Contrasting responses of above- and below-ground herbivore communities along elevation

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Abstract

Above- and below-ground herbivory are key ecosystem processes that can be substantially altered by environmental changes. However, direct comparisons of the coupled variations of above- and below-ground herbivore communities along elevation gradients remain sparse. Here, we studied the variation in assemblages of two dominant groups of herbivores, namely, above-ground orthoptera and belowground nematodes, in grasslands along six elevation gradients in the Swiss Alps. By examining variations of community properties of herbivores and their food plants along montane clines, we sought to determine whether the structure and functional properties of these taxonomic groups change with elevation. We found that orthoptera decreased in both species richness and abundance with elevation. In contrast with aboveground herbivores, the taxonomic richness and the total abundance of nematode did not covary with elevation. We further found a stronger shift in above- than below-ground functional properties along elevation, where the mandibular strength of orthoptera matched a shift in leaf toughness. Nematodes showed a weaker pattern of declined sedentary behavior and increased mobility with elevation. In contrast to the direct exposure of aboveground organisms to the surface climate, conditions may be buffered belowground, which together with the influence of edaphic factors on the biodiversity of soil biota, may explain the differences between elevational patterns of above- and below-ground communities. Our study emphasizes the necessity to consider both the above- and below-ground compartments to understand the impact of current and future climatic variation on ecosystems, from a functional perspective of species interactions.

Keywords Environmental gradient · Species richness · Herbivory · Functional traits · Ecological interactions

Introduction

The study of community properties along environmental gradients is a fundamental and necessary step toward better predictions of the functioning of natural systems (Garnier et al. 2016; Mayor et al. 2017). In most of the taxonomic groups and trophic levels studied so far, biodiversity shifts along elevation gradients have been generally associated with the variation in climate that exists between low- and...
highlands (Rahbek 1995; Hodkinson 2005; McCain and Grytnes 2010; McCain and Grytnes 2010; Guo et al. 2013). One of the most accurate climatic correlates of elevation is temperature, and the linear decrease of temperature with elevation represents a major constraint for species distribution along elevation gradients (Peters et al. 2016). Temperature may stimulate metabolic and diversification rates, fostering more species in warm environments (Clarke & Fraser 2004; Allen et al. 2006; Hatfield and Prueger 2015), while cold extremes may generate a stress effect, filtering the composition of communities that can occupy the alpine belt (Körner 2003; Sierra-Almeida et al. 2009; Hoiss et al. 2012; Buckland and Huey 2016). To date, our understanding of the variation of species assemblages along montane gradients is biased toward aboveground organisms (McKenzie et al. 2013), while the ecological signature of elevation on soil communities remains largely unknown (Martiny et al. 2006; Pellissier et al. 2014a).

Above- and below-ground communities both influence important ecological processes, including nutrient cycling and trophic interactions (Bardgett et al. 1998; Adams and Wall 2000; Wardle et al. 2004). Their common role in mediating some of the most important aspects of ecosystem functioning calls for more comprehensive studies on the coupled variation of both systems along environmental gradients. Climatic variation along elevation clines may have different effects on below- compared to above-ground organisms (Adams and Wall 2000) as surface conditions can be buffered in the soil compartment (Beyens et al. 2009), which may result in different species turnover rates between both systems along environmental gradients. For instance, a general decline in species richness along elevation or latitude has been documented for multiple aboveground taxonomic groups of plants and animals (Hodkinson 2005; Sharma et al. 2009; McCain and Grytnes 2010; Guo et al. 2013; Descombes et al. 2017a, b). In contrast, the few studies conducted on soil systems thus far have not demonstrated a similar association between elevation and taxonomic richness and/or abundance (Margesin et al. 2009; Fierer et al. 2011; Jarvis et al. 2015; Kergunteuil et al. 2016; van den Hoogen et al. 2019) or even found an increase of these community indices with elevation. For instance, Pellissier et al. (2014a) reported a higher fungal richness and phylogenetic diversity at lower temperatures and higher moisture conditions. Similarly, Kergunteuil et al. (2016) found that the abundance and metabolic footprint of soil nematodes increased when moving from forested low-elevation sites up to Alpine grasslands. If such contrasting patterns of biodiversity generalize to others components of trophic networks, such as major herbivore groups, it would lead to a shift and decoupling of ecosystem processes, from above- to below-ground, depending on where the system is along the climatic gradient (Adams and Wall 2000; Hooper et al. 2000).

The species interactions that drive ecosystem processes are mediated by functional traits. Reciprocally, functional characteristics modulate ecological interactions and can be used together with community structure indices to understand community responses to environmental gradients (Diaz et al. 1998). The choice of relevant functional traits that mediate species–environment interactions through abiotic and biotic mechanisms presupposes a good knowledge of the taxa under study and their ecology (Mlambo 2014; Nock et al. 2016). Studies on the variation of functional traits with elevation are common for plant traits such as height (Moles et al. 2009), leaf structure (Reich et al. 1998), or leaf resource content (Shipley and Vu 2002). For insects, a decrease in body size, in wing length, or shifts in coloration associated with heat absorption have also been related to the elevation gradient (Hodkinson 2005). In contrast, the characterization of functional strategies for soil communities along elevation remains overly marginal (Kergunteuil et al. 2016), although some recent studies have attested to the functional response of soil organisms to the elevation gradient (Bond-Lamberty et al. 2016; Looby et al. 2016). Yet, as underground systems are less exposed to climatic conditions (Adams and Wall 2000), we expect that functional trait composition and species turnover along elevation is dictated by different structural forces than those acting on the surface.

Climatic variation shapes functional trait composition at the community level (Diaz et al. 1998), in turn, potentially modifying the mode and strength of species interactions (Hillyer and Silman 2010), and ultimately, ecosystem processes (Tylianakis et al. 2008). For instance, plant–herbivore interactions are modulated by the coupling of plant resistance and feeding-related traits (Moles et al. 2013). In this regard, shifts in the composition and functional identity of herbivores along elevation gradients (Hodkinson 2005) has been shown to reduce the intensity of herbivory (Rasmann et al. 2014), and should, therefore, change the investment of plant defences along elevation (Pellissier et al. 2012). As a consequence, variation in plant and herbivore functional traits along environmental gradients should modify species interactions within ecosystems. Consequently, studying both above- and below-ground community composition and functional traits could provide valuable information on the plant–herbivore relationships within the two sub-systems along elevation gradients and enable a stratified characterization of ecosystem functioning.

