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Winter climate change in alpine tundra: plant responses to changes in snow depth and snowmelt timing

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Abstract Snow is an important environmental factor in alpine ecosystems, which influences plant phenology, growth and species composition in various ways. With current climate warming, the snow-to-rain ratio is decreasing, and the timing of snowmelt advancing. In a 2-year field experiment above treeline in the Swiss Alps, we investigated how a substantial decrease in snow depth and an earlier snowmelt affect plant phenology, growth, and reproduction of the four most abundant dwarf-shrub species in an alpine tundra community. By advancing the timing when plants started their growing season and thus lost their winter frost hardiness, earlier snowmelt also changed the number of low-temperature events they experienced while frost sensitive. This seemed to outweigh the positive effects of a longer growing season and hence, aboveground growth was reduced after advanced snowmelt in three of the four species studied. Only *Loiseleuria procumbens*, a specialist of wind exposed sites with little snow, benefited from an advanced snowmelt. We conclude that changes in the snow cover can have a wide range of species-specific effects on alpine tundra plants. Thus, changes in winter climate and snow cover characteristics should be taken into account when predicting climate change effects on alpine ecosystems.

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1 Introduction

The winter snow cover is one of the fastest changing climate features under current climate change (IPCC 2007). In mountains this is especially evident as an upward shift of the snow line, and hence, a thinner snow cover of shorter duration at low and medium elevations (Latenser and Schneebeil 2003; Lopez-Moreno 2005; Mote et al. 2005). The timing, depth and duration of the snow cover are of special importance to upland ecosystems, as they define the start and duration of the potential growing season (Inouye and Wielgolaski 2003), as well as the temperatures under the snow (Pomeroy and Brun 2001). Although winter precipitation may increase in the future, the proportion falling as rain instead of snow will rise (IPCC 2007), leading to a thinner and less insulating snow cover, which will develop later and melt earlier. Thus, mountain ecosystems may face colder winters, but longer and warmer growing seasons.

In alpine and arctic ecosystems there is a close association between the spatial patterns of snow distribution and the occurrence of vegetation types or plant species (e.g. Evans et al. 1989; Walker et al. 1993; Odland and Munkejord 2008), illustrating the importance of snow depth, snowmelt and season length in structuring communities. Changes in snowmelt timing in alpine ecosystems can immediately affect plant growth (Walker et al. 1993; Galen and Stanton 1995; Wipf et al. 2006; Siffi 2007) and, in the longer term, vegetation composition (Galen and Stanton 1995; Seastedt and Vaccaro 2001; Welch et al. 2005; Wipf et al. 2005). Due to its effects on plant and soil temperatures and soil freezing processes, the snow depth controls biochemical, microbial and plant processes in winter (e.g. Campbell et al. 2005; Sturm et al. 2005), the effects of which can persist well into the growing season (Jones et al. 1998; Weih and Karlsson 2002). Thus, changes in the winter climate are likely to affect ecosystems in seasonally snow covered regions in various ways.

In a field experiment in alpine tundra just above timberline, we factorially reduced the snow depth in winter and advanced snowmelt in spring over 2 years to explore their individual and combined effects on ecosystem processes. The thinner snow cover decreased soil temperatures, and the advanced snowmelt increased the duration of the vegetation period and the temperature sum over summer, but induced lower temperatures just after snowmelt and a higher number of frost days. During the summers following these two winter manipulations, we measured the response in phenology, growth and reproduction of the four most abundant and dominant tundra plant species. We used correlations to explore and interpret the mechanisms leading to changes in plant growth, especially the relationships between growth, phenology, frost occurrence, and temperature.

We hypothesised that changes in snowmelt would have stronger short-term effects on plant performance than changes in snow depth. In particular, we expected an earlier snowmelt to advance the timing of phenological development and thus, to increase plant fitness due to higher temperature sums and a longer snow-free period available for growth and resource allocation. However, we expected the responses to differ between species, which could help to identify the potential winners and losers under long-term climate change.

2 Methods

2.1 Study site, experimental design and treatments

The study was conducted at the treeline research site of Stillberg, Davos (Central Alps, Switzerland, 47°28' N 7°30' E at 2,200 masl). The local climate (means 1975 to 2004 at 2,090 masl) is characterised by a mean annual precipitation sum of 1,150 mm, 48% of which falls between June and September. Temperatures average 2.0°C over the year, with February as the coldest, and July and August as the warmest months (means = -4.9°C and 9.9°C, respectively). Snow can fall at any time, but lays continuously from 18 October to 26 May on average, reaching a mean maximum depth of 143 cm. Over these 30 years, the fraction of precipitation falling as snow (i.e. at $\leq 0^\circ\text{C}$) decreased by 3% per decade, and the first snow-free day became earlier by 4 days per decade (linear regressions, $F_{1,28} = 4.8$ and 5.2 , $P = 0.04$ and 0.03).

