

Intra and inter-annual climatic conditions have stronger effect than grazing intensity on root growth of permanent grasslands

Catherine Picon-Cochard, Nathalie Vassal, Raphaël Martin, Damien Herfurth, Priscilla Note, Frédérique Louault

▶ To cite this version:

Catherine Picon-Cochard, Nathalie Vassal, Raphaël Martin, Damien Herfurth, Priscilla Note, et al.. Intra and inter-annual climatic conditions have stronger effect than grazing intensity on root growth of permanent grasslands. Peer Community Journal, 2021, 1, pp.e43. 10.24072/pcjournal.54 . hal-03997770

HAL Id: hal-03997770 https://hal.inrae.fr/hal-03997770

Submitted on 20 Feb 2023 $\,$

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

Peer Community Journal

Section: Ecology

RESEARCH ARTICLE

Published 2021-11-30

Cite as

Catherine Picon-Cochard, Nathalie Vassal, Raphaël Martin, Damien Herfurth, Priscilla Note and Frédérique Louault (2021) Intra and inter-annual climatic conditions have stronger effect than grazing intensity on root growth of permanent grasslands, Peer Community Journal, 1: e43.

Correspondence

catherine.picon-cochard@inrae.fr

Peer-review

Peer reviewed and recommended by PCI Ecology, https://doi.org/10.24072/pci. ecology.100073



This article is licensed under the Creative Commons Attribution 4.0 License.

Intra and inter-annual climatic conditions have stronger effect than grazing intensity on root growth of permanent grasslands

Catherine Picon-Cochard^{®,1}, Nathalie Vassal^{®,2}, Raphaël Martin^{®,1}, Damien Herfurth², Priscilla Note³, and Frédérique Louault^{®,1}

Volume 1 (2021), article e43

https://doi.org/10.24072/pcjournal.54

Abstract

Understanding how direct and indirect changes in climatic conditions, management, and species composition affect root production and root traits is of prime importance for the delivery of carbon sequestration services of grasslands. This study considers the effects of climatic variability and gradients of herbage utilisation by grazing on root production over the course of two years. The root and leaf traits of the plant communities were determined to detect their capacity to predict above- and below-ground net primary production, ANPP and BNPP, respectively. A long-term field experiment was used to compare the effects of abandonment and low (Ca-) and high (Ca+) grazing intensities (resulting in mean residual plant heights of 15.2 cm and 7.7 cm, respectively) induced by grazing rotations on upland fertile grasslands after 10 years of treatment application. Ingrowth cores and exclusion cages were used to measure, respectively, the root and shoot mass production several times each year and at an annual scale. The root and leaf traits of the communities were measured near the vegetations peak growing season. We observed strong seasonal root production across treatments in both a wet and a dry year, but the response to grazing intensity was hardly observable within growing seasons. In the abandonment treatment, the spring and autumn root growth peaks were delayed by approximately one month compared to the two cattle treatments, possibly due to a late plant canopy green-up induced by lower soil temperatures and an accumulation of litter. The BNPP was slightly lower in the abandonment treatment compared to the cattle treatments only during the dry year, whereas a decline of the ANPP in the abandonment treatment compared to the Ca+ treatment was observed during the wet year. In response to drought, which occurred during the second year, the root-to-shoot biomass ratio was stable in the cattle treatments but declined in the abandonment treatment. The higher allocation to root mass could benefit plant communities under drier conditions. Rotational grazing pressures and climatic condition variabilities had limited effects on root growth seasonality, although drought had stronger effects on the BNPP than on the ANPP. The stability of the root-to-shoot biomass ratio during the dry year evidenced a higher resistance to drought by grazed versus abandoned grassland communities.

¹Université Clermont Auvergne, INRAE, VetAgro Sup, UREP, 63000 Clermont-Ferrand, France, ²Université Clermont Auvergne, VetAgro Sup, 63370 Lempdes, France, ³INRAE, UE1414, Herbipôle, 63122 Saint-Genès-Champanelle, France



Peer Community Journal is a member of the Centre Mersenne for Open Scientific Publishing http://www.centre-mersenne.org/



Introduction

Permanent grasslands provide many services that connect to human activities through livestock products but also contribute to regulate greenhouse gas emissions because their soils accumulate large amounts of carbon (C) in organic matter fractions. Root activity (growth, exudation, and turnover) contributes to C and nitrogen (N) inputs and to absorb both nutrients and water, which are essential to fix atmospheric carbon dioxide (CO₂) and produce plant biomass. The intensification of management practices may affect these services as well as climate variability (Conant et al., 2001; Jones & Donnelly, 2004; Soussana & Duru, 2007). Thus, improving our understanding of grassland roots dynamics under different management and climatic conditions may help to identify management options to maintain the forage production and C sequestration abilities of this ecosystem and thus its sustainability.

Different management practices modify forage production and the amount of soil C and N through both the direct effects of defoliation, fertilisation, or returns of excreta to soil on root growth and soil abiotic factors and the indirect effects of species composition changes (Bardgett & Wardle, 2003; Dawson et al., 2000; Soussana et al., 2004). In mown grasslands, the root mass production is generally lower when grasses are frequently mown and fertilised (Leuschner et al., 2013; Picon-Cochard et al., 2009). This result may be explained by changes in the root-to-shoot allocation with an increase of above-ground growth to maximise light capture. The complexity of these phenomena in grazed grasslands is greater than in mown systems owing to animals' selective defoliation of plant species and because returns to soil are spatially heterogeneous (Rossignol et al., 2011). In addition, the level of soil fertility may buffer the degree of root response to defoliation in grazed grasslands as plants exhibit specific responses to defoliation in fertile and unfertile grasslands (Duru et al., 1998). Overall, the complexity of phenomena being direct and indirect effects on plants could explain why no clear trend has been found for the effects of grazing on above- and below-ground production (e.g., refer to the syntheses of Milchunas & Lauenroth [1993] and McSherry & Ritchie [2013]), although two meta-analyses have emphasised the negative effect of grazing intensity on above- and below-ground C stocks compared to ungrazed systems (Zhou et al., 2017; Li et al., 2018). Furthermore, repeated defoliations induced by grazing or mowing grasslands can simultaneously increase soil temperatures and moisture (Moretto et al., 2001; Pineiro et al., 2010; Smith et al., 2014). Soil moisture can also be modified by high stocking rates through changes in the soil bulk density due to soil compaction and changes in the leaf area index after defoliation (Pineiro et al., 2010). These direct effects of grazing on soil abiotic factors should affect the root growth of grazed grasslands, although these phenomena have not been well documented in field conditions.

Species composition change induced by management is also an important determinant of above- and below-ground responses in grazed grasslands. Intensive practices (e.g., high grazing intensity or fertilisation) generally favour the development of fast-growing species (plant exploitative strategy), whereas the opposite extensive practices (e.g., low grazing intensity or the absence of fertilisation) favour slow-growing species (plant conservative strategy; Klumpp et al., 2009; Louault et al., 2005; Soussana & Lemaire, 2014; Wardle et al., 2004). The root-to-shoot biomass allocation, as well as the functional traits (used as proxies of ecosystem properties such as the ANPP or BNPP; e.g., Laliberté & Tylianakis, 2012), are thus likely to change in response to an intensification of practices, such as from ungrazed to intensively grazed temperate grasslands, alpine meadows, steppes or desert steppes (Klumpp & Soussana, 2009; Zeng et al., 2015). According to Ziter & MacDougall (2013), the uncertainty surrounding nutrient-defoliation responses makes it difficult to predict whether C storage would be higher in managed versus unmanaged grasslands. Thus, soil fertility should be considered when comparing different grazing intensities in grasslands (Louault et al., 2005).

Increased climate variability is another source of response uncertainty in managed ecosystems. As more frequent and longer periods of drought associated with heat waves may threaten and shape the long-term dynamics of perennial ecosystems such as grasslands (Brookshire and Weaver, 2015), it is important to understand how above- and below-ground compartments respond to climate variability. However, the data on above- and below-ground biomass responses to drought for grasslands are limited (Byrne et al., 2013; Wilcox et al., 2015; Li et al., 2018), although some evidence shows that resource conservation strategy is associated with drought tolerance (Pérez-Ramos et al., 2012; Reich, 2014). Changes in root morphology and functioning may thus be important determinants in plants' adaptive strategies to drought

and have been less studied than above-ground plant responses (Biswell & Weaver, 1933; Dawson et al., 2000; McInenly et al., 2010). However, there are not enough data to make generalisations about the combined impacts of management and climatic condition variabilities, such as precipitation reduction, on root and shoot biomass production and plant traits defining plant strategies related to resource use and grazing intensity.

This study comprised of a long-term field experiment for which controlled grazing intensity was applied for 10 years. We compared grazing abandonment and two levels of herbage utilisation by grazing based on five rotations per year. In two consecutive years, the ingrowth core method was used to measure the monthly root biomass production and calculate the BNPP. The ANPP was measured using grazing exclusion cages, and the community-weighted mean (CWM) leaf and root traits were assessed the first year. We tested the following hypotheses: (i) high grazing intensity increases above-ground mass at the expense of root production as a result of the direct negative effect of defoliation on root growth, whatever the climatic conditions; (ii) inter-annual climatic conditions modulate the above- and below-ground biomass production response to grazing intensity as a consequence of the higher presence of defoliation-tolerant and drought-sensitive species (*Lolium perenne* or *Trifolium repens*) in the high grazing intensity treatment; (iii) root traits respond to treatments and are a determinant of the BNPP, as observed for leaf traits and the ANPP.