In this study, we explore structural properties and functional constituents of above- and below-ground herbivore communities along six elevational transects in the Swiss Alps. We compare the assemblage of two groups of herbivores: orthoptera that feed on leaves, and herbivore nematodes that feed on roots. Orthoptera is among the most influential aboveground insect herbivores in open habitats, removing high percentages

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of plant biomass in natural meadows (up to 30%, Blumer and Diemer 1996). Herbivory pressure exerted by orthoptera has been reported to decrease at high elevation, as species richness and abundance are reduced at that level (Scheidel and Bruelheide 2001; Hodkinson 2005; Descombes et al. 2017a). Nematodes are belowground organisms characterized by high functional, trophic, and taxonomic diversity. Herbivore nematodes are capable of substantial uptakes of plant root biomass (Ingham and Detling 1990; Hodda et al. 2009) and are distinguished into functional feeding groups including five plant root-feeding strategies (sedentary endoparasites, migratory endoparasites, semi-endoparasites, ectoparasites, and epidermal cells and root-hair feeders; Yeates et al. 1993). Responses of herbivore nematodes to elevation have seldom been described with reports of decrease (Dong et al. 2017) or increase (Kergunteuil et al. 2016) in taxonomic richness and abundance at a higher elevation. To date, concomitant investigations of both orthoptera and nematode communities, their functional responses and those of local plant communities along elevation, are still lacking. In this study, we address these questions with the following expectations:

1. Species richness and total abundance should decline more above- than below-ground along elevation because strong aboveground temperature shifts are buffered belowground. Differential rates of change in communities along elevation may lead to decoupled above- and below-ground community composition.

2. The functional traits of aboveground herbivore communities should vary more along elevation than the abundance of the functional feeding groups of belowground herbivores nematodes (e.g. epidermal/root hair feeders, ectoparasites) because of attenuated abiotic variations. In particular, we expect that feeding strategies, reflected by the above-mentioned functional properties, should be more stable among below- compared to above-ground herbivores.

3. Variations in herbivore traits should match those of plant traits along elevation. In particular, we expect the mandibular strength of orthoptera to increase toward high elevation in response to increasing plant leaf toughness in more stressful environmental conditions. We further expect different responses to the elevation between specialists (e.g. sedentary and migratory endoparasitic nematodes) and generalist feeders due to change in plant traits and edaphic conditions.

Methods

Study sites

To study above-belowground plant–herbivore interaction along elevation gradients, we selected six elevation transects spanning the major macro-climatic and environmental conditions (i.e., climate and bedrock type) of the Central Alps (see Fig. S1). We selected eight study sites per elevation transect in open, non-woody areas with elevations ranging from 578 m to 2417 m, and an average elevation difference between sites of 240 m. Study sites were located in semi-natural grasslands characterized by low impact from agricultural practices in land-use and pasture. Most of the low to medium elevation sites corresponded to dry meadows and pastures in the Swiss inventory for national protected areas (Federal Act on the Protection of Nature and Cultural Heritage (LPN), status as of 1 January 2017; Article 18. Protection of animal and plant species, https://www.admin.ch), while high elevation sites were situated in alpine meadows with no mowing and low grazing pressure. Herbivores and plant surveys took place during the summers of 2016 and 2017 within a square area of 10 m × 10 m. Study plots were positioned within the study zone to represent the dominant vegetation type of the surrounding natural environment and were set a minimum three meters away from forest edges when present. Plant, orthoptera and nematode herbivore inventories were compiled when communities reached maximal species richness and abundance, gradually surveying low to high elevation sites, between early June and the end of August.

Herbivore surveys

Among orthoptera, the Caelifera suborder includes only strictly vegetarian species, while Ensifera are omnivorous, but largely feed on plant material (Ingrisch and Köhler 1998; Baur et al. 2006). We, therefore, included both suborders in the subsequent analyses. Orthoptera surveys were performed under optimal weather conditions for insect activity, between 10 a.m. and 5 p.m. on days with maximal sunshine levels. Species determination was done through visual and auditory identification using the reference work for Swiss Orthoptera (Baur et al. 2006). We estimated the abundance of each species following a “Z” sampling pattern across the 100 m² study area by counting all adult orthoptera specimens that were visually detected without distinguishing sex. Nematode sampling consisted in a random sampling of 15–20 soil cores (2 cm diameter, 10–25 cm depth) within a 2 m × 2 m area located in the
middle of the 100 m² study area to obtain 1 kg of soil after the removal of all rock pieces greater than 2 cm in diameter. The bulk soil was then mixed homogenously, and a sample of 300 g was collected and stored at 4 °C before nematode extraction. Twenty-five grams of fresh soil were air-dried and used to calculate soil moisture. Two-hundred grams of fresh soil were used to extract nematodes from each sample using a modification of the sieving and Baermann funnel method (Barker 1985). Once extracted, all nematodes were counted under a binocular microscope. After counting, nematodes were concentrated and mounted alive in a slide and at least 150 nematodes were identified to the genus or family level according to Bongers (1989). Nematodes were identified at 200 × or 400 × under an inverted microscope, and nematode abundances expressed as no. of nematodes standardized to 100 g of dry soil.

**Plant surveys**

The vegetation inventories were first conducted in a circular subplot of 9 m² positioned in a floristically homogenous area within the 100 m² plot, across which we further searched for additional rare species. Plant species determination was done following Swiss Floras (Lauber et al. 2012; Eggenberg and Möhl 2013). We visually estimated the relative cover of each plant species according to a 9-level scale (< 0.25, 0.25–0.5, 0.5–1, 1–5, 5–15, 15–25, 25–50, 50–75 and > 75%). Low abundance classes (i.e. < 0.25 and 0.25–0.5) were used to accurately account for the difference in the relative abundance of rare plant species. The median values of these classes were used in all subsequent statistical analyses.

**Herbivore functional trait measurements**

To measure orthoptera incisive strength, we collected insect specimens in falcon tubes that were killed with cold treatment at − 20 °C. Mandibular trait measurements were performed following the approach described by Ibanez et al. (2013), for 90% of the study species using three specimens of each species and sex. When a species was observed in more than one study site, collection points were selected within and between the study areas to cover the full extent of elevation and geographical range of each species. After the extraction of the left mandible, we took photographs of each mouthpiece in triplicate using a high-resolution measuring digital microscope (Leica DVM6, Leica Microsystems, GmbH, Wetzlar, Germany), which, together with the high-resolution photo stacking option available in Leica Application Suite X (LAS X) and Leica Map Premium software (Leica Microsystems), maximizes photographic resolution to increase measurement accuracy. The mandibular incisive strength \( F_I \) was obtained \( F_I = F_A L_A / R_1 \) using the formula, where \( F_A \) is a proxy for the mandibular section area, \( L_A \) is the adductor muscle lever, \( L_I \) is the incisive lever and \( R_1 \) is the incisive region length (Ibanez et al. 2013). Values were measured on mandible photographs using the ImageJ image processing program (Rueden et al. 2017). For nematodes, we used the Nematode INdicator Joint Analysis system (NINJA, Sieriebriennikov et al. 2014) to assign feeding groups to each nematode taxon, and extracted only the genus corresponding to plant-parasitic nematodes. The functional classification within herbivore nematodes includes (a) epidermal/root hair feeders, which generally induce low phytopathological damage, (b) ectoparasites, which feed on superficial root tissues, (c) semi-endoparasites, which partially introduce their body into the root tissue, (d) migratory endoparasites, which migrate through the root tissue, and (d) sedentary endoparasites, which complete their life cycle inside roots, thus establishing a permanent feeding site and inducing the formation of root galls or cysts.