The vegetation, growing on a 5–20 cm thick organic layer on Rankers over siliceous gneiss, consists of alpine tundra dominated by deciduous and evergreen dwarf shrubs, with few scattered herbaceous and graminoid species. At sites with visually homogeneous vegetation we established eight blocks, each consisting of four plots of 1 m² at approx. 4 m distance. In the winters 2003/2004 and 2004/2005, one combination of the following two treatments was assigned to each plot of a block, resulting in a fully factorial design: (1) reduced snow depth over winter and (2) advanced snowmelt in spring. To reduce the snow cover, and thus, its temperature insulation capacity over winter, we kept its depth at approx. 40 cm by manually removing snow after each snowfall period. This treatment was stopped towards spring, and snow depths levelled out during the next snowfalls. In spring we advanced snowmelt on half of the plots by carefully reducing the remaining snow from approx. 1 m to 30 cm, and letting those melt naturally. Due to an unusual warm spell in late spring 2005, the snow cover melted very rapidly, and some plots were snow-free before the snowmelt manipulations even started (they were excluded from analysis). When we reduced the snow cover on the still snow covered early-melting plots on 25 May 2005, we also added 30 to 50 cm of snow to the remaining control plots to generate significant difference in snowmelt dates between treatments. For simplicity we still call them controls throughout the manuscript.

Over winter, we measured temperatures in 3 h intervals at the snow/soil interface with temperature loggers (Thermochron iButtons, Maxim Integrated Products, Inc., Sunnyvale, CA, USA) that we installed at soil surface in each plot before snowfall. Thus we were also able to record snow-free periods as days when temperature fluctuations by $>5^\circ\text{C}$ or night temperatures of $>0^\circ\text{C}$ were measured. Soil temperatures were reduced by the snow removal, especially in the first year (minimum temperatures under the snow: -2.9°C vs. -4.8°C in mid-December 2004, -4.5°C vs. -5.1°C in mid-December 2005; $F_{1,2} = 67.9$, $P = 0.01$; unmanipulated snow depth at these times was approx. 40 and 15 cm). The snowmelt manipulations advanced snowmelt by 11.3 days in 2004 and by 6.2 days in 2005 ($F_{1,7} = 52.6$, $P < 0.001$; Table 1), corresponding to a scenario for 2020 to 2040 based on the rate snowmelt was advancing at the site in previous decades (see above). The snowmelt treatment

prolonged the potential growing season (between melt-out and first heavy fall frosts with $> -5^{\circ}\text{C}$ temperatures) by 11% in 2004 and by 6% in 2005, and increased the number of frost days (reaching $<0^{\circ}\text{C}$) during that period by 45% in 2004 and 16% in 2005 (Table 1). Soil moisture during the snow free season (measured at five dates in 2005) was not affected by the snow manipulation treatments, probably due to lateral water flow during snowmelt (data not shown).

2.2 Plant responses

We marked four shoots per plot of each of the four most abundant and dominant species, namely the evergreen dwarf shrubs *Empetrum nigrum* L. ssp. *hermaphroditum* (Lange ex Hagerup) Böcher and *Loiseleuria procumbens* (L.) Desv. and the deciduous dwarf shrubs *Vaccinium myrtillus* L. and *V. uliginosum* L. ssp. *gaultierioides* Bigelow. Between snowmelt and leaf senescence, we visited the plots two to three times weekly to record the number of living shoots, their phenological state, and the number of live flower buds, flowers, and fruits of each marked shoot. We recorded the phenological development as the dates when a shoot entered the following vegetative and reproductive phenophases (=stage of development): (1) shoot snow free, (2) greening up, i.e. reduction of anthocyanins in evergreens, or first leaf parts visible in deciduous species, (3) start of shoot growth, (4) onset of fall coloration, (5) burst of first flower buds, (6) first flowers open, and (7) first flowers senescent. Fruit ripening and *L. procumbens* reproduction did not occur in sufficient numbers to be analysed. From the daily mean temperatures of the nearby weather station we calculated the growing degree days accumulated after greening-up (GDD, i.e. sum of daily mean temperatures above 0°C between greening-up and first heavy fall frosts). In 2005, we recorded the shoots with visible frost damages (aborted leaf and flower buds and blackened leaf tissue) and calculated the abortion rate of flower buds. Each fall we harvested, dried and weighed the ripe berries of the

Table 1 Effects of the snow depth and snowmelt manipulation treatments in 2004 and 2005 on the environmental variables snowmelt date, number (n) of frost days (i.e. minimum temperatures below freezing), and growing degree days (GDD) above 0°C between snowmelt date and the end of the growing season

Treatments	N	Snowmelt date	n frost days	GDD $> 0^{\circ}\text{C}$
		Mean (SE)	Mean (SE)	Mean (SE)
2004				
Overall	32	156.7 (1.4)	11.5 (0.7)	941 (7.3)
Deep snow	16	157.6 (2.0)	11.3 (0.9)	933 (11.2)
Shallow snow	16	155.8 (2.1)	11.7 (1.2)	949 (9.1)
Early melt	16	151.1 (1.8)	13.6 (1.1)	970 (6.3)
Control melt	16	162.4 (1.0)	9.4 (0.6)	913 (8.3)
2005				
Overall	26	149.3 (0.6)	8.3 (0.2)	931 (5.3)
Deep snow	14	149.6 (0.9)	8.3 (0.4)	929 (7.6)
Shallow snow	12	148.9 (0.9)	8.3 (0.3)	933 (7.6)
Early melt	14	146.4 (0.2)	8.9 (0.3)	949 (5.1)
Control melt	12	152.6 (0.4)	7.7 (0.3)	910 (5.4)

Snowmelt date given as day of year

N number of plots per treatment

marked shoots and measured the leaf lengths of approx. 20 leaves per species and plot. As a measure of belowground productivity, we quantified root production in 4 blocks (=16 plots) by burying root ingrowth cores, i.e. root-free organic soil cores (depth 10 cm, diameter 3 cm) in a nylon mesh (mesh size 1 mm) in October 2003, and quantifying root biomass grown into the cores by 30 August 2004. In October 2005 we harvested all marked shoots and measured their above ground growth as the annual growth increments of the last 2 years.

