

Onset of autumn senescence in High Arctic plants shows similar patterns in natural and experimental snow depth gradients

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Abstract

Predicted changes in snow cover and temperature raise uncertainties about how the beginning and the end of the growing season will shift for Arctic plants. Snowmelt timing and temperature are known to affect the timing of bud burst, but their effects on autumn senescence are less clear. To address this, researchers have examined senescence under natural and experimental environmental gradients. However, these approaches address different aspects of plant responses and the extent to which they can be compared is poorly understood. In this study, we show that the effect of snowmelt timing on the timing of autumn senescence in High Arctic plants is the same between a natural and an experimental gradient in three out of four studied species. While the two approaches mostly produce comparable results, they give in combination greater insight into the phenological responses to predicted climate changes. We also showed that a short warming treatment in autumn delayed senescence by 3.5 days in *D. octopetala*, which is a 10 % extension of the growing season end for this species. Warming treatments have commonly been applied to the whole growing season, but here we show that even isolated autumn warming can be sufficient to affect plant senescence.

Keywords: senescence, phenology, natural gradient, experimental gradient, warming

Introduction

Climate is changing most rapidly in the Arctic with mean annual temperatures rising twice as fast as the global average (IPCC 2013). However, the seasons are not affected equally and the autumn is warming disproportionately more than the summer (Chylek et al. 2009; Serreze et al. 2009). In addition, higher temperatures in late winter and spring may lead to earlier snowmelt, while the increase in snowfall predicted for some areas of the Arctic may delay snowmelt, with large regional variations (McBean et al. 2005; Räisänen 2008). This could affect the timing of bud burst and shift the end of the growing season for plants (Semenchuk et al. 2016). These changes in temperature and snow cover raise uncertainties about how the end of the growing season will shift for Arctic plants, with implications for multiple key ecosystem processes (Cooper 2014).

Due to the short growing season in the Arctic tundra, the timing of autumn senescence is important for maximising the period for plant growth and reproduction and allowing sufficient time for the development of frost hardiness and nutrient resorption. Early and sudden frost events can inhibit the development of frost hardiness, causing premature leaf fall, loss of nutrients due to incomplete resorption and damage to shoots, with possible consequences for plant fitness (May and Killingbeck 1992). Despite its importance, autumn senescence is still far less studied than spring phenology, especially in tundra ecosystems (Gallinat et al. 2015). It is currently unclear what drives the timing of autumn senescence in Arctic plants, but indirect effects of snowmelt timing (Keenan and Richardson 2015; Semenchuk et al. 2016) and direct effects of photoperiod and temperature have been suggested (Barnes et al. 1998; Marchand et al. 2004).

In order to study the effect of snowmelt timing on plant phenology, both natural (Kudo 1991; Kudo and Suzuki 1999; Sedlacek et al. 2015; Wheeler et al. 2015; Carbognani et al. 2016; Gehrman et al. 2017) and experimental gradients of snowmelt timing (Henry and Molau 1997; Walker et al. 1999; Wipf and Rixen 2010; Cooper et al. 2011) have been used. However, both of these methods record responses at different temporal scales and feature their own set of advantages and limitations,

which complicates inter-comparison among methods and ultimately impedes development of a robust consensus on likely phenological responses to climate change to inform environmental policy and species distribution models (Dunne et al. 2004).

Natural gradients in snowmelt timing give information about the range of conditions plant populations can cope with under the current climate and this may give insight into the resilience of the population in the future (Gehrmann et al. 2017, 2020). Observations reflect the equilibrated responses of biotic communities, which have adapted over a long period to the local temperature and moisture regime. However, this method assumes that the variation over space elicits the same responses in plants as change over time and that biotic interactions remain the same as the climate changes, which is unlikely to be the case. Further, it remains difficult to completely remove, and/or control for, the confounding effects of other environmental factors, which often co-vary with the factors of interest along the gradient (Dunne et al. 2004; Fukami and Wardle 2005).

Experimental manipulations of snowmelt timing and temperature, such as snow fences which passively accumulate snow during winter (Cooper et al. 2011), or open top chambers (OTCs) which passively warm the vegetation throughout the whole growing season, allow the establishment of possible causal relationships between the plant response and the treatment because specific controls are available. However, such broad manipulations alone cannot determine the role of specific primary drivers of autumn phenology, such as soil temperature under the snow or short-term, selective warming of the autumn season. Rather, they show the net effect of these climate conditions (Dunne et al. 2004). Furthermore, the initial response of plants to climatic changes may not correspond to the equilibrated response to long-term, sustained changes in climate. Therefore, short-term experiments may over- or under-predict the plant responses to climate change so that observations and experiments often give different results and lead to different conclusions (Leuzinger et al. 2011; Wolkovich et al. 2012; Blume-Werry et al. 2016a). Few studies have directly compared natural with experimental gradients, so it is difficult to predict how plant responses to gradients caused by natural environmental

variations or experimental manipulations differ. Which approach is more relevant to understanding the impacts of anthropogenic climate change on plants depends on the speed and magnitude with which the climate will change. Overall, the limitations of each of these study methods imply that a combination of approaches is necessary to better understand plant responses to a changing climate.

Here, we combine natural and experimental gradients of snowmelt timing and selective warming during autumn senescence in the same site to study the long- and short-term effects of changes in snowmelt timing and the short-term effects of elevated temperature on the timing of autumn senescence. The experimental snowmelt gradient has been in place since 2006 whereas the natural snowmelt gradient is based on the local microtopography, providing significantly contrasting timescales of exposure. Furthermore, our selective experimental passive warming in autumn only allows us to examine the effects of elevated temperature specifically in the autumn period as observed and predicted across the Arctic (Chylek et al. 2009; Serreze et al. 2009), instead of passive warming across the whole growing season, as has been most commonly studied (e.g. Henry and Molau 1997).

We hypothesise that

1. Experimental and natural gradients yield different conclusions on the effect of snowmelt timing on autumn senescence attributable to the different timescales and processes (adaptation vs. plasticity) involved.
2. Passive warming in autumn, rather than whole-growing season warming, is sufficient to delay autumn senescence.

Methods

Site description

The field site is located in Adventdalen valley (78°10'N, 16°06'E) on the High Arctic island of Spitsbergen in the Svalbard archipelago, where the polar day (no sunset) lasts from April 19th to August 25th. At the nearest weather station (Svalbard airport), the annual mean temperature during

the normal period (1991-2020) was $-3.9\text{ }^{\circ}\text{C}$ and mean temperatures during the coldest and warmest month were $-12\text{ }^{\circ}\text{C}$ (March) and $7\text{ }^{\circ}\text{C}$ (July), respectively (www.eklima.no). The mean annual precipitation during the same period is 217 mm with over 60 % falling as snow, assuming precipitation falls as snow during October to May when the mean monthly temperature is below $0\text{ }^{\circ}\text{C}$ (www.eklima.no). This constitutes a $2.8\text{ }^{\circ}\text{C}$ rise in mean annual temperature and a 27 mm rise in mean annual precipitation compared to the previous normal period (1961-1990, www.eklima.no). Locally, snow can be almost absent on ridges or reach up to 3 m depth in hollows due to the effect of wind and microtopography on snow deposition. Approximately 40 % of Svalbard is covered by continuous permafrost which has an active layer depth of 0.8 to 2.5 m (Humlum et al. 2003). The vegetation zones on Svalbard are polar desert and, to a lesser extent, Arctic tundra dominated by dwarf shrubs such as *Cassiope tetragona* (L.) D. Don, *Salix polaris* Wahlenb. and *Dryas octopetala* L. as well as graminoids. The study was carried out in two vegetation types which were found to have different soil moisture during the growing season (Cooper et al. 2011): dry heath (“Heath”) and mesic meadow (“Meadow”). The Heath site was located at the foot of a mountain slope and was faster draining, while the Meadow site was located near a river basin and retained higher soil moisture until mid-July (Morgner et al. 2010; Cooper et al. 2011). Further description of the site can be found in Cooper et al. (2011).

Snowmelt gradients

Six experimental and six natural gradients of snowmelt timing were set up in the study area (Figure 1a).

Three snow fences previously erected in each of two vegetation types (Heath and Meadow sites) were used to create the six experimental gradients. The wooden fences were 1.5 m tall and 6.2 m long and were established in 2006 (Cooper et al. 2011). They were placed perpendicular to the prevailing wind direction (south-east) which caused an accumulation of snow on the lee side of the fence with different

snow regimes (Figure S1a). In the following, “snow regime” is referred to the combination of environmental conditions caused by the different snow depths, which includes variations in snowmelt timing, soil moisture, soil temperature and nutrient availability. Snow regime was therefore treated here as an “umbrella” environmental factor which combines the effects of these other conditions. Three 75 x 75 cm plots were marked between 2 m and 14 m on the leeward side of the fence in the area with the deepest snow accumulation (“*Deep_{exp}*”). An additional three plots were marked between 11 m and 26 m on the leeward side of the fence with medium accumulation of snow (“*Medium_{exp}*”). Three plots with ambient snow regime (“*Low_{exp}*”) were marked in the vicinity of each fence, either on the windward side of the fence or next to the fences beyond the area of snow accumulation.