**Plant functional trait measurement**

Leaf functional traits were collected for 79% of the plant species occurring in the plant surveys. We measured four leaf functional traits that were initially considered as relating to climatic disturbance, but could also confer resistance to herbivory since they reflect leaf toughness and resource acquisition: specific leaf area (SLA), leaf dry matter content (LDMC), punch strength, and carbon-to-nitrogen ratio (C/N). We sampled a minimum of three individuals per species and site, however, when the species occurred at more than one study site along the elevation gradient, we accounted for potential intra–species variation by increasing the number of replicates across different elevations to a maximum of twelve by multiples of three for each additional site. We selected well-developed and healthy leaves that were moisturized directly after collection and stored at 4 °C with additional moisture for a maximum of 24 h before trait measurement. Measurements of SLA (calculated as the area of a fresh leaf divided by the dry weight and expressed in mm² mg⁻¹) and LDMC (the ratio of the leaf dry mass to the water-saturated weight in mg g⁻¹) were performed following standard procedures (Cornelissen et al. 2003; Vaieretti et al. 2007). Dry weight was measured after oven-drying the fresh leaves at 55 °C for a minimum of 72 h. The leaf force to punch, which is considered to be the trait that best captures leaves’ mechanical properties that are relevant for herbivory (Sanson et al. 2001; Ibanez et al. 2013), was measured using a digital force gauge that records the force required to pierce the leaf lamina (IMADA CO., LTD. Toyohashi, Japan). Measurements were taken on fresh leaves with the measurement point selected to avoid leaf veins. Values are expressed in MN m⁻² and were corrected for leaves with widths of less than the diameter of the gauge.
pin (2 mm). Leaf width was measured using a digital caliper gauge (0.01 mm precision). Total organic carbon (C) and nitrogen (N) amounts were determined for 89% of all species by dry combustion of ground leaf material using a CN elemental analyzer (NC-2500 from CE Instruments, Wigan, Lancashire, United Kingdom).

**Taxonomic richness and abundance along elevation**

Species richness of plants and orthoptera insects, and the genera richness of herbivore nematodes were calculated as the sum of the number of taxa identified at each study site. Abundance was estimated using the total specimen count at each site for the orthoptera and no. of individuals per 100 g of dry soil for nematodes assemblages. The abundance of nematodes for one particular site that differed from others by a particularly thick layer of dead plant organic matter exceeded the median value of the total number of nematodes by a factor of ten. This likely indicates a bias in the sampling process such as the accidental inclusion of soil containing a non-representative high level of plant organic matter resulting in the extremely high value of nematodes abundance. This site was, therefore, removed from subsequent analyses. Relationships between elevation and response variables were tested using regression models that included the transect identity as a random factor using functions within the lme4 and lmerTest R packages (Bates 2008, Kuznetsova et al. 2017). All analyses were conducted using R (R Core Team 2019). The variation of orthoptera and nematode taxonomic richness and orthoptera abundance along elevation was tested using generalized linear mixed-effects models for the Poisson distribution of count data (glmer) and we use a second-degree polynomial regression when the shape of the relationship was not linear. We also tested the variation of species richness and abundance along the elevation for both orthoptera suborders independently using the same model types. The nAGQ parameter was set to 0 for orthoptera and nematode models to ensure model convergence. The variation of the plant taxonomic richness was tested using a second-degree polynomial regression in a generalized linear model with a negative binomial family to avoid overdispersion of the residuals. We related nematode abundance data to elevation using a linear mixed effect model for Gaussian distribution data with a logarithmic transformation (log + 1). Overdispersion of the residuals of the models was verified by simulating standardized residuals from the fitted models in DHARMa R package (Hartig 2020).

**Co-inertia analyses and temperature variation along elevation**

We conducted a co-inertia analysis to test the coupling of above- and below-ground herbivore communities. For each transect, we first performed a PCA on abundance matrices of orthoptera and nematodes independently using the function “dudi.pca” in the ade4 package (Thioulouse et al. 2018). We then used the first two factorial axes of the PCAs to apply the co-inertia criterion procedure and quantify the co-variance between the two tables (co-inertia function) using the RV coefficient. Values close to one indicate maximal co-variance between matrices. To further investigate how surface climate might be buffered in the soil, we compared the variation of above- vs. below-ground temperature along the elevation using soil temperature measured with data loggers (DS1921G-F5 HomeChip, Farnell, Zug, Switzerland) from October 2017 to September 2018 and air temperature data supplied by MeteoSwiss (2 m aboveground; 2020, Swiss Federal Office of Meteorology and Climatology).

**Functional traits along elevation**

The functional classification of herbivore nematodes was used to reflect the functional aspects of belowground communities. We summed the abundance of each herbivore group to quantify their abundance at each site and performed a logarithmic transformation of the data (log + 1) to fulfil model assumptions. The mean community values (CM) of functional traits for orthoptera (i.e., incisive mandibular strength) and plants (i.e., SLA, LDMC, punch strength, C/N) were obtained by averaging the sum of the trait values by the total number of species. The community weighted means (CWM) accounting for species abundance were computed for the same traits and each site using where R is the number of species, Pi is the relative abundance of the species i, and ti is the mean trait value of the species i. While the two metrics are based on means, CM reflects the average of life-strategies occurring within a community weighting all species equally, and CWM accentuates the ecological role of dominant species (Garnier et al. 2004). We used linear mixed-effects models with Gaussian data distributions to test whether abundances of the nematode functional groups were associated with the elevation gradient. For the variation of orthopteran functional trait metrics along elevation (i.e., CM and CWM), we fitted linear mixed-effects models for Gaussian data distribution. The weighted and unweighted CM of plant functional traits along the elevation were analyzed using the same models. We excluded the three lowest elevation sites of the Salgesch transect since they were more similar.
to steppic environments, with plant traits strongly biased toward functional responses that are typical of extremely dry environmental conditions (Volaire 2008; Delarze et al. 2015).

Results

Taxonomic richness and abundance along elevation

From vegetation surveys, we identified 526 plant species belonging to 251 genera and 69 families. From orthoptera surveys, 48 species including 19 Ensifera and 29 Caelifera taxa were determined. In total, we identified 55 nematode genera, comprising 14 herbivore genera that corresponded to five different plant-parasitic types (i.e., ectoparasites, epidermal/root hair feeders, semi-endoparasites, migratory, and sedentary endoparasites). The nematode taxa identified, their functional classification, and percentage contribution of herbivore functional groups to the nematode community are indicated in Table S1. We found that the species richness of orthoptera and plant significantly decreased with elevation following linear and hump-shaped relationships, respectively (Fig. 1a, c, Table 1). When tested independently for Caelifera and Ensifera, the same declining trends were found, except for the variation of Caelifera species richness along

Fig. 1 Illustration of the relationships between the elevation and community structure indices using linear and generalized linear mixed effects models with the taxonomic richness of (a) orthoptera, (b) herbivore nematodes, and (c) plants, and the specimen abundance of (d) orthoptera communities and (e) herbivore nematodes (log-transformed). Regression lines of fitted values and standard error intervals are only displayed for relationships that are statistically significant. A significant decrease in taxonomic richness was found for orthopteran and plant but not for herbivore nematode communities. The abundance of orthoptera also decreased with the elevation, a signal that was not retrieved for herbivore nematodes

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elevation (Fig. S2, Table S2). This result contrasts with the nematode distribution patterns that presented no variation along the transects for the genus richness and the abundance (Fig. 1b, e, Table 1). Orthoptera abundance displayed a significant polynomial relationship with elevation, with the highest values found at mid-elevation (Fig. 1d, Table 1).

