herbivory at high plant species richness. We could show that plant-individual herbivory decreases with increasing plant species richness, partially confirming our first hypothesis (H: plant species richness and the resource acquisition strategies of the plant community affect individual plant herbivory). Using partial-least-squares path-modelling, we uncovered the relationships between plant diversity, soil community composition, plant metabolomes, and above-ground herbivory, thus supporting our second hypothesis (H: plant species richness and plant community resource acquisition traits directly or indirectly, via the soil biota, relate to the plant’s metabolome and thereby may explain variation in herbivory). Compared to previous studies (Scherber et al., 2010; van Dam & Heil, 2011) our study yields novel insights by highlighting how below-ground communities may shape plant metabolomes, thereby becoming a significant driver of above-ground herbivory.

The abundance, diversity and community structure of soil biota are commonly determined by the species identity and traits of individual plants as well as the plant community diversity (Bezemer et al., 2010; Lange et al., 2015; Strecker et al., 2016). Accordingly, our path model showed that plant species richness and variation in resource acquisition-related functional traits can explain variation in soil microbial and nematode community composition. The relationship between plant species richness and microbial community composition is likely due to an increased and more diverse influx of organic matter in the form of rhizodeposits (Eisenhauer et al., 2017; Lange et al., 2015; Steinauer et al., 2016). The observed negative relationship between resource acquisition traits and most functional nematode guilds was mainly driven by community-weighted plant height, growth starting time, and flowering onset. This suggests that the abundance and seasonality of resource influx from the plant community into the soil determines nematode community structure (Yeates, 1999). In contrast, rooting depth and root length density were positively correlated with phytophagous nematodes, suggesting that phytophagous nematodes can also be affected by root architecture (Yeates, 1999). Changes in soil community composition were related to significant changes in plant metabolomes. Specifically, the abundance of bacterial feeders, predators and phytophagous nematodes positively correlated to the concentration of defence-related metabolites in individual plants. Bacterial-feeding nematodes contribute to the mineralization of nitrogen in the soil, which supports plant growth and potentially the synthesis of defence-related metabolites (Freckman & Caswell, 1985). Predatory nematodes control plant parasitic nematodes, thus also indirectly supporting plant growth (Freckman & Caswell, 1985). In contrast, phytophagous nematodes can induce systemic defence responses, which can explain the positive correlation between nematode abundance and defensive secondary metabolites in leaves (van Dam & Heil, 2011; Wondafrash et al., 2013). Interestingly, we identified several phenolic compounds that are produced via the shikimic acid pathway. Salicylic acid, which is involved in the plant’s systemic response to root feeding nematodes, is a product of this pathway (Dempsey et al., 2011; Wondafrash et al., 2013). Our path models all contained significant relationships between plant diversity and soil biota community composition, which may have affected the shoot metabolome via the systemic induction of metabolites (Agrawal & Weber, 2015; van Dam & Heil, 2011). Such an induction of metabolites can affect herbivore resistance and may explain the significant link between nematode community composition, the composition of the plant metabolome, and herbivory in our model (van Dam & van der Meiijden, 2011). This is supported by earlier findings reporting effects of plant diversity on soil community composition, especially plant growth facilitators and plant antagonists, and on plant metabolomes and thereby herbivory (Bezemer & van Dam, 2005; Hol et al., 2010; Kos et al., 2015; Ristok et al., 2019; Wurst et al., 2010).
Similarly, also root herbivory may impact shoot metabolomes (Bezemer & van Dam, 2005).

While an aim of our study was to test if the plant's metabolome can explain variation in herbivory, we could not disentangle potential effects of herbivory on the plant's metabolome. Because we analysed field plants, it is likely that the metabolomes result from several (a)biotic interactions. If anything, this adds realism to our results, as in nature above-ground herbivores often encounter plants induced by other interactors (van Dam & Heil, 2011). In all models, the relationship between biomass and metabolome was maintained. The negative relationships found in the full and grasses model, may support the hypothesis that larger plants produced less defence, because they can tolerate biomass loss to herbivory and prioritize growth over defence production (de Jong, 1995). In forbs, however, there was a positive relationship between biomass and metabolome. This might point to the fact that flowering forbs are commonly larger and produce more and different metabolites to protect their reproductive organs (McKey, 1979). Additional experiments are necessary to test the hypothesis that the growth-defence trade-off varies with ontogeny and between plant functional groups.