In July 2016, six natural gradients (three in each vegetation type) of three snow regimes with different snow depth and snowmelt timing were defined according to the local microtopography within approximately 50 m of the fences. Along microelevational gradients in the landscape (*ca.* 1-3 m vertical height), three plots were set up at the top of the slope where little snow accumulates and therefore melts early (“*Low_{nat}*”), three plots were installed along the slopes with medium snow accumulation and moderate snowmelt timing (“*Medium_{nat}*”) and three plots were placed at the foot of the slopes where snow accumulates the most and therefore melts late (“*Deep_{nat}*”; Figure S1b). The *Low_{nat}* plots in two of the natural gradients in the Heath site were the same as the *Low_{exp}* plots in the two closest experimental gradients. The third natural gradient had its own set of *Low_{nat}* plots.

On April 4th 2017, snow depth behind each snow fence of the experimental gradient was measured along three parallel 18 m long transects in 2 m intervals starting at 1.5 m away from the fences. The approximate snow depth in each *Deep_{exp}* and *Medium_{exp}* plot was then estimated from their known location behind the fence. Snow depth over the winter period 2016-2017 in the *Low_{exp}* plots of the experimental gradient and all plots of the natural gradient was measured at marking sticks which had been left in the ground during the snow-free period near the plots. Ground ice was common

in the area especially in places with little snow cover and was estimated to be up to 10 cm in some locations.

Open top chamber set up

In one of the three natural gradients in the Heath site, three five-sided open top chambers (OTCs; height = 0.5 m, surface area = 2.56 m²) made of Plexiglas® (Henry and Molau 1997; Marion et al. 1997) were installed in each snow regime (*Low_{nat}*, *Medium_{nat}* and *Deep_{nat}*), with each OTC adjacent to one of each of the natural gradient plots (see above), on July 20th 2016 and removed at the end of the experiment on August 7th 2016. The *Low_{nat}*, *Medium_{nat}* and *Deep_{nat}* plots of the natural snowmelt gradient (see above) were used as control plots. The OTC installation took place shortly before the beginning of visible leaf colouration associated with autumn senescence which coincides with a sharp decline in biomass production (Blume-Werry et al. 2016b), so we selectively increased the temperature at the end of the growing season.

Temperature and soil moisture measurements

Air temperatures at approximately 20 cm above the ground were measured at hourly intervals from July 23rd 2016 until August 7th 2016 with iButtons® (DS1922L-F5 thermochrons; Homechip Ltd., Milton Keynes, UK) in one or two plots in each snow treatment of the twelve individual gradients, as well as in the centre of six of the OTC chambers. Air temperature loggers were shaded from direct insolation with white plastic cups (9 cm tall and 7.5 cm diameter at the widest end) placed on top of the loggers. Several cuts of 6 cm length and spaced 2 cm apart were made lengthways into the cups to improve ventilation.

Soil surface temperatures were measured at hourly intervals from July 23rd 2016 until summer 2017 with iButtons® (DS1922L-F5 thermochrons; Homechip Ltd., Milton Keynes, UK). Of the soil temperature loggers, 47% stopped functioning over the subsequent winter, so the closest functioning

loggers were used as a temperature estimate for the determination of snowmelt timing. In different locations of the experimental gradient, the soil surface temperature had been logged additionally with Tinytag (Gemini Data Loggers, Chichester, UK) temperature loggers at hourly intervals for several years before and including the present study.

Snowmelt timing in 2017 was estimated from both types of loggers in the experimental gradient and from the iButtons in the natural gradient as the day on which the soil surface temperature started fluctuating above 0 °C. The correlation of snowmelt timing as estimated from the two loggers in the experimental gradient ($R=0.91$) in 2017 was used to estimate snowmelt timing in the experimental gradient in 2016. The correlation of snowmelt timing between the experimental and natural gradient as estimated from the iButtons ($R=0.8$) was used to estimate snowmelt timing in the natural gradient in 2016.

Soil moisture was measured on the same days as observations of phenology and NDVI measurements (see next sections) with a Theta ML 2x probe (Delta-T Devices Ltd, Cambridge, UK) in three locations per plot.

Observation of autumn senescence

Observations of autumn phenology were made on the dwarf shrub *Salix polaris*, the evergreen dwarf shrub *Dryas octopetala* and the herbaceous species *Bistorta vivipara* (L.) Gray and *Oxyria digyna* (L.). In each plot, five individual shoots per species were loosely marked with a cable tie or string. *Bistorta vivipara* was present in all marked plots and *S. polaris* in all plots but one. *Oxyria digyna* was not sufficiently present in the Meadow site, so only data from the Heath site was included in this study. *Oxyria digyna* and *D. octopetala* were lacking in about 20 % of all studied plots, but there was only one of the six natural and one of the six experimental gradients in which they were lacking completely from a whole snowmelt group.

The onset of autumn senescence was defined as the first day on which any amount of autumn colouration (yellow in *S. polaris* and *D. octopetala*, brown or yellow in *B. vivipara* and red in *O. digyna*) was visible on at least one leaf (the top leaf in *D. octopetala*) of the marked shoot. In this study, we chose the onset of senescence as the first and most distinguishable sign of senescence. It is also the easiest to consistently and objectively compare between different observers. The shoots were observed every other day from July 19th 2016 (in OTCs: July 20th and July 23rd, then every other day) to August 6th 2016 and the date of onset of autumn senescence was noted down for each marked shoot. Additionally, the timing of autumn senescence was calculated as the number of days after snowmelt (DAS) until the onset of senescence by subtracting the day of year (number of days from January 1st; DOY) from day of year of snowmelt. No noticeable amount of leaf colouration was seen in the study area at the beginning of the observation period. By the end of the study period, 90 % of all individuals had started to senesce. The remaining individuals were excluded from the analysis in order to focus the analysis on when senescence occurred rather than if it occurred during the study period.

NDVI measurement

On the same days as the phenological observations, the average NDVI index (normalized difference vegetation index, calculated from the red and near-infrared light reflected by vegetation) was measured on each plot with a GreenSeeker handheld optical crop sensor (Trimble Inc. ©, Sunnyvale, US) as a measure of vegetation greenness. The GreenSeeker was held at approximately 120 cm height and moved horizontally across each plot to give an average reading of a 50 cm oval area on the ground.

Statistical analysis

The effects of vegetation type (Heath or Meadow), snow regime (Low_{nat} , $Medium_{nat}$ or $Deep_{nat}$ in the natural gradient; Low_{exp} , $Medium_{exp}$ or $Deep_{exp}$ in the experimental gradient) and OTC treatment (control or OTC) on the DOY of autumn senescence and on the NDVI, was analysed separately for each gradient type (natural or experimental) and separately for each species (in the DOY analysis) with a mixed effects linear model using the *nlme* package (Pinheiro et al. 2018, version 3.1-137) in R (R Core Team 2020, version 3.5.1). In the snowmelt gradient study, the DAS was additionally analysed as a response variable. The fixed effects were snow regime and vegetation type and the random effect Plot nested in Fence. The sample size in the DAS analysis was lower since the day of year of snowmelt was not available for some plots in the natural gradient. No analysis was carried out for DAS in *O. digyna* in the natural gradient due to the low sample size. In the OTC study, the fixed effects were snow regime and OTC treatment and the random effect was Plot, since only a single gradient was used. Model selection was carried out using the maximum likelihood test and significance of predictor variables ($p \leq 0.05$) was determined with F-tests. Residuals were checked visually for compliance with assumptions of normality of the residual distribution, homogeneity of residual variance and independency of observations. Pairwise comparisons were carried out with the *emmeans* package in R (Lenth 2018, version 1.3.0). When pairwise comparisons were significant, the p -value was adjusted using the Holm's method.

The difference in mean soil moisture between Vegetation types and Snow regime, and their interaction, was tested for significance ($p \leq 0.05$) with a linear mixed model using Plot nested in Fence as a random effect. Residuals were checked visually for compliance with ANOVA assumptions.

Figures 2-7 were prepared in R (version 3.6.2) and figure 1 was prepared in Powerpoint.

Results

Comparison of experimental and natural snowmelt gradients

Snow depth was greater and more variable along the experimental compared to the natural gradient (Table 1). In the experimental gradient, *Deep_{exp}* plots accumulated 1.1 ± 0.04 m (mean \pm SE) of snow and *Medium_{exp}* plots 0.6 ± 0.07 m. *Low_{exp}* plots were covered by between 0.1 and 0.5 m of snow (measured approximately). In the natural gradient, the snow depth in *Low_{nat}* plots was approximately 0-0.1 m, in *Medium_{nat}* plots 0.1-0.2 m and in *Deep_{nat}* plots 0.3-0.5 m. Hence, when considering snow depth, *Medium_{exp}* plots in the experimental snow gradient were more similar to *Deep_{nat}* plots, rather than *Medium_{nat}* plots, in the natural snow gradient.

In the experimental treatment, *Low_{exp}* plots melted out on 24th-26th May, *Medium_{exp}* plots on 4th-5th June and *Deep_{exp}* plots on 10th-12th June 2016. Despite the differences in snow depth between the experimental and the natural snowmelt gradients, snowmelt timing in the experimental gradient was similar in the corresponding plots of the natural gradient (Table 1), i.e. *Low_{exp}*-*Low_{nat}*, *Medium_{exp}* - *Medium_{nat}* and *Deep_{exp}* - *Deep_{nat}* had similar snowmelt timing.

Soil moisture varied significantly with snowmelt timing (Snow_{exp} : $F_{2,1568}=10.34$, $p<0.0001$; Snow_{nat} : $F_{2,1597}=45.61$, $p<0.0001$) but not with Vegetation type, so Vegetation type did not represent different soil moisture conditions during the autumn. In the experimental gradient, soil moisture was lowest in *Low_{exp}* and highest in *Deep_{exp}* (Table 2, Figure 2). In the natural gradient, it was highest in *Deep_{nat}* and lowest in *Medium_{nat}*.

