2 3 4

5

6

7

8

11

12

13

14

15

17

18

19

20

21

Arctic Science Downloaded from cdnsciencepub.com by 86.158.198.15 on 05/21/21 For personal use only. This Just-IN manuscript is the accepted manuscript prior to copy editing and page composition. It may differ from the final official version of record.

Intra-specific variation in phenology offers resilience to climate change for *Eriophorum vaginatum*

Thomas C. Parker^{1*}, Steven L. Unger², Michael L. Moody³, Jianwu Tang⁴ & Ned Fetcher⁵

¹Biological and Environmental Sciences, School of Natural Sciences, University of Stirling, Stirling, FK9 4LA UK.

9 Email: t.c.parker@stir.ac.uk

10 *Corresponding Author

²Florida International University, 11200 SW 8th Street, Miami FL 33199, USA

Email: sunge001@fiu.edu

³Biological Sciences, University of Texas at El Paso, El Paso, TX, USA

16 Email: mlmoody@utep.edu

⁴The Ecosystems Center, Marine Biological Laboratory, Woods Hole, MA 02543, USA

Email: jtang@mbl.edu

⁵Institute for Environmental Science and Sustainability, Wilkes University, Wilkes-Barre, PA

22 18766, USA

23 Email: ned.fetcher@wilkes.edu

Publisher policy allows this work to be made available in this repository. The original publication is available at: https://doi.org/10.1139/AS-2020-0039

Arctic Science Downloaded from cdnsciencepub.com by 86.158.198.15 on 05/21/21 personal use only. This Just-IN manuscript is the accepted manuscript prior to copy editing and page composition. It may differ from the final official version of record. For 1

24 <u>Abstract:</u>

25 Phenology of arctic plants is an important determinant of the pattern of carbon uptake and may 26 be highly sensitive to continued rapid climate change. *Eriophorum vaginatum* has a 27 disproportionate influence over ecosystem processes in moist acidic tundra, but it is unclear 28 whether its growth and phenology will remain competitive in the future. We asked whether 29 northern tundra ecotypes of E. vaginatum could extend their growing season in response to direct 30 warming and transplanting into southern ecosystems. At the same time, we asked whether 31 southern ecotypes could adjust their growth patterns in order to thrive further north, should they 32 disperse quickly enough. Detailed phenology measurements across three reciprocal transplant 33 gardens and two years showed that some northern ecotypes were capable of growing for longer 34 when conditions were favourable, but their biomass and growing season length was still shorter 35 than the southern ecotype. Southern ecotypes retained large leaf length when transplanted north and mirrored the growing season length better than the others, mainly due to immediate green-up 36 37 after snowmelt. All ecotypes retained the same senescence timing, regardless of environment, indicating a strong genetic control. E. vaginatum may remain competitive in a warming world if 38 39 southern ecotypes can migrate north.

41

42

40

Key Words: Phenology, Tundra, Eriophorum vaginatum, Ecotype, Climate Change

Page 3 of 34

55

43 Introduction

44 The Arctic is warming at twice the global average, resulting in profound changes in not only 45 temperature but also precipitation and growing season lengths (Mudryk et al. 2019). Biological 46 processes in the Arctic are closely tuned to environmental cues and as such are showing signals 47 of change in response to a changing climate (Post et al. 2009). This is important because living 48 organisms hold critical control over biogeochemical, energy, and hydrological fluxes with huge 49 potential to further exacerbate climate change (Wookey et al. 2009). Plant communities across 50 the Arctic have shown particularly striking changes in response to warming as they grow taller 51 (Bjorkman et al. 2018), increase cover, and undergo shifts in dominance with mosses often in 52 decline and deciduous shrubs in ascendance (Elmendorf et al. 2012b). Natural observations of 53 change are supported by experimental evidence that shows that there are clear winners and losers 54 in the plant community as the climate continues to change (Elmendorf et al. 2012a).

56 The direct effects of warming on arctic plant community composition and growth have been 57 well studied through a circumpolar network of open-top chamber (OTC) experiments 58 (Elmendorf et al. 2012a). These generally show that plant growth increases with warming and 59 that deciduous shrubs increase in dominance but also that responses are mediated by site conditions such as local climate and soil moisture (Elmendorf et al. 2012a). Wider observation 60 networks are detecting 'greening' signals with increases in height and cover at the plot level 61 62 (Bjorkman et al. 2018) and increases in Normalised Difference Vegetation Index (NDVI) at 63 satellite levels (Epstein et al. 2012). Observations of the expansion of deciduous shrub cover are 64 consistent with these trends (Myers-Smith et al. 2011). One of the key findings is that certain 65 groups in the community such as mosses decrease in cover as the community responds to 66 warming, while the response of other groups such as sedges is mixed (Elmendorf et al. 2012a). It 67 is important to understand how all constituents of the plant community will change in the future

because they all contribute significantly to ecosystem processes such as primary productivity,
reflectance, and phenology, among others (Myers-Smith et al. 2019).