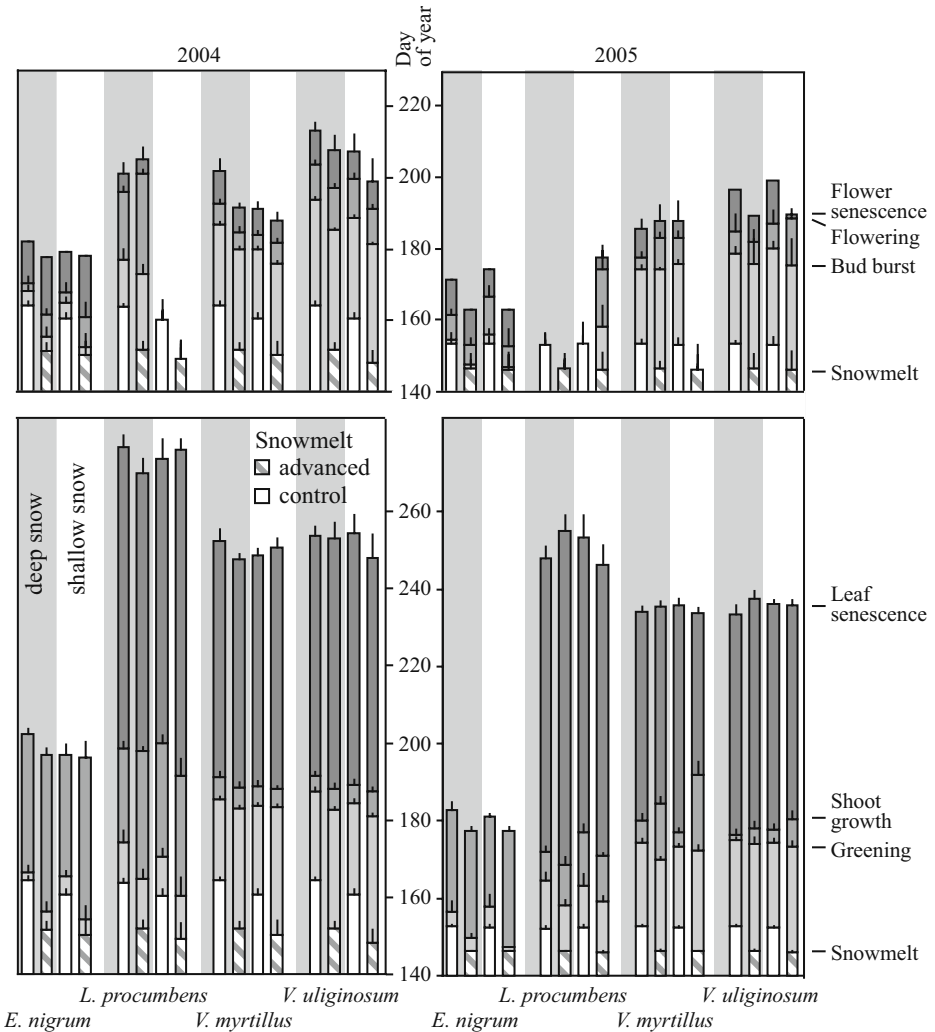


Fig. 1 Phenology of the vegetative (*above*) and reproductive (*below*) development of four alpine dwarf shrub species subject to manipulations in snow depth and snowmelt timing in 2004 and 2005. The phenological stages recorded after snowmelt were: leaf greening, start of shoot growth, start of leaf senescence/fall coloration, bud break of flower buds, start of flowering, and start of flower senescence. Error bars = ± 1 SE. For full species names see text