The effect of snow regime on autumn senescence

Oxyria digyna senesced first (DOY= 206 ± 0.3 [mean \pm SE]), followed by *B. vivipara* (DOY= 209 ± 0.2), *S. polaris* (DOY= 210 ± 0.2) and lastly *D. octopetala* (DOY= 212 ± 0.3) (Figure 3).

The effect of snow regime on the timing of senescence was the same in the experimental and the natural gradient in *S. polaris*, *B. vivipara* and *O. digyna* and different in *D. octopetala*. In *S. polaris*, snow regime had no significant effect on the timing of senescence onset in either gradient (Table 3, Figure 3a). In *B. vivipara*, senescence was significantly delayed by one day in *Medium_{nat}*

plots compared to Low_{nat} and $Deep_{nat}$ plots (Low_{nat} - $Medium_{nat}$ $p = 0.04$, $Deep_{nat}$ - $Medium_{nat}$ $p = 0.04$) in the natural gradient and by two days in $Medium_{exp}$ plots compared to Low_{exp} and $Deep_{exp}$ plots in the experimental gradient (Low_{exp} - $Medium_{exp}$ $p = 0.0428$, $Deep_{exp}$ - $Medium_{exp}$ $p = 0.0428$; Figure 3b). In *D. octopetala*, the snow regime only altered senescence timing significantly in the experimental gradient in the Meadow site where senescence occurred four to five days later in $Medium_{exp}$ plots compared to Low_{exp} plots ($p = 0.03$) or $Deep_{exp}$ plots ($p = 0.01$; Figure 3c). Senescence timing did not differ along the natural gradient in this species. *Oxyria digyna* was only present in both the experimental and the natural gradient in the Heath site where its senescence timing was not significantly affected by snow regime (Figure 3d).

Vegetation type only had a significant effect on senescence in *S. polaris* growing in the natural gradient where it senesced 1.6 days earlier in the Meadow compared to the Heath site (Table 3). Data for senescence timing along the snow regimes are therefore presented as means across both vegetation types in Figure 3.

The effect of snow regime and vegetation type on the time from snowmelt until the onset of senescence (DAS) was similar among *S. polaris*, *B. vivipara* and *D. octopetala* (Figure 4a, b, c). In these species, later snowmelt timing always significantly decreased the DAS (Table 4). This effect was significantly influenced by Vegetation type in the experimental gradient, where DAS was similar in Low_{exp} and $Medium_{exp}$ plots in the Heath site but Low_{exp} was significantly higher than $Medium_{exp}$ in the Meadow site (Table 4). In *O. digyna*, no clear trend in the effect of Snow or Vegetation on DAS could be seen (Figure 4d).

The effect of passive autumn warming on the timing of autumn senescence

The OTC chambers caused an average increase in air temperature of 1 °C between 23rd July (DOY 205) and 6th August 2016 (DOY 2019) (Figure S2). At night, temperatures between OTC and control plots were similar, but during daytime, the OTC plots were heated by up to 8 °C compared to the

temperature in the control plots. This is likely due to additional heating effects from direct insolation or rising warm air from the soil, as the actual temperature during those periods of very high warming greatly exceeded the temperature measured at the weather station (Appendix A). Removing any temperatures above 18 °C (considering the maximum weather station temperature was 13.4 °C) yielded a maximum temperature increase of 5 °C for OTC plots. It is worth noting that although wind reduction inside OTCs has previously not been found to be the driver of air temperature (Bokhorst et al. 2013), the affect it has on the boundary layer of plants can cause large differences in plant tissue surface temperature inside OTCs (De Boeck et al. 2012). The passive warming effect was stronger in the *Low_{nat}* than in *Medium_{nat}* and *Deep_{nat}*. The soil surface temperature within the OTCs was colder in *Low_{nat}* and *Deep_{nat}* and warmer in *Medium_{nat}* compared to the control plots (Figure S2b, Appendix A). Both cooling and warming effects of soil surface temperature due to OTCs have been reported previously and can be due to water logging, plant cover and microsite placement of the loggers (Hollister et al. 2006). In future studies, more temperature loggers and their more careful placement could achieve clearer results.

The OTC chambers caused an overall delay of 1.5 days in the autumn senescence across all species ($F_{1,14}=8.3$, $p=0.012$; Figure 5). This was mostly due to *Dryas octopetala*, which senesced 3.7 days later in plots with the OTC treatment ($p=0.0014$), while the other species had no significant delay in the timing of senescence. The effect of the OTC chambers was the same across all snowmelt regimes. The first signs of senescence in the tagged shoots appeared later in the OTC treatment compared to the control and most individuals senesced at the end of the observation period compared to the control plants for which there was a peak in senescence on DOY 213 (Figure 6).

The effect of snow regime and temperature on NDVI

In both the experimental and the natural gradient, there was a significant decline in NDVI (DOY_{exp}: $F_{1,528} = 49.39$, $p < 0.0001$; DOY_{nat}: $F_{1,530} = 50.0$, $p < 0.0001$) over the late summer growing season

measurement period, as would be expected for the onset of autumn senescence (Figure 7a, b). Snow regime also had a significant effect in both gradients (Snow_{exp}: $F_{2,528} = 80.82, p < 0.0001$; Snow_{nat}: $F_{2,530} = 6.17, p = 0.0022$). In the natural gradient, the NDVI was the same in *Low_{nat}* and *Deep_{nat}* and significantly lower in *Medium_{nat}* (Table 5, Fig. 7a). In the experimental gradient, *Low_{exp}* also had the highest NDVI in both vegetation types, while *Medium_{nat}* was lowest in the Heath site and *Deep_{exp}* was lowest in the Meadow site (Table 5, Fig. 7b). The NDVI was initially higher and declined at a faster rate over time in the Heath site than in the Meadow site (Fig 7a, b; Vegetation_{exp}: n.s.; Vegetation_{nat}: $F_{1,4} = 11.5, p = 0.0275$).

The OTC treatment significantly decreased the NDVI value by 0.03 compared to the control in all snow regimes (Figure 7c; Treatment: $F_{1,166} = 24.584, p < 0.0001$).

Discussion

Timing of senescence in experimental and natural snowmelt gradients

Our results show that experimental and natural gradient studies lead to similar conclusions regarding the response of plants to changes in snow conditions in three out of the four studied species (*S. polaris*, *B. vivipara* and *O. digyna*). In our study, the snowmelt timing in the experimental gradient was the same as the corresponding snow regime in the natural gradient, which could imply that the timing of autumn senescence in *B. vivipara* is linked to snowmelt timing. This species senesces the latest in *Medium_{exp}* and *Medium_{nat}* snow regime plots (Fig. 3b), indicating the presence of the most optimal conditions for a longer growing season. The senescence timing in *S. polaris* and *O. digyna* was not affected by snow regime after 10 years of experimentally modified snowmelt timing. This response persisted along the long-term, natural gradient, implying that the mechanism for responding to changes in snowmelt timing is the same during short-term and long-term climate variations. In *S. polaris*, the DAS decreased with later snowmelt. According to Gehrman et al. (2018), phenophases

which occur synchronously but require fewer DAS along a snow gradient may be under photoperiodic control or other environmental regulation such as temperature changes. In our study location, day length was 24 hours during the entirety of the study period, so it is unlikely to control senescence onset in *S. polaris*. Changes in spectral composition have been suggested to affect autumn senescence of deciduous trees (Brelsford et al. 2019) but the effect on tundra shrubs is unknown. The compensation response of a decrease in DAS with later snowmelt could instead be due to faster development of plants in later melting areas, as temperatures are higher compared to earlier melting plots upon melt-out. Oberbauer et al. (2013) found that the timing of senescence in Arctic plants is advancing with higher spring and summer temperatures, suggesting that senescence timing could be controlled by growing season temperatures.

In *D. octopetala*, the results from the natural gradient suggest that increased snow depth and delayed snowmelt timing will not change the timing of autumn senescence. Projected increases in winter precipitation would therefore lead to a shorter growing season, as this would delay the onset of growth while not extending the end of the growing season. The experimental gradient, on the other hand, suggests that medium snow depth in mesic sites delays senescence in *D. octopetala*, thereby compensating for the delayed start in spring to some extent. On the other hand, the NDVI measurements, which were made at the plot level, suggest that little snow cover (as in the *Low_{nai}* and *Low_{exp}* plots) delays the beginning of autumn senescence the most and that the rate of autumn browning is strongly dependent on vegetation type (Figure 7). NDVI values are strongly dependent on both the type of vegetation and the amount of vegetation coverage, both of which are altered by the snow manipulation (Riedel et al. 2005; Cooper et al. 2019). While this could explain the divergence in results between different methods, it also highlights the need to employ a variety of methods in order to understand both species- and community-scale responses.

Although we saw similar responses in the timing of autumn senescence to experimental and natural snowmelt gradients, we were able to simultaneously address different aspects of plant

responses to climate with this combination of approaches. Short-term (a few years to a few decades) experiments, such as this snow manipulation, show whether plant phenology is plastic. This plasticity enables plants to respond immediately to a changing climate before genetic changes can take place. Observations along natural gradients, on the other hand, show the long-term adaptation to climate change, which can include plasticity in the phenology or genetic differentiation of phenological timing. Such a combination approach remains rare even though it is critical to estimate the contributions of phenotypic plasticity and evolutionary adaptation (Shaver et al. 2000; Dunne et al. 2004).