70

71 Climate change in the Arctic is multifaceted and will affect aspects of plant performance in 72 different ways (Post et al. 2009; Box et al. 2019). For example, summer growing seasons are extending in the Arctic due to reductions in snow cover duration (SCD) (Box et al. 2019). Model 73 74 projections indicate that SCD over much of the Arctic will decline by about 10-30% by the end of this century as a consequence of delayed onset of snow cover as well as earlier snowmelt 75 76 (Brown et al. 2017). The projected decrease in SCD implies that the potential growing season should lengthen, as found by Park et al. (2016), who used the normalized difference vegetation 77 index (NDVI) to analyze changes in growing season length in boreal and arctic vegetation. 78 79 Broadly speaking, plant phenology in the Arctic has been shown to be sensitive to abiotic 80 conditions (Assmann et al. 2019; Prevéy et al. 2017). At the beginning of the growing season, earlier snowmelt should result in earlier green-up, as abundant sunshine and the disappearance of 81 82 snow produces good growing conditions. Many studies have documented the importance of 83 snowmelt timing for controlling the phenology of arctic plants with earlier snowmelt, which usually results in earlier onset of growth (Høye et al. 2007; Bjorkman et al. 2015; Khorsand Rosa 84 et al. 2015; Semenchuk et al. 2016; May et al. 2020). Once the growing season is underway, it is 85 less clear whether higher average temperatures will affect plant phenology in part because of 86 87 interactions with snowmelt timing (Oberbauer et al. 2013). Geographical patterns in phenology 88 further complicate the response of arctic plants to climate change. Across the Arctic, phenology 89 of plants from more northern sites exhibited greater sensitivity to warming temperatures than 90 plants from sites at more southern latitudes (Prevéy et al. 2017).

91

92 Increasing temperatures in autumn (Box et al. 2019) may offer an opportunity to plant 93 communities to grow for longer, but it is difficult to forecast the effect of mid- and late-season growing conditions on phenology in the autumn. If autumn temperatures increase, it is not clear 94 95 that arctic plants will respond by extending their growing season (Parker et al. 2017). Many 96 species start to turn yellow in August when temperatures are still warm (Shaver and Laundre 97 1997). This may be because, in the Arctic, harsh winter conditions may appear suddenly, which 98 could result in the loss of valuable resources through frost damage to live aboveground biomass 99 that hasn't fully senesced. Some functional groups, notably some graminoids, may be able to 100 delay senescence in response to warming conditions, while other functional groups may have 101 fixed leaf life spans which are correlated with average growing season lengths (Oberbauer et al. 102 2013). Manipulation of the timing of green-up by removing snow or adding it with snow fences 103 has shown that the length of phenological stages such as growth, flowering, or seed setting 104 remained invariant even though the dates of start-up varied greatly (Khorsand Rosa et al. 2015; Semenchuk et al. 2016). Semenchuk et al. (2016) concluded that a range of herbaceous and shrub 105 106 species in their study are periodic, meaning that the duration of phenological periods is 107 genetically fixed. By extension, therefore, even if the end of season environment is suitable for 108 continued growth, tundra plants may senesce early if their green-up was early.

While many studies have focused on variation at the species and community level of organization, few studies have looked at intraspecific variation in phenology of tundra plants. Since most arctic plants have widespread distributions, local adaptations are likely to be important for many species (Linhart and Grant 1996). Local adaptation is widespread in plant populations, especially those with many individuals covering a wide geographic range (Leimu and Fischer 2008; Hereford et al. 2009). Wagner and Simons (2009) reported differences between arctic and alpine populations in phenology of the annual *Koenigia islandica*, where the

117 arctic population flowered earlier than the alpine population. Bjorkman et al. (2017) reported that southern populations of the arctic plants Oxvria digyna and Papaver radicatum were slower to 118 119 leaf out and to initiate senescence than northern (local) populations. Likewise, Parker et al. 120 (2017) showed that senescence of *Eriophorum vaginatum* grown in a common garden occurs 121 later for populations from the southern portions of a latitudinal gradient in the Alaskan Arctic. 122 Although growth rates were the same, the southern populations were able to accumulate more 123 biomass because of the longer growing season (Parker et al. 2017). Thus, it is important to base models of phenology on not only a generalized phenotype but also to consider the variation 124 125 within species across their range where local dynamics may vary, although the assemblage remains the same. 126

128 Many arctic plant species are distributed along the latitudinal gradient from Low to High 129 Arctic, which provides ample scope for locally adapted populations or ecotypes. Strong adaptation to local climates may render arctic plants vulnerable to rapid climate change in their 130 131 locales if they are not able to respond quickly enough (McGraw et al. 2015). The degree of 132 phenotypic plasticity of ecotypes of arctic plants may determine their potential to take advantage of, or survive, warmer conditions. Eriophorum vaginatum is a foundational species of moist 133 134 acidic tundra, meaning that it strongly dictates the system's physical structure as well as its 135 process rates (Chapin and Shaver 1985). E. vaginatum demonstrates clear ecotypic 136 differentiation in phenotypes (Shaver et al. 1986; Fetcher and Shaver 1990) and gene expression 137 (Mohl et al. 2020) across its South-North distribution in Alaska which reflects a wide range in 138 growing season conditions. McGraw et al. (2015) showed that the optimal environment for 139 tussock survival and tiller population growth in E. vaginatum had shifted northwards, meaning 140 that this important species may suffer from 'adaptational lag' and not keep pace with current 141 rates of climate change. To address the lag in the performance of local populations, they may

127

- Caro	niu.
ر در بر	1110
5	
	16131
- 	CIAI
:550	
5001	111141
4	
fuo m	
1 ff2#	IICI
21/2	ay u
05/	11
5 on	1011.
98.1	IIRUU
58.1	, IIIO
86.]	Jago
m by	4 nin
b.col	ŝ
cepu	
cien	(don
cdns	2) T
rom	n pur
ded f	dine
nload	Iallu
Dow	
nce I	rchr
Scie	IC ar
ctic	n et 1
IA:	, cup
51140	Iallu
ž	
Tangt	-len(
1:1	г . т
90 O	20.01
5.10	lal u
0000	10612
5	54. 5
Ľ	-

