Response of temperate grasslands at different altitudes to simulated summer drought differed but scaled with annual precipitation

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Abstract. Water is an important resource for plant life. Since climate scenarios for Switzerland predict an average reduction of 20% in summer precipitation until 2070, understanding ecosystem responses to water shortage, e.g. in terms of plant productivity, is of major concern. Thus, we tested the effects of simulated summer drought on three managed grasslands along an altitudinal gradient in Switzerland from 2005 to 2007, representing typical management intensities at the respective altitude. We assessed the effects of experimental drought on above- and below-ground productivity, stand structure (LAI and vegetation height) and resource use (carbon and water). Responses of community above-ground productivity to reduced precipitation input differed among the three sites but scaled positively with total annual precipitation at the sites ($R^2=0.85$). Annual community above-ground biomass productivity was significantly reduced by summer drought at the alpine site receiving the least amount of annual precipitation, while no significant decrease (rather an increase) was observed at the pre-alpine site receiving highest precipitation amounts in all three years. At the lowland site (intermediate precipitation sums), biomass productivity significantly decreased in response to drought only in the third year, after showing increased abundance of a drought tolerant weed species in the second year. No significant change in below-ground biomass productivity was observed at any of the sites in response to simulated summer drought. However, vegetation carbon isotope ratios increased under drought conditions, indicating an increase in water use efficiency. We conclude that there is no general drought response of Swiss grasslands, but that sites with lower annual precipitation seem to be more vulnerable to summer drought than sites with higher annual precipitation, and thus site-specific adaptation of management strategies will be needed, especially in regions with low annual precipitation.

1 Introduction

Water availability is among the strongest limitations to plant productivity globally, even in temperate or boreal regions (Lambers et al., 1998). Annual above-ground productivity of vegetation strongly depends on mean annual precipitation (for grassland: Paruelo et al., 1999; Knapp and Smith, 2001). Water stress for plants can arise from low precipitation inputs, high rates of water loss due to high atmospheric vapour pressure deficit (VPD) or from drying soil, all mechanisms ultimately reducing plant productivity. Such effects might become more pronounced since water stress is predicted to increase in the future in certain regions due to climate change, e.g. in Central Europe, for which decreasing summer precipitation is expected (Christensen et al., 2007). Projections for Switzerland indicate that by 2070 the mean decrease of summer precipitation (June through August) might be around 20% compared to 1990 with a maximum decrease of 40% (Frei et al., 2006). Thus, we expect ecosystem processes, from microbial activities to plant performance, to be strongly affected. At the same time, the response to drought of terrestrial ecosystems might vary dependent on vegetation composition and local environmental conditions.

While research on drought effects on grassland species has often been carried out under controlled conditions (e.g. Arp et al., 1998; Karsten and MacAdam, 2001), research at the ecosystem level in the field used two approaches: (1) naturally occurring droughts and their impact on the long-term field trials (Weaver et al., 1935; Gibbens and Beck, 1988;
2.1 Experimental sites and setup

The study was conducted at three different temperate grassland sites across Switzerland, representing Swiss grassland systems at the respective altitudes (Table 1). The experiment was established by the end of June 2005 at the two lower sites, Chamau and Früebüel, and in July 2006 at the alpine site, Alp Weissenstein, and was continued until 2007, thus during three and two growing seasons, respectively. At each site, we installed five portable rain shelters and excluded rainfall in spring/summer to simulate a pronounced drought. In 2006, two additional rain shelters and control plots were installed at Chamau and one each at Früebüel. The tunnel-shaped rain shelters had an area of $3 \times 3.5$ m and were about 2.1 m high at the highest point. During the entire drought

<table>
<thead>
<tr>
<th>Political community</th>
<th>Chamau</th>
<th>Früebüel</th>
<th>Alp Weissenstein</th>
</tr>
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<tbody>
<tr>
<td>Coordinates WGS84</td>
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<td>Bergün, GR</td>
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<td>May to mid-October</td>
<td>mid-June to September</td>
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<tr>
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<td>gleyso$^b$</td>
<td>humus sand loam$^b$</td>
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<tr>
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<td>&gt;120 cm$^a$</td>
<td>30 cm</td>
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<tr>
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<td>7.7°C</td>
<td>2.3°C</td>
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<td>Vegetation type (dominant species)</td>
<td>artificial grass-legume mixture ($Trifolium repens, Lolium perenne, Poa pratensis$)</td>
<td>permanent managed pasture ($Dactylis glomerata, Alopecurus pratensis, Taraxacum officinale$)</td>
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<td>35</td>
<td>around 20</td>
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<td>silage/hay; cattle grazing in autumn</td>
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$^a$ Roth (2006)  
$^b$ Schärer (2003)  
$^c$ data from Zeeman (2008) adapted with data from MeteoSwiss  
$^d$ Sautier (2007)  
$^e$ Keller (2006)
2.2 Micrometeorological measurements