Species-specific differences in senescence timing along snow depth gradients

We found species-specific differences in the response of the onset of senescence to the timing of spring snowmelt and the depth of winter snow. Semenchuk et al. (2016) measured senescence timing over several years in three of the four species (*B. vivipara*, *D. octopetala*, *S. polaris*) investigated here in the same snow fence experiment and reported on the mid-point of senescence of the vegetation plots, rather than onset as we present here. They found that senescence was delayed by approximately ten days in *Deep_{exp}* compared to *Low_{exp}* in all of these species. This difference is greater than the delay (or lack of delay) in phenology in the species in the present study. However, the study by Semenchuk et al. (2016) only measured phenology once a week and therefore could not reliably detect shifts in phenology of only a few days. They also found that the shift in senescence timing along the snow gradient is not the same between different years. While some yearly variation in plant responses can be expected due to a certain degree of phenotypic plasticity, both our study and Semenchuk et al. (2016) showed a similar direction in the plant responses, i.e. no advance in senescence. Yearly variation and greater measurement intervals may explain the difference between our results and those from the literature and act as modifiers on the plant responses. Species-specific differences in the response of phenology to snow regimes are common in the literature (e.g. Wipf et

al. 2009; Wipf and Rixen 2010; Cooper et al. 2011) and some studies suggest that growth form or temporal niche of a species determines their response to snowmelt timing or snow depth (Wipf 2010; Petraglia et al. 2014; Khorsand Rosa et al. 2015; Livensperger et al. 2016), but our study could not confirm this. It has been found that genetic adaptation could account for 20 %-75 % of changes in flowering time in response to climate change in *Boechnera stricta* (Graham) Al-Shehbaz, with larger changes calculated for shorter generation times (Anderson et al. 2012). Short-lived, herbaceous species may be more strongly controlled by genetic adaptation than long-lived woody species. These effects of stronger genetic adaptation and shorter generation times could play a role in the stronger variation in senescence timing we see in the herbaceous species *B. vivipara* compared to the woody species *S. polaris*. Senescence timing can also be affected by bud burst timing. Bud burst responds strongly to snowmelt timing, but many Arctic species retain the same duration for some phenophases, so that shifts in senescence timing can be a lag effect of shifted bud burst (Semenchuk et al. 2016). In our study, budburst data was not available.

In our study, snow regime and temperature increase did not alter senescence timing in *O. digyna*, so the onset of senescence is likely controlled by another factor in this species. It was recently shown that the timing of leaf-out and senescence of *O. digyna* follow a clear latitudinal trend, suggesting that day length and/or spectral composition could be an important phenological control for this species (Bjorkman et al. 2017). This would support our result that senescence timing was the same along the snow gradients, as all plants experienced the same photoperiod.

Lower soil moisture has previously been found to be correlated with earlier onset of senescence (Livensperger et al. 2019), but since it is confounded by snowmelt timing its effect is difficult to separate from snowmelt. In our study, we did not find significant differences in the mean soil moisture for the period 19th July to 6th August between vegetation types (Fig. 2). This is likely because the autumn in 2016 had relatively little precipitation, so even the poorer draining Meadow site did not accumulate as much moisture. Higher soil moisture has previously been measured in the

Meadow compared to the Heath site especially shortly after snowmelt but the difference decreased in July (Cooper et al. 2011, 2019). Higher snow depth commonly led to higher soil moisture in this study site, although this difference is more pronounced shortly after snowmelt and not always significant when considering the whole-season mean of soil moisture (Cooper et al. 2019; Mörsdorf et al. 2019). While this study was not established to specifically test gradients in soil moisture, we additionally examined our data for a significant correlation between the mean soil moisture of the study period and the DOY (Appendix B). We found that DOY in *S. polaris* was significantly affected by soil moisture with opposing effects in the natural and the experimental gradient (Figure B1, Table B1).

Increased nitrogen was measured in the soil and leaves of *S. polaris* in *Deep_{exp}* plots in the snow experiment in previous studies (Semenchuk et al. 2015; Mörsdorf et al. 2019). In warmed winter soil under *Deep_{exp}* snow, vegetation type affects the availability of the increase in nutrients (Semenchuk et al. 2015), but it is currently not clear how nutrient availability affects senescence. Senescence is strongly linked to nutrient cycling since nutrient resorption takes place during that time, so it may indirectly be affected by soil moisture. Therefore, we may not have seen strong effects of vegetation type on senescence timing in this study due to the similar autumn soil moisture.

Overall, data on senescence timing in Arctic plants are still scarce and studies use different stages in the senescence process to define senescence onset, such as 1%, 50% or 100% leaf colouration (Abbandonato 2014). Furthermore, the progress of senescence from the onset (i.e. 1 % colouration) to the end (i.e. 100 % colouration) can take as long as 6 weeks (Abbandonato 2014; Semenchuk et al. 2016) and is therefore not directly comparable to measurements of a single senescence stage. These discrepancies make it difficult to directly compare study results and implies that a common measure of senescence timing is needed (Gallinat et al. 2015).

Passive autumn warming

In our study, *D. octopetala* was the only species for which the onset of senescence was significantly affected by the presence of the OTCs. A delay of four days in the leaf colouration of *D. octopetala* exposed to the same method of passive warming was also found by Welker et al. (1997). They suggest that the extension of the active period is possible because *D. octopetala* is a “wintergreen” plant, meaning it retains its leaves throughout the winter and greens up rapidly in spring during a leaf lifespan of approximately 2 years. They also found a higher C:N ratio in *D. octopetala* leaves from warmed plots in the autumn, which could represent a greater resorption efficiency from leaves due to internal N cycling. This is in contrast to the other species in our study, which senesced at the same time regardless of temperature conditions. In a meta-analysis of the international tundra experiment (ITEX), there was generally no treatment effect after four years of OTC treatment on the timing of senescence in various species in up to six different arctic and alpine sites (Arft et al. 1999). A more recent meta-analysis including studies with up to 20 years of OTC treatment found a mean delay in senescence across 61 species (Collins et al. 2021), which could indicate that there is a lag of several years in the response of autumn senescence to warming. A recent review of ITEX found that the timing of senescence is overall not sensitive to temperature increases across time or space when considering multiple species across the Arctic (Prevéy et al. 2017). However, most data in these syntheses is from the sub-Arctic and low Arctic as senescence data from the High Arctic is rare. Due to the different light conditions in the High Arctic we therefore do not know if we can extrapolate along this latitudinal gradient. While these studies have employed OTC chambers throughout the whole growing season, our study shows that a selective passive warming of only the end of the growing season can already lead to the extension of senescence onset, at least for some species.

Shifts in senescence due to warming may be small and undetectable if summarised across several species, but our results show that substantial changes can be seen in individual species. The growing season length for *D. octopetala* in this location is only about 5.5 weeks (Cooper et al. 2011), so even a small lengthening of the growing season by four days, which we found here, is equivalent

to a 10 % increase of time available for growth and other essential processes. It has been proposed that as little as a 1-day increase in the length of the growing season increases the annual GPP by 0.6 % (Piao et al. 2007). However, autumn warming increases respiration more than photosynthesis in northern ecosystems (Piao et al. 2008), so a delay in senescence caused by warming could severely impact net ecosystem exchange of carbon, potentially resulting in increased carbon gain. This is especially true in heath communities, where evergreen dwarf shrubs make a major contribution to net ecosystem exchange (Strimbeck et al. 2019). To fully evaluate the changes in carbon dynamics caused by growing season shifts, the effect of climate changes during each season has to be evaluated separately. Furthermore, the progression of senescence needs to be considered from the onset (first colouration) to the end (leaf fall), as each phase during senescence can be differentially influenced by environmental factors and can vary greatly across habitats, years and phylogeny (Abbandonato 2014; Panchen et al. 2015).

Conclusions

Climate change will not affect all seasons equally and changes in one season can have a legacy effect on plant responses in another season. We have shown here that spring snowmelt and passive autumn warming both affect the timing of autumn senescence. The timing of autumn senescence is critical for plant ecology and ecosystem cycling because the length of the photosynthetic activity in plants determines plant productivity, nutrient and water cycling and carbon uptake and release. To understand how these processes will be affected by climate change, we should study the plant responses using a variety of methods and scales. Our results showed that combining different study methods, such as experiments and natural gradients, and different scales, such as the plant level observations and plot level NDVI measurements, can provide complementary insights into the short- and long-term plant responses. Applying multiple approaches in the same study helps to overcome the limitations inherent in individual approaches, to generate more robust scientific conclusions. In

the future, we need to aim to combine different methods at overlapping temporal and spatial scales to develop a better understanding of how plants will be affected by climate change.

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Figure 1. Plot and sampling design of the study. a) Three snow fences and three natural gradients were used in each vegetation type (Heath or Meadow). In each gradient, three plots were marked in a location with *Low* snow depth, three plots in a location with *Medium* snow depth and three plots in a location with *Deep* snow depth. In the experimental gradients, the *Low_{exp}* plots were located at either at the windward side of the fence or next to the fence beyond the area of snow accumulation, while *Medium_{exp}* and *Deep_{exp}* were leeward. In each plot, five shoots per species were marked for observations (black dots). b) In one of the natural gradient slopes, three OTCs (pentagons) were set up in each snow regime adjacent to the control plots (squares).

Figure 2. Volumetric water content (VWC %) in an experimental (*Low_{exp}*=solid line, *Medium_{exp}*=dotted line, *Deep_{exp}*=dashed line) and natural (*Low_{nat}*=solid line, *Medium_{nat}*=dotted line, *Deep_{nat}*=dashed line) snowmelt gradient at two different vegetation types (Heath and Meadow). VWC was measured every other day from 19th July to 7th August 2016 (n=9).