142	need to be supplemented by gene flow from the south (McGraw et al. 2015). Performance of the
143	northern ecotypes of E. vaginatum is less flexible than the southern ecotypes in both net
144	ecosystem exchange (NEE) (Curasi et al. 2019) and leaf growth (Fetcher and Shaver 1990). But
145	as previously stated, changes in growing season length offer new opportunities to grow for
146	longer and remain competitive in their environment.
147	
148	Here we investigate the role of genetic background and environmental conditions as they
149	affect the phenology of E. vaginatum growing in a reciprocal transplant experiment in northern
150	Alaska. We use this system to ask:
151	1. Can the phenology of <i>E. vaginatum</i> ecotypes match growing conditions when
152	transplanted into warmer ecosystems with longer growing seasons?
153	2. Do southern populations retain their growth patterns when transplanted north?
154	3. Do local ecotypes increase growth and growing season length when experimentally
155	warmed <i>in situ</i> ?
156	4. Does <i>E. vaginatum</i> exhibit a fixed periodicity in its phenology, i.e. if it starts growing
157	early will it senesce early?
158	
159	
160	Materials and methods.
161	Site description and experimental design.
162	Eriophorum vaginatum L. (Cyperaceae) is a tussock-forming sedge that has a strong
163	influence on tundra microclimate and carbon cycling potential (Chapin et al. 1979, Curasi et al.
164	submitted). It covers large areas of northeastern Siberia (Walker et al. 2005) and is also found in

165 wetlands and moorlands throughout the circumpolar region (Wein 1973). In Alaska, full-sized

adult tussocks can consist of 300–600 live tillers (Fetcher and Shaver 1982). Tussocks can live
for well over 100 years (Mark et al. 1985) and can vary widely in size (Shaver et al. 1986).

Three common gardens of reciprocally transplanted tussocks of E. vaginatum were 169 170 established at Sagwon (SG; 69.42°N, 148.72°W, elev. 300 m), Toolik Lake (TL; 68.63°N, 149.36°W, elev. 760 m) and Coldfoot (CF; 67.26°N, 150.17°W, elev. 331 m) along the Dalton 171 Highway in Alaska, USA. CF is approximately 4 °C warmer than the other sites during the 172 summer months of June and July and average temperature stays above freezing for 2 more 173 174 months during Spring and Autumn, resulting in more thawing degree days (Fig S1, Supplementary Figure S2). Although SG is further north, it is at a lower elevation than TL, 175 176 resulting in similar overall temperature regimes (Fig S1, Supplementary Figure S2). Tussocks of E. vaginatum dominate all three sites with deciduous (Betula nana L., Salix spp., and Vaccinium 177 uliginosum L.) and evergreen shrubs (Vaccinium vitis-idaea L, Rhododendron tomentosum 178 179 Harmaja), mosses, and lichens growing in between the tussocks. The northern ecotypes of E. 180 vaginatum are found at Sagwon, which is on the northern edge of moist acidic tundra and may 181 not have been glaciated during the Pleistocene, and at Toolik Lake, which is in moist acidic 182 tundra near the Brooks Range and was most recently glaciated in the Late Wisconsinian (~20,000 yr BP) (Hamilton 2003, Kaufman and Manley 2004, Kaufman et al. 2011, Walker et al. 183 184 2005). One of the southern ecotypes is found at Coldfoot, which is in muskeg with encroaching 185 trees (Picea mariana (Mill.) Britton, Sterns & Poggenb.) that were not present in 1982 when 186 previous common gardens were established (Shaver et al. 1986). Coldfoot was glaciated during 187 the Early Wisconsinian (~70,000 - ~40,000 yr BP), but probably not during the Late 188 Wisconsinian (Kaufman and Manley 2004, Kaufman et al. 2011). The three sites were likely 189 colonised by E. vaginatum at different times and therefore were differentiating as ecotypes for

different amounts of time, nonetheless they have all had at least 20,000 years to potentiallydevelop traits that reflect their home environments.

192

193 In August 2014 mature tussocks were transplanted between the three sites with tussocks 194 from each home site transplanted into their home site to act as controls according to methods specified in Bennington et al. (2012) and Schedlbauer et al. (2018). Briefly, a serrated knife was 195 196 used to sever the rhizomes from roots and soil at a tussock's base and remove it from the tundra. 197 Tussocks were then placed in the vacant positions at the common garden where local tussocks 198 had been removed. This method has a high success rate because of E. vaginatum's deciduous 199 rooting habit; although roots are severed during transplanting, new roots grow in each 200 subsequent year, restoring full root function (Bennington et al. 2012). Tussocks were planted in 201 clusters of three, approximately 0.5 m apart from each other. Clusters were paired at SG and TL 202 where one cluster of each pair was passively warmed using open-top chambers (OTC) 203 (Schedlbauer et al. 2018). Ten pairs of clusters of the three populations were arranged in an 204 approximately 25 m x 30 m grid. Open-top chambers were placed on the selected clusters from 205 11 July until 28 August in 2015, from 2 June until 28 August in 2016, and from 30 May until 206 26 August in 2017 causing a mean hourly air temperature increase of 1.16°C and 1.04°C at 207 Sagwon in 2016 and 2017, respectively. At Toolik Lake, the respective temperature increases 208 were 0.60°C and 1.01°C. At CF, where there was no warming treatment, clusters were arranged 209 as singletons in a smaller grid (25 m x 15 m). At each site, 10 non-transplanted tussocks were 210 identified next to the transplant garden in order to assess the effect of transplanting on measured 211 response variables.

212 Leaf Measurements

Through the growing seasons of 2016 and 2017 (early June - mid-September), leaf growth and senescence were monitored on transplanted tussocks. Growing season air temperatures at each transplant garden during the measurement years were representative of typical climatic
conditions of each site (Supplementary Table S1). A tiller from one tussock of each cluster was
tagged and monitored according to Shaver and Laundre (1997) and Parker et al. (2017). A small
zip tie was secured around the base of the tiller, so as to include all leaves with any visible green
portions while excluding any previously senesced leaves from previous growth. The total leaf
length and the length of the green portions were measured to the nearest 5 mm approximately
once a week for each leaf in a tiller, from oldest to youngest.