Materials and Methods

Site characteristics

The experiment transpired at the long-term observatory network (ACBB, <u>https://www.anaee-france.fr/en/infrastructure-services/in-natura-experimentation/agrosystem/acbb</u>) located in St-Genès-Champanelle, France (45° 43'N, 03° 01'E, 880 m a.s.l.). The local climate is semi-continental with oceanic influences and a mean annual temperature and precipitation of 8.5°C and 784 mm, respectively (Table 1). The site supports mesotrophic multi-specific permanent grasslands dominated by species with high Ellenberg indicator values for N (Schaffers and Sykora, 2000), indicating a high level of fertility for the site (Table S1; Louault et al., 2017). The soil is a cambisol with a sandy loam texture developed on a granitic bedrock. Differences in the local soil composition and profile led us to consider two distinct blocks characterised by an eutric cambisol (54% sand, 26% silt, 20% clay, 7.0% organic matter, and pH: 5.9) and a colluvic cambisol (50% sand, 26% silt, 24% clay, 7.4% organic matter, and pH: 6.0), including some volcanic materials.

Management

Prior to the installation of this experiment in 2005, the study area had been used for intensive hay and silage production (combining grazing, mowing, and fertilisation) with mineral fertilisation, and two years preceding the start of the experiment (2003 and 2004), the grassland site was mown three times per year without fertilisation. From 2005, the grassland was managed for 10 years with a gradient of grazing intensity resulting from three treatments: abandonment, low herbage utilisation (Ca-), and high herbage utilisation (Ca+). The Ca- and Ca+ treatments were obtained by modifying the stocking density (6.9 and 13.8 livestock units (LSU) ha⁻¹, respectively) with five grazing rotations each year occurring at mid-April, late May, early July, September, and November and lasting on average 9.6, 9.0, 10.7, 8.6, and 2.1 days, respectively. The two cattle treatments corresponded to two levels of herbage utilisation by grazing and had on average a residual plant height of 15.2 cm \pm 0.5 (mean \pm SE) for Ca- and 7.7 cm \pm 0.2 for Ca+ at the end of each grazing rotation. For each treatment, two replicate plots were created per block, resulting in four replicates per treatment and a total of 12 plots (2 blocks x 2 plots x 3 treatments). The average distance between the two blocks was approximately 230 m, and all the treatments were randomised within each block. The size of the plots differed according to the treatment: 2,200 m² for the two cattle treatments and 400 m² for the abandonment treatment.

Climatic and edaphic conditions

The daily precipitation (mm) and air temperature (°C) were measured for the two years and recorded with an onsite meteorological station. A climatic water balance was calculated as precipitation minus the potential evapotranspiration (P - PET, mm) with the Penman-Monteith equation. The daily soil temperature

(°C) was measured with thermocouple sensors (homemade copper-constantan sensors) inserted at a 20cm depth in each plot and recorded with an HOBO data logger (U12-014, Onset Instruments, MA, USA). The daily volumetric soil water content (SWC; m³ m⁻³) of each plot was measured with two probes (ECHO-10, Decagon, USA) inserted horizontally at a 20-cm depth and connected to data loggers (EM5 and EM50, Decagon, USA). From January 2014 to November 2015 (DOY 132–326), the SWC was measured every 30 min and averaged at a daily scale. For each plot, the average values of the two probes were used. The daily relative soil water content (RSWC) data are shown and calculated as the ratio:

 $RSWC = \frac{SWC - SWCmin}{SWCmax - SWCmin}$, where SWC is the soil moisture at a given day, SWCmin is the minimum value of soil moisture, and SWCmax is the maximum value of soil moisture, all observed during the two years. For the soil temperature and RSWC, the values were averaged according to the root growth timescale.

Root growth and root mass

Six months before the start of the experiment, shallow (0-20 cm) soil was collected on each of the two blocks, sieved (using a 5 mm mesh size) to remove stones and coarse organic matter, and then left unused outside covered under a shelter and protected from direct sunlight. This air-dried soil was subsequently used to fill the ingrowth cores each month. In December 2013, soil cores were collected with an auger (which had an 8-cm diameter and 0-20 cm depth) for each of the 12 plots at four locations representative of the plant community in the treatment. On average, the mean distance between locations was 19.8 m \pm 0.2 for Ca+, 21.7 m \pm 0.1 for Ca-, and 17.2 m \pm 0.2 for Ab (mean \pm standard deviation; refer to Figure S1). After the core harvest, each hole was filled with an 8-mm mesh sized plastic net containing a fixed volume of air-dried sieved soil (called thereafter ingrowth core) collected six months beforehand. For the next two years and approximately each month (2 x 10 times), the ingrowth cores, containing soil and the root and rhizome material that had grown therein, were extracted and then replenished with another fixed volume of air-dried sieved soil. Thus, the monthly and annual root production (BNPP, g m⁻² y⁻¹) were measured from February 2014 to December 2015. The root production period ranged on average 36.5 days, but with longer periods in the winter and shorter periods in the spring and summer (Table 1). In periods without precipitation, a fixed volume of water was added to adjust soil humidity to field conditions. After collection, the ingrowth cores were transported to the laboratory and immediately stored at 4°C before being processed in the next five days. The roots were washed under tap water with a 200-µm sieve and then oven dried for 48 h at 60°C.

To measure the root mass stock, the soil cores were collected three times (December 2013 and March and June 2014) with the same auger and near the ingrowth core locations. These samples were stored in the freezer at -18°C; after defrosting, the roots were washed with the same procedure as that used for the ingrowth cores and then oven dried for 48 h at 60°C.

Root traits

Subsamples of the washed roots collected with the ingrowth cores in June 2014 were fresh weighed and then frozen at -18°C before performing a morphology analysis. After defrosting, the roots were stained with methylene blue (5 g L⁻¹) for approximately 5–10 minutes, rinsed in water, spread in a transparent glass box containing a thin layer of water, and covered with a transparent plastic sheet. High resolution images were recorded with a double light scanner (800 dpi, perfection V700, Epson, JA) and analysed using WinRhizo software (PRO 2012b, Regent Instruments, CA) with the automatic procedure. Two scans per location were recorded and separately analysed to measure the root length (m), root volume (cm³), root surface area (m²), average root diameter (mm), and root length by class diameter (there are 13 classes: 11 with a 0.1-mm interval and 2 with a 0.5-mm interval). The specific root length (SRL; m g⁻¹), root tissue density (RTD; g cm⁻³), and specific root area (SRA; m² g⁻¹) were calculated for fine roots as in Picon-Cochard et al. (2012).

Botanical composition

The species contribution (%) was visually observed on a 20-cm diameter circle around each ingrowth core location in April (cattle treatments) and May (abandonment treatment) 2014. For each zone, a score on a 10-point scale was allocated to the species present according to their volume occupancy, and the

percentage of each species was calculated at the plot scale by averaging the values of the four zones. Table S2 lists the species and their relative contributions.

Above-ground biomass production

On each plot and on each sampling date, four fenced sampling areas $(0.6 \times 0.6 \text{ m})$ were used to measure the accumulation of above-ground biomass after the above-ground standing biomass was clipped to 5.5 cm, oven dried, and weighed. Measurements were made five times over the year, once before each grazing event in the Ca+ and Ca- treatment plots and three times (spring, summer, and autumn) in the abandonment plots. The sampling areas were moved within the plot at each measurement date during the year. The annual above-ground net primary production (ANPP, g m⁻² y⁻¹) was calculated as the sum of the successive biomass accumulations throughout the year.

Leaf traits

The CWM trait values of the leaf dry matter content (LDMC), specific leaf area (SLA) and reproductive plant height were calculated for each ingrowth core zone using both the relative contribution of the dominant species to the community (i.e., species that account for at least 85% of the cumulated species contribution to the community) measured in 2014 and the leaf trait measurements made at plot scale in 2006 and 2007. The traits were measured on 10 vegetative plants using standard protocols (refer to the methods in Louault et al., 2005). The reproductive plant height was measured on mature plants located in fenced zones to allow full plant development. The CWM is expressed with the following equation: CWM = $\sum p_i \times \text{trait}_i$, where p_i is the relative contribution of species *i* to the community and trait_i is the trait of species *i*.

Statistical analyses

For a given date, data averages of the root mass and root traits collected at each of the four locations (four ingrowth cores in each plot) were used to obtain a single value for each of the 12 plots and test for the effect of the treatment and dates. Before variance analysis (ANOVA), the normality of residuals was inspected with quantile-quantile plots of model residuals, and variance homogeneity was confirmed by checking the plots of model residuals versus the fitted values. Data were transformed if they deviated from the ANOVA assumptions (square root, In, reciprocal). Linear mixed effects models as available in the R 'nlme' package (Pinheiro et al., 2015) were used to perform repeated measure ANOVAs to test the effects of treatments and dates, as well as their interactions on values of root growth, soil temperature, RSWC, and root mass stock, with plots nested within block as a random factor accounting for temporal pseudoreplication (S4). For the root growth dynamics, soil temperature, and RSWC (Figure 1; Table S3), the dates correspond to 20 dates, whereas for the root mass stock, the dates correspond to three harvest dates (Table 2). For the BNPP, ANPP, and root-to-shoot ratio (BNPP/ANPP), the data were analysed using a nested mixed model procedure with the treatments and year used as fixed factors and with plot nested within block used as random factors. For the leaf and root traits data, the treatments were used as fixed factors with plots nested within block used as a random factor. Post-hoc tests were performed to compare the significance levels across fixed factors with a Tukey test ('Ismeans' package). Principal component analyses (PCAs) were performed for each year to analyse the relationships between the leaf and root traits, soil temperature, RSWC, root mass stock, ANPP, and BNPP measured at the plot level; the treatments were considered supplementary categories ('FactoMineR' package). This statistics approach allows sets of trait and property relationships to be compared to detect response and effect traits and to analyse multiple dimensions of trait relationships, which is not possible with pairs of correlations. All statistical analyses were performed in an R environment (Version 3.5.2, R Core Team, 2012) using RStudio (Version 1.1.463). The scripts are shown in S4.