**Co-inertia analyses and temperature variation along elevation**

Along the co-inertia analysis performed on each transect individually, the PCAs’ first two axes together explained more variance in nematode abundance than orthoptera (Table 3), accounting, on average, for 41% and 27% of the variance in orthoptera, and 79% and 15% in nematodes, respectively. The analyses suggested a partial decoupling between above- and below-ground herbivore communities with RV coefficients ranging from 0.20 to 0.44 (see Table 3). When comparing above vs. below-ground temperature along the elevation, we found that the monthly average and minimum temperature are generally higher belowground than at the surface (Fig. S3a & S3b) and that the temperature variance considerably decreased in the soil (Fig. S3c).

**Orthoptera and nematode traits along elevation**

Among the five herbivore nematode functional groups, we only found that sedentary parasites decreased significantly in abundance with elevation (Fig. 2d, Table 2). While no clear trend in abundance variation was visible for ectoparasites, epidermal/root hair feeders, or semi-endoparasites (Fig. 2a, b, e, Table 2), the abundance of migratory endoparasites was significantly higher at high elevation (Fig. 2c, Table 2). The CM of the incisive mandibular strength of orthoptera increased with elevation for males, while for females, values were stable along the gradient (Fig. 2f, Table 2) and largely outreached those of males. The CWM of this trait followed the same pattern (Fig. S4, Table S3).

**Plant traits along elevation**

The CM of SLA and LDMC showed a significant negative relationship with elevation (Fig. 3a, b, Table 2), whereas a positive relationship was found for punch strength and C/N (Fig. 3c, d, Table 2), the latter being concurrently explained by a decrease in nitrogen and an increase in carbon. The results of the regression models applied to CWM generally indicate similar trends to those found for CM (Fig. S5, Table S3).

**Discussion**

Variations in above- and below-ground herbivore communities along elevation gradients have rarely been directly compared to date. Here, we have focused on two dominant groups of herbivores in grasslands, aboveground orthoptera and belowground nematodes, and studied variation in assemblages along six elevation gradients in the Swiss Alps. We examined variations in community taxonomic properties and the functional traits of above- and below-ground herbivores, and their food plants along montane gradients. We have shown that variations in community indices along elevation considerably differ between above- and below-ground organisms. The evenness of nematode taxonomic richness along the elevation gradient contrasted with the decrease in orthoptera species richness at the highest elevation and may be related to the tempering effect of stressful abiotic conditions that limits environmental filtering in the soils. The response of above- and below-ground communities to elevation also differed as regard to abundance. We found no relationship for nematodes contrasting with an upward decline of orthoptera along the gradient. We have also

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**Table 1** Slope coefficients (slope estimate), P values and standard errors (SE) for linear and generalized linear mixed effects models testing the relationships between elevation and community indices

| Species richness | Linear model | 2nd degree polynomial model |
|------------------|--------------|---------------------------|
|                   | Slope Estimate | P value | SE | Slope Estimate | P value | SE |
| Orthoptera       | −0.0003       | <0.001        | 0.0001 |
| Herbivore nematodes | −0.0001   | 0.584       | 0.0001 |
| Plants           | 0.240         | 0.16        | 0.17 | −0.696      | <0.001 | 0.169 |
| Abundance        | Orthoptera    | −0.273       | 0.006 | 0.10 | −1.053 | <0.001 | 0.099 |
| Herbivore nematodes | 0.0004   | 0.245       | 0.0003 |

Values are provided for linear and 2nd degree polynomial models (when applicable) that included the transect identity as a random factor to quantify the variation of the species richness of orthoptera, herbivore nematodes, plants, and the species abundance of orthoptera and herbivore nematodes along the elevation. Overdispersion of the residuals was verified by simulating standardized residuals from the fitted models in DHARMa R package (Hartig 2020)
found a stronger shift in above- than below-ground functional properties along elevation. The mandibular strength of orthoptera matched a shift in leaf punch strength along elevation, while nematodes showed a pattern of elevational variation only through a decline in sedentary endoparasites and an increase of migratory endoparasites with elevation.
Abundance of nematode functional groups

Table 2  Slope coefficients (slope estimate) and \(P\) values and standard errors (SE) for linear mixed effects models testing the relationships between the elevation and the abundance of herbivore nematode functional groups using a logarithm transformation, male and female orthoptera incisive mandibular strength, and plant functional traits

| Functional groups                | Slope Estimate | \(P\) value | SE  |
|----------------------------------|----------------|-------------|-----|
| Ectoparasites                    | 0.0002         | 0.56        | 0.0004 |
| Epidermal root hair feeders      | 0.0002         | 0.66        | 0.0003 |
| Migratory endoparasites          | 0.001          | < 0.05      | 0.0004 |
| Sedentary parasites              | − 0.001        | < 0.001     | 0.0003 |
| Semi-endoparasites               | 0.0004         | 0.32        | 0.0004 |
| Orthoptera mandibular strength   | Male           | 0.00003     | < 0.001 | 0.00001 |
|                                  | Female         | 0.00001     | 0.55  | 0.00002 |
| Plant functional traits          | SLA            | − 0.01      | 0.01  | 0.0003 |
|                                  | LDMC           | − 0.01      | < 0.01 | 0.003  |
|                                  | Punch strength | 0.0002      | < 0.001 | 0.00004 |
|                                  | C/N            | 0.001       | < 0.001 | 0.0004 |

The abundance of the functional groups of belowground herbivore nematodes displayed weaker variation with elevation than the functional traits of aboveground herbivore communities

We discuss our results in terms of abiotic conditions that act above- and below-ground, and examine their implications for shifts in plant–insect herbivore interactions along elevation gradients.