Overall, sampling campaign had a strong effect on plant-individual herbivory, the richness and Shannon diversity of secondary metabolites, and the relationship among these variables as well as with resource acquisition traits. While this potential seasonal effect is certainly interesting, our experimental and sampling design does not allow for mechanistic or causal interpretation. Our data are limited because (1) we have no repeated seasonal measurements, (2) we sampled at the end of one growing season and at the beginning of the next growing season, which could have led to differences in herbivore community and herbivory between sampling campaigns (Meyer et al., 2017), and (3) leaf traits and the plant's metabolome can vary within and between years (Peters et al., 2018). Dedicated experiments that repeat sampling throughout the season (e.g. Marr et al., 2021) and in multiple consecutive years are necessary to analyse the seasonal variation in the relationship between the plant's metabolome, resource acquisition traits, and herbivory.

We also discovered that the metabolite diversity in grasses and forbs varied differently to changes in resource acquisition-associated community-weighted traits. These contrasting responses are likely due to differences in defensive strategies. Grasses possess silica crystals providing mechanical protection from herbivory (Massey & Hartley, 2009), while forbs invest in carbon-based defences, such as phenolics (Cooke & Leishman, 2012; Larson, 1988). Moreover, grasses and forbs differ in their associations with soil biota, such as the symbiosis with mycorrhizal fungi, which can contribute to diverging metabolomic responses (Chialva et al., 2018; Ristik et al., 2019). While the difference between grasses and forbs was not the focus of our study and we only analysed a small subset in each functional group, our results stress the importance of including functional group identity to improve predictive models analysing plant-herbivore interactions.

While the present experiment provides novel insights into above-below-ground relationships, additional experiments should manipulate and disentangle the individual and interactive roles of plant and soil biodiversity in driving changes in plant metabolomes and herbivory rates (Peters et al., 2018). Such studies should be conducted in the presence and absence of above-ground herbivores (Seabloom et al., 2017), to assess if above-ground herbivory modulates plant and soil biodiversity effects on the metabolome, for example via induced responses (Peters et al., 2018). Preferably, these studies should include specialist and generalist herbivores as well as different feeding types (Mithöfer & Boland, 2008).

Taken together, the present study provides support for the existence of tight relationships between plant diversity, soil biota communities, plant metabolomes, and above-ground herbivores. Our results especially suggest that the plant metabolome is an important functional trait (Walker et al., 2022) that can aid to explain more variation (22%) in herbivory than commonly used morphological and physiological traits (on average 12.7%; van der Plas et al., 2020). By including metabolomic analyses, we advanced our knowledge on the potential mechanisms linking plant diversity and herbivory rates via changes in plant metabolomes (Peters et al., 2018). In addition, we highlight that the soil nematode and microbial communities shape above-ground interactions and that season and plant functional group identity should be considered when analysing such relationships. Our study creates a framework for future experimental research which can further illuminate the underlying mechanisms through targeted and independent manipulation of the plant, soil biota, and herbivore community. It thereby expands our capability to better characterize the complex nature of multitrophic interactions above and below the ground.

**AUTHOR CONTRIBUTIONS**

Christian Ristik, Alexander Weinhold, Nico Eisenhauer and Nicole M. van Dam conceived the study. Christian Ristik, Christiane Roscher, Fredd Vergara and Alexander Weinhold collected the data. Marcel Ciobanu identified and computed the nematode indices. Christian Ristik and Yvonne Poeschl analysed the data. Christian Ristik, Alexander Weinhold, Nico Eisenhauer and Nicole M. van Dam interpreted the data. Christian Ristik wrote the manuscript under guidance of Alexander Weinhold, Nico Eisenhauer and Nicole M. van Dam. All authors contributed critically to the drafts and gave final approval for publication.

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CONFLICT OF INTEREST
The authors declare no conflict of interest. Nicole van Dam is an Associate Editor for Journal of Ecology, but took no part in the peer review or decision-making process for this paper.

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Data available from the Dryad Digital Repository https://doi.org/10.5061/dryad.d51c5b046 (Ristok et al., 2022).

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