Results

Climatic conditions during the experiment

Compared with the average long-term climatic data for the site, the first and second years of the experiment received higher (+92 mm) and lower (-199 mm) precipitation, respectively (Table 1). The PET in the second year was also higher than the long-term average (a 73-mm difference), leading to a negative

annual climatic water balance (P – PET = -181 mm and a deficit of 271 mm compared to the long-term average). The annual temperature in the two experimental years was similar and approximately 0.8°C higher than the long-term average for the site (Table 1). At a monthly timescale and during part of the growing season (March to September), in comparison with the first year, the second year had a cumulated water deficit difference of -266 mm and warmer temperatures by an average of 1.9°C. Larger differences between the two years occurred in June and July with an average higher temperature (+6°C), higher water deficit (P – PET = -152.6 mm), and less precipitation (-81%) in the second year.

Table 1. Air temperature (°C), precipitation (P, mm), potential evapotranspiration (PET) and climatic water balance: cumulated (P – PET, mm) and calculated for the 28 y period 1986-2013, mean values ± SD) and measured for the 10 dates in 2014 and 2015 corresponding to measurements of root growth and averaged (temperature) or summed (P, PET, P – PET) at annual scale.

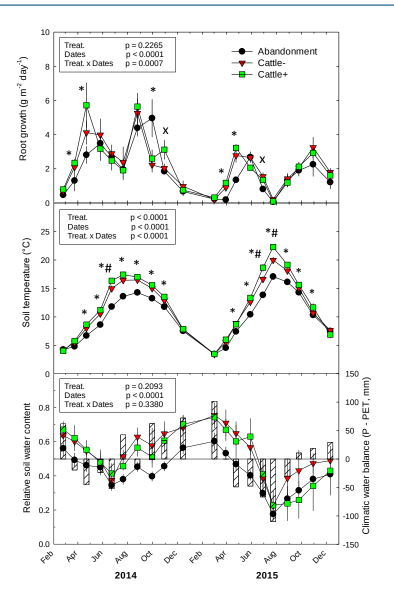
Year	Dates	Air temperature	Precipitation	PET	P-PET
	Annual long-term average	8.5 ± 0.6	784 ± 1376	693 ± 96	91 ± 195
	December 12 – February 23	3.7	98	37.5	60.5
	February 24 – March 23	5.3	27	46.3	-19.3
	March 24 – April 21	7.2	23.5	68.7	-45.2
	April 22 – May 25	9.2	79.5	103.1	-23.6
2014	May 26 – June 22	14.2	58	110.2	-52.2
2014	June 23 –July 20	15.1	136.5	93.9	42.6
	July 21 – August 24	14.4	90.5	100.5	-10
	August 25 – September 29	13.7	141.8	79.5	62.3
	September 30 – October 29	11.7	69	36.3	32.7
	October 30 – December 14	5.3	111	10.9	72.1
	Annual	9.2	876	691	157.7
	December 15 – March 1	1.3	132.5	31	101.5
	March 2 – March 29	4.5	36.5	36.8	-0.3
	March 30 – April 23	8.5	17.5	66.4	-48.9
	April 24 – May 28	11.0	66	113.6	-47.6
	May 29 – June 28	15.5	62.5	129.1	-66.6
2015	June 29 –July 23	21.1	26	136	-110
	July 24 – August 27	16.4	94.5	124.6	-30.1
	August 28 – September 24	12.8	77	66.3	10.7
	September 25 – October 29	7.8	55	36.1	18.9
	October 30 – December 11	7.0	54.5	25.1	29.4
	Annual	9.4	585	766	-180.9

Dynamics of soil temperature and relative soil water content

The soil temperature was significantly affected by the treatment, date, and treatment × date (Figure 1; Table S3). For most of the dates (February to October), the abandonment treatment corresponded with a lower soil temperature (1.76° C on average) than the cattle treatments, whereas the Ca- treatment soil temperature was significantly lower (0.64° C) than the Ca+ treatment. However, this difference was only observed to be significant for a limited number of dates in early summer of both years. The RSWC fluctuated from 0.6-0.7 at the beginning of spring to 0.38 in June during the wet year and 0.2 during the dry year, which accords with the variation of the climatic water balance (P – PET). In the case of the dry year, from summer until autumn, the RSWC remained lower than 0.4 and P - PET was negative.

6

Figure 1. The root growth (g m⁻² day⁻¹), soil temperature (°C), relative soil water content, and climatic water balance (P - PET, mm; hashed bars) dynamics measured over two years for the abandonment, low (Cattle-), and high (Cattle+) grazing intensity treatments. The vertical bars correspond to 1 SE (n = 4). The insets indicate P values from the repeated measure two-tailed ANOVA, where treat. is the treatment; dates, and interaction for the main treatments. *: P < 0.05; x: $P \le 0.1$. For soil temperature, *# corresponds to significant differences between all treatments (Abandonment < Cattle- < Cattle+).



Root growth dynamics

Root growth was affected by the date and treatment × date interaction (Figure 1). Each year, the root growth peak occurred twice, in spring and autumn, and growth was markedly reduced in summer and winter. Only in the second year did growth cease in summer. Second year root growth was significantly lower than the first year. Regarding the treatment effect, the abandonment treatment showed significant lower root growth than the two cattle treatments for the spring period in both years and for the autumn period of the second year. In autumn 2014 (end of September), a delay of growth peaks was also observed, which led to a twofold higher root growth for the abandonment treatment versus the two cattle treatments. The two cattle treatments experienced similar root growth across the years and seasons.

Seasonal root mass stock, below- and above net primary production, and root-to-shoot biomass ratio

The stock of root mass did not change throughout the seasons or across treatments (Table 2). The BNPP, ANPP, and root-to-shoot biomass ratio (R/S) were significantly lower during the second year, with a stronger effect on the BNPP (-44% on average) than the ANPP (-24% on average; Figure 2; Table 3). Only the abandonment treatment maintained its ANPP value in the second year, which led to a 48% decline in the R/S (significant treatment × year, P < 0.01; Table 3). Accordingly, a treatment effect was only observed for the BNPP during the second year, with a decline of 24% for the abandonment treatment compared to the cattle treatments. For the ANPP during the first year, the Ca+ treatment had 22% and 68% higher values, respectively, than the Ca- and abandonment treatments, while the Ca- treatment had a 38% higher ANPP than the abandonment treatment.

Table 2. a) Repeated measure ANOVA is shown for treatment, date (December 2013, March 2014, June 2014) and interaction effects on root mass (g m-2). Numerator (num), denominator (den) of degree of freedom (DF) and F values are shown. b) Root mass (g m-2) of abandonment, low (Cattle-) and high (Cattle+) stocking density treatments measured in winter (December 12 2013), spring (March 20 2014), summer (June 20 2014) and averaged across the three dates. Means ± SE are shown, n = 4. Superscripts ns correspond to P > 0.05.

a)	num/den DF	F-value	
Treatment	2/8	1.151 ^{ns}	
Date	2/18	2.027 ^{ns}	
Treatment × date	4/18	1.340 ^{ns}	
b) Date	Abandonment	Cattle-	Cattle+
December 2013	636.4 ± 133.1	403.3 ± 66.4	496.5 ± 20.6
March 2014	559.1 ± 166.2	609.2 ± 45.3	719.8 ± 47.5
June 2014	574.2 ± 84.8	482.2 ± 38.6	591.2 ± 101.7
3 dates average	589.9 ± 99.9	498.2 ± 43.6	602.5 ± 44.4

Table 3. Repeated measure ANOVA is shown for treatment, year and interaction effects on annual root production (BNPP, g m⁻² y⁻¹), annual above-ground production (ANPP, g m⁻² y⁻¹) and root to shoot ratio (R/S). Numerator (num), denominator (den) of degree of freedom (DF), *F* values are shown. Superscripts ^{ns, *, **, ***} correspond to P > 0.05, P < 0.05, P < 0.01, P < 0.001, respectively.

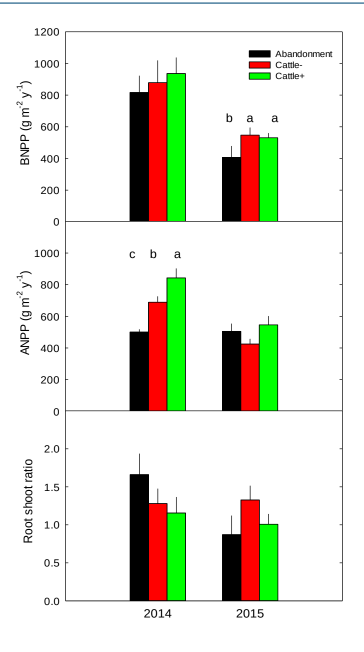
		BNPP	ANPP	R/S
	num/den	F-value	F-value	F-value
Treatment	2/8	2.51ns	8.10*	0.46ns
Year	1/9	70.72***	83.77***	13.09**
Treatment × Year	2/9	3.83ns	22.21**	9.52**

Species composition, leaf and root traits

The abandonment treatment was characterised by the dominance of tall grass species (76% in total) with 27.2% of Alopecurus pratensis, 18.8% of Elytrigia repens, 11.3% of Poa pratensis, and 10.3% of Arrhenatherum elatius. Some forbs (19%) were also present, whereas legumes were absent (Table S2). The two cattle treatments differed from the abandonment treatment by the equal presence of Taraxacum officinale (18% on average) and Trifolium repens (17% on average). Differences also existed concerning grass species (56% in total), with the dominance of Dactylis glomerata (22.2%), A. pratensis (7.6%), and Festuca arundinacea (5.6%) for the Ca- treatment and Lolium perenne (13.6%), D. glomerata (9.1%), and Poa trivialis (7.2%) for the Ca+ treatment. Thus, the Ca+ treatment had a higher percentage of L. perenne than the Ca- treatment (Table S2).