Microclimatic variables were continuously monitored close to the centre of two drought and control plots per site, starting in autumn 2005 at Früebüel, spring 2006 at Chamau and summer 2006 at Alp Weissenstein. Soil temperature (Precision IC Temperature Transducer AD592AN, Analog Devices, Norwood, MA, USA), soil moisture at three soil depth (5, 15 and 30 cm) using 20 cm long ECH2O probes (EC-20, Decagon Devices, Inc., Pullman, WA, USA) and soil heat flux (HF01, Hukseflux Thermal Sensors B.V., Delft, Netherlands) close to the soil surface were recorded. We also monitored air temperature at 60 and 160 cm (using the same sensors as for soil temperature but with ventilation) and photosynthetically active radiation (PAR LITE, Kipp & Zonen B.V., Delft, Netherlands). The air temperature sensors were disconnected in winter and spring 2006/2007 due to technical modifications in the setup. After mid-October 2007, the air temperature sensors were not ventilated any longer to save battery lifetime. Measurements were made every 10 s while ten minute averages were logged with a CR10X data logger (Campbell Scientific Inc., Logan, UT, USA).

In addition, we assessed soil moisture by taking additional measurements of gravimetric soil water content in 2007. We therefore sampled 15 cm deep soil cores from the centre of the plots and divided them into three 5 cm pieces. Samples were stored in tightly sealed plastic bags and fresh weight was measured immediately after returning from the field. The soil cores were dried to weight constancy at 100 °C (some days) and dry weight was measured afterwards. Gravimetric soil water content was then calculated as the difference of the fresh and the dry weight (weight of the water) divided by the dry weight (weight of the soil).

2.3 Above-ground productivity

Above-ground biomass was harvested at the cutting dates of the surrounding farm, i.e. six times per year at Chamau (three times in 2005), two times per year at Früebüel (once in 2005) and once at the end of the growing season (end of September) at Alp Weissenstein. Biomass was collected using 20×50 cm frames that were randomly placed on the plots in 2005 and installed at fixed locations starting spring 2006. Cutting height of the vegetation was approximately 7 cm above the soil according to the common management practice on the farms. Two samples per plot were taken and then pooled for the analyses (representing 0.2 m²). Biomass was stored in plastic bags at 4 °C for a maximum of one week.
until it could be separated into species (plant functional types for Alp Weissenstein) and dried at 60°C until weight constancy. Dry matter was then determined for each species sample, and the sum of all samples from the same plots was used as an estimate for community above-ground biomass productivity. Each species was assigned a plant functional type (grass, forb (i.e. non-leguminous forb) or legume). The dead biomass (often also referred to as necromass) was considered its own plant functional type and was never separated into species (i.e. biomass of the other three functional types was alive by definition). It was however included in community above-ground productivity estimates (often also called phytomass).

2.4 LAI and vegetation height

During the growing seasons 2006 and 2007 (only 2007 at Alp Weissenstein), leaf area index (LAI) was measured approximately 7 cm above soil (cutting height) using an LAI-2000 Plant Canopy Analyzer (LI-COR Biosciences, Lincoln, NE, USA). Five measurements distributed over the plot were averaged to represent the plot LAI. A 270° view cap was used to reduce plots size required for measurements, i.e. only 90° of the sensor view were used for measurements. Vegetation height was estimated at the same dates as LAI using a stick and a falling styrofoam plate (0.5×0.5 m, 1 cm thick). Two to four measurements were averaged for each plot.