The species richness and abundance of orthoptera decreased with elevation (Fig. 1a, d), while for nematodes, no variation was observed (Fig. 1b, e). A decrease in aboveground biodiversity is generally associated with a shift in climatic conditions along elevation although such strong climatic gradients may not extend to belowground communities (Bryant et al. 2008; Fierer et al. 2011). If environmental conditions apply similarly above- and below-ground, we should find an equivalent variation in taxonomic richness and abundance for above- and below-ground assemblages. Contrary to this expectation, the co-inertia analyses indicated generally low congruence between orthoptera and nematode communities (Table 3), suggesting that the environmental gradients shaping soil communities are different from those acting on the surface. Among the environmental conditions that shift strongly with elevation, the temperature gradient and in particular, thermal variability may be buffered for soil biota (Fig. S3), differentially shaping the structure of above-compared to below-ground communities (Beyens et al. 2009; Ryalls et al. 2013). We found a monotonical decrease in orthoptera species richness that directly follows the gradual decrease in temperature with elevation (Barry 2008), while orthoptera abundance peaks at mid-elevation, where plant species richness is highest (Fig. 1c; Haddad et al. 2001; Descombes et al. 2017b). In contrast, the taxonomic richness and the abundance of nematodes did not vary with elevation (Fig. 1b, e). This may indicate the limited influence of temperature decline and variability along the elevation gradient on soil communities. It also suggests that edaphic factors (e.g., soil fertility and humidity) may vary along montane clines to shape soil communities, which, together with buffered surface conditions, may result in patterns that strongly diverge from those observed aboveground (Kergunteuil et al. 2016; van den Hoogen et al. 2019). In agreement with Kergunteuil et al. (2016) who analyzed nematode community properties along elevation transects, our study found that species diversity and abundance were higher in alpine meadows. Hence, indices of community structure that respond to elevation should fundamentally differ depending on whether the study compartment is located above or below-ground. These results also parallel biodiversity patterns occurring at a wider biogeographic scale. The distribution of orthopteran diversity was found to be higher in Southern Europe (Hochkirch et al. 2016) whilst the abundance and diversity of belowground biota were shown to increase in temperate regions (van den Hoogen et al. 2019). Alongside variation in taxonomic richness and abundance, we also expected a weaker variation of functional properties in above-compared to below-ground communities.

Both orthoptera and nematode communities showed some degree of functional changes with elevation. The CM of mandibular strength for male orthoptera increased with increasing elevation, while the community values for females were steady and systematically greater. Morphological differences between sexes have been documented for this insect group (Laiolo et al. 2013), and we showed here that sexes responded differently to elevation. Because mandibular strength influences plant ingestion, higher values found for females could underscore the greater importance for this sex to bypass plant mechanical barriers at any elevation. This would ensure a nutrient intake that is capital for the production of eggs (Hochkirch and Gröning 2008). For males, although a minimum nutrient intake is necessary to guarantee survival, they may invest less in their feeding ability. Hence, the constant and high mandibular strength of females, and the increase for males with elevation possibly represents a response to plant physical resistance at high elevation where greater mandibular strength would help the ingesting of tougher plants (Ibanez et al. 2013). With regards to nematodes, we found a significant increase in abundance for migratory endoparasitic nematodes and a significant decrease for sedentary endoparasites, with no variation along elevation for nematodes characterized as ectoparasites, semi-endoparasites, or epidermal/root hair feeders. Both sedentary (Meloidogyne, Heterodera) and migratory (Pratylenchus) endoparasitic nematodes are able to feed on large numbers of plant species, being extremely...
polyphagous (Jones and Fosu-Nyarko 2014; Truong et al. 2015). Plant-parasitic nematodes are seldom studied in natural systems, and little information is available on the effects of elevation on parasitic nematodes. The limited information available, however, reported increased \textit{Pratylenchus} and low \textit{Meloidogyne} abundances at high elevations in tropical areas (Fogain 2001; Gaidashova et al. 2009; Avelino et al. 2009; Kamira et al. 2013). Our results showed that low-specialized belowground herbivory is equally distributed along elevation transects, while the most specialized herbivore conditions, represented by sedentary and migratory endoparasitism (Perrine-Walker 2019), present peak abundances at different elevations, respectively at 578 m and 2346 m. Competition among \textit{Pratylenchus} and \textit{Meloidogyne} has been detected in crops under experimental and natural conditions (Avelino et al. 2009; Fontana et al. 2015), and similar processes have been described in natural systems between \textit{Pratylenchus} and other endoparasitic nematodes such as \textit{Heterodera} (Brinkman et al. 2005). Besides plant functional traits and other ecological factors, such competition may play a role in structuring herbivore nematode communities across environmental gradients.

If functional traits govern the trophic interplay between plants and herbivores, and reflect community responses to environmental changes, they also reshape plant–insect herbivore interactions along elevation gradients. Influenced or not by abiotic constraints, the abundance of herbivores is also involved in modulating the functional interface between herbivore feeding abilities and plant defence (War et al. 2012). We have investigated a set of plant functional traits that represent physical resistance against herbivory. Among the four plant functional traits studied, we found that the CM of SLA and LDMC decreased with increasing elevation, while punch strength and the C/N ratio showed opposite relationships. All the tested plant functional traits displayed a significant relationship with the elevation, that was positive for the SLA and LDMC and negative for the C/N and the punch strength.

![Fig. 3](image-url) Relationships between plant functional trait community means (CM) and elevation using linear mixed effects models for (a) SLA, (b) LDMC, (c) punch strength, and (d) C/N. Regression lines of fitted values and standard error intervals are only shown for significant relationships.
Transect edaphic factors govern to shape soil communities. Surface climatic conditions might be buffered belowground where and below-ground herbivore abundance matrices suggesting that the PCA matrices and RV coefficients of the co-inertia analyses performed on axes of the PCAs applied to orthoptera and nematode abundance sect are given the percentage of variance explained by the first two Table 3 should be studied accordingly. The decline in herbivore biotic forces that are unique to the study compartment, and to those belowground, are controlled by a set of abiotic and herbivore relationships in aboveground systems, in contrast communities. These findings suggest that the plant–insect functional responses is of greater amplitude for aboveground matrices are partially decoupled and that the variation of in the soil. Our study also found that herbivore abundance along elevation gradients, plant defences in roots may not show the same decline as documented for leaves (Pellissier et al. 2012; Callis-Duehl et al. 2017). We, therefore, propose that plant defence and herbivory relationships of above- vs. below-ground compartments react differently to environmental change, which calls for a greater effort to document belowground plant–herbivore interactions. However, the factors that operate in the structuring of plant–herbivore interactions remain unexplored, particularly for soil systems. Given the specificities of above- and below-ground systems, we believe that a line of research that considers both community types, in a functional and network perspective, is required to identify the drivers of species interaction, and to anticipate how climate change will affect distinct ecosystem compartments and functioning.

| Transect | Orthoptera | Nematodes | RV coefficient |
|----------|------------|-----------|----------------|
|          | Axis 1     | Ax i s 2  | Axis 1         | Axis 2         |
| Bex      | 30.7       | 29.7      | 87.4           | 8.7            | 0.35 |
| Calanda  | 55.3       | 21.7      | 93.8           | 5.1            | 0.25 |
| Salgesch | 38.8       | 32.6      | 75.3           | 24.4           | 0.20 |
| Grindelwald | 44.7 | 26.5      | 69.4           | 18.0           | 0.20 |
| Martigny | 37.3       | 25.6      | 54.9           | 29.6           | 0.44 |
| Faido    | 38.6       | 24.5      | 94.6           | 4.2            | 0.25 |