The CWM leaf traits were significantly modified by the treatments. The plant height and LDMC were significantly higher (P < 0.05 and P < 0.0001, respectively) and the SLA was lower (P < 0.05) in the abandonment treatment than in the two cattle treatments (Table 4). Unlike the leaf traits, the root traits were only slightly affected by the treatments. The SRL (P < 0.1) and SRA (P < 0.05) were lower in the abandonment treatment than in the Ca- treatment, but not than in the Ca+ treatment. For other root traits (diameter, RTD, and root length percentage by class diameter), no between-treatment differences were observed (Table 4).

Figure 2. The annual root biomass production (BNPP, g m⁻² y⁻¹), annual above-ground biomass production (ANPP, g m⁻² y⁻¹), and root-to-shoot biomass ratio measured in 2014 and 2015 for the abandonment, low (Cattle-), and high (Cattle+) grazing intensity treatments. The vertical bars correspond to 1 SE (n = 4). Within a year, different letters correspond to significant differences at P < 0.05.

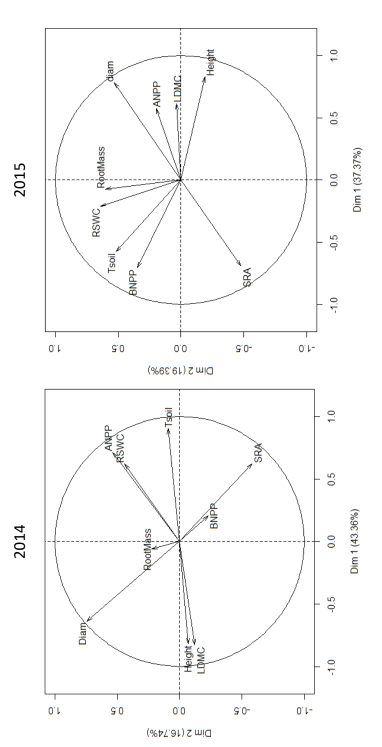


Co-variation of traits and production

The two main axes of the standardised PCA explained 60.1% and 56.8% of the community trait and production variation in 2014 and 2015, respectively (Figure 3). For the first year, the first PCA axis (PC1), which accounted for 43.4% of the total variation, was significantly related to the leaf and root traits, ANPP, and soil temperature. The soil temperature, SRA, and ANPP had positive loadings, while the diameter, plant height, and LDMC had negative loadings (Table 5). The second PCA axis (PC2), which accounted for 16.7% of the total variation, was significantly and positively related to the root diameter and negatively related to the SRA. For the second year, the PC1 accounted for 37.4% of the total variation and was significantly related to the leaf and root traits, ANPP, and BNPP. The BNPP and SRA had negative loadings, whereas the root diameter, plant height, and ANPP had positive loadings (Table 5). The PC2, which accounted for 19.4% of the total variation, was significantly and positively related to the RSWC and the stock of root mass averaged across three dates. The abandonment treatment was significantly related to the PC1s, with negative loadings for the first and second years, respectively.

Table 4. Root traits measured from ingrowth core collected in June 2014 and leaf traits measured from botanical observation in abandonment (May 2014), Ca- and Ca+ (April 2014) treatments. Diameter: root diameter (mm); SRL: specific root length (m g⁻¹); RTD: root tissue density (g cm⁻³); SRA: specific root area (m² g⁻¹); % 0-0.1 mm: percentage of root length in the class diameter 0-0.1 mm; % 0.1-0.2 mm: percentage of rot length in the class diameter 0.1-0.2 mm; % 0.2-0.3 mm: percentage of root length in the class diameter 0.1-0.2 mm; % 0.2-0.3 mm: percentage of root length in the class diameter > 0.3 mm; Community-weighted mean (CWM) Height: plant height (cm); SLA: specific leaf area (cm² g⁻¹); LDMC: leaf dry matter content (g g⁻¹). Means ± SE are shown (n = 4). num/den DF: numerator and denominator of degree of freedom. Superscripts ^{ns, +, *, ***} correspond to P > 0.1, P ≤ 0.1, P < 0.05, P < 0.01, P < 0.001, respectively. For SRL, SRA, height, SLA and LDMC, different letters correspond to significant differences between treatments.</p>

	num	F-	Abandonmen	Cattle-	Cattle+
	/den DF	value	t	Cattle-	
Root traits					
Diameter	2/8	1.61 ^{ns}	0.240 ± 0.015	0.210 ± 0.006	0.222 ± 0.015
SRL	2/8	3.71+	237.2 ± 26.3 b	332.7 ± 30.4 a	277.8 ± 23.8 a
RTD	2/8	0.55 ^{ns}	0.099 ± 0.007	0.095 ± 0.003	0.102 ± 0.007
SRA	2/8	4.96*	0.137 ± 0.011	0.182 ± 0.008 a	0.155 ± 0.01 a
% 0-0.1	2/8	1.28 ^{ns}	28.5 ± 1.1	32.9 ± 5.5	28.8 ± 2.6
% 0.1-0.2	2/8	0.46 ^{ns}	37.7 ± 4.4	37.7 ± 2.2	39.1 ± 1.8
% 0.2-0.3	2/8	0.30 ^{ns}	16.6 ± 1.2	16.2 ± 2.4	17.1 ± 1.9
% > 0.3	2/8	1.22 ^{ns}	17.2 ± 5.0	13.2 ± 1.3	15.1 ± 2.1
Leaf traits					
CWM_Hei	2/8	8.45*	93.0 ± 3.5 a	72.8 ± 7.0 b	68.6 ± 3.8 b
CWM_SLA	2/8	5.30*	205.1 ± 5.7 b	231.8 ± 7.3 a	225.5 ± 7.1 ab
CWM_LD	2/8	11.22*	0.261 ± 0.008	0.227 ± 0.007 b	0.213 ± 0.010



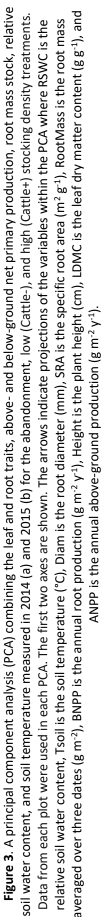


Table 5. Contribution of the different variables to the first two axes of the principal component analysis (PCA) calculated for 2014 and 2015. Variables used in the PCA were annual relative soil water content (RSWC), annual soil temperature (Tsoil, °C), root diameter (Diam, mm), specific root area (SRA, m² g⁻¹), root mass averaged over three dates (RootMass, g m⁻²), annual root production (BNPP, g m⁻² y⁻¹), plant height (Height, cm), leaf dry matter content (LDMC, g g⁻¹), annual above-ground production (ANPP, g m⁻² y⁻¹). Treatments were added as supplementary categories. Contribution in bold indicates significant correlation of the variables on the PCA axis (P < 0.05).

	2014		2015	
Variable	Axis 1 (43.4 %)	Axis 2 (16.7 %)	Axis 1 (37.4 %)	Axis 2 (19.4 %)
RSWC	0.62	0.44	-0.21	0.64
Tsoil	0.91	0.09	-0.58	0.52
Diam	-0.64	0.75	0.78	0.53
SRA	0.62	-0.58	-0.69	-0.48
RootMass	-0.06	0.22	-0.07	0.60
BNPP	0.21	-0.23	-0.71	0.35
Height	-0.82	-0.07	0.83	-0.19
LDMC	-0.83	-0.12	0.61	0.03
ANPP	0.71	0.54	0.57	0.20
Suppl. Categories				
Abandonment	-2.62	-0.24	2.04	-0.27
Cattle-	1.07	-0.55	-1.21	-0.62
Cattle+	0.70	0.18	-0.83	0.90

Discussion

Ten years of contrasted management strongly modified the functional diversity and above-ground production of this fertile upland grassland as previously shown for the same site (Herfurth et al., 2015; Louault et al., 2017). Accordingly, we expected that the above-ground biomass patterns would be mirrored below ground, especially during the periods of grazing. In this section, we first discuss the within-year differences of root growth, followed by the inter-annual variation responses to grazing intensity and climatic condition variabilities between the two contrasting years. Finally, we analyse the relationships between traits and the above- and below-ground production.