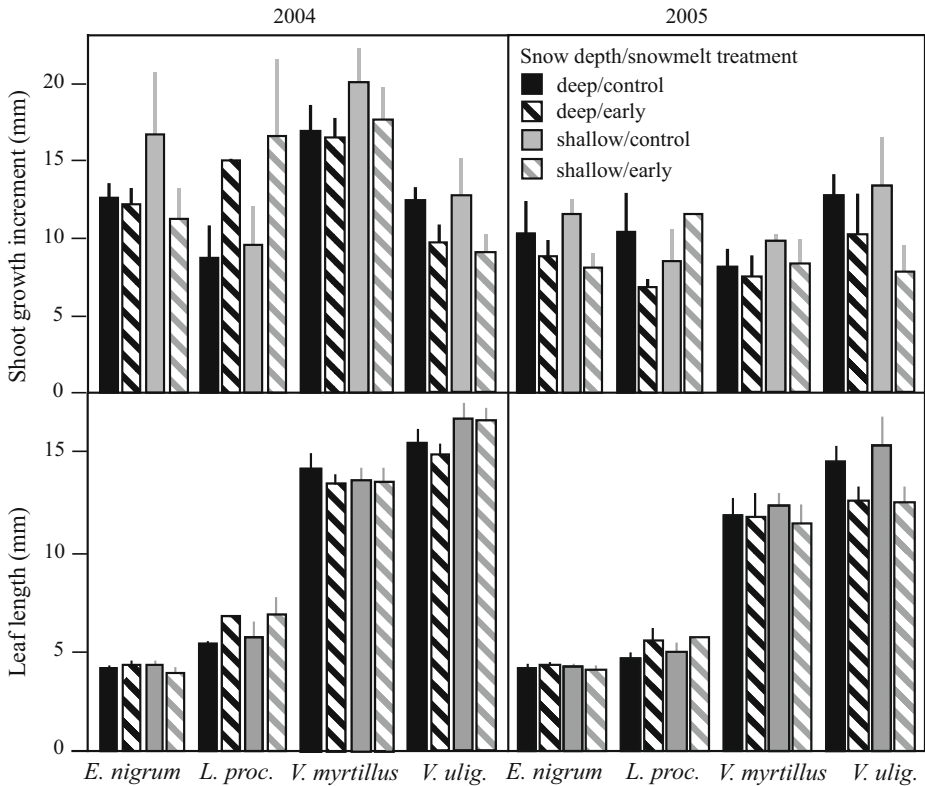
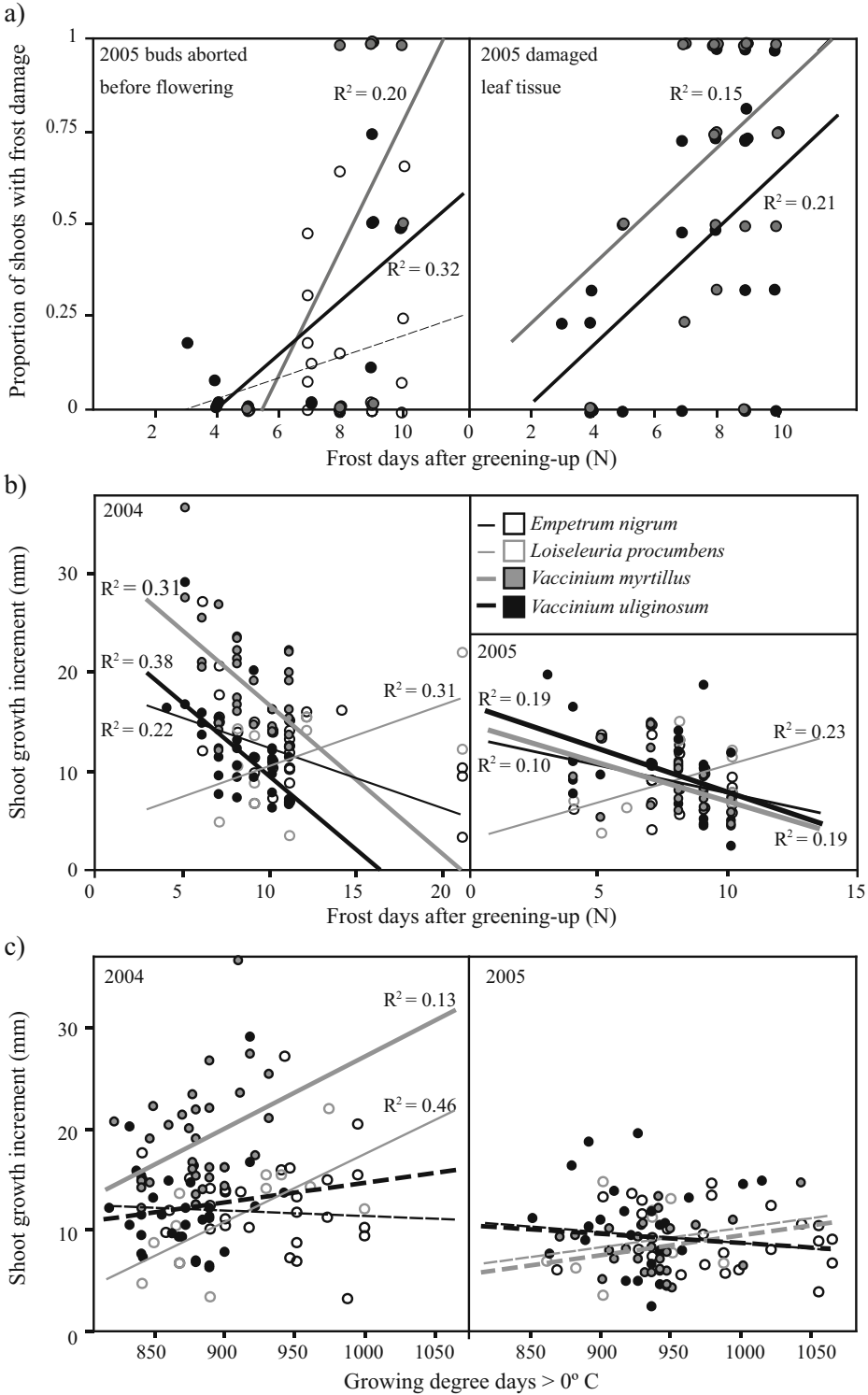


Fig. 2 The growth response of shoots (*above*) and leaves (*below*) of four alpine dwarf shrubs species to manipulations in snow depth and snowmelt timing in 2004 and 2005. Error bars = ±1 SE

2.3 Statistical analyses

All data per species and plot were pooled and analysed for the effects of the snow manipulation treatments with ANOVAs with Type I Sums of Squares. We analysed whether the response variables differed between years, snowmelt and snow depth treatments, and tested for interactions between these factors. Due to the blockwise structure of the experiment, we defined year, snowmelt and snow depth treatments as fixed and block as random factors. Residuals were visually checked and variables transformed where necessary to meet the assumptions of ANOVA (Appendix). Analyses were conducted in SPSS 11.0 (SPSS Inc 2001).