222 Tiller Phenology Data Processing

The senesced portions of leaves were fragile and sometimes broke off; since this occurred 223 224 after leaves had reached their full length, the total length was corrected to match the last 225 measurement of the unbroken leaf. Where lengths of single leaves were missing for a time point 226 due to human error, they were replaced with the mean of the previous and following time points. 227 Only leaves that were growing during the season of measurement were measured, thereby excluding leaves that were grown in the previous year and were senescing as well as leaves that 228 229 had been initiated for the next year but were not elongating. A double logistic phenology model 230 (Busetto et al. 2010) was fit using non-linear least squares regression to green leaf growth pattern 231 over the growing season on every tiller in each year (See Fig.1 for example fits):

$$G(d) = (G_{Min} + (G_{Max} - G_{Min})) \left(\frac{1}{1 + e^{-mS(d-S)}} + \frac{1}{1 + e^{-mA(d-A)}} - 1\right)$$
 Equation 1

where G(t) is the green leaf length (cm) at day of the year (d), G_{Max} is the maximum green leaf length observed, G_{Min} is the minimum green leaf length over the year (here set to 1 cm because *E. vaginatum* retains a small amount of green biomass over winter (Shaver & Laundre, 1997)), *mS* is the spring growth rate, and *mA* is the autumn senescence rate at time-points *S* and *A*, which are found halfway on the increase and decrease curves, respectively.

232

238 Phenology metrics specified by Busetto et al. (2010) as significant points on the phenology 239 curve were extracted from each curve (Fig. 1). S1, S5, A1, and A5 are the time points at which 240 changes in curvature are at their maximum or minimum (Busetto et al. 2010). S2, S4, A2, and 241 A4 are dates at which the double logistic curve transitions from linear to non-linear (or vice 242 versa), and S3 and A3 are the points of maximum increase or decrease of the curve (Busetto et al. 2010). The tiller growing season (S1A1) was calculated as the number of days between 243 244 metrics S1 and A1, which represents the period between the beginning of peak growth rate and the end of peak biomass (before senescence) and therefore when the majority of primary 245 246 productivity takes place.

Poorly fitting models for individual tillers were removed from the dataset if they made 247 248 biologically unrealistic estimates of Spring (onset of growth (S1) before April 1st, growth rate 249 (mS) above 0.4 cm day ⁻¹) and peak growing season (S5A1) phenology (metric A1-S5 less than 250 0). Additionally, if the any phenology model had a particularly poor fit to the extent that it was an outlier compared to other fit models (RMSE higher than 95 % percentile of all model fits), it 251 252 was discarded. After this process, 130 curves from the three gardens could be analyzed in 2016 253 and 113 in 2017 (20 total removed). The curves were split relatively evenly between populations 254 (SG, TL, or CF), sites (SG, TL, or CF) and treatments (OTC or control), resulting in even 255 replication across all combinations (Supplementary Figure S1).



Figure 1: Example of double logistic model (Equation 1) fit to the growth pattern of a tiller over a growing season and metrics that can be calculated from this curve (in red, left panel) and fit to two other example datasets (in black, right panel).

262 Environmental data processing

263 Air temperatures for Coldfoot and Sagwon were extracted and calculated from daily average 264 data from the SNOTEL database (http://www.wcc.nrcs.usda.gov/snow/) and from the Toolik Field Station Environmental Data Center (EDC) of the University of Alaska, Fairbanks 265 266 (Environmental Data Center Team) for the Toolik Lake site. Snowmelt timing was extracted 267 from the SNOWTEL database for Sagwon and Coldfoot and from the Environmental Data 268 Center for the Toolik site (Environmental Data Center, University of Alaska, Fairbanks; 269 https://toolik.alaska.edu/edc/index.php). The end of the growing season was defined as the first 270 day in autumn that the prior seven-day running average minimum daily air temperature returned 271 to 1°C; consequently, the potential growing season length was determined as the number of days 272 between snowmelt and a return to consistently low temperatures, We used a seven-day running

average temperature because cold-snaps can happen at any time in the season and we chose 1 °C because the seven-day running average of 0 °C did not occur until long after all plant activity had ceased (October). Note that this was the authors' judgement of a 'potential growing season length' for the purpose of this paper, to our knowledge there is no recognised definition in this system. Late season temperature was defined as the average air temperature at each site between 1 August and 14 August in any given year. This was deemed a period of time when plants are green but potentially receptive to phenological cues for senescence.

280 Statistical Analysis

281 Linear mixed effects models (Pinheiro et al. 2017) were used to test whether phenology 282 variables (onset of growing season (S1), onset of senescence (A1), and growing season length (S1-A1)) were significantly affected by fixed effects: population source, common garden site, or 283 sampling year using the 'nlme' package in R (R Development Core Team, 2016, Pinheiro et al. 284 285 2017). The tussock ID was used as a random intercept term. Models were simplified by removing interaction terms if they did not have a significant effect in order to get best estimates 286 of main fixed effects (Crawley 2007). The effect of each factor in the final model was assessed 287 288 relative to the null model (intercept only) by ANOVA (Crawley 2007). Linear mixed effects 289 models (tussock ID as random intercept term) were used to assess the effect of population and 290 environmental factors: potential growing season length and the effect of snowmelt date and late 291 season temperature on the onset of growth, and onset of senescence, respectively. The number of 292 days between the onset of growth and onset of senescence was used to determine actual growing 293 season length. All analyses were carried out with R v3.3.3 (R Development Core Team, 2016).