Seasonality of root growth in relation to grazing intensity and climatic conditions

As expected, the permanent grassland root growth was affected by seasons and the spring and autumn peaks (Garcia-Pausas et al., 2011; Pilon et al., 2013; Steinaker and Wilson, 2008), but unexpectedly, grazing pressure applied by rotations and climatic condition variabilities had very limited effects on this seasonality. Accordingly, at the below-ground level, the plant community behaviour was not affected by either the rotational grazing management or the climatic condition variabilities, although a severe drought occurred in summer of the second year. Only the abandonment treatment showed a delayed root growth peak in spring. This delay was probably the result of a slower shoot budburst and a reduced capacity to produce new green leaves in the dense litter canopy, especially at the beginning of the growing season in spring (data not shown). Moreover, the tall and dense canopy of the abandonment treatment strongly modified the soil temperature; as expected, the soil conditions were cooler in the abandoned vegetation (Picon-Cochard et al., 2006; Zhou et al., 2017; Zhu et al., 2016). As shown in some studies, light, soil, water, and nutrient availability (Edwards et al., 2004; Garcia-Pausas et al., 2011; Steinaker and Wilson, 2008) are other abiotic factors determining root growth dynamics in grasslands, for example root peaks were observed before the summer soil temperature peak when a negative climatic water balance occurred, especially in the second year. Nevertheless, plants growing in the abandonment treatment offset their slower root growth by producing a similar annual root biomass, especially during the wet year. The presence of tall grass species, such as A. pratensis, A. elatius, and E. repens, having plant trait syndromes related to both disturbance and resource conservation strategies (i.e. lower SLA and SRL and higher plant height and root depth; Pagès & Picon-Cochard, 2014) might explain the tall grass capacity to produce higher root biomasses before the canopy senescence onset. Additionally, pre-existing soil fertility can be maintained in conditions of very low levels of herbage utilisation (near-abandonment) because of the absence of biomass exportation and increased internal recycling of N within senescent plants, both of which contribute to an increase in the total N available for plant growth (Loiseau et al., 2005).

The similar root growth dynamics of the two cattle treatments was unexpected since infrequent defoliation and moderate excreta returns to the soil might increase the root biomass production at the expense of the shoot biomass production (Klumpp et al., 2009). The absence of effect on the root growth and BNPP indicates that the grazing applied to the plant communities by rotations was too short but sufficient to observe the effect on the ANPP in wet conditions.

Different methods exist globally to manage grasslands by grazing (Huyghe et al., 2014), including rotational or permanent grazing options with different stocking densities, durations, and types of herbivores. In general, grazing management creates high spatial heterogeneity within the plots due to animals' selective defoliation of plant species and because returns to soil are spatially heterogeneous. Thus, in grazed grasslands, disturbances create patches of vegetation, which should affect the local root growth and below-ground biomass of the plant communities if the intensity of grazing is sufficient. The complexity of these phenomena in grazed grasslands is greater than in mown systems (Rossignol et al., 2011).

Notwithstanding, the confounding effect of soil fertility and defoliation may mask a clear response of the below-ground compartment in grazed grasslands. In consequence, we postulate that the root growth in the Ca+ treatment was favoured by the higher soil temperature compensating for the negative effects of frequent defoliation on root growth, while the cooler soil conditions encountered in the Ca- treatment might have slowed the root growth. Soil moisture level is a main determinant of plant growth and can be affected by cattle treatments. Some studies have shown an increase of soil moisture in grazed compared ungrazed treatments due to the lower leaf area index in the grazed conditions (Moretto et al., 2001; Pineiro et al., 2010) or an absence of effect in others studies (LeCain et al., 2002; Smith et al., 2014). The presence of herbivores can increase soil bulk density and consequently modify soil moisture. In our field conditions after 10 years of treatment applications, the absence of effect on soil moisture could be due to several of these factors: absence of change of soil density, due to the rotational grazing as cattle spend less time in the plots than in continuous grazing systems; the functional composition of the community regarding both their response to defoliation and their water-use strategies; the monthly-based temporal scale used, which could buffer shorter-term responses.

We should also consider the level of soil fertility and species composition as drivers of root growth and trait plasticity (Dawson et al., 2000). The soil fertility of our site, reflected by the nitrogen nutrition index (NNI; Lemaire & Gastal, 1997), was similar across our grazing intensity gradient (Table S1), at least in 2014.

Thus, our site provided us with the opportunity to compare grazing intensity effects at equivalent soil fertility levels. Because root trait plasticity generally shows larger differences with respect to soil fertility than from cutting or defoliation (Leuschner et al., 2013; Picon-Cochard et al., 2009), we could expect grazing intensity to produce a less pronounced effect on root growth under similar soil fertility conditions. Indeed, the higher presence of defoliation-tolerant species, characterised by a shorter stature and root system (e.g., *L. perenne* and *P. trivialis*) but a higher shoot and root growth capacity after defoliation and higher rhizosphere activity (Dawson et al., 2000), probably compensated for the negative effect of defoliation in the Ca+ treatment. The sampling depth might also have had an effect as we expect that due to species-specific differential root depth distribution, harvesting root systems deeper than 20 cm should have resulted in a more contrasting root growth response across the two cattle treatments according to the grass species composition (Xu et al., 2014). The results indicate that the high soil temperatures, high soil fertility, and species composition moderated the root growth response across our grazing intensity gradient. The difficulty of assigning species composition in root mixtures, however, makes it difficult to draw firm conclusions.

Responses of ANPP, BNPP, and root-to-shoot biomass production ratio across the grazing intensity gradient and climate variability

According to meta-analyses and recent results (McSherry & Ritchie, 2013; Zeng et al., 2015; Zhou et al., 2017; Li et al., 2018), grazing intensity generally negatively affects the above- and below-ground biomass of grasslands regardless of the climatic conditions or vegetation type, although these effects can be modulated by the level of grazing intensity. The study results do not confirm these findings because the ANPP and BNPP increased in the wet and dry year, respectively, in response to grazing intensity compared to the abandonment treatment. Methodology issues for estimating the ANPP and BNPP should thus be considered for studies comparison. Some papers report the use of biomass stock or fluxes measured once at peak of growth or at several periods (Scurlock et al., 2002), other report estimation of the BNPP from indirect measurements (e.g., Zeng et al., 2015). The root mass, based on stock, provides a snapshot of plant functioning that generally includes mixtures of living and senescent tissues, thus depending on abiotic factors and plant growth, whereas measurements based on new shoot and root biomass reflect the growth potential of grasslands. Although these methods are very different, the BNPP measured with ingrowth cores produced similar results to the root mass stock assessed at three seasons. Another point to consider is the number of samples used to compare treatments and detect significant differences. In grasslands, the coefficient of variation of the root dry weight in auger samples is generally high, i.e. between 30 and 50% (Bengough et al., 2000). According to these authors, we can expect that our sampling protocol (with 16 samples) was adapted to detect at least 35% differences between treatments, whereas to detect less than 10% differences, more than 100 replicates should be collected. While it is possible that collecting more samples would have highlighted significant differences across the treatments, we had to find a compromise between performing more frequent samplings (20 dates) to study the seasonal dynamics of root growth and collecting more samples at the plot level but less frequently.

Negative climatic water balance (P - PET) conditions had stronger effects on the ANPP and BNPP than grazing intensity because severe drought had a direct negative effect on plant growth. In comparison with another experiment located along side ours, 80% of canopy senescence was reached for P – PET of -156 mm (Zwicke et al., 2013). This index reached -303 mm from March to August, confirming that a severe drought occurred in the second year of our experiment explaining the root growth cessation during the summer. At an annual scale, the ANPP of the two cattle treatments showed lower resistance to increased aridity (resistance is defined as ANPP_{year2}/ANPP_{year1} and equalled 0.63) than the abandonment treatment (ratio = 1). For the BNPP, the results were inversed, leading to a lower resistance for the root-to-shoot biomass ratio in the abandonment treatment than in the two cattle treatments. The absence of root growth modification by grazing at an annual scale during the wet year reflects the slight change in root-to-shoot biomass allocation. Other processes, such as root turnover, are expected to change in grazed versus ungrazed grasslands. For our study, Herfurth et al. (2015) observed a similar root mass stock along a grazing disturbance gradient, but by using a simplified C flux model, these authors showed that the Ca+ treatment tended to accelerate C cycling in plant communities, resulting in a higher quantity of C allocated to the soil organic matter continuum. Taken together, these results suggest that the slight BNPP increase

grazing may occur with an increase in rhizodeposition because the root turnover, calculated as the BNPP to root mass stock ratio was not different across the treatments (data not shown; Lauenroth & Gill, 2003).

Furthermore, our results suggest that cattle treatments slow the negative effect of aridity on the rootto-shoot biomass ratio, underling that these treatments seem better adapted to buffering the negative effect of drought on grassland root production than abandonment. This finding is consistent with previous work showing that moderate grazing could be more beneficial than no grazing for drought resistance and the recovery of the ANPP and BNPP (Frank, 2007; Xu et al., 2012). The BNPP has also been shown to have greater resistant than the ANPP to changes in precipitation (Yan et al., 2013). Other studies observed no prevalent effects of grazing, drought, or fire on grassland production in North America or South Africa (Koerner & Collins, 2014). Nevertheless, further research is needed to determine whether grazing pressure has additive or combined effects on the drought response of grasslands (Ruppert et al., 2015).

Community-weighted mean leaf and root traits as predictors of the ANPP and BNPP

As shown by some studies (e.g., Diaz et al., 2007; Laliberté & Tylianakis, 2012; Louault et al., 2017; Zheng et al., 2015), disturbances induced by grazing pressure profoundly affect plant community and functional traits by selecting defoliation-tolerant species, such as *L. perenne*, *P. trivialis*, or *T. repens*, with possible cascading effects on multiple ecosystem functions. Given their capacity to regrow quickly after defoliation, these species generally exhibit high values of SLA and low values of LDMC and plant height. They contrast with species that are adapted to fertile soil but exhibit a slower regrowth capacity after defoliation, such as *D. glomerata* or *F. arundinacea*, with opposite leaf trait values. In abandonment, competition for light tends to select plants with trait syndromes related to disturbance and conservative strategies (e.g., high plant height, low SLA, and high LDMC values). Thus, the CWM traits of a community depend on the balance between these species groups, which are expected to affect the ANPP and BNPP (Klumpp et al., 2009; Milchunas & Lauenroth, 1993). Although species that are tolerant and intolerant to defoliation were present in both cattle treatments, the leaf trait values were similarly and positively related to the ANPP and only differed from the traits of species present in the abandonment treatment. This means that the cessation of grazing strongly differentiated the plant communities, whereas within the two cattle treatments, differences were slighter.