Figure 3. The mean day of year (DOY \pm SE) of the onset of senescence in a) *Salix polaris*, b) *Bistorta vivipara*, c) *Dryas octopetala* and d) *Oxyria digyna* along an experimental (*Low_{exp}*, *Medium_{exp}*, *Deep_{exp}*) and natural (*Low_{nat}*, *Medium_{nat}*, *Deep_{nat}*) snowmelt gradient. Different letters above the data points indicate significant different means between snow regimes within the same species and snowmelt gradient. Vegetation type only had a significant effect on senescence in *S. polaris* growing in the natural gradient. Data for senescence timing along the snow regimes are therefore presented as means across both vegetation types.

Figure 4. The mean number of days after snowmelt (\pm SE) of the onset of senescence in a) *Salix polaris*, b) *Bistorta vivipara*, c) *Dryas octopetala* and d) *Oxyria digyna* along an experimental (*Low_{exp}*, *Medium_{exp}*, *Deep_{exp}*) and natural (*Low_{nat}*, *Medium_{nat}*, *Deep_{nat}*) snowmelt gradient in two

different vegetation types (Heath=circle, Meadow=triangle). Different letters denote significant differences between snow regimes (lower case letters for the Heath site and upper case letters for the Meadow site). No statistical testing was carried out for *O. digyna* due to the low sample size.

Figure 5. The mean day of year (\pm SE) of the onset of autumn senescence in four arctic species (BV= *Bistorta vivipara*, DO=*Dryas octopetala*, OD= *Oxyria digyna*, SP= *Salix polaris*) along a snowmelt gradient in control (filled symbols) and OTC (open symbols) treatment plots. Different letters denote statistically different means.

Figure 6. The accumulated number of marked senescing *Dryas octopetala* shoots for each observational day of year in control (black line) and OTC (grey line) plots.

Figure 7. Mean NDVI values and modelled regression with 95% confidence intervals for a) natural snowmelt gradient (Low_{nat} =solid line/circles, $Medium_{nat}$ =dotted line/squares, $Deep_{nat}$ =dashed line/triangles) and an b) experimental snowmelt gradient (Low_{exp} =solid line/circle, $Medium_{exp}$ =dotted line/squares, $Deep_{exp}$ =dashed line/triangles) in two vegetation types as well as for c) control and OTC treatment plots along one natural snowmelt gradient (Low_{nat} =solid line/circles, $Medium_{nat}$ =dotted line/squares, $Deep_{nat}$ =dashed line/triangles) in Heath.

1 Table 1. Estimated date of snowmelt (day of year in parentheses) in 2016 and snow depth on
 2 April 4th 2017 along an experimental and a natural snowmelt gradient in two different
 3 vegetation types (n=9).
 4

Gradient	Vegetation	Low	Medium	Deep
Experimental	Heath	21 May 2016 (142) 0.1 - 0.5 m	3 June 2016 (155) 0.7 ± 0.2 m	8 June 2016 (160) 1.2 ± 0.2 m
	Meadow	23 May 2016 (144) 0.1 - 0.5 m	04 June 2016 (156) 0.5 ± 0.3	12 June 2016 (164) 1.1 ± 0.2
Natural	Heath	22 May 2016 (143) 0 - 0.1 m	29 May 2016 (150) 0.1 - 0.2 m	9 June 2016 (161) 0.3 - 0.5 m
	Meadow	22 May 2016 (143) 0 - 0.1 m	3 June 2016 (155) 0.1 - 0.2 m	4 June 2016 (156) 0.3 - 0.5 m

5
 6 Table 2. Pairwise comparisons of mean volumetric water content during the measuring period
 7 (19th July – 7th August 2016) between snow regimes in an experimental and a natural snowmelt
 8 gradient. The effect size, standard error, degrees of freedom and p-value of comparisons
 9 between plots with different snow regimes is shown.

Comparison	Effect size	SE	df	t ratio	p value
Experimental gradient					
Low-Medium	-2.13	0.826	2, 1568	-2.529	0.0115
Low-Deep	-4.06	0.827	2, 1568	-4.827	<.0001
Deep-Medium	1.94	0.818	2, 1568	2.326	0.0201
Natural gradient					
Low-Medium	2.94	0.719	2, 1597	4.086	<.0001
Low-Deep	-3.91	0.719	2, 1597	-5.434	<.0001
Deep-Medium	6.85	0.719	2, 1597	9.520	<.0001

10 Table 3 The effect of Snow regime (*Low, Medium, Deep*) and Vegetation type (Heath, Meadow)
 11 on the DOY (day of year) of the onset of autumn senescence in four High Arctic plant species
 12 growing along an experimental and a natural snowmelt gradient. The results of the F-statistics
 13 from the ANOVA are shown. NA signifies lack of statistical testing due to insufficient
 14 representation of plants within study plots.

	Snow	Vegetation	Snow*Vegetation
Species			
<i>S. polaris</i>			
Experimental gradient	n.s.	n.s.	n.s.
Natural gradient	n.s.	$F_{1,4}=9.5, p=0.037$	n.s.
<i>B. vivipara</i>			
Experimental gradient	$F_{2,46}=3.7, p=0.0332$	n.s.	n.s.
Natural gradient	$F_{2,46}=3.5, p=0.0386$	n.s.	n.s.
<i>D. octopetala</i>			
Experimental gradient	n.s.	n.s.	$F_{2,31}=3.15, p=0.0571$
Natural gradient	n.s.	n.s.	n.s.
<i>O. digyna</i>			
Experimental gradient	n.s.	NA	NA
Natural gradient	n.s.	NA	NA

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26 Table 4 The effect of Snow regime (*Low, Medium, Deep*) and Vegetation type (Heath,
 27 Meadow) on the DAS (days after snowmelt) until the onset of autumn senescence in four
 28 High Arctic plant species growing along an experimental and a natural snowmelt gradient.
 29 The results of the F-statistics from the ANOVA are shown. NA signifies lack of statistical
 30 testing due to missing or low sample size.

	Snow	Vegetation	Snow*Vegetation
Species			
<i>S. polaris</i>			
Experimental gradient	$F_{2,43}=25.06, p<0.0001$	n.s.	$F_{2,43}=6.33, p=0.0039$
Natural gradient	$F_{2,29}=14.75, p<0.0001$	n.s.	n.s.
<i>B. vivipara</i>			
Experimental gradient	$F_{2,44}=32.65, p<0.0001$	n.s.	$F_{2,44}=10.57, p=0.0002$
Natural gradient	$F_{2,27}=35.73, p<0.0001$	n.s.	$F_{2,27}=7.29, p=0.0029$
<i>D. octopetala</i>			
Experimental gradient	$F_{2,138}=54.27, p<0.0001$	n.s.	$F_{2,138}=18.23, p<0.0001$
Natural gradient	$F_{2,128}=76.41, p<0.0001$	n.s.	$F_{2,128}=16.76, p<0.0001$
<i>O. digyna</i>			
Experimental gradient	$F_{2,16}=7.2, p=0.0059$	NA	NA
Natural gradient	NA	NA	NA

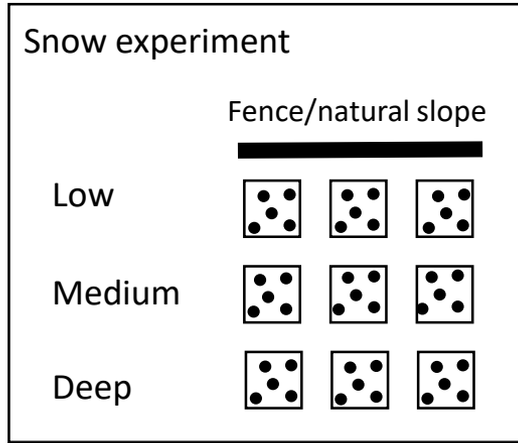
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41 Table 5 Pairwise comparisons of NDVI between snow regimes in an experimental and a natural
 42 snowmelt gradient in two different vegetation types (Heath and Meadow). The effect size,
 43 standard error, degrees of freedom and p-value of comparisons between plots with different
 44 snow regimes is shown (n=9). P values were adjusted using the Holm's method

Comparison	Heath					Meadow				
	Effect size	SE	df	t ratio	p value	Effect size	SE	df	t ratio	p value
Natural gradient										
Low-Medium	0.02	0.006	5, 530	3.369	0.0032	0.02	0.006	5, 530	3.369	0.0032
Low-Deep	0.005	0.006	5, 530	0.819	0.4	0.005	0.006	5, 530	0.819	0.4
Medium-Deep	0.015	0.006	5, 530	2.550	0.02	0.015	0.006	5, 530	2.550	0.02
Experimental gradient										
Low-Medium	0.055	0.009	4, 528	6.411	0.0006	0.055	0.009	4, 528	6.437	0.0006
Low-Deep	0.024	0.009	4, 528	2.849	0.0046	0.124	0.009	4, 528	14.467	0.0006
Medium-Deep	0.031	0.009	4, 528	3.562	0.0008	-0.069	0.009	4, 528	-8.03	0.0006

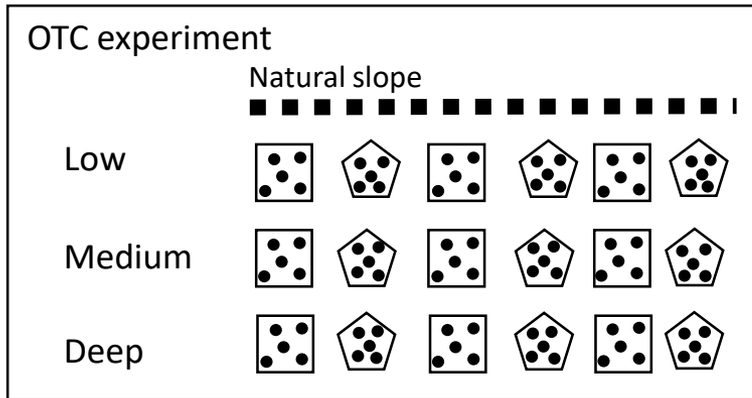
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a)