The co-inertia analyses indicated a partial decoupling between above- and below-ground herbivore abundance matrices suggesting that the surface climatic conditions might be buffered belowground where edaphic factors govern to shape soil communities trends. The increase in C/N and punch strength are indicators of a greater physical resistance of alpine plant communities. This is in line with the review of Moreira et al. (2018), who concluded that plants adapted to stressful environmental conditions tend to invest more in constitutive defence, which include some chemical but mostly physical defences. Changes in plant leaf traits correspond to the increase in mandibular strength in male orthoptera with increasing elevation. However, we lack data on root properties to properly assess changes in nematode communities. Due to the technical difficulties involved in measuring root functional traits, the imprint of herbivore abundance on root defence is relatively unexplored, with only a few studies focusing on chemical responses to root herbivory (Kaplan et al. 2008; Rasmann et al. 2011). Without knowledge of root defensive traits against herbivory, it remains difficult to connect nematodes and plant functional responses. However, since root biomass generally increases with elevation (WeiLing et al. 2010), plants may support constant herbivory pressures along elevation gradients based on tolerance rather than defensive response. Although this requires further analyses aiming at tracking different defence/tolerance trade-offs in above- and below-ground compartments, our study showed that at the surface, herbivore abundance declines with elevation while no variation or opposite patterns are observed in the soil. Our study also found that herbivore abundance matrices are partially decoupled and that the variation of functional responses is of greater amplitude for aboveground communities. These findings suggest that the plant–insect herbivore relationships in aboveground systems, in contrast to those belowground, are controlled by a set of abiotic and biotic forces that are unique to the study compartment, and should be studied accordingly. The decline in herbivore abundance along the elevation gradient documented for aboveground insect herbivores (Reynolds and Crossley 1997; Pellissier et al. 2014b; Descombes et al. 2017a) may not exist belowground. As a result of the increase in nematode abundance along elevation gradients, plant defences in roots may not show the same decline as documented for leaves (Pellissier et al. 2012; Callis-Duehl et al. 2017). We, therefore, propose that plant defence and herbivory relationships of above- vs. below-ground compartments react differently to environmental change, which calls for a greater effort to document belowground plant–herbivore interactions. However, the factors that operate in the structuring of plant–herbivore interactions remain unexplored, particularly for soil systems. Given the specificities of above- and below-ground systems, we believe that a line of research that considers both community types, in a functional and network perspective, is required to identify the drivers of species interaction, and to anticipate how climate change will affect distinct ecosystem compartments and functioning.

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Compliance with ethical standards Conflict of interest The authors declare that they have no conflict of interest.

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References

Adams GA, Wil DH (2000) Biodiversity above and below the surface of soils and sediments: linkages and implications for global change. Bioscience 50:1043–1048. https://doi.org/10.1641/0006-3568(2000)050[1043:BAABTSI]2.0.CO;2

Allen AP, Gillooly JF, Savage VM, Brown JH (2006) Kinetic effects of temperature on rates of genetic divergence and speciation. Proc Natl Acad Sci 103:9130–9135. https://doi.org/10.1073/pnas.0906155103

Avelino J, Bouvet ME, Salazar L, Cilas C (2009) Relationships between agro-ecological factors and population densities of Meloidogyne exigua and Pratylenchus coffeae sensu lato in coffee roots, in Costa Rica. Appl Soil Ecol 43:95–105. https://doi.org/10.1016/j.apsoil.2009.06.006

Barker KR (1985) Nematode extraction and bioassays. In: Barker KR, Carter CC, JN (eds) An Advanced Treatise on Meloidogyne, Vol 2 Methodology. North Carolina State University Graphics, Raleigh, pp 19–35

Bardgett RD, Wardle DA, Yeates GW (1998) Linking above-ground and below-ground interactions: how plant responses to foliar herbivory influence soil organisms. Soil Biol Biochem 30:1867–1878. https://doi.org/10.1016/S0038-0717(98)00069-8

Barker KR (1985) Nematode extraction and bioassays. In: Barker KR, Carter CC, JN (eds) An Advanced Treatise on Meloidogyne, Vol 2 Methodology. North Carolina State University Graphics, Raleigh, pp 19–35

Barry RG (2008) Mountain weather and climate, 3rd edn. Cambridge University Press, Cambridge

Bates D, Mächler M, Bolker B, Walker S (2008) Fitting mixed-effects models using the lme4 package in R. J Stat Softw. https://doi.org/10.18637/jss.v067.i01

Baur B, Baur H, Roesti C, Roesti D (2006) Sauterelles, grilloins et criquets de Suisse. Haupt, Bern

Beyens L, Ledeganck P, Graae BJ, Nijs I (2009) Are soil biota buffered against climatic extremes? An experimental test on testate amoebae in arctic tundra (Qeqertarsuaq, West Greenland). Polar Biol 32:453–462. https://doi.org/10.1007/s00300-008-0540-y

Blythe P, Diemer M (1996) The occurrence and consequences of grasshopper herbivory in an alpine grassland. Swiss central Alps Arct Alp Res 28:435. https://doi.org/10.2307/1551854

Bond-Lamberty B, Bolton H, Fansler S et al (2016) Soil respiration and bacterial structure and function after 17 years of a reciprocal soil transplant experiment. PLoS ONE 11:e0150599. https://doi.org/10.1371/journal.pone.0150599

Bongers T (1989) The nematodes of the Netherlands. Wageningen Agricultural University, Wageningen

Brinkman EP, Duyts H, Van Der Putten WH (2005) Competition and bacterial structure and function after 17 years of a reciprocal soil transplant experiment. PLoS ONE 11:e0150599. https://doi.org/10.1371/journal.pone.0150599

Borgers T (1989) The nematodes of the Netherlands. Wageningen Agricultural University, Wageningen

Bryant J, Lamanna C, Morlon H et al (2008) Microbes on mountainsides: contrasting elevational patterns of bacterial and plant diversity. Proc Natl Acad Sci 105:11505–11511. https://doi.org/10.1073/pnas.0906155103

Buckley LB, Huey RB (2016) Temperature extremes: geographic patterns, recent changes, and implications for organismal vulnerabilities. Glob Chang Biol 22:3829–3842. https://doi.org/10.1111/gcb.13313

Callis-Duehl K, Vittoz P, Defossez E, Rasmann S (2017) Community-level relaxation of plant defenses against herbivores at high elevation. Plant Ecol 218:291–304. https://doi.org/10.1007/s11258-016-0688-4

Clarke A, Fraser KPP (2004) Why does metabolism scale with temperature? Funct Ecol 18:243–251. https://doi.org/10.1111/j.0269-8463.2004.00841.x

Cornelissen JHC, Lavorel S, Garnier E et al (2003) A handbook of protocols for standardised and easy measurement of plant functional traits worldwide. Aust J Bot 51:335. https://doi.org/10.1071/BT02124

Delarze R, Gontzly S, Eggenberg S et al (2015) Guide des milieux naturels de Suisse: écologie, menaces, espèces caractéristiques. Rossoli, Lausanne, Switzerland

Descombes P, Marchon J, JnN P et al (2017) Community-level plant palatability increases with elevation as insect herbivore abundance declines. J Ecol 105:142–151. https://doi.org/10.1111/1365-2745.12664

Descombes P, Vittoz P, Giusan A, Pellissier L (2017) Uneven rate of plant turnover along elevation in grasslands. Alp Bot 127:53–63. https://doi.org/10.1007/s00035-016-0173-7

Diaz S, Cabido M, Casanoves F (1998) Plant functional traits and environmental filters at a regional scale. J Veg Sci 9:113–122. https://doi.org/10.2307/3237229