For the below-ground compartment, we expected the above-ground differences to be mirrored by the root growth and traits, assuming that higher root diameter values and lower SRL and SRA values would be associated with a lower BNPP in the abandonment treatment compared with the two cattle treatments. Although the root response to grazing (mainly through defoliation) is generally reported to result in a reduction of root mass or root length (Dawson et al., 2000), our study did not confirm these assumptions. The contrasting results are possibly due to the variable abundance of defoliation-tolerant species or the combined effects of both defoliation and the level of soil fertility on the roots of grazed grasslands (Leuschner et al., 2013; Picon-Cochard et al., 2009; Yan et al., 2013; Ziter & McDougall, 2013). Thus, root growth reductions associated with grazing may have a greater impact in locations where the grazer-mediated N return is spatially decoupled from defoliation (McInenly et al., 2010). Furthermore, the higher SRA observed in the Ca- treatment than in the abandonment or Ca+ treatment should reflect a higher presence of species with fine roots, such as *D. glomerata* or *H. lanatus* (Picon-Cochard et al., 2012), because the soil fertility approximated by the NNI was similar across treatments.

Conclusions

Rotational grazing pressures and climatic condition variabilities had limited effects on root growth seasonality, although drought had stronger effects on the BNPP than on the ANPP. The nearly identical functional traits of the plant communities and similar soil fertility across the two cattle treatments explains the absence of changes in the root mass production for these treatments. Our site disentangled the combined effects of fertility and defoliation on root production, which most previous studies have not addressed. Thus, our results suggest the prevalence of a soil fertility effect on the root production response rather than a defoliation effect. The stability of the root-to-shoot biomass ratio during the dry year evidenced a higher resistance to drought by grazed versus abandoned grassland communities. Moreover, the strong effect of climatic condition variabilities on the ANPP and BNPP observed in the short term could increase in the future as more frequent climatic extremes are expected. It is thus necessary to improve our

knowledge at a larger timescale of the grazing practices that would increase grasslands resilience to more frequent and intense climatic events, such as drought and heat waves.

Acknowledgements

We thank staff from INRAE UREP: V. Guillot and E. Viallard for their technical expertise in field measurements, D. Colosse and S. Toillon for the soil temperature database, and S. Revaillot, A. Bartout, L. Bulon and S. Sauvat and M Mattei (VetAgro Sup) for their help in root sample measurements, and the staff of INRAE UE1414 Herbipôle. Data of the weather station are coming from the platform INRAE CLIMATIK (<u>https://intranet.inrae.fr/climatik/</u>, in French) managed by the AgroClim laboratory of Avignon, France.

A previous version of this article has been peer-reviewed and recommended by Peer Community In Ecology (<u>https://doi.org/10.24072/pci.ecology.100073</u>)

Funding

The experiment is part of the SOERE-ACBB project (<u>http://www.soere-acbb.com/</u>) funded by Allenvi and the French National Infrastructure AnaEE-F through ANR-11-INBS-0001. DH received a doctoral fellowship from VetAgro Sup and DGER pole "ESTIVE". The present work falls within the thematic area of the French government IDEX-ISITE initiative 16-IDEX-0001 (CAP 20-25).

Conflict of interest disclosure

The authors declare that they have no financial conflict of interest with the content of this article.

Data, script and code availability

Data are available online: https://zenodo.org/record/4034903#.YA129-fjJPZ

```
R scripts used in the paper

1) Root growth, soil temperature and relative soil water content

> lme1 <- lme (variable ~ trait*date, random = ~ 1 | bloc/ID,

+ data=IGC_ID, method="ML")

> anova(lme1)
```

plot(lme1)

- > par(mfrow=c(2,2))
- > plot(residuals(lme1))
- > qqPlot(residuals(lme1))
- > hist(residuals(lme1))
- > Ime1.lsmeans <-lsmeans(Ime1, pairwise ~ trait | date)</p>

```
> print(lme1.lsmeans[[2]])
```

```
2) Root mass stock measured at three dates
> Ime1 <- Ime (StockRac ~ trait*date, random = ~ 1 | bloc/ID,</li>
+ data=Stock_ID, method="ML")
```

```
> anova(lme1)
```

```
plot(lme1)
```

- > par(mfrow=c(2,2))
- > plot(residuals(lme1))
- > qqPlot(residuals(lme1))
- > hist(residuals(lme1))

```
> Ime1.lsmeans <-lsmeans(Ime1, pairwise ~ trait | date)</p>
   > print(lme1.lsmeans[[2]])
   3) Root mass stock: averaged of the three dates, root and leaf traits
   > Ime1 <- Ime (variable ~ trait, random = ~ 1 | bloc/ID,</p>
               data=Stock3, method="ML")
   > anova(lme1)
   plot(lme1)
   > par(mfrow=c(2,2))
   > plot(residuals(lme1))
   > qqPlot(residuals(lme1))
   > hist(residuals(lme1))
   > Ime1.lsmeans <-lsmeans(Ime1, pairwise ~ trait)</p>
   > print(lme1.lsmeans[[2]])
   BNPP, ANPP, Root shoot mass ratio, NNI
   > Ime1 <- Ime (variable ~ trait*year, random = ~ 1 | bloc/ID,
               data=BNPP, method="ML")
> anova(lme1)
plot(lme1)
> par(mfrow=c(2,2))
> plot(residuals(lme1))
```

```
> qqPlot(residuals(lme1))
```

```
> hist(residuals(lme1))
```

> lme1.lsmeans <-lsmeans(lme1, pairwise ~ trait | year)

> print(lme1.lsmeans[[2]])

Supplementary information

Table S1. Nitrogen nutrition index (NNI %, Lemaire & Gastal 1997, Cruz et al., 2006) measured on forage regrowth of May in 2014 and 2015 on the non-leguminous part to assess the effect of treatments on N availability according to grazing intensity. When legumes were below 4.5% in the herbage mass, NNI was assessed using the procedure defined by Cruz et al. (2006) based on the total forage and the legume contribution. The P-values are associated with a nested mixed model: treatment used as fixed factor with plots nested in blocks as random factors. Mean ± SE is shown (n = 4). For each year, different letters

correspond to significant differences at P < 0.05.

Year	P-value	Abandonment	Cattle-	Cattle+
2014	0.146	65.64 ± 3.10 a	59.54 ± 1.78 a	63.72 ± 2.86 a
2015	0.018	69.72 ± 1.19 a	61.71 ± 1.53 b	69.25 ± 2.09 a

References

Lemaire G, Gastal F (1997) N uptake and distribution on plant canopy. In: Lemaire, G (ed.) Diagnosis of the nitrogen status in crops, pp. 3-43. Springer-Verlag, Berlin, DE.

Cruz P, Jouany C, Theau J-P, Petibon P, Lecloux E, Duru M (2006) L'utilisation de l'indice de nutrition azotée en prairies naturelles avec présence de légumineuses. Fourrages, 187,369-376.

Group	Species	P-value	Abandonment	Cattle-	Cattle+
Grasses	Agrostis capillaris	ns	0.0 ± 0.0	0.6 ± 0.6	1.7 ± 1.2
	Arrhenatherum elatius	ns	10.3 ± 6.8	2.2 ± 2.2	2.5 ± 2.5
	Alopecurus pratensis	**	27.2 ± 7.9 a	7.8 ± 3.3 b	3.3 ± 1.7 b
	Dactylis glomerata	*	3.1 ± 2.7 b	22.2 ± 9.8 a	9.1 ± 3.8 ab
	Elytrigia repens	*	18.8 ± 9.9 a	2.8 ± 1.8 b	3.8 ± 2.7 b
	Festuca arundinacea ^x	ns	5.0 ± 2.3	5.6 ± 2.1	6.3 ± 2.2
	Holcus lanatus	*	0.0 ± 0.0 b	4.7 ± 1.6 a	3.4 ± 1.9 a
	Lolium perenne	***	0.0 ± 0.0 b	0.9 ± 0.9 b	13.6 ± 3.8 a
	Poa pratensis	ns	11.3 ± 2.2	3.1 ± 1.5	3.4 ± 2.5
	Poa trivialis	*	0.0 ± 0.0 b	5.0 ± 2.5 a	7.2 ± 2.4 a
	Trisetum flavescens	ns	0.0 ± 0.0	2.2 ± 1.3	0.6 ± 0.4
Forbs	Achillea millefolium	ns	1.3 ± 0.9	3.8 ± 2.4	3.1 ± 2.3
	Anthriscus sylvestris	ns	2.5 ± 2.1	0.0 ± 0.0	0.0 ± 0.0
	Cerastium fontanum	ns	0.0 ± 0.0	1.3 ± 0.9	0.0 ± 0.0
	Cerastium	ns	0.0 ± 0.0	0.0 ± 0.0	0.3 ± 0.3
	Cirsium arvense	ns	5.0 ± 3.5	0.0 ± 0.0	0.0 ± 0.0
	Hypocheris radicata	ns	0.0 ± 0.0	0.9 ± 0.9	0.0 ± 0.0
	Ranunculus acris	ns	0.0 ± 0.0	0.0 ± 0.0	3.8 ± 3.8
	Stellaria graminea	ns	0.6 ± 0.6	0.6 ± 0.4	0.0 ± 0.0
	Taraxacum officinale	**	0.0 ± 0.0 b	17.5 ± 1.8 a	19.1 ± 6.0 a
	Urtica dioïca	*	9.7 ± 4.9 a	0.0 ± 0.0 b	0.0 ± 0.0 b
	Veronica serpyllifolia	ns	0.0 ± 0.0	0.3 ± 0.3	0.0 ± 0.0
Legumes	Lathyrus pratensis	ns	0.0 ± 0.0	0.3 ± 0.3	0.3 ± 0.3
	Trifolium pratense	ns	0.0 ± 0.0	0.0 ± 0.0	0.3 ± 0.3
	Trifolium repens	***	0.0 ± 0.0 b	16.3 ± 4.0 a	17.7 ± 2.5 a

Table S2. Species contribution (%) in the community present around the ingrowth core measured in
April and May 2014 for Cattle-, Cattle+ and Abandonment, respectively. Mean ± SE is shown (n = 4). For
each species, different letters correspond to significant differences at *: P < 0.05; **: P < 0.01; ***: P <
0.001; ns: P > 0.05.