2.5 Root biomass productivity

Root biomass productivity at the community level over the complete growing season was determined at Chamau and Früebüel in 2007 and over nearly two growing seasons (2006 and 2007) at Alp Weissenstein using ingrowth cores (4.4 cm diameter, 30 cm length). Cores were positioned in an angle of approximately 45°. All ingrowth cores were filled with root-free, sieved (2 mm) soil from the corresponding site. In-growth cores were installed from 13 March 2007 to 1 December 2007 at Chamau (263 days), from 15 December 2006 to 1 December 2007 at Früebüel (351 days), and from 14 July 2006 to 25 September 2007 at Alp Weissenstein (438 days). One ingrowth core per plot was buried at Alp Weissenstein while two cores per plot were used at the two other sites. After removal from the soil, cores were stored at 4°C until further analysis. The ingrown roots were washed from the soil cores in the laboratory, roots were dried to weight constancy at 60°C and the dry weight was determined. Where two ingrowth cores had been installed and recovered, root biomass of two cores was pooled to calculate root productivity of the plot. In some cases, part of the soil was lost when removing the ingrowth cores from the soil. Therefore, the actual length of the soil core was used to calculate the amount of roots per unit soil depth. As the ingrowth cores remained in the field for different time periods at the three sites, the root weight was divided by the number of days that roots had been allowed to grow into the cores and then multiplied by 365 to represent the root mass per m² and year.

2.6 Carbon isotope and nitrogen concentration measurements

To determine carbon isotope ratios (δ¹³C) and nitrogen concentrations, the most abundant species were sampled at Chamau (Agrostis stolonifera L., Alopecurus pratensis L., Dactylis glomerata L., Lolium multiflorum LAM., Phleum pratense L. AGG., Poa pratensis L. AGG., Poa trivialis L. S.L., Rumex obtusifolius L., and Trifolium repens L.) and Früebüel (Agrostis capillaris L., A. stolonifera, A. pratensis, Anthoxanthum odoratum L., P. pratense, P. pratensis, P. trivialis, Rumex acetosa L., R. obtusifolius, and T. repens), while the four plant functional types were analysed at Alp Weissenstein. Biomass was ground to a fine powder and analysed using a Flash EA 1112 Series elemental analyser (Thermo Italy, former CE Instruments, Rhodano, Italy) coupled to a Finnigan MAT DeltaPlus XP isotope ratio mass spectrometer (Finnigan MAT, Bremen, Germany) via a 6-port valve (Brooks et al., 2003), a ConFlo III (Werner et al., 1999) and an additional Nafion-trap backed by a conventional Mg(Clo4)2-trap followed by a 4-port valve (Werner, 2003) between reduction tube and GC column. Post-run offline calculations (blank, offset and possibly drift corrections) were performed to assign the final δ-values on the V-PDB and AIR-N₂ scales according to Werner and Brand (2001). The long-time precision for the lab’s quality control standard tyrosine (~2.5 years) was 0.05‰ for δ¹³C. Precision for N concentrations was 0.05%. For statistical analyses, isotope and nitrogen data were weighted by biomass to calculate a vegetation and functional group mean for each plot.

2.7 Statistical analysis

Due to the rather small number of replicates (n=5–7), all plots were included in the statistical analyses of biomass, vegetation height, δ¹³C and N concentrations, i.e. no outliers were excluded. LAI values were excluded only if the standard error of the five measurements per plot was higher than a third of the mean LAI of the plot or if MTA (mean tilt angle) was below 30% (criteria used for 2007 since this information was not available for 2006).

Statistical analyses were performed using R 2.5.0 (R Development Core Team, 2007). Community above-ground and functional type biomass, LAI, vegetation height and isotope signatures were generally all analysed with the same ANOVA model. In a first step, the full datasets were analysed (all sites over all years), using an ANOVA model considering site, harvest date, treatment (control vs. drought treatment) and all interactions, including the triple interaction. In a second step, datasets were analysed for each year separately using the same model. In a third step, the dataset was analysed for each site separately and finally for each harvest.
Fig. 1. Micrometeorology during the experiment at Chamau: daily precipitation sum (a), difference of daily average air temperature at 160 cm between drought and control plots (b), proportion of total photosynthetically active radiation (PAR) relative to the control plots (c), difference of daily average soil temperature at 5 cm depth between drought and control plots (d), and relative development of soil moisture at 15 cm depth (relative to the soil moisture at the beginning of the drought treatment in 2006; e). Solid lines: control plots, dashed lines: drought plots. Measurements of gravimetric soil water content in soil cores from 10–15 cm are given as points (open symbols for drought plots). The periods of drought treatment are shaded in grey. Averages of two plots per treatment are shown.

3 Results

3.1 Micrometeorological measurements

Annual precipitation sums were quite similar in 2005 (1170 mm and 1481 mm at Chamau and Früebüel, respectively) and 2006 (1136 mm, 1649 mm and 867 mm at Chamau, Früebüel and Alp Weissenstein, respectively), but around 100 mm higher at all three sites in 2007 (1232 mm, 1765 mm and 969 mm at Chamau, Früebüel and Alp Weissenstein, respectively, Zeeman et al., 2009, adjusted with data from nearby MeteoSwiss stations). While in 2006, July was naturally dry, spring 2007 started with very low precipitation but was followed by a rather wet summer 2007 (Fig. 1a).