Dong K, Moroenyane I, Tripathi B et al (2017) Soil nematodes show a mid-elevation diversity maximum and elevational zonation on Mt. Norikura, Japan. Sci Rep 7:13028. https://doi.org/10.1038/s41598-017-03655-3

Eggenberg S, Möhl A (2013) Flora vegetativa: ein Bestimmungsbuch für Pflanzen der Schweiz im blütenlosen Zustand, 2nd edn. Haupt, Bern

Fierer N, McCain CM, Meir P et al (2011) Microbes do not follow the elevational diversity patterns of plants and animals. Ecology 92:797–804. https://doi.org/10.1890/10-1170.1

Fogain R (2001) Nematodes and weevil of bananas and plantains in Cameroon: occurrence, importance and host susceptibility. Int J Pest Manag 47:201–205. https://doi.org/10.1080/0967078010021108

Fontana LF, Dias-arieira CR, Mattei D et al (2015) Competition between Pratylenchus zeae and Meloidogyne incognita on sugarcane. Nematropica 45:1–8

Gaidashova S, Van Asten P, De Waele D, Delvaux B (2009) Relationship between soil properties, crop management, plant growth and vigour, nematode occurrence and root damage in East African Highland banana-cropping systems: a case study in Rwanda. Nematology 11:883–894. https://doi.org/10.1111/j.1365-2745.2010.01730.x

Garnier E, Cortez J, Billès G et al (2004) Plant functional markers capture ecosystem properties during secondary succession. Ecology 85:2630–2637. https://doi.org/10.1890/03-0799

Garnier E, Navas M-L, Griulius K (2016) Plant functional diversity: Organism traits, community structure, and ecosystem properties. Oxford University Press, Oxford

Guo Q, Kelt DA, Sun Z et al (2013) Global variation in elevational diversity patterns. Sci Rep 3:3007. https://doi.org/10.1038/srep03007

Haddad NM, Tilman D, Haarstad J et al (2001) Contrasting effects of plant richness and composition on insect communities: a field experiment. Am Nat 158:17–35. https://doi.org/10.1086/320866

Hartig F (2020) DHARMa: Residual Diagnostics for Hierarchical Regression Models. In: Compr. R Arch. Netw.

Hatfield JL, Prueger JH (2015) Temperature extremes: effect on plant growth and development. Weather Clim Extrem 10:4–10. https://doi.org/10.1016/j.wace.2015.08.001

Hillyer R, Silman MR (2010) Changes in species interactions across a 2.5 km elevation gradient: effects on plant migration in response to climate change. Glob Chang Biol 16:3205–3214. https://doi.org/10.1111/j.1365-2486.2010.02206.x

Hochkirch A, Gröning J (2008) Sexual size dimorphism in Orthoptera (sens. str.)—a review. J Orthoptera Res 17:189–196. https://doi.org/10.1665/1365-2486.2010.02206.x

Hochkirch A, Nieto A, Criado MG et al (2016) European red list of grasshoppers, crickets and bush-crickets. Publications Office of the European Union, Luxembourg. https://doi.org/10.2779/60944
Hodkinson ID (2005) Terrestrial insects along elevation gradients: species and community responses to altitude. Biol Rev 80:489–493. https://doi.org/10.1017/S1464793105006767
Hoiss B, Krauss J, Potts SG et al (2012) Altitude acts as an environmental filter on phylogenetic composition, traits and diversity in bee communities. Proc R Soc B Biol Sci 279:4447–4456. https://doi.org/10.1098/rspb.2012.1581
Hooper DU, Bignell DE, Brown VK et al (2000) Interactions between aboveground and belowground biodiversity in terrestrial ecosystems: patterns, mechanisms, and feedbacks. Bioscience 50:1049–1061. https://doi.org/10.1641/0006-3568(2000)050[1049:iaabt2.0.co;2]
Ibanez S, Lavorel S, Puijalon S, Moretti M (2013) Herbivory and interspecific variation in sexual size dimorphism in mountain grasshoppers. J Evol Biol 26:2171–2183. https://doi.org/10.1111/jeb.12213
Ingrisch S, Köhler G (1998) Die Heuschrecken Mitteleuropas. Westarp Wissenschaften, Magdeburg
Jarvis SG, Woodward S, Taylor AFS (2015) Strong altitudinal partitioning in the distributions of ectomycorrhizal fungi along a short (300 m) elevation gradient. New Phytol 206:1145–1155. https://doi.org/10.1111/nph.13315
Jones MGK, Fosu-Nyarko J (2014) Molecular biology of root lesion nematodes (Pratylenchus spp.) and their interaction with host plants. Ann Appl Biol 164:163–181. https://doi.org/10.1111/aab.12105
Kamira M, Hauser S, Van Asten P et al (2013) Plant parasitic nematodes associated with banana and plantain in eastern and western Democratic Republic of Congo. Nematropica 43:216–225. https://doi.org/10.13140/2.1.970.4969
Kaplan I, Halitschke R, Kessler A et al (2008) Physiological integration of roots and shoots in plant defense strategies links above- and belowground heribvory. Ecol Lett 11:841–851. https://doi.org/10.1111/j.1461-0248.2008.01200.x
Kurganteuil A, Campos-Herrera R, Sánchez-Moreno S et al (2016) The abundance, diversity, and metabolic footprint of soil nematodes is highest in high elevation alpine grasslands. Front Ecol Evol. https://doi.org/10.3389/fevo.2016.00084
Körner C (2003) The alpine plant life: functional plant ecology of high mountain ecosystems, 2nd edn. Springer, Berlin
Kuznetsova A, Brockhoff PB, Christensen RHB (2017) lmerTest package: tests in linear mixed effects models. J Stat Softw. https://doi.org/10.18637/jss.v082.i13
Laiolo P, Illera JC, Obeso JR (2013) Local climate determines intra- and interspecific variation in sexual size dimorphism in mountain grasshopper communities. J Evol Biol 26:2171–2183. https://doi.org/10.1111/jeb.12213
Labber K, Wagner G, Gygax A (2012) Flora Helvetica : flore illustrée de Suisse, 4th edn. Haupt, Bern
Looby CI, Maltz MR, Treseder KK (2016) Belowground responses to elevation in a changing cloud forest. Ecol Evol 6:1996–2009. https://doi.org/10.1002/ece3.2025
Margesin R, Jud M, Tscherko D, Schinner F (2009) Microbial communities and activities in alpine and subalpine soils. FEMS Microbiol Ecol 67:208–218. https://doi.org/10.1111/j.1574-6941.2008.00620.x
Martiny JBH, Bohannan BJM, Brown JH et al (2006) Microbial biogeography: putting microorganisms on the map. Nat Rev Microbiol 4:102–112. https://doi.org/10.1038/nrmicro1341
Mayor JR, Sanders NJ, Classen AT et al (2017) Elevation alters ecosystem properties across temperate treelines globally. Nature 542:91–95. https://doi.org/10.1038/nature21027
Mccain CM, Grytnes JA (2010) Elevational gradients in species richness. In: Encyclopedia of Life Sciences. John Wiley & Sons
McKenzie SW, Hentley WT, Hails RS et al (2013) Global climate change and above- belowground insect herbivore interactions. Front Plant Sci 4:412. https://doi.org/10.3389/fpls.2013.00412
Mlamo BC (2014) Not all traits are ‘functional’: insights from taxonomy and biodiversity-ecosystem functioning research. Biodivers Conserv 23:781–790. https://doi.org/10.1007/s10531-014-0618-5
Moles AT, Peco B, Wallis IR et al (2013) Correlations between physical and chemical defences in plants: tradeoffs, syndromes, or just many different ways to skin a herivorous cat? New Phytol 198:252–263. https://doi.org/10.1111/nph.12116
Moles AT, Warton DI, Warman L et al (2009) Global patterns in plant height. J Ecol 97:923–932. https://doi.org/10.1111/j.1365-2745.2009.01526.x
Moreira X, Petry WK, Mooney KA et al (2018) Elevational gradients in plant defences and insect herbivory: recent advances in the field and prospects for future research. Ecography (Cop) 41:1485–1496. https://doi.org/10.1111/ecog.03184
Nock CA, Vogt RJ, Beisner BE (2016) Functional Traits. eLS. John Wiley and Sons, Chichester, pp 1–8
Pellissier L, Fiedler K, Nidrié C et al (2012) Shifts in species richness, herbivore specialization, and plant resistance along elevation gradients. Ecol Evol 2:1818–1825. https://doi.org/10.1002/eece.3296
Pellissier L, Niculita-Hirzel H, Dubuis A et al (2014) Soil fungal communities of grasslands are environmentally structured at a regional scale in the Alps. Mol Ecol 23:4274–4290. https://doi.org/10.1111/mec.12854
Pellissier L, Roger A, Bilat J, Rasmann S (2014) High elevation Plantago lanceolata plants are less resistant to herbivory than their low elevation conspecifics: is it just temperature? Ecography (Cop) 37:950–959. https://doi.org/10.1111/ecog.00833
Perrine-Walker F (2019) Interactions of endoparasitic and ectoparasitic nematodes within the plant root system. Funct Plant Biol 46:205. https://doi.org/10.1071/FP18176
Peters MK, Hemp A, Appelhans T et al (2016) Predictors of elevational biodiversity gradients change from single taxa to the multi-taxa community level. Nat Commun 7:13736. https://doi.org/10.1038/ncomms13736
R Core Team (2019) R: A language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria. URL https://www.r-project.org/.
Rahbek C (1995) The elevational gradient of species richness: a uniform pattern? Ecography (Cop) 18:200–205. https://doi.org/10.1111/j.1600-0587.1995.tb00341.x
Rasmann S, Erwin AC, Halitschke R, Agrawal AA (2011) Direct and indirect root defences of milkweed (Asclepias syriaca): trophic cascades, trade-offs and novel methods for studying subterranean herbivory. J Ecol 99:16–25. https://doi.org/10.1111/j.1365-2745.2010.01713.x
Rasmann S, Pellisser L, Defossez E et al (2014) Climate-driven change in plant-insect interactions along elevation gradients. Funct Ecol 28:46–54. https://doi.org/10.1111/1365-2435.12135
Reich PB, Ellsworth DS, Walters MB (1998) Leaf structure (specific leaf area) modulates photosynthesis-nitrogen relations: evidence from within and across species and functional groups. Funct Ecol 12:948–958. https://doi.org/10.1046/j.1365-2435.1998.00274.x
Reynolds BC, Crossley DA (1997) Spatial variation in herbivory by forest canopy arthropods along an elevation gradient. Environ Entomol 26:1232–1239. https://doi.org/10.1093/ee/26.6.1232