×: new species name: Schedonorus arundinaceus

Table S3. Repeated measure ANOVA is shown for root growth (g m⁻² day⁻¹), soil temperature (Tsoil, °C) and relative soil water content (RSWC) responses to treatment, dates (d1 to d20) and interaction effects. Numerator (num), denominator (den) of degree of freedom (DF) and *F* values are shown. Superscripts ^{ns, **, ***} correspond to P > 0.05, P < 0.001, P < 0.0001, respectively.

Variables	Treatment		Dates Treat. x Date			es	
	num/den	F-	num/den	F-	num/den	F-	
Root	2/8	1.80 ⁿ	19/171	50.40*	38/171	2.09	
Tsoil	2/8	33.9	19/166	** 944.83	38/166	-** 9.75	
RSWC	2/8	1.91	19/163	25.287	38/163	** 1.09	



Figure S1: Scheme of the plots and blocks on the experimental site

References

- Bardgett RD, Wardle DA (2003) Herbivore-mediated linkages between aboveground and belowground communities. *Ecology*, **84**, 2258-2268. <u>https,//doi.org/10.1890/02-0274</u>
- Bengough AG, Castrignano A, Pagès L, van Noordwijk M (2000) Sampling Strategies, Scaling, and Statistics. In, Smit AL, Bengough AG, Engels C, van Noordwijk M, Pellerin S, van de Geijn SC (eds) Root Methods. Springer, Berlin, Heidelberg. <u>https://doi.org/10.1007/978-3-662-04188-8_5</u>
- Biswell H, Weaver JE (1933) Effect of frequent clipping on the development of roots and tops of grasses in prairie sod. *Ecology*, **14**, 368-390. <u>https://doi.org/10.2307/1932660</u>
- Brookshire ENJ, Weaver T (2015) Long-term decline in grassland productivity driven by increasing dryness. *Nature Communications*, **6**, 7148. <u>https://doi.org/10.1038/ncomms8148</u>
- Byrne KM, Lauenroth WK, Adler PB (2013) Contrasting Effects of Precipitation Manipulations on Production in Two Sites within the Central Grassland Region, USA. *Ecosystems*, **16**, 1039-1051. <u>https,//doi.org/10.1007/s10021-013-9666-z</u>
- Conant RT, Paustian K, Elliott ET (2001) Grassland management and conversion into grassland, effects on soil carbon. *Ecological Applications*, **11**, 343-355. <u>https://doi.org/10.1890/1051-0761(2001)011[0343,GMACIG]2.0.CO;2</u>
- Dawson LA, Grayston SJ, Paterson E (2000) Effects of grazing on the roots and rhizosphere of grasses. In, Lemaire G, Hodgson J, de Moraes A, Nabinger C, De F. Carvalho PC (eds) Grassland ecophysiology and grazing ecology. CABI Publishing, Wallingford, pp 61–84
- Diaz S, Lavorel S, McIntyre S, Falczuk V, Casanoves F, et al. (2007) Plant trait responses to grazing-a global synthesis. *Global Change Biology*, **13**, 313-341. <u>https://doi.org/10.1111/j.1365-2486.2006.01288.x</u>
- Duru M, Balent G, Gibon A, Magda D, Theau JP, Cruz P, Jouany C (1998) Fonctionnement et dynamique des prairies permanentes. Exemple des Pyrénées centrales. *Fourrages*, **153**, 97-113.
- Edwards EJ, Benham DG, Marland LA, Fitter AH (2004) Root production is determined by radiation flux in a temperate grassland community. *Global Change Biology*, **10**, 209-227. <u>https://doi.org/10.1111/j.1365-2486.2004.00729.x</u>
- Frank DA (2007) Drought effects on above-and belowground production of a grazed temperate grassland ecosystem. *Oecologia*, **152**, 131–139. DOI 10.1007/s00442-006-0632-8
- Garcia-Pausas J, Casals P, Romanyà J, Vallecillo S, Sebastià M-T (2011) Seasonal patterns of belowground biomass and productivity in mountain grasslands in the Pyrenees. *Plant and Soil*, **340**, 315–326. https,//doi.org/10.1007/s11104-010-0601-1
- Herfurth D, Vassal N, Louault F, Alvarez G, Pottier J, Picon-Cochard C, Bosio I, Carrère P (2015) How does soil particulate organic carbon respond to grazing intensity in permanent grasslands? *Plant and Soil*, **394**, 239-255. <u>https,//doi.org/10.1007/s11104-015-2528-z</u>
- Huyghe C, De Vliegher A, van Gils B, Petters A (2014) Grasslands and herbivore production in Europe and effects of common policies. In, Huyghe C, De Vliegher A, van Gils B, Petters A eds, Quæ editions, Versailles, France, 287pp.
- Jones MB, Donnelly A (2004) Carbon sequestration in temperate grassland ecosystems and the influence of management, climate and elevated CO₂. *New Phytologist*, **164**, 423-439. <u>https,//doi.org/10.1111/j.1469-8137.2004.01201.x</u>
- Klumpp K, Fontaine S, Attard E, Le Roux X, Gleixner G, Soussana JF (2009) Grazing triggers soil carbon loss by altering plant roots and their control on soil microbial community. *Journal of Ecology*, **97**, 876–885. <u>https://doi.org/10.1111/j.1365-2745.2009.01549.x</u>
- Klumpp K, Soussana JF (2009) Using functional traits to predict grassland ecosystem change, a mathematical test of the response-and-effect trait approach. *Global Change Biology*, **15**,2921-2934. https,//doi.org/10.1111/j.1365-2486.2009.01905.x
- Koerner SE, Collins SL (2014) Interactive effects of grazing, drought, and fire on grassland plant communities in North America and South Africa. *Ecology*, **95**, 98-109. <u>https://doi.org/10.1890/13-0526.1</u>
- Laliberté E, Tylianakis JM (2012) Cascading effects of long-term land-use changes on plant traits and ecosystem functioning. *Ecology*, **93**, 145–155. <u>https://doi.org/10.1890/11-0338.1</u>