The rain shelters had no substantial effect on daily average and minimum air temperatures at 160 cm height (shown for Chamau; Fig. 1b): at Chamau, the average temperature change during the drought treatment was $-0.03^\circ C$, at Früebüel $0.2^\circ C$ and at Alp Weissenstein $0.1^\circ C$. The rain shelters increased daily maximum temperatures at Chamau and Früebüel (by around $0.8^\circ C$) while no effect was seen at Alp Weissenstein. The plastic foils reduced average daily PAR sums by 20% at Chamau and Früebüel and by 26% at Alp Weissenstein (Fig. 1c). The effect of the shelters on soil temperature was also small: in general, there was an increase of less than $1^\circ C$ (but a decrease of $0.3^\circ C$ at Alp Weissenstein) in soil temperature under the shelters compared to unsheltered periods (Fig. 1d). However, the rain shelters had the
Drought also affected vegetation composition and species richness. At Chamau, the average number of species was only slightly affected in the first two years (2005 and 2006), but significantly reduced by drought from 7.3 to 5.3 in 2007. At Alp Weissenstein, the average species number was significantly reduced in 2006 but not in 2007, while at Früebüel, no significant impact on species richness was found (data not shown). Focusing on the drought responses of the four plant functional types (PFT) separately (Table 4, upper part) revealed significant differences among sites (except for forbs that grew similar amounts of biomass at all sites) but only slight differences of annual PFT above-ground productivities between treatments. Thus, over the three-year experiment, site was the most important factor influencing annual above-ground biomass productivities of the community as well as of individual PFTs. In comparison, impacts of the drought treatment significantly only affected the annual amount of dead biomass in 2006 ($P=0.05$) and grass biomass in 2007 ($P=0.03$), while annual above-ground productivities of forbs

desired effect on soil moisture. Maximum reduction of soil moisture was around 83% at Chamau, with soil moisture being reduced by 20 to 60% during the drought treatment. Additional measurements of gravimetric soil water content showed a similar pattern as the continuous measurements (Fig. 1e). Due to lack of rain in the weeks prior to the start of the treatment, soil moisture dropped to very low levels in both, drought and control plots before the start of the drought treatment in 2007. At Früebüel, the effect of the shelters on soil moisture was smaller than at Chamau because the site was generally much wetter: soil moisture was reduced on average by 30% at 30 cm soil depth during the drought treatment. At Alp Weissenstein, no data from continuous measurements were available due to technical problems, but drought significantly reduced gravimetric soil water content in 0–5 cm depth at the end of the drought treatment (absolute reduction of 41%, $P<0.001$).

### 3.2 Above-ground productivity

Overall, annual community above-ground biomass differed significantly among the three sites ($P<0.001$). At Chamau, annual community above-ground productivities were similar in 2005 and 2006 for both treatments but drought tended to decrease community above-ground productivity in 2007. No treatment effect was found at Früebüel in any of the three years, while annual community above-ground productivity at Alp Weissenstein was significantly decreased by drought in both years (Table 3). However, above-ground productivities at Chamau were strongly influenced by one weed species, *Rumex obtusifolius*, which gained competitive advantage over the other species in 2006 (Gilgen et al., 2010). We therefore excluded *R. obtusifolius* biomass at Chamau from all further above-ground productivity analyses. As a result, the overall drought effect across all sites and all three years became significant ($P=0.02$ when including year in the analysis, data not shown). We found a strong relationship of the average annual community above-ground biomass response with annual precipitation sum ($R^2=0.73$, $P=0.004$; without *Rumex*: $R^2=0.85$, $P<0.001$; Fig. 2a), which was much stronger than the relationship with amount of precipitation excluded ($R^2=0.33$, $P=0.08$; without *Rumex*: $R^2=0.60$, $P=0.01$; Fig. 2b). No such relationship was found when relating above-ground biomass response to the fraction of annual precipitation excluded by the shelters ($R^2<0.001$, $P=0.89$) or to growing degree days ($R^2<0.001$, $P=0.5$).