	Heath		Meadow	
	Experimental	Natural	Experimental	Natural
No. fences/ natural slopes	3	3	3	3
No. plots				
Low	9	9	9	9
Medium	9	9	9	9
Deep	9	9	9	9

b)



	Natural - Control	Natural - OTC
No. plots		
Low	3	3
Medium	3	3
Deep	3	3

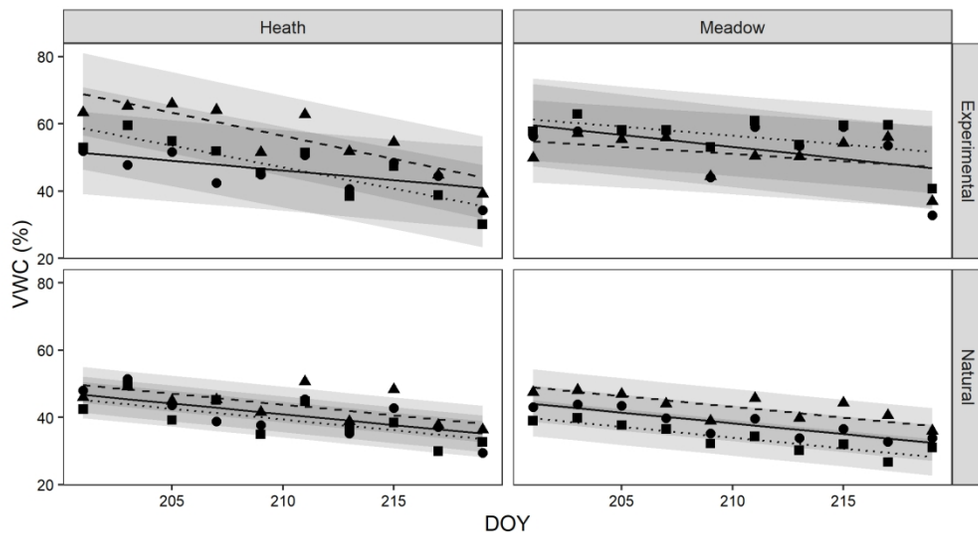


Figure 2. Volumetric water content (VWC %) in an experimental (Lowexp=solid line, Mediumexp=dotted line, Deepexp=dashed line) and natural (Lownat=solid line, Mediumnat=dotted line, Deepnat=dashed line) snowmelt gradient at two different vegetation types (Heath and Meadow). VWC was measured every other day from 19th July to 7th August 2016 (n=9).

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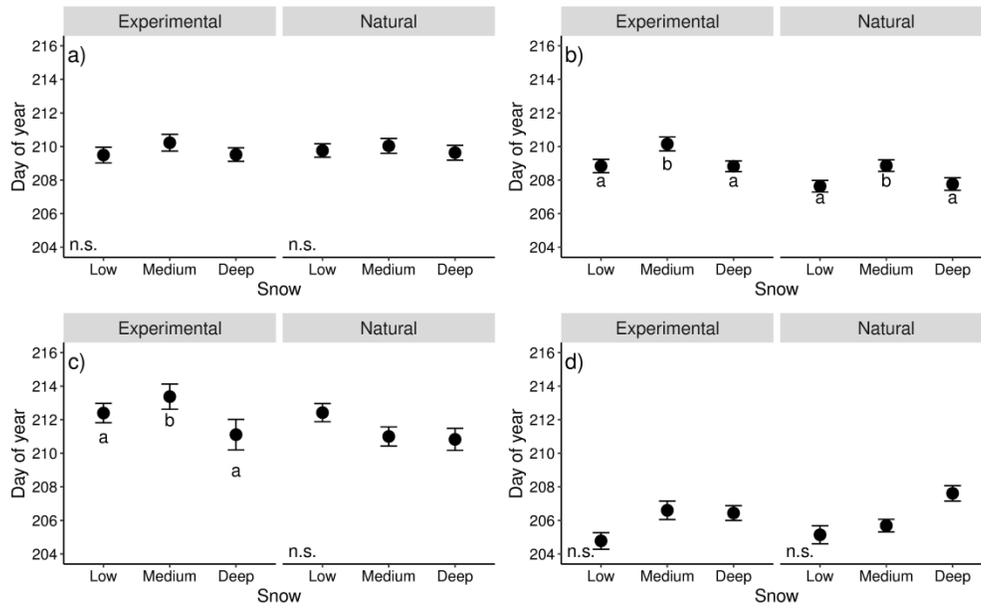


Figure 3. The mean day of year (DOY \pm SE) of the onset of senescence in a) *Salix polaris*, b) *Bistorta vivipara*, c) *Dryas octopetala* and d) *Oxyria digyna* along an experimental (Lowexp, Mediumexp, Deepexp) and natural (Lownat, Mediumnat, Deepnat) snowmelt gradient. Different letters above the data points indicate significant different means between snow regimes within the same species and snowmelt gradient. Vegetation type only had a significant effect on senescence in *S. polaris* growing in the natural gradient. Data for senescence timing along the snow regimes are therefore presented as means across both vegetation types.

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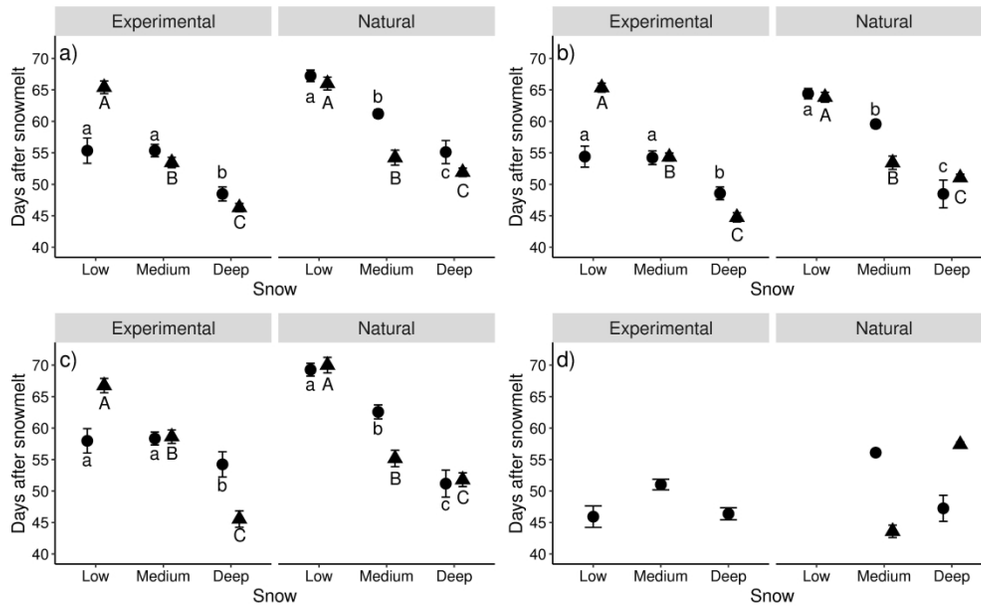


Figure 4. The mean number of days after snowmelt (\pm SE) of the onset of senescence in a) *Salix polaris*, b) *Bistorta vivipara*, c) *Dryas octopetala* and d) *Oxyria digyna* along an experimental (Lowexp, Mediumexp, Deepexp) and natural (Lownat, Mediumnat, Deepnat) snowmelt gradient in two different vegetation types (Heath=circle, Meadow=triangle). Different letters denote significant differences between snow regimes (lower case letters for the Heath site and upper case letters for the Meadow site). No statistical testing was carried out for *O. digyna* due to the low sample size.

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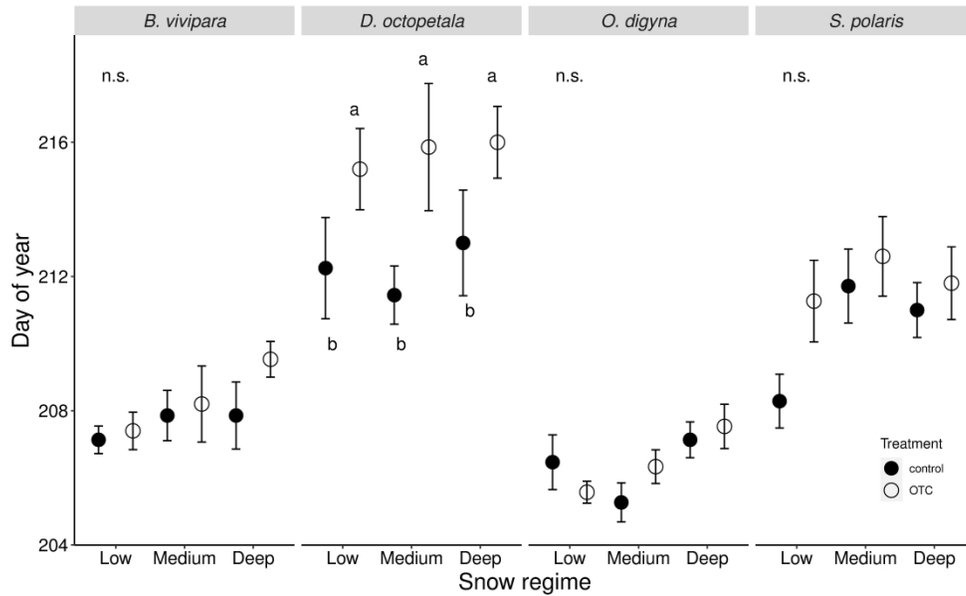


Figure 5. The mean day of year (\pm SE) of the onset of autumn senescence in four arctic species (BV= *Bistorta vivipara*, DO= *Dryas octopetala*, OD= *Oxyria digyna*, SP= *Salix polaris*) along a snowmelt gradient in control (filled symbols) and OTC (open symbols) treatment plots. Different letters denote statistically different means.