Fig. 3 The relationship between the number of frost days experienced by four dwarf shrub species after starting their life cycle in spring, **a** the occurrence of frost damaged flower buds and leaves and **b** shoot growth, and **c** the relationship between growing degree days and shoot growth. Significant correlations are indicated in *solid lines* and with R^2 values, non-significant ones with *dashed lines*



3 Results

The phenological development of all species was clearly influenced by changes in the snow cover (Fig. 1; Table 2). The greening-up, which is the first sign of seasonal plant activity, was advanced by earlier snowmelt in all four species, and also advanced by reduced snow in both *Vaccinium* species. An earlier snowmelt also advanced the start of shoot growth of *E. nigrum* (marginally significant), *L. procumbens*, and *V. uliginosum* (in 2004 only), but delayed that of *V. myrtillus*, especially in 2005 and in plots with reduced snow cover. In 2004, leaf senescence of both *Vaccinium* species tended to start earlier after advanced snowmelt. Early snowmelt advanced all reproductive phenophases of all three species it was analysed for, but not all effects were statistically significant (Fig. 1; Table 2). A reduced winter snow cover further advanced flowering and flower senescence, but only significantly so in *V. uliginosum*.

Although an earlier snowmelt advanced the start of all species' life cycle, and thus, increased the growing degree days they accumulated, the annual growth increments and leaf lengths of *E. nigrum*, *V. myrtillus* and *V. uliginosum*, were generally decreased after early snowmelt (but not in all cases significantly; Fig. 2; Appendix). *L. procumbens* differed from this prevalent pattern: its shoots and leaves grew larger after earlier snowmelt, especially in 2004 (Fig. 2). The reduced winter snow depth did not significantly alter aboveground growth, but decreased the total root mass in root ingrowth cores by 55% (from 34.7 to 15.7 mg, $F_{1,9} = 5.4$, $P = 0.05$). A positive effect of temperature (i.e. increased growing degree days) on shoot growth was only found in *L. procumbens* and *V. myrtillus* in 2004, but not in the other species and years (Fig. 3). The proportion of *Vaccinium* sp. shoots with visible frost damage in 2005 was more than doubled by advanced snowmelt (from 46.5% to 93.3% in *V. myrtillus*, $F_{1,6} = 19.3$, $P = 0.005$, from 23.6% to 64.3% in *V. uliginosum*, $F_{1,6} = 6.0$, $P = 0.05$) and was correlated with the number of frosts that shoots experienced after greening-up (Fig. 3). The more frost days *E. nigrum* and *Vaccinium* shoots experienced after greening-up, the less they grew in that year, however, the correlation between frost occurrence and *L. procumbens* growth was a positive one (Fig. 3).

An advanced snowmelt and a higher number of frost days after greening-up resulted in a higher proportion of buds of both *Vaccinium* species killed before flowering in 2005 (Fig. 3). The fecundity (=proportion of shoots with ripe fruits) in these two species did not differ between treatments. The fecundity of *L. procumbens* was decreased with reduced snow cover over winter (from 2.1% to 0%; $F_{1,6} = 5.7$, $P = 0.05$). Shallow snow cover reduced fecundity in *E. nigrum*, but only in 2005 (from 23.1% to 6.9%; snow depth \times year: $F_{1,6} = 13.3$, $P = 0.01$). *E. nigrum* berries, however, were heavier in plots with earlier snowmelt, especially in 2005 (24.8 vs. 20.6 mg, snowmelt treatment: $F_{1,7} = 4.3$, $P = 0.08$; snowmelt treatment \times year: $F_{1,5} = 9.0$, $P = 0.03$).

4 Discussion

Our field experiment is to our knowledge the first to manipulate factorially the presence and depth of the alpine snow cover and to study simultaneously the role of two of the key variables that control winter and spring conditions experienced by alpine plants. By disentangling their effects on the phenology and fitness of four

Table 2 Response in phenology of four alpine dwarf shrub species to two winters of manipulations of snow depth and snowmelt timing