294

Results

Across all populations, tillers of *E. vaginatum* initiated growth earlier at Coldfoot than the other two sites (Table 1, Fig. 2) but there was no significant difference between populations across all gardens (P = 0.195, Table 1). However, the CF population responded to differences in site growing conditions more than the other populations, resulting in a significant interaction between populations and site (Table 1). The CF population started to senesce later than the northern ecotypes as represented by the TL and SG populations at all of the sites (Fig. 2). But the onset of growth at Toolik Lake and Sagwon was significantly delayed after snowmelt in 2016, which had low temperatures in early June (Table 1, Supplementary Figure S1). Thus, early June temperatures appeared to exert some control on the initiation of growth.



Figure 2: Summary of growing season of Coldfoot (yellow) Toolik Lake (green) and
Sagwon (blue) populations across all three common gardens and over two years. Points on
the left signify mean (+/- 1 standard error) onset of growing season (metric S1) and points
on the right signify mean (+/- 1 standard error) onset of senescence (metric A1). The
number of days between these (colored line) signifies the length of the growing season
(metric A1S1). Statistics testing the effect of population, transplant garden (site) and year
are found in Table 1, for model details, see Supplementary Table S5.

Page 15 of 34

Arctic Science Downloaded from cdnsciencepub.com by 86.158.198.15 on 05/21/21 For personal use only. This Just-IN manuscript is the accepted manuscript prior to copy editing and page composition. It may differ from the final official version of record.

314	Table 1: Test statistics from linear mixed effects models showing the effect of fixed factors
315	on growth patterns in <i>E. vaginatum</i> . Data are divided into an analysis of transplant only
316	tussocks (no warming treatment, three common gardens) and an analysis that includes the
317	effect of warming with open-top chambers (Sagwon and Toolik sites only). For model
318	details, see Supplementary Table S5.

details, see Supplementary Table S5.

Non-warmed transplanted tussocks

Response variable Fixed effect			d.f	F	Р
S1 (Onset of Growth)	Population	2,	81	1.7	0.195
	Site	2,	81	21.6	< 0.001
	Year	1,	57	7.0	0.011
	Population x Site	4,	81	4.0	0.005
A1 (Onset of Senescence)	Population	2,	85	32.4	< 0.001
	Site	2,	85	3.4	0.038
	Year	1,	57	40.7	< 0.001
S1A1 (Growing Season)	Population	2,	81	12.5	< 0.001
	Site	2,	81	13.1	< 0.001
	Year	1,	57	42.3	< 0.001
	Population x Site	4,	81	3.7	0.008
Maximum Green length	Population	2,	87	19.2	< 0.001
	Site	2,	87	10.1	< 0.001
	Year	1,	68	33.9	< 0.001

Warmed and non-warmed transplant tussocks

Response variable	Fixed effect	d.f		F	Р
A1 (Onset of Senescence)	Population	2,	112	18.4	< 0.001
	Site	1,	112	0.4	0.552
	Year	1,	79	58.4	< 0.001
	Warming	1,	112	2.4	0.125
S1A1 (Growing Season)	Population	2,	112	10.6	< 0.001
	Site	1,	112	3.7	0.056
	Year	1,	79	38.3	< 0.001
	Warming	1,	112	6.0	0.016
Maximum Green length	Population	2,	115	13.4	< 0.001
	Site	1,	115	10.8	0.001
	Year	1,	91	38.3	< 0.001

1.8

0.184

1,

Warming

Arctic Science Downloaded from cdnsciencepub.com by 86.158.198.15 on 05/21/21 For personal use only. This Just-IN manuscript is the accepted manuscript prior to copy editing and page composition. It may differ from the final official version of record.

319

The southern ecotype had significantly longer leaves than the northern ecotypes (P < 0.001, Table 1), although this difference was less pronounced at Toolik Lake than at the other two sites (Table 1, Supplementary Figure S3). Warming with OTCs had no effect on leaf length (Table 1), but on average, over the two years, it did result in a significantly (P < 0.05) longer growing season, defined by the number of days between S1 and A1 (Table 1, Fig. 2). Warming did not affect spring phenology or autumn phenology in a statistically detectable way, but the combined effects may have increased the overall season length slightly.

327 The effect of transplanting was analysed by comparing tussocks that were transplanted into 328 their 'home' site with non-transplanted 'control' tussocks. Across all sites transplanting did not 329 affect onset of growth (metric S1 (Supplementary Figure S4, Supplementary Figure S5, Supplementary Table S3)) but on average make onset of senescence marginally earlier (metric 330 A1 (P = 0.066, Supplementary Figure S4, Supplementary Figure S5, Supplementary Table S3)), 331 332 although this pattern was not consistent and depended on population. Growing season length was 333 not affected by transplanting (metric S1A1 (Supplementary Figure S4, Supplementary Figure S5, 334 Supplementary Table S3)), but transplanting did significantly reduce maximum green length compared to non-transplanted controls (P = 0.01, Supplementary Figure S4, Supplementary 335 336 Figure S5, Supplementary Table S3).

There was no significant relationship between the date of growth onset and the date of senescence onset across populations and no interaction between onset of growth and population (Fig. 3, Supplementary Table S4). CF populations consistently senesced later than the others, but this was unrelated to the onset of growth. Over the whole growing season, the actual growing season of leaves (Metric S1A1) of all populations was positively affected by potential growing length but the CF tussocks responded particularly strongly. This resulted in a statistically significant effect of potential growing season, population origin (CF was highest on average),

344 and an interaction between the two (Fig. 4, Supplementary Table S4). The onset of growth in 345 spring was positively related to the day of snowmelt across all populations but the CF population 346 was particularly responsive, with initiation of growth closely tracking the loss of snow at any 347 given site (Fig. 5, Supplementary Table S4). In autumn, none of the populations in either year 348 were responsive to differences in late season environmental conditions, in this case, temperature 349 in the first half of August. Instead, the populations maintained a significant difference in 350 senescence timing regardless of the garden they were present in with CF senescing particularly late (Fig. 6, Supplementary Table S4). There were significant differences between years with 351 352 most tussocks senescing later in 2017 than 2016.