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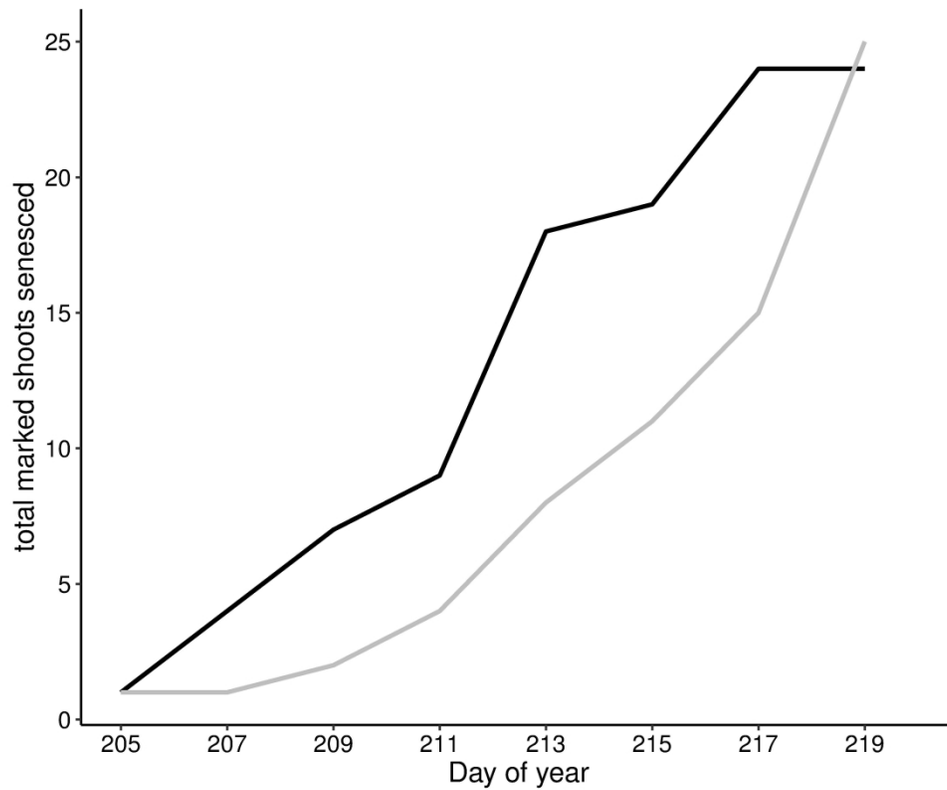


Figure 6. The accumulated number of marked senescing *Dryas octopetala* shoots for each observational day of year in control (black line) and OTC (grey line) plots.

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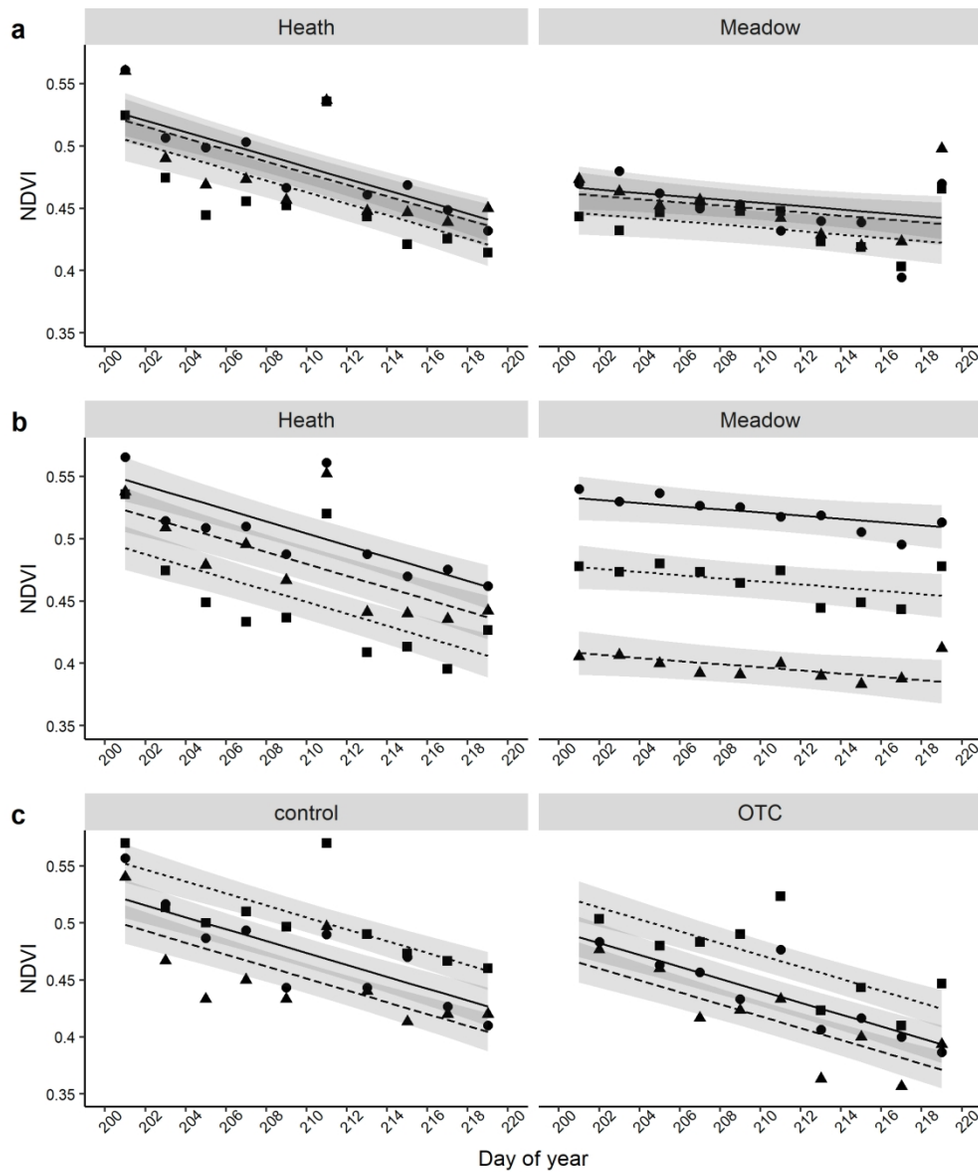


Figure 7. Mean NDVI values and modelled regression with 95% confidence intervals for a) natural snowmelt gradient (Lownat=solid line/circles, Mediumnat=dotted line/squares, Deepnat=dashed line/triangles) and an b) experimental snowmelt gradient (Lowexp=solid line/circle, Mediumexp=dotted line/squares, Deepexp=dashed line/triangles) in two vegetation types as well as for c) control and OTC treatment plots along one natural snowmelt gradient (Lownat=solid line/circles, Mediumnat=dotted line/squares, Deepnat=dashed line/triangles) in Heath.

182x215mm (300 x 300 DPI)