Year	Treatments	N	Leaves green		Vegetative growth		Leaf senescence		N	Flowering		N	Flower senescence	
			Mean (SE)	Mean (SE)	Mean (SE)	Mean (SE)	Mean (SE)	Mean (SE)		Mean (SE)	Mean (SE)			
<i>Empetrum nigrum</i>														
2004	Overall	30	161 (1.5)	198 (1.2)		26	160 (1.8)	29	165 (1.1)	29	179 (0.7)			
	Deep snow	14	161 (2.2)	199 (1.1)		12	161 (2.6)	13	166 (1.7)	13	180 (1.0)			
	Shallow snow	16	160 (2.1)	197 (2.1)		14	159 (2.6)	16	164 (1.5)	16	179 (1.0)			
	Early melt	15	155 (2.0)	197 (2.0)		13	154 (2.4)	15	161 (0.9)	15	178 (0.9)			
2005	Control melt	15	166 (1.1)	199 (1.4)		13	166 (1.5)	14	169 (1.4)	14	180 (1.0)			
	Overall	24	152 (1.2)	180 (0.8)		21	151 (0.9)	21	157 (1.3)	19	165 (1.2)			
	Deep snow	13	153 (1.7)	180 (1.3)		11	151 (1.3)	11	156 (1.4)	11	165 (1.5)			
	Shallow snow	11	151 (1.9)	179 (0.9)		10	150 (1.5)	10	157 (2.3)	8	165 (2.0)			
2005	Early melt	13	148 (0.5)	178 (0.7)		11	147 (0.2)	11	152 (0.4)	10	161 (0.8)			
	Control melt	11	157 (1.8)	182 (1.2)	Y***; M**	10	154 (1.2)	10	162 (1.5)	9	169 (1.2)			
					Y***; M(*)		Y***; M**		Y***; M***; Y × D(*)		Y***; M***; Y × M*			
<i>Loiseleuria procumbens</i>														
2004	Overall	27	167 (1.8)	197 (1.3)										
	Deep snow	13	170 (2.2)	198 (0.9)										
	Shallow snow	14	165 (2.7)	196 (2.4)										
	Early melt	14	163 (2.5)	195 (2.1)										
2005	Control melt	13	172 (1.9)	199 (1.4)										
	Overall	20	161 (1.1)	172 (1.0)										
	Deep snow	11	161 (1.5)	170 (1.4)										
	Shallow snow	9	161 (1.7)	174 (1.3)										
2005	Early melt	11	159 (1.1)	170 (1.3)										
	Control melt	9	164 (1.6)	174 (1.3)	Y***; M**									
					Y***; M**									

Table 2 (continued)

Year	Treatments	N	Leaves green		Vegetative growth		Leaf senescence		Flower buds burst		Flowering		Flower senescence	
			Mean (SE)	N	Mean (SE)	N	Mean (SE)	N	Mean (SE)	N	Mean (SE)	N	Mean (SE)	N
<i>Vaccinium myrtillus</i>														
2004	Overall	32	184 (0.3)	189 (0.7)	250 (1.2)	16	180 (0.9)	16	184 (1.0)	16	191 (1.5)			
	Deep snow	16	184 (0.6)	190 (1.1)	250 (1.9)	7	181 (1.4)	7	186 (1.8)	7	193 (2.3)			
	Shallow snow	16	184 (0.4)	188 (0.7)	249 (1.4)	9	178 (1.2)	9	183 (1.0)	9	190 (2.1)			
	Early melt	16	183 (0.4)	188 (0.8)	249 (1.3)	11	179 (1.0)	11	183 (0.9)	11	190 (1.5)			
2005	Control melt	16	184 (0.5)	190 (1.0)	251 (2)	5	182 (2.0)	5	186 (2.3)	5	194 (3.6)			
	Overall	26	173 (0.6)	184 (1.6)	235 (0.7)									
	Deep snow	14	172 (0.8)	182 (1.3)	235 (1.0)									
	Shallow snow	12	173 (0.9)	187 (3.2)	235 (0.9)									
Control melt	Early melt	14	171 (0.8)	189 (2.2)	235 (0.9)									
	Control melt	12	174 (0.7)	179 (1.2)	235 (1.0)									
			Y***; M**;	Y**; M**;	Y***; M**;									
		M**;	Y × M**;	Y × M(*)										
		M × D*	Y × D(*)											
			M × D*											

widespread tundra species, our experiment contributes to the understanding of these species' sensitivities to different abiotic factors and of the consequences of winter climate change on alpine plant communities.

Our results confirm that both the depth and the timing of disappearance of the snow cover have important implications for plant phenology, growth and reproduction, but that snowmelt timing has stronger effects than snow depth. The phenological development was generally advanced after earlier snowmelt, which could potentially foster plant fitness by extending the period available for growth and resource allocation (Galen and Stanton 1993; Stinson 2004). Three out of four plant species, however, were not able to turn the longer growing season into increased growth and reproduction, which is in line with other snow removal studies in upland ecosystems (Saavedra 2002; Wipf et al. 2006; Siffi 2007). The one species that responded positively to advanced snowmelt was *Loiseleuria procumbens*, which tended to produce longer shoots and leaves. *L. procumbens* is a specialist of windswept ridges with little or lacking winter snow cover (Grabherr 1980) and is noted for its extreme winter frost hardiness (Koerner 1999). Although it is much more frost sensitive in summer, at ca. -7°C it is still more frost hardy than the other species studied here (Ulmer 1937; Taschler and Neuner 2004), and thanks to its prostrate life form, it may profit from some protection by other species (Jordan and Smith 1995). In the following year 2006 without major frost events, however, *V. myrtillus* growth increased after early snowmelt (subsample of this experiment, C. Schwoerer, unpublished data). These findings, together with the negative relationship between frost occurrence and growth, suggest that frosts might play an important role for the survival, growth and reproduction of alpine plant species in a changing climate (see Inouye 2000 for a review on this topic). The positive correlation of frost occurrence and *L. procumbens* shoot growth even suggests that this prostrate plant might benefit from the frost damage to the other, larger growing species.