Rueden CT, Schindelin J, Hiner MC et al (2017) Image J2: imageJ for the next generation of scientific image data. BMC Bioinform. https://doi.org/10.1186/s12859-017-1934-z

Ryalls JMW, Riegler M, Moore BD et al (2013) Effects of elevated temperature and CO2 on aboveground-belowground systems: a case study with plants, their mutualistic bacteria and root/shoot herbivores. Front Plant Sci 4:445. https://doi.org/10.3389/fpls.2013.00445

Sanson G, Read J, Aranwela N et al (2001) Measurement of leaf biomechanical properties in studies of herbivory: opportunities, problems and procedures. Austral Ecol 26:535–546. https://doi.org/10.1046/j.1442-9993.2001.01154.x

Scheidel U, Bruelheide H (2001) Altitudinal differences in herbivory on montane Compositae species. Oecologia 129:75–86. https://doi.org/10.1007/s004420100695

Sieriebriennikov B, Ferris H, de Goede RGM (2014) NINJA: an automated calculation system for nematode-based biological monitoring. Eur J Soil Biol 61:90–93. https://doi.org/10.1016/J.EJSOB.2014.02.004

Sierra-Almeida A, Cavieres LA, Bravo LA (2009) Freezing resistance varies within the growing season and with elevation in high-Andean species of central Chile. New Phytol 182:461–469. https://doi.org/10.1111/j.1469-8137.2008.02756.x

Thioulouse J, Dufour AB, Jombart T et al (2018) Multivariate analysis of ecological data with ade4. Springer, New York, New York

Truong NM, Nguyen CN, Abad P et al (2015) Function of root-knot nematode effectors and their targets in plant parasitism. Adv Bot Res 73:293–324. https://doi.org/10.1016/bs.abr.2014.12.010

Tylianakis JM, Didham RK, Bascompte J, Wardle DA (2008) Global change and species interactions in terrestrial ecosystems. Ecol Lett 11:1351–1363. https://doi.org/10.1111/j.1461-0248.2008.01250.x

Vairetti MV, Diaz S, Vile D, Garnier E (2007) Two measurement methods of leaf dry matter content produce similar results in a broad range of species. Ann Bot 99:955–958. https://doi.org/10.1007/10.1093/aob/mcm022

van den Hoogen J, Geissen S, Routh D et al (2019) Soil nematode abundance and functional group composition at a global scale. Nature 572:194–198. https://doi.org/10.1016/41586-019-1418-6

Valoire F (2008) Plant traits and functional types to characterise drought survival of pluri-specific perennial herbaceous swards in Mediterranean areas. Eur J Agron 29:116–124. https://doi.org/10.1016/J.EJA.2008.04.008

War AR, Paulraj MG, Ahmad T et al (2012) Mechanisms of plant defense against insect herbivores. Plant Signal Behav 7:1306–1320. https://doi.org/10.4161/psb.21663

Wardle DA, Bardgett RD, Klironomos JN et al (2004) Ecological linkages between aboveground and belowground biota. Science 304:1629–1633. https://doi.org/10.1126/science.1094875

WeiLing M, PeiLi S, WenHua L et al (2010) Changes in individual plant traits and biomass allocation in alpine meadow with elevation variation on the Qinghai-Tibetan Plateau. Sci China Life Sci 53:1142–1151. https://doi.org/10.1007/s11427-010-4054-9

Yeates GW, Bongers T, De Goede RG et al (1993) Feeding habits in soil nematode families and genera-an outline for soil ecologists. J Nematol 25:315–331