- Lauenroth WK, Gill R (2003) Turnover of root systems. In, de Kroon H, Visser EJW (eds) Root Ecology, Ecological Studies (Analysis and Synthesis), vol 168. Springer, Berlin, Heidelberg. https,//doi.org/10.1007/978-3-662-09784-7 3
- LeCain DR, Morgan JA, Schuman GE, Reeder JD, Hart RH (2002) Carbon exchange and species composition of grazed pastures and exclosures in the shortgrass steppe of Colorado. *Agriculture, Ecosystems & Environment*, **93(1)**, 421-435. <u>https://doi.org/10.1016/S0167-8809(01)00290-0</u>
- Lemaire G, Gastal F (1997) N uptake and distribution on plant canopy. In, Lemaire, G (ed) Diagnosis of the nitrogen status in crops, Springer-Verlag, Berlin, pp 3-43. <u>https://doi.org/10.1007/978-3-642-60684-7_1</u>
- Leuschner C, Gebel S, Rose L (2013) Root trait responses of six temperate grassland species to intensive mowing and NPK fertilisation, a field study in a temperate grassland. *Plant and Soil*, **373**, 687-698. <u>https,//doi.org/10.1007/s11104-013-1836-4</u>
- Li W, Li X, Zhao Y, Zheng S, Bai Y (2018) Ecosystem structure, functioning and stability under climate change and grazing in grasslands, current status and future prospects. *Current Opinion in Environmental Sustainability*, **33**, 124-135. <u>https://doi.org/10.1016/j.cosust.2018.05.008</u>
- Loiseau P, Louault F, Le Roux X, Bardy M (2005) Does extensification of rich grasslands alter the C and N cycles, directly or via species composition? *Basic and Applied Ecology*, **6**, 275-287. https,//doi.org/10.1016/j.baae.2004.07.006
- Louault F, Pillar VD, Aufrere J, Garnier E, Soussana JF (2005) Plant traits and function types in response to reduced disturbance in a semi-natural grassland. *Journal of Vegetation Science*, **16**, 151–160. <u>https://doi.org/10.1111/j.1654-1103.2005.tb02350.x</u>
- Louault F, Pottier J, Note P, Vile D, Soussana JF, Carrère P (2017) Complex plant community responses to modifications of disturbance and nutrients availability in productive grasslands. *Journal of Vegetation Science*, 28, 538-549; <u>https,//doi.org/10.1111/jvs.12509</u>
- McInenly LE, Merrill EH, Cahill JF, Juma NG (2010) Festuca campestris alters root morphology and growth in response to simulated grazing and nitrogen form. *Functional Ecology*, **24**, 283-292. <u>https,//doi.org/10.1111/j.1365-2435.2009.01642.x</u>
- McSherry ME, Ritchie ME (2013) Effects of grazing on grassland soil carbon, a global review. *Global Change Biology*, **19**, 1347–1357. <u>https://doi.org/10.1111/gcb.12144</u>
- Milchunas DG, Lauenroth WK (1993) Quantitative effects of grazing on vegetation and soils over a global range of environments. *Ecological Monographs*, **63**, 327-366. <u>https://doi.org/10.2307/2937150</u>
- Moretto AS, Distel RA, Didoné NG (2001) Decomposition and nutrient dynamic of leaf litter and roots from palatable and unpalatable grasses in a semi-arid grassland. *Applied Soil Ecology*, **18**, 31–37. <u>https,//doi.org/10.1016/S0929-1393(01)00151-2</u>
- Pagès L, Picon-Cochard C (2014) Modelling the root system architecture of Poaceae. Can we simulate integrated traits from morphological parameters of growth and branching? *New Phytologist*, **204**,149-158. <u>https://doi.org/10.1111/nph.12904</u>
- Pérez-Ramos IM, Roumet C, Cruz P, Blanchard A, Autran P, Garnier E (2012) Evidence for a 'plant community economics spectrum' driven by nutrient and water limitations in a Mediterranean rangeland of southern France. *Journal of Ecology*, **100**, 1315-1327. <u>https://doi.org/10.1111/1365-2745.12000</u>
- Picon-Cochard C, Coll L, Balandier P (2006) The role of below-ground competition during early stages of secondary succession, the case of three-year-old Scots pine (*Pinus sylvestris* L.) seedlings in an abandoned grassland. *Oecologia*, **148**, 373-383. <u>https://doi.org/10.1007/s00442-006-0379-2</u>
- Picon-Cochard C, Pilon R, Revaillot S (2009) Plasticity of grass root functional traits and root mass in response to cutting frequency and N fertilisation. Proceedings of the 7th ISRR Symposium, Root Research and Applications (RootRAP), Vienne September 2–4, 2009. 4pp. <u>https://hal.inrae.fr/hal-02758074</u>
- Picon-Cochard C, Pilon R, Tarroux E, Pagès L, Robertson J, Dawson L (2012) Effects of species, root branching order and season on root traits of 13 perennial grass species. *Plant and Soil*, **353**,47-57. https://doi.org/10.1007/s11104-011-1007-4
- Pilon R, Picon-Cochard C, Bloor JMG, Revaillot S, Kuhn E, Falcimagne R, Balandier P, Soussana JF (2013) Grassland root demography responses to multiple climate change drivers depend on root morphology. *Plant and Soil*, **364**, 395-408. <u>https://doi.org/10.1007/s11104-012-1371-8</u>

- Pineiro G, Paruelo JM, Oesterheld M, Jobbagy EG (2010) Pathways of grazing effects on soil organic carbon and nitrogen. *Rangeland Ecology and Management*, **63**, 109–119. <u>https://doi.org/10.2111/08-255.1</u>
- Pinheiro J, Bates D, Debroy S, Sarkar D and R Core Team (2015) nlme, linear and non linear mixed effect models. R Package Version 3.1-119. <u>https://CRAN.R-project.org/package=nlme</u>.
- R Core Team (2012) R, a language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria.
- Reich PB (2014) The world-wide 'fast-slow' plant economics spectrum, a traits manifesto. *Journal of Ecology*, **102**, 275-301. <u>https://doi.org/10.1111/1365-2745.12211</u>
- Rossignol N, Chadoeuf J, Carrère P, Dumont B (2011) A hierarchical model for analysing the stability of vegetation patterns created by grazing in temperate pastures. *Applied Vegetation Science*, **14**, 189-199. https://doi.org/10.1111/j.1654-109X.2010.01106.x
- Ruppert JC, Harmoney K, Henkin Z, Snyman HA, Sternberg M, Willms W, Linstädter A (2015) Quantifying drylands' drought resistance and recovery, the importance of drought intensity, dominant life history and grazing regime. *Global Change Biology*, **21**, 1258-1270. <u>https://doi.org/10.1111/gcb.12777</u>
- Schaffers A, Sykora K (2000) Reliability of Ellenberg indicator value for moisture, nitrogen and soil reaction, a comparison with field measurements. *Journal of Vegetation Science*, **11**, 225-244. https,//doi.org/10.2307/3236802
- Scurlock JMO, Johnson K, Olson RJ (2002) Estimating net primary productivity from grassland biomass dynamics measurements. *Global Change Biology*, **8**, 736-753. <u>https://doi.org/10.1046/j.1365-2486.2002.00512.x</u>
- Smith SW, Woodin SJ, Pakeman RJ, Johnson D, van der Wal R (2014) Root traits predict decomposition across a landscape-scale grazing experiment. *New Phytologist*, **203**, 851-862. <u>https,//doi.org/10.1111/nph.12845</u>
- Soussana JF, Loiseau P, Vuichard N, Ceschia E, Balesdent J, Chevallier T, Arrouays D (2004) Carbon cycling and sequestration opportunities in temperate grasslands. *Soil Use and Management*, **20**, 219-230. https,//doi.org/10.1111/j.1475-2743.2004.tb00362.x
- Soussana JF, Duru M (2007) Grassland science in Europe facing new challenges, biodiversity and global environmental change. CAB Reviews, Perspectives in Agriculture, Veterinary Science, Nutrition and Natural Resources, **272**, 1-12.
- Soussana JF, Lemaire G (2014) Coupling carbon and nitrogen cycles for environmentally sustainable intensification of grasslands and crop-livestock systems. *Agriculture, Ecosystems and Environment*, **190**, 9-17. <u>https://doi.org/10.1016/j.agee.2013.10.012</u>
- Steinaker DF, Wilson SD (2008) Phenology of fine roots and leaves in forest and grassland. *Journal of Ecology*, **96**, 1222-1229. <u>https://doi.org/10.1111/j.1365-2745.2008.01439.x</u>
- Wardle DA, Bardgett RD, Klironomos JN, Setala H, van der Putten WH, Wall DH (2004) Ecological linkages between aboveground and belowground biota. *Science*, **304**, 1629-1633. <u>DOI</u>, <u>10.1126/science.1094875</u>
- Wilcox KR, von Fischer JC, Muscha JM, Petersen MK, Knapp AK (2015) Contrasting above- and belowground sensitivity of three Great Plains grasslands to altered rainfall regimes. *Global Change Biology*, **21**, 335-344. <u>https://doi.org/10.1111/gcb.12673</u>
- Xu X, Sherry RA, Niu S, Li D, Luo Y (2012) Net primary productivity and rain-use efficiency as affected by warming, altered precipitation, and clipping in a mixed-grass prairie. *Global Change Biology*, **19**, 2753-2764. <u>https://doi.org/10.1111/gcb.12248</u>
- Xu X, Luo Y, Shi Z, Zhou X, Li D (2014) Consistent proportional increments in responses of belowground net primary productivity to long-term warming and clipping at various soil depths in a tallgrass prairie. *Oecologia*, **174**, 1045-1054. <u>https://doi.org/10.1007/s00442-013-2828-z</u>
- Yan L, Zhou G, Zhang F (2013) Effects of different grazing intensities on grassland production in China, A meta-analysis. *PLoS ONE*, **8**, e81466. <u>https://doi.org/10.1371/journal.pone.0081466.s002</u>
- Zeng C, Wu J, Zhang X (2015) Effects of grazing on above- vs. below-ground biomass allocation of alpine grasslands on the northern tibetan plateau. *PLoS ONE*, **10**, e0135173. <u>https,//doi.org/10.1371/journal.pone.0135173</u>
- Zheng S, Li W, Lan Z, Ren H, Wang K (2015) Functional trait responses to grazing are mediated by soil moisture and plant functional group identity. *Scientific Reports*, **5**, 18163. <u>https,//doi.org/10.1038/srep18163</u>

- Zhou G, Zhou X, He Y, Shao J, Hu Z, Liu R, Zhou H, Hosseinibai S (2017) Grazing intensity significantly affects belowground carbon and nitrogen cycling in grassland ecosystems, a meta-analysis. *Global Change Biology*, 23, 1167-1179. <u>https://doi.org/10.1111/gcb.13431</u>
- Zhu J, Zhang Y, Liu Y (2016) Effects of short-term grazing exclusion on plant phenology and reproductive succession in a Tibetan alpine meadow. *Scientific Reports*, **6**, 27781. <u>https://doi.org/10.1038/srep27781</u>
- Ziter C, MacDougall AS (2013) Nutrients and defoliation increase soil carbon inputs in grassland. *Ecology*, **94**, 106-116. <u>https://doi.org/10.1890/11-2070.1</u>
- Zwicke M, Alessio GA, Thiery L, Falcimagne R, Baumont R, Rossignol N, Soussana JF, Picon-Cochard C (2013) Lasting effects of climate disturbance on perennial grassland above-ground biomass production under two cutting frequencies. *Global Change Biology*, **19**, 3435–3448. <u>https://doi.org/10.1111/gcb.12317</u>