The interaction between snowmelt timing and phenology can control the amount of frost damage in two ways. First, the start of the life cycle and loss of frost hardiness occurs earlier after advanced snowmelt and thus, plants experience a higher number of frost events while in a relatively frost sensitive stage. Second, by starting the formation of apical meristems and buds earlier, plants also have “more to lose”, should low temperature extremes occur. Although alpine and arctic plants are generally able to replace frost aborted shoots and leaves within the same growing season (e.g. *V. myrtillus* in our experiment), frost damage severely reduces plant growth and fecundity in the short term (Molgaard and Christensen 1997; Price and Waser 1998; Inouye et al. 2002), and may even become lethal if occurring several years in a row, as plant resources may be depleted (Molau 1997). A strategy to reduce frost damages in a fluctuating or changing climate is photoperiodism, i.e. a plant's ability to time its development according to a certain day length. This allows many alpine species to remain frost hardy until later in summer, when growing conditions are more favourable (Koerner 1999; Keller and Korner 2003). In the absence of frosts, on the other hand, this mechanism inhibits plants from benefiting from warmer temperatures and longer growing periods.

By removing snow without increasing spring temperatures (unlike e.g. Harte and Shaw 1995 or Oberbauer et al. 1998, who applied additional warming treatments), our experiment caused the seemingly paradox climate scenario of a colder spring in a warmer world. In snow-rich regions, the amount of snow does not only control the

plant and soil temperatures in winter, but also the temperatures that the plants first experience in spring. Due to the interaction of a shallower snow cover and warmer spring temperatures, snowmelt can shift to an earlier, naturally colder period of the year. Accordingly, long-term data from the Colorado Rocky Mountains, a region with a snow and temperature regime comparable to ours, show that while the amount of snow decreased and snowmelt timing advanced, the amount of frost damage on alpine plants strongly increased, regardless of an overall warming trend (Inouye et al. 2002; Inouye 2008). Hence, whether frost events will increase or decrease depends on the balance between warmer spring temperatures (which generally decrease the occurrence of frost events) and earlier snowmelt (which increases the risk that plants experience frosts while not covered in snow).

The decreased snow depth, which induced colder soil temperatures throughout most of the winter, had little direct effect on plant phenology and aboveground growth. Our snow reduction treatment did not expose plants during winter, thus, rather than aboveground plant parts, we would expect snow depth manipulations to affect belowground processes. In fact the snow reduction treatment did decrease root production in the following summer in our study. This contrasts with another snow manipulation experiment, where decreased snow cover stimulated root turnover (Tierney et al. 2001), probably because soil freezing and the associated soil heaving damaged the fine roots and stimulated their replacement growth in the following spring. Whether the different results are caused by the different methods used in these studies or represent a biological pattern remains unclear. Our experiment (Bauer, unpublished data) as well as other studies (Fitzhugh et al. 2001; Groffman et al. 2001) found that reduced snow depth and winter temperatures increased mobile soil nitrogen, which can be used by microbes, but also by plants in spring (Bilbrough et al. 2000). Hence, if reduced snow depth and soil freezing can affect aboveground plant growth at all, the extent may depend on the balance between the negative (root damage) and positive effects (increased nutrient availability).

In conclusion, our study shows that plant responses to winter climate change and modifications of the snow cover strongly differ among species. The fact that plants respond differently to changes in the same abiotic variable suggests that the interaction between plant species (Wipf et al. 2006), but also between plants and their pollinators, pests or pathogens, may be disrupted (Roy et al. 2004). Such mechanisms have a great potential to mediate or amplify effects of climate change on community composition. The important role of frost events on the outcome of our experiment underlines the importance of incorporating extreme nonlinear climate factors into analyses and predictions of climate change effects. The ongoing snow manipulations of our experiment will show whether the initial changes in species fitness and reproduction found after the first 2 years will proceed over the longer term and will eventually result in significant changes in community composition and structure.

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Appendix

Table 3 F statistics of ANOVAs analysing the response of four alpine dwarf shrub species to two winters of manipulating snowmelt timing and snow depth