### Table 3. Effect of drought on community (i.e. sum of dead and alive), alive and dead annual above-ground biomass productivity at Chamau (including *Rumex obtusifolius*), Früebüel and Alp Weissenstein in g m$^{-2}$. Means and standard errors are given ($n=5–6$). Significantly different means ($P \leq 0.05$) are given in bold, marginally significant differences ($0.1 \geq P > 0.05$) in italics.

<table>
<thead>
<tr>
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<th>Chamau</th>
<th>Früebüel</th>
<th>Alp Weissenstein</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>control</td>
<td>drought</td>
<td>control</td>
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<td>2005$^a$</td>
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<td>425±60</td>
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<td></td>
<td>alive</td>
<td>457±55</td>
<td>398±60</td>
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<tr>
<td></td>
<td>dead</td>
<td>20±4</td>
<td>26±5</td>
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<td>community</td>
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<td>alive</td>
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<td></td>
<td>dead</td>
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<td>33±4</td>
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<td>2007</td>
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<td>1211±134</td>
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<td>alive</td>
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<td>dead</td>
<td>52±6</td>
<td>67±6</td>
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</tbody>
</table>

$^a$ Annual biomass productivity sum based on only three (instead of six) and one (instead of two) harvests at Chamau and Früebüel, respectively.
and legumes were not significantly affected in any of the three years.

Community above-ground regrowth (i.e. biomass grown between cutting dates) across all sites and all three years of the experiment (Fig. 3, Table 4, lower part) was affected by site and harvest date ($P<0.001$) as well as by the drought treatment ($P=0.003$). A significant site × treatment interaction ($P=0.001$) indicated differing grassland responses to drought at the three sites, as already observed for the annual community above-ground productivities. Including year in the analysis did not change significance levels and was thus omitted. When analysing the three years separately, site and harvest date explained a large fraction of the variation in productivity in all three years (2005–2007), while the treatment effect was insignificant in 2005, it became marginally significant in 2006 ($P=0.06$) and stayed significant in 2007 ($P=0.01$). In 2007, the drought treatment showed highly significant effects on above-ground productivities at Chamau ($P=0.001$) and Alp Weissenstein ($P<0.001$), but still no effects at Fruebuel ($P=0.5$; Fig. 3). These annual patterns could also be seen in the response of the single harvests that were unaffected by the drought treatment at Fruebuel and in the two first years at Chamau (except for the fifth harvest in 2006), but significantly reduced at Chamau in the third year (first, third and fourth harvest) and also strongly reduced at Alp Weissenstein (Fig. 3). The date × treatment interaction term was never significant, indicating that the direction of the drought responses of the respective grasslands was stable over time (Table 4, lower part).

Focusing on the different PFTs clearly showed that grasses resembled the drought response of the community above-ground productivity best (independent of Rumex). The drought treatment clearly decreased grass regrowth above-ground ($P=0.001$). Although this response differed among sites (drought strongly decreased grass productivities at Chamau and Alp Weissenstein but no effect was observed at Fruebuel; site × treatment interaction: $P<0.001$), the response of grasses to drought was stable over the growing season (harvest date × treatment interaction: not significant). Furthermore, similar patterns for community and grass above-ground productivity were also found when analysing the years separately (except site effect in 2005): grass productivity differed among the three sites and grass regrowth changed during the growing season (site and harvest date effects: $P<0.001$). Forbs reacted to drought with decreased regrowth (overall: $P=0.007$), particularly in 2006. Unlike their annual productivities, forb regrowth differed between sites and also during the season (site and harvest date effects: $P<0.001$), but the direction of the drought response was unaffected by site and harvest date (overall site × treatment and harvest date × treatment interactions: $P>0.6$). Furthermore, legumes were generally positively affected by the drought treatment (overall: $P=0.005$), particularly in the year 2007, but this was driven by the positive drought response at Chamau (data not shown).

**3.3 LAI and vegetation height**

LAI measurements during 2006 and 2007 (Fig. 4) represented above-ground biomass regrowth patterns in higher temporal resolution than community above-ground productivity and varied significantly during the growing seasons ($P<0.001$). Similar to productivity measurements, no significant drought effect on LAI was found at Chamau in 2006 (except very early in the season), probably confounded by Rumex abundance. In contrast, the drought response of LAI in 2007 was stronger than that of above-ground productivity, with drought lowering LAI values significantly ($P<0.001$). At Fruebuel, LAI was negatively affected by drought in
Table 4. Results of the ANOVA models for annual sums of above-ground biomass and above-ground harvest yields of community (comm.=sum of dead and alive biomass) and plant functional type (grass, forb (excluding Rumex obtusifolius at Chamau), legume; only alive biomass) as well as dead above-ground biomass. Main factors are site (Chamau, Fruebuel or Alp Weissenstein), harvest date and treatment (drought vs. control). Significant $P$ values ($P \leq 0.05$) are given in bold, marginally significant values ($0.1 \geq P > 0.05$) in italics; $n=5–6$.

<table>
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<td>0.3</td>
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<td>0.09</td>
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<td>0.03</td>
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Fig. 3. Effect of summer drought on community above-ground biomass production (i.e. dead and alive plant material) at Chamau (excluding R. obtusifolius biomass; a–c), Fruebuel (d–f) and Alp Weissenstein (g–h) during the experiment. Bars represent biomass production per harvest (i.e. biomass regrown between cutting dates). Means and standard errors are given ($n=5–7$). Periods of drought treatment are shaded in grey. * $0.05 \geq P > 0.01$, ** $0.01 \geq P > 0.001$. 

both years ($P < 0.001$) although there was no such effect on community above-ground biomass. At Alp Weissenstein, drought reduced LAI values ($P = 0.02$), but despite the large decrease in biomass under drought conditions this trend was not significant at the different measurement dates. No relationship of LAI before the cut with harvested community above-ground biomass was found at any of the three sites ($R^2 \leq 0.1$, $P > 0.2$ but $P < 0.001$ for Chamau).

Vegetation height (data not shown) developed similar to LAI but was not as strongly affected by the drought treatment as LAI (except for Alp Weissenstein where vegetation height was significantly reduced by drought). Vegetation height before the cuts was not related to community above-ground biomass at Chamau and Alp Weissenstein, in contrast to Fruebuel ($R^2 = 0.4$, $P < 0.001$).

### 3.4 Root biomass

Root biomass productivity differed significantly among sites ($P = 0.01$), with productivities being almost twice as high at Alp Weissenstein compared to Fruebuel and Chamau (Fig. 5). However, below-ground productivity was not affected by the drought treatment ($P = 0.9$). In addition, the site × treatment interaction term was not significant ($P = 0.9$), indicating that the response of below-ground productivity to drought was similar across all three sites.

### 3.5 Carbon isotope and nitrogen concentration measurements

Bulk above-ground carbon isotope ratios ($\delta^{13}C$) ranged between $-31.3\%e$ and $-26.7\%e$ for the grassland vegetation (Fig. 6), with significant differences among sites ($P < 0.001$).
Fig. 5. Effect of summer drought on annual community above- (dead and alive plant material) and below-ground biomass productivity at the three sites in 2007. Below-ground biomass was estimated using ingrowth cores. Means and standard errors are given (n=3–6). *( ) 0.1 ≥ P > 0.05, ** 0.01 ≥ P > 0.001. No significant treatment effects on the below-ground biomass productivity were found.

δ¹³C values were highest for Alp Weissenstein and varied over the course of the growing season (P < 0.001). In addition, we found significant effects of the drought treatment on δ¹³C at all sites (P < 0.001). While drought increased δ¹³C in 2005 and 2007, there was only a delayed response in 2006 at Chamau (date × treatment interaction: P < 0.001; Fig. 6a–c). This was controlled by the drought response of grasses that contributed most to community biomass. Legumes on the other hand did not react to drought in 2005 and 2007 but instead significantly decreased their δ¹³C values in 2006 (data not shown). At Fruëbuel, the drought response of vegetation δ¹³C varied throughout the experiment but a nonsignificant harvest date × treatment interaction indicated no change in the direction of the drought effect (Fig. 6d–e). Grasses were the only functional group displaying an overall positive drought response but forbs also showed slightly increasing δ¹³C values in response to drought in 2007. The drought response of vegetation δ¹³C at Alp Weissenstein was mainly driven by a very strong positive drought effect in the first year (significant date and date × treatment interaction effects; Fig. 6g–h), dominated by grasses that showed highest increases in δ¹³C in response to drought as well as by the positive drought effect on δ¹³C of forbs in 2006. Interestingly, δ¹³C values of legumes were not significantly affected by drought, although legume biomass had been reduced significantly.

On the single species level, no clear patterns could be detected. Grass species showed clear drought responses but no species differed strikingly from the others. The only outstanding drought response was observed in R. obtusifolius at Chamau (excluded from all analyses, for detailed results see Gilgen et al., 2010). Community nitrogen concentrations varied between 1.1% and 4.9%, with highly significant differences among sites and harvest dates (P < 0.001 each). The drought treatment increased overall N concentrations in above-ground biomass significantly by around 0.15% (P = 0.003). However, analysing the three years separately revealed that the drought effect was only significant in 2006. At Chamau, drought significantly increased N concentrations in above-ground biomass (P < 0.001). Independent of the treatment, N concentrations increased from around 2.5% in 2005 to more than 3.5% in autumn 2007. At Fruëbuel, no effect of drought on N concentrations was detected. At Alp Weissenstein, drought decreased average N concentrations at Alp Weissenstein, but this effect was only significant in the first year (P = 0.001), not in the second year (P = 0.3) of the drought treatment.

4 Discussion

4.1 Above-ground productivity and N supply

Drought typically reduces above-ground biomass productivity in grasslands (Hopkins, 1978; Bollinger et al., 1991; Kahmen et al., 2005). In contrast to these findings and rather unexpected, no consistent decrease in community above-ground biomass under drought was observed in our experiment for all sites in all years. One might suspect that especially at Fruëbuel the plants were not really water stressed. However, measurements of pre-dawn leaf water potential, assimilation rate and stomatal conductance show that plants of all three plant functional types (grasses, herbs, legumes) were negatively affected by the reduced precipitation input (Signarbieux and Feller, 2008; Signarbieux, 2009). As environmental and management characteristics differed considerably among the three sites (i.e. site explained most of the differences in community above-ground biomass productivity) and with time (i.e. significant effect of the harvest date), the sites seemed to react rather site- and year-specific. However, there was a strong relationship of average annual community above-ground biomass response with annual precipitation. Sites with lower annual precipitation thus seem to be more vulnerable to summer drought than sites with higher annual precipitation. Moreover, if annual precipitation is very high (above 1500–2000 mm), decreased precipitation input can even generate a beneficial effect on annual above-ground biomass productivity. The partial pressure of oxygen in wet soils is reduced due to decreased oxygen diffusion from the atmosphere and thus less oxygen is available for
Fig. 6. Effect of summer drought on $\delta^{13}$C of vegetation (alive plant parts only) at Chamau (excluding *Rumex obtusifolius*; a–c), Früebüel (d–f) and Alp Weissenstein (g–h) during the experiment. Means and standard errors are given (n=5–7). Periods of drought treatment are shaded in grey. * $0.05 \geq P > 0.01$, ** $0.01 \geq P > 0.001$, *** $P \leq 0.001$.

root respiration. Also the availability of nutrients can be limited in wet conditions due to changes in redox potentials and microbial activities which will also negatively affect plant performance (Nilsen and Orcutt, 1996). The lacking relationship of average annual community above-ground biomass response with the fraction of annual precipitation excluded by the shelters or with growing degree days clearly indicates that long-term annual precipitation rather than temperature controlled the drought response of these grasslands. A dependence of biomass production on soil moisture at drier sites but not at wetter sites was found for shrublands (Penuelas et al., 2007).

Grasses contributed most to community biomass (generally more than 50%, except for Chamau in 2006), thus reflecting the response of the community well or rather shaping the response of the community. Also Grime et al. (2000) had found that perennial grasses were particularly vulnerable to climate change in a limestone grassland. This is supported by findings from a companion study showing that drought had a strong negative impact on the physiological performance of grasses (especially at Chamau, Signarbieux and Feller, 2008). On the other hand, we found only slight effects of drought on forbs or legumes (except for *R. obtusifolius* at Chamau in 2006, Gilgen et al., 2010). Although *Trifolium repens*, the most abundant legume species at our sites, has been shown to be drought sensitive (Foulds, 1978; Stevenson and Laidlaw, 1985), in our study, the fraction of *T. repens* of community biomass was not significantly affected by drought (except at Alp Weissenstein in 2006). Furthermore, the amount of dead biomass was slightly increased under drought, with the strongest effect again at the driest site, Alp Weissenstein, in 2006. This increase in
dead biomass in drought plots was probably caused by earlier senescence as has been reported in other studies as well (Volaire, 2002; Monti et al., 2007).

Although other factors than water can also limit plant productivity, such as N (Harpole et al., 2007), we can fairly assume that the three grasslands studied here were not N limited throughout the experiment. This was shown in a long-term biodiversity experiment in Germany where biomass productivity was not affected by the lack of fertilisation after six years (Marquard et al., 2009). Prior to our experiment, the plots at the two lower sites were regularly fertilised with manure, according to Swiss regulations. At Chamau, this high fertilisation was even shown to control the C cycle more than climate (Zeeman et al., 2009). Increasing N concentrations in above-ground biomass in drought plots at Chamau and no change in N concentrations at Früebüel indicate an appropriate N supply. Even at the alpine site where drought decreased N concentrations in 2006, we can still assume there was no N limitation since biomass productivity was higher in the second year than in the first year of the experiment and N concentrations in above-ground biomass were unaffected by the treatment in the second year.

4.2 Below-ground productivity

Below-ground productivity can also react to drought: plants are able to sense the water available in soils and when the soil dries out, they produce abscisic acid (ABA) which was shown to maintain or increase root growth while shoot growth is inhibited (Saab et al., 1990; Davies and Zhang, 1991; Tardieu et al., 1992). Indeed, increased root growth under drought was often found in other studies (Jupp and Newman, 1987; Field et al., 1992; Kalapos et al., 1996; van den Boogaard et al., 1996; Kahmen et al., 2005). However, in our study, no change in below-ground biomass productivity was observed at any of the three sites. One reason could be that roots of different grass species reacted differently to drought (Molyneux and Davies, 1983; Bessler et al., 2009) and could thus compensate each other to hold community root biomass constant. Another explanation could be timing. Since our ingrowth cores remained in the field for a whole season or longer and not only during the drought treatment, the untreated conditions during most of the season might have dampened any drought effect. Still, on an annual basis, constant below-ground compared to decreasing above-ground productivity resulted in a higher root/shoot ratio of total plant biomass, thus indicating an increased allocation of resources to root growth.

4.3 Carbon isotope measurements

According to theory, $\delta^{13}C$ of plants can be used as an estimate for water use efficiency (integrated WUE) because the $^{13}C$ signature depends on the ratio of intercellular to ambient CO$_2$ concentrations ($c_i/c_a$, Farquhar et al., 1982) and an increase in $\delta^{13}C$ is thus coupled to a decrease in $c_i/c_a$, which can either be caused by higher photosynthetic fixation or decreasing stomatal conductance (Farquhar et al., 1989). Although this simplified view has recently been discussed rather critically (Seibt et al., 2008), our data still met the predictions made based on those earlier assumptions, i.e. an increase in $\delta^{13}C$ under drier condition due to decreased stomatal conductance (Signarbieux and Feller, 2009). Vegetation $\delta^{13}C$ was more positive in drought plots than in control plots at all sites in all years, although other physiological processes than just photosynthesis, e.g. cell elongation and protein synthesis, might affect growth even more strongly than photosynthesis (Lambers et al., 1998). Nevertheless, vegetation $\delta^{13}C$ was mainly controlled by grass $\delta^{13}C$ values that were also higher in drought plots than in control plots while forb and legume $\delta^{13}C$ values were unaffected by drought indicating no change in WUE efficiency in these two plant functional types.

4.4 Perspectives

Extrapolating results of the present study to the future might be difficult, since not only the amount of summer precipitation is changing under climate change but also of winter precipitation as well as timing and variability of rainfall, both important for grassland productivity (Nippert et al., 2006; Heisler-White et al., 2008). Future winters are predicted to be wetter than today in Central Europe (Frei et al., 2006). Higher winter precipitation will therefore probably recharge water reservoirs regularly, thereby buffering any summer drought effects. In addition, it has been shown that winter precipitation had no effect on biomass productivity, which was rather controlled by summer precipitation of the previous and the current year (Morecroft et al., 2004). This indicates that even if community above-ground biomass productivity recovered quickly after the removal of the rain shelters in our experiment, a long-term effect could still have been preserved.

Under future climate change drier summers will also be accompanied by higher atmospheric CO$_2$ concentrations. These have been shown to lead to water savings due to a reduced stomatal opening (Campbell et al., 1997; Volk et al., 2000). However, the relatively strong reductions of biomass productivity at Alp Weissenstein and at Chamau in 2007 will probably not be offset by the typically small water saving effects under increasing CO$_2$ as observed in Swiss forest (Leuzinger et al., 2005) and grassland sites (Volk et al., 2000).

In conclusion, results from this study indicate that there is no uniform grassland response to drought in Switzerland. Different grassland types as well as different plant functional types differed in their response to drought. However, the drought response scaled with total annual precipitation, thus sites with high annual precipitation seem to be better buffered against disturbance by summer drought than sites with low
annual precipitation. If this is also true in other parts of Europe remains to be tested, but it already now emphasises the strong needs for adapted management strategies in the drier parts of Switzerland.

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