356 vaginatum across all gardens populations in 2016 (squares) and 2017 (triangles). A linear

mixed effects model (Supplementary Table S4) showed no effect of onset of growth on onset of senescence (P = 0.388) but significant effect of population (P < 0.001). Modelled marginal effects (with 95 % confidence intervals) of onset of growth and population on the onset of senescence are displayed, for model details, see Supplementary Table S5.



Potential growing season (Days)

Figure 4: Relationship between the potential growing season (snowmelt - return of cold
temperatures) and the actual growing season (S1A1) for three populations in 2016
(squares) and 2017 (triangles). See Supplementary Table S4 for linear mixed effects models
showing a significant effect of population (P < 0.001), a significant positive effect of
potential growing season (P < 0.001) and a significant interaction between the two (P =
0.002). Modelled marginal effects (with 95 % confidence intervals) of potential growing

362

369 season and population on actual growing season are displayed, for model details, see

370 Supplementary Table S5.

371



372

Arctic Science Downloaded from cdnsciencepub.com by 86.158.198.15 on 05/21/21 For personal use only. This Just-IN manuscript is the accepted manuscript prior to copy editing and page composition. It may differ from the final official version of record.





Figure 6: Relationship between late season temperature and onset of senescence (A1) for three populations in 2016 (squares) and 2017 (triangles). See Supplementary Table S4 for linear mixed effects models showing a significant effect of population (*P* < 0.001) but no effect late season temperature (P = 0.575). Modelled marginal effects (with 95 % confidence intervals) of late season temperature and population on onset of senescence are displayed, for model details, see Supplementary Table S5.

389 <u>Discussion</u>

390 Response of northern ecotypes to warming

391 Climate change is progressing rapidly in arctic ecosystems so it is essential for tundra 392 plants, which are adapted to cold environments, to respond in kind. For the foundational species 393 of moist acidic tundra, Eriophorum vaginatum, there is evidence that these long-lived species are 394 already growing outside of their optimal climates (McGraw et al. 2015). To better understand 395 how this species will respond to climate change, we measured phenology and growth in reciprocal transplant experiment combined with warming using OTCs. Firstly, we asked whether 396 397 the phenology of the northern ecotypes can match longer growing seasons when transplanted 398 south. The Sagwon population did grow for longer when transplanted south to Coldfoot, 399 managing to take advantage of earlier snowmelt, followed by warm temperatures. Although the 400 length of the growing season of the Toolik population did not change when moved southward, 401 the initiation of growth following snowmelt was earlier in 2017 compared to its home site. When 402 experimentally warmed with OTCs, there was a general pattern across all populations to slightly 403 increase their growing season length, but they did not grow any larger. Taken together, 404 phenology of the northern ecotypes showed some responsiveness to climate change simulation, 405 but the effects were mixed and relatively small. This is consistent with the lack of change in tiller size when northern populations from Sagwon, Toolik Lake, and Prudhoe Bay were moved south 406 407 in an earlier experiment (Fetcher and Shaver 1990, Souther et al. 2014).

408

409 Effects of transplanting on phenology of Eriophorum vaginatum

By comparing phenology leaf growth of tussocks transplanted into their 'home' site with non-transplanted tussocks, we show that there is a minimal effect of the physical disturbance on measured phenology traits (Supplementary Figure S4, Supplementary Figure S5). This is an effect that often assumed in such experiments (Parker et al. 2017, Curasi et al. 2018, Walker et Arctic Science Downloaded from cdnsciencepub.com by 86.158.198.15 on 05/21/21 For personal use only. This Just-IN manuscript is the accepted manuscript prior to copy editing and page composition. It may differ from the final official version of record.

424

414 al. 2018) but rarely tested. One of the advantages of E. vaginatum for reciprocal transplants is its 415 deciduous root system and lack of mycorrhizal symbiosis, which means that tussocks can be 416 transplanted without disturbing the belowground environment (Shaver et al 1986, Parker et al. 417 2017, Schedlbauer et al. 2108). To this feature, we can now add the relative lack of response of 418 phenology to transplanting. The timing of onset of growth and the length of the growing season 419 was not different. The timing of senescence was overall marginally earlier, and the maximum 420 length of leaves was slightly shorter which perhaps reflects a less well established connection between rhizomes and the soil resulting in less effective nutrient uptake. These data further 421 422 underline the usefulness of E. vaginatum as a model species to study ecotypic variation of traits 423 in mature plants.

425 Tussock competitiveness in a warming tundra

426 Eriophorum vaginatum will need to contend with changing temperature regimes as well as increasing competition from plant functional types that respond well under warming. The 427 428 Arctic is warming rapidly (Mudryk et al. 2019) and there is only limited evidence as to how well 429 E. vaginatum will fare in these warmer conditions. Parker et al. (2017) were not able to detect a response to simulated warming but Sullivan and Welker (2005) showed that warming to similar 430 431 levels as in our experiments initiated early season growth of E. vaginatum in the tundra. Our 432 study showed that tussocks across all populations stayed green for 3.76 days longer in response 433 to direct warming but little else was responsive, therefore it is not clear whether the response of 434 *E. vaginatum* to a gradual temperature increase will have tangible ecosystem effects. OTC 435 experiments have recently shown that tundra plant communities (including moist acidic tundra, 436 dominated by *E. vaginatum*) extend their growing season when warmed (May et al. 2020). This 437 suggests that contemporary plant communities can take advantage of milder growing conditions, 438 at least in the short term. In the long term, however, the future success of E. vaginatum may

439 depend more on the performance of its fellow community members than on its own. Many areas 440 of tundra are becoming more productive and taller in response to climate change (Bjorkman et al. 441 2018) and deciduous shrubs are often the plants that increase growth the most as climate warms 442 (Elmendorf et al. 2012a). If deciduous shrubs overgrow tussocks, which are more limited in their 443 ability to grow taller, then the foundation species of moist acidic tundra may suffer declines. 444 However, if tussocks remain green for longer in extended growing seasons (Park et al. 2016), 445 and extend beyond that of shrubs, they may retain an important place in northern ecosystems. E. vaginatum remains photosynthetically active as long as it holds green leaves into August (Curasi 446 447 et al. 2019), hence tussocks that can delay senescence may continue to accumulate carbon later into the season, after other species have dropped their leaves. 448

449

450

Plasticity and competitiveness of southern ecotypes

451 If the northern populations of E. vaginatum have only limited potential to respond to 452 climate warming, can tussock tundra be maintained if southern populations or their genes move 453 northward? In our study, the Coldfoot ecotype from the warmer site, south of the treeline (CF) 454 did show plasticity in spring because the timing of green-up varied with the time of snowmelt. At 455 the same time, senescence of the southern ecotype occurred later than that of the northern 456 ecotypes across all environments, resulting in an apparent plasticity of growing season length in 457 the CF ecotype. Parker et al. (2017) showed, using a single common garden in moist acidic 458 tundra, that the southern ecotype grows later into the season and suggested that this trait is driven 459 by adaptation to their warmer home site in the south. We show here that the southern ecotype 460 maintains green leaves on average longer than the northern ecotypes (16 days longer than SG 461 and 9 days longer than TL), regardless of which common garden they are growing in (700 462 thawing degree days difference between CF and SG gardens). Green leaves in E. vaginatum 463 retain active photosynthetic tissue late into August (Curasi et al. 2019), therefore if southern

464 ecotypes can migrate north in sync with climate warming they may increase the fitness of a 465 species which is currently suffering in situ (McGraw et al. 2015). Southern ecotypes grow taller leaves (Fetcher and Shaver 1990) and maintain green tissue later in the season than northern 466 467 ecotypes, therefore they may have a greater capacity for C fixation (Shaver et al. 2007). 468 Dependent on dispersal by wind and suitable ecosystem disturbance for establishment (McGraw 469 et al. 2015), northward migration of southern ecotypes could potentially have ecosystem-level 470 impacts by influencing net ecosystem exchange. This needs to be tested by explicitly considering tussocks (transplanted and non-transplanted) in ecosystem analyses. Further, studies that measure 471 472 ecosystem processes are currently limited by peak season-only measurements (Souther et al. 473 2014, Walker et al. 2018, Curasi et al. 2019), to integrate ecotypes into ecosystem gas exchange, 474 the whole growing season needs to be considered.

475 The pattern of greater spring phenological plasticity in the southernmost E. vaginatum 476 population stands in contrast to results from that of the community-wide large-scale synthesis of 477 phenology (Prevéy et al. 2017). The results from this synthesis extended over 21 degrees of 478 latitude and over 10 degrees further north than our most northern site (SG). At very high 479 latitudes, in the harsh growing conditions of High Arctic desert, the benefit of earlier spring 480 green-up may outweigh the risk of damage by variation in early season weather (cold-snaps) (Prevéy et al. 2017), thus more northern sites had higher plasticity. In the present experiment, the 481 482 southern ecotypes showed more plasticity in the timing of green-up. At Coldfoot, there may be 483 less environmental risk to greening-up as soon as the snow melts, whereas in the tundra there is a 484 high risk that harsh growing conditions will return post snowmelt (Supplementary Table S2, 485 Parker et al. 2017).

486

487 Environmental vs genetic controls on phenology

488 Because snowmelt in the tundra is getting earlier and causing earlier plant green-up (Park 489 et al. 2016; Assmann et al. 2019), one of the next questions is how does this affect biological 490 processes later in the growing season and how does this compare with other important controls 491 on late season phenology? Our Ouestion 4 arose from the hypothesis that some arctic plant 492 species are periodic (Semenchuk et al. 2016), Under this hypothesis, early green-up would result 493 in early senescence due to genetic control over the length of E. vaginatum's growing season. We 494 found no evidence to support this hypothesis. In contrast to other authors (Khorsand Rosa et al. 2015; Semenchuk et al. 2016) we found no relationship between timing of early season and late 495 496 season phenology. Instead, we find that timing of senescence is best predicted by the population 497 origin of the tussock. This is postulated as the result of genetic adaptations to past environmental 498 conditions, which genotype-environment association studies support as a driver in forming 499 population structure conditions and patterns of E. vaginatum in north central Alaska (glaciation 500 (Elizabeth Stunz, pers. comm.). Senescing at the right time is particularly important in the Arctic where the abrupt start of winter can be harsh and damaging to exposed tissues (McGraw and et 501 502 al. 1983). Therefore, it is plausible that genetic control over average timing of the return of cold 503 temperatures at each of the home sites has a role in shaping the observed phenology patterns in 504 the three populations studied.

505 While previous selection pressure clearly has a part to play in shaping contemporary late 506 season phenology, it is important to consider the plastic response of phenology to environmental 507 factors. Cold-snaps and frost can cause senescence in multiple species (McGraw et al. 1983) and 508 a warm late season can delay senescence (May et al. 2017). We observed later senescence at the 509 northern sites in 2017 compared with 2016 when late season temperatures were significantly 510 warmer (Fig. 6) but more years of measurements at the same sites would be needed to start to determine the driving factors behind this variation. Temperatures towards the end of the growing 511 512 season are quite variable, but photoperiod or the quality of light could be a more reliable cue for

Arctic Science Downloaded from cdnsciencepub.com by 86.158.198.15 on 05/21/21 For personal use only. This Just-IN manuscript is the accepted manuscript prior to copy editing and page composition. It may differ from the final official version of record.

513 triggering senescence. The length of photoperiod was shown to be critical for growth cessation in 514 Salix pentandra, with northern populations requiring a shorter dark period to stop growth 515 (Juntilla and Kaurin 1985). This is consistent with the need to cease growing before an early 516 frost occurs, which is more likely at higher latitudes. Another potentially important but equally 517 understudied light cue for senescence in arctic plants is the ratio of red:far red light as monitored 518 by the phytochrome photoreceptors (Buchanan et al. 2015). As discussed by Parker et al. (2017) 519 research into the sensitivity of tundra plants to light quality should remain a priority for research in the future. 520

521

522 Conclusions

523 In the tundra ecosystem dominated by E. vaginatum, the growing season of the northern 524 ecotypes had a limited response to longer, more favorable growing conditions when transplanted 525 south and therefore limited the potential to sequester C during warm shoulder seasons. The Arctic is warming faster than any other biome on Earth (Park et al. 2018) and warming is 526 527 particularly pronounced at the shoulder seasons, resulting in longer growing seasons (Park et al. 528 2016). Our results support a hypothesis that the southern ecotype of E. vaginatum is better 529 adapted to take advantage of this warming through having more plasticity in phenological 530 response, but the northern ecotypes are more constrained, therefore would be unable to take 531 advantage of a lengthened growing season (Fetcher and Shaver 1990). If the Arctic continues to 532 become more productive (Epstein et al. 2012) and grow taller (Bjorkman et al. 2019), locally 533 adapted ecotypes lacking the ability extend growth in height or phenology in the northern part of 534 their range, as found for E. vaginatum, may need to rely on gene flow from southern populations 535 to maintain a competitive balance in the ecosystem. At the present rate of climate change, the 536 northern ecotypes already appear to be poorly adapted if the climate warms to the temperatures 537 currently encountered by the southern ecotype (McGraw et al. 2015). Given the slow growth and

	538	longevity of tussock forming plants in the arctic, the chances for establishment of southern
51 10	539	ecotypes in the north seem unlikely, outside of assisted gene flow or migration (Borrell et al.
	540	2020).
	541	
	542	Acknowledgments:
	543	Thanks to Alana Thurston, Darrell Dech, David Heinz, Elizabeth Fortin, Jon Gewirtzman, Myra
	544	Melendez, Sophia Iglesia, and Stephen Turner for field assistance and to Gus Shaver for advice.
11 11149	545	We would also like to acknowledge the helpful comments of two anonymous reviewers.
.IIOme	546	
-combr	547	Competing interests
iu pago	548	The authors declare there are no competing interests
ung an	549	
upy cu	550	Contributors statement
	551	NF, MLM and JT and TCP designed the experiment and measurement routine, TCP, NF and SU
nd idn	552	took the measurements. All Authors contributed to writing the manuscript.
Sentiali	553	
r pridre	554	Funding
חור מרר	555	Funding for this research was provided through the National Science Foundation (NSF/PLR
er idriv	556	1418010 to NF, NSF/PLR 1417645 to MLM, and NSF/PLR 1417763 to JT). Logistic support
Sentipin	557	came from Toolik Field Station and the Arctic LTER (NSF/PLR 1637459).
- NTT-Jer	558	
r entr	559	Data availability statement
. YIII		

560	Raw leaf length data that underpin the paper are supplied in Supplemental file 2
561	Literature Cited
562 563 564 565	Arft, A. M., Walker, M. D., Gurevitch, J., Alatalo, J. M., Bret-Harte, M. S., Dale, et al. 1999. Responses of tundra plants to experimental warming: Meta-analysis of the International Tundra Experiment. Ecological Monographs, 69(4): 491–511. doi: 10.1890/0012- 9615(1999)069[0491:rotpte]2.0.co;2
566 567 568 569	Assmann, J.J., Myers-Smith, I.H., Phillimore, A.B., Bjorkman, A.D., Ennos, R.E., Prevéy, J.S., et al. 2019. Local snow melt and temperature—but not regional sea ice—explain variation in spring phenology in coastal Arctic tundra. Global Change Biology, 25 : 2258–2274. doi:10.1111/gcb.14639.
570 571 572 573	Bennington, C. C., Fetcher, N., Vavrek, M. C., Shaver, G. R., Cummings, K. J., and McGraw, J. B. 2012. Home site advantage in two long-lived arctic plant species: results from two 30-year reciprocal transplant studies. Journal of Ecology, 100 (4): 841–851. http://dx.doi.org/10.1111/j.1365-2745.2012.01984.x
574 575 576 577	Bjorkman, A. D., Elmendorf, S. C., Beamish, A. L., Vellend, M., and Henry, G. H. R. 2015. Contrasting effects of warming and increased snowfall on Arctic tundra plant phenology over the past two decades. Global Change Biology, 21(12): 4651–4661. doi: 10.1111/gcb.13051
578 579 580	Bjorkman, A. D., Vellend, M., Frei, E. R., and Henry, G. H. R. 2017. Climate adaptation is not enough: warming does not facilitate success of southern tundra plant populations in the high Arctic. Global Change Biology, 23(4): 1540–1551. doi: 10.1111/gcb.13417
581 582 583	Bjorkman, A.D., Myers-Smith