	N		Year		Snowmelt treatment		Snow depth treatment		Year × snowmelt		Year × snow depth		Snowmelt × snow depth					
	df	F	P	df	F	P	df	F	P	df	F	P	df	F	P			
Leaf greening-up																		
<i>E. nigrum</i>	54	1, 6	< 0.001*	1, 7	24.1	0.002*	1, 7	0.7	ns	1, 6	3.3	ns	1, 6	0.0	ns	1, 5	1.6	ns
<i>L. procumbens</i>	47	1, 5	0.02*	1, 6	13.2	0.01*	1, 6	1.0	ns	1, 5	2.9	ns	1, 5	0.8	ns	1, 5	0.3	ns
<i>V. myrtillus</i>	58	1, 6	< 0.001*	1, 7	25.8	0.001*	1, 7	0.0	ns	1, 6	1.5	ns	1, 6	1.1	ns	1, 7	10.7	0.01*
<i>V. uliginosum</i>	56	1, 6	< 0.001*	1, 7	12.1	0.01*	1, 7	9.4	0.02*	1, 6	1.6	ns	1, 6	0.7	ns	1, 6	0.3	ns
Start of vegetative growth																		
<i>E. nigrum</i>	54	1, 6	< 0.001*	1, 7	4.1	0.08(*)	1, 7	0.8	ns	1, 6	1.0	ns	1, 6	0.1	ns	1, 5	0.5	ns
<i>L. procumbens</i>	47	1, 5	< 0.001*	1, 6	23.7	0.003*	1, 6	0.0	ns	1, 5	0.0	ns	1, 5	4.2	0.10(*)	1, 5	1.3	ns
<i>V. myrtillus</i>	58	1, 6	0.01*	1, 7	4.8	0.07(*)	1, 7	2.1	ns	1, 6	3.0	0.001*	1, 6	4.1	0.09(*)	1, 7	8.1	0.02*
<i>V. uliginosum</i>	56	1, 6	< 0.001*	1, 7	0.6	ns	1, 7	0.0	ns	1, 6	5.2	0.06(*)	1, 6	2.8	ns	1, 6	0.1	ns
Start of leaf senescence																		
<i>L. procumbens</i>	42	1, 5	0.002*	1, 6	0.8	ns	1, 6	0.1	ns	1, 3	0.7	ns	1, 4	1.0	ns	1, 3	0.0	ns
<i>V. myrtillus</i>	58	1, 6	< 0.001*	1, 7	0.8	ns	1, 7	0.2	ns	1, 6	4.8	0.07(*)	1, 6	0.0	ns	1, 7	2.1	ns
<i>V. uliginosum</i>	56	1, 6	< 0.001*	1, 7	0.2	ns	1, 7	0.1	ns	1, 6	5.1	0.07(*)	1, 6	0.1	ns	1, 6	0.2	ns
Bud burst of flower buds																		
<i>E. nigrum</i>	47	1, 5	< 0.001*	1, 7	22.4	0.002*	1, 6	0.0	ns	1, 4	9.4	0.04*	1, 5	0.0	ns	1, 3	0.0	ns
<i>V. myrtillus</i>	21	1, 2	0.01*	1, 2	1.6	ns	1, 2	3.7	ns	not enough samples								
<i>V. uliginosum</i>	46	1, 6	< 0.001*	1, 7	8.3	0.02*	1, 7	1.3	ns	1, 4	2.7	ns	1, 3	2.6	ns	1, 3	0.1	ns

Start of flowering																			
<i>E. nigrum</i>	50	1, 5	123	<0.001*	1, 7	50	<0.001*	1, 7	0.0	ns	1, 5	0.4	ns	1, 5	5.2	0.07(*)	1, 4	0.1	ns
<i>V. myrtillos</i>	23	1, 2	112	0.01*	1, 3	1.3	ns	1, 3	5.3	ns	not enough samples								
<i>V. uliginosum</i>	45	1, 6	190	<0.001*	1, 7	2.6	ns	1, 7	2.8	ns	1, 4	6.6	0.05*	1, 3	11.3	0.04*	1, 2	0.2	ns
Start of flower senescence																			
<i>E. nigrum</i>	48	1, 5	454	<0.001*	1, 7	22.1	0.002*	1, 7	0.4	ns	1, 5	10.4	0.02*	1, 4	0.3	ns	1, 4	0.3	ns
<i>V. myrtillos</i>	22	1, 2	6.4	ns	1, 3	7.7	0.07(*)	1, 2	2.3	ns	not enough samples								
<i>V. uliginosum</i>	33	1, 5	457	<0.001*	1, 5	4.9	0.08(*)	1, 6	4.5	0.08(*)	1, 1	0.9	ns	1, 1	445	0.03*	1, 0		
Length of shoot increment (log-transformed)																			
<i>E. nigrum</i>	53	1, 6	22.0	0.003*	1, 7	4.3	0.08(*)	1, 7	0.0	ns	1, 6	0.0	ns	1, 6	1.0	ns	1, 7	7.4	0.05*
<i>L. procumbens</i>	23	1, 3	1.5	ns	1, 3	6.9	0.07(*)	1, 2	0.0	ns	1, 1	5.0	ns	1, 1	23.0	ns	1, 1	0.8	ns
<i>V. myrtillos</i>	58	1, 6	895	<0.001*	1, 7	1.2	ns	1, 7	1.5	ns	1, 6	0.6	ns	1, 6	0.2	ns	1, 7	1.4	ns
<i>V. uliginosum</i>	56	1, 6	4.2	0.09(*)	1, 7	6.1	0.04*	1, 7	1.6	ns	1, 6	0.7	ns	1, 6	1.3	ns	1, 6	0.0	ns
Length of leaves																			
<i>E. nigrum</i>	54	1, 6	0.7	ns	1, 7	1.3	ns	1, 7	1.0	ns	1, 6	1.5	ns	1, 6	1.3	ns	1, 5	2.9	ns
<i>L. procumbens</i>	20	1, 3	13.9	0.03*	1, 1	20.3	ns	1, 2	0.1	ns	1, 1	200	0.04*	1, 1	23.0	ns	1, 1	0.0	ns
<i>V. myrtillos</i>	58	1, 6	87	<0.001*	1, 7	0.5	ns	1, 7	0.2	ns	1, 6	0.1	ns	1, 6	0.5	ns	1, 7	0.2	ns
<i>V. uliginosum</i>	56	1, 6	28.5	0.002*	1, 7	12.8	0.009*	1, 7	1.4	ns	1, 6	6.6	0.04*	1, 6	1.9	ns	1, 7	0.0	ns

For details of the analysis see “Methods” section

df degrees of freedom (numerator, denominator), ns not significant

* $P < 0.05$, significant; (*) $P < 0.1$, marginally significant

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