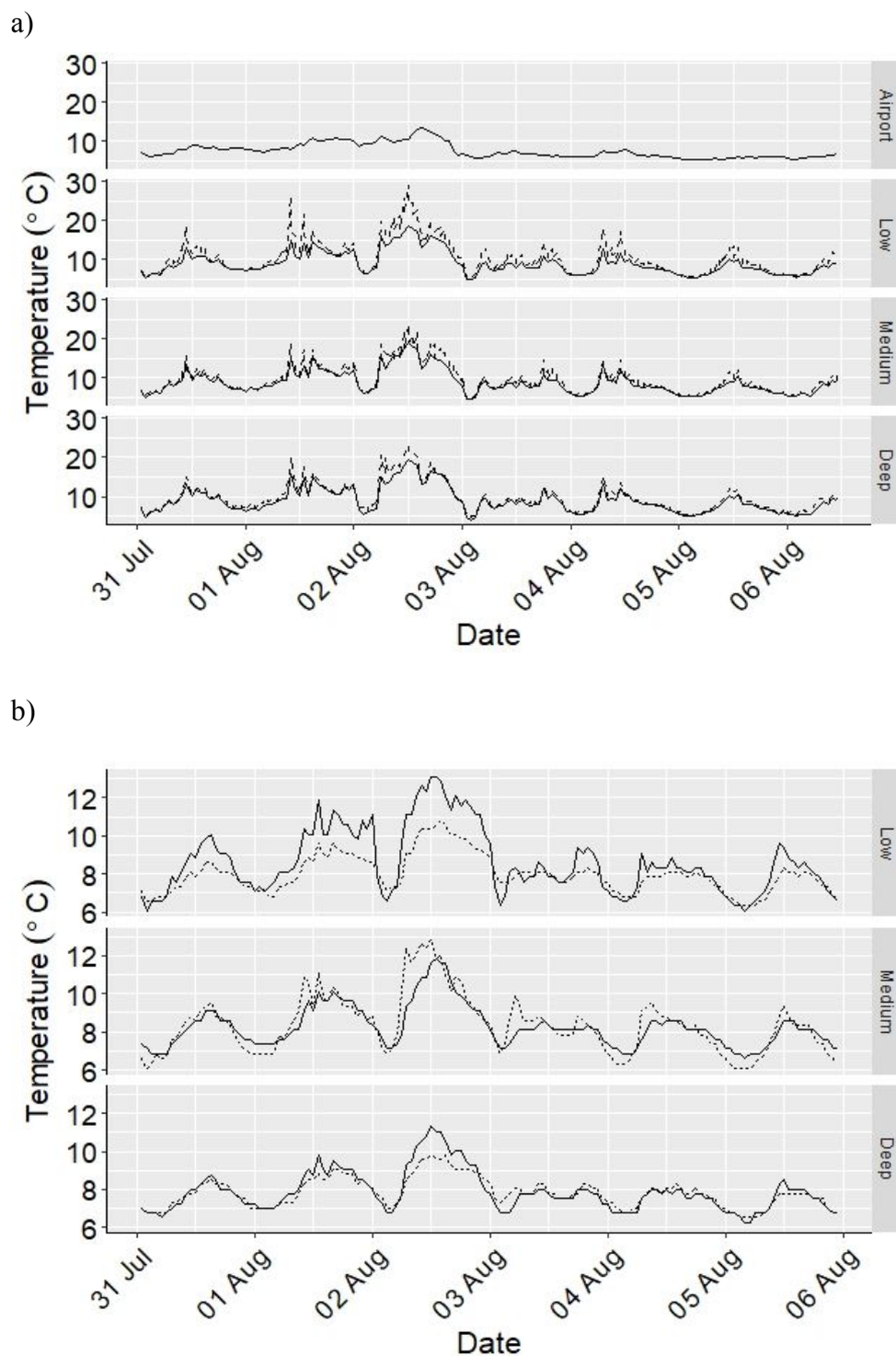
Appendix

Onset of autumn senescence in High Arctic plants show similar patterns in natural and experimental snow depth gradients.

Friederike Gehrman^{1,2}, Camille Ziegler^{3,4}, Elisabeth J. Cooper⁵

Appendix A

Figure A1. a) Air temperature and b) soil temperature measured during the period 31st July to 6th August in control plots (solid line) and OTC chambers (dashed lines) along a natural snow depth gradient with Low, Medium or Deep snow regimes, as well as the temperature measured at the nearest weather station (Airport) at Svalbard airport.



Appendix B

Method

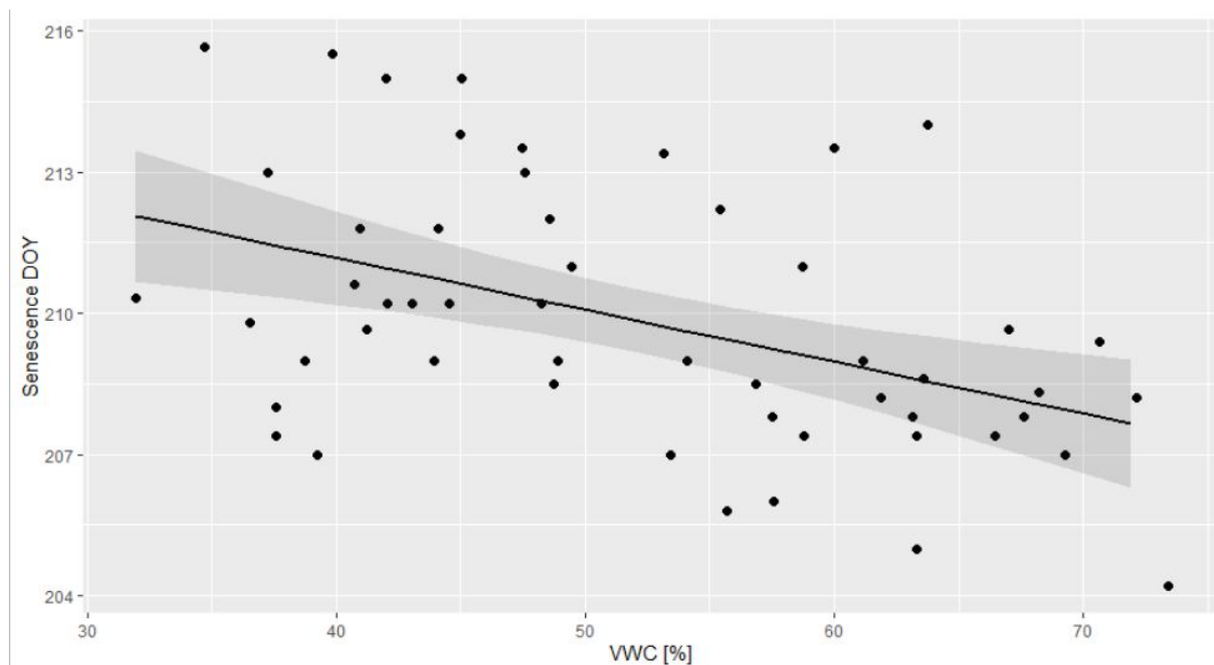
We additionally examined our data for the effect of soil moisture on the DOY of senescence. This was done by first calculating the mean soil moisture (VWC, %) of the whole study period (19th July to 6th August 2016) for each plot ("SM_Season") as a representation of the relative wetness of each plot. Then, DOY was modelled as a function of Snow, Vegetation and SM_Season in a linear mixed-effects model with Site and Plot as random effects. The model was fitted for each species and each gradient type (natural or experimental) separately.

Results

The effect of mean soil moisture on the DOY was only significant in *S. polaris*. In the experimental gradient, higher soil moisture during the study period was correlated with an earlier DOY (Fig. A2, Table A2). In the natural gradient, the correlation was the opposite (Fig. A2, Table A2). Additionally, individuals in Meadow senesced overall 1.4 days earlier than in Heath and 1.9 days earlier in Meadow-Medium than in Meadow-Low.

Figure B1. The mean day of year (DOY) of the onset of senescence in *S. polaris* in an a) experimental and a b) natural snowmelt gradient in two different Vegetation types (Heath, Meadow) and plots with differing mean autumn soil moisture (SM_Season). Only significant relationships between DOY and explanatory variables are displayed. Mean autumn soil moisture was calculated as the mean soil moisture during the study period for each individual plot.

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b)

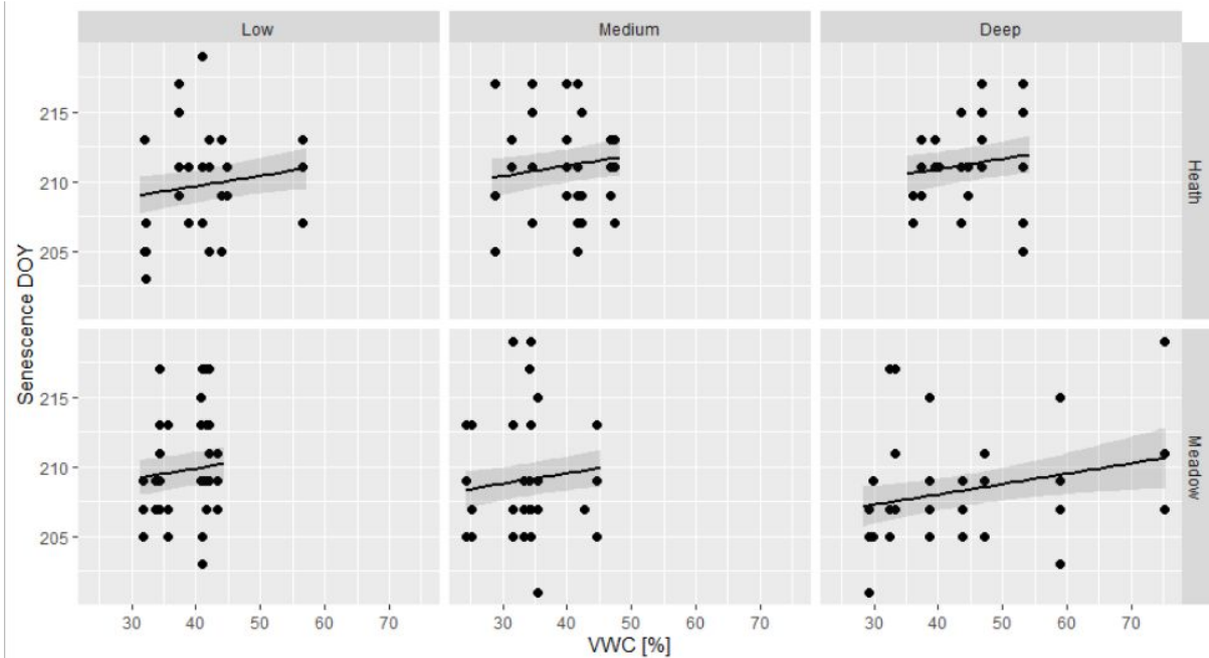


Table B1. Analysis of variances results of linear mixed-effects models testing the effect of mean soil moisture during the study period (SM_Season), snow regime (Snow) and vegetation type (Vegetation) on the DOY of senescence in *S. polaris* in a natural and an experimental snowmelt gradient.

	SM_Season	Snow	Vegetation	Snow*Vegetation
Natural gradient	$F_{1,43}=6.8, p=0.012$	n.s.	$F_{1,4}=8.1, p=0.047$	$F_{2,43}=3.4, p=0.0437$
Experimental Gradient	$F_{1,46}=13.1, p<0.001$	n.s.	n.s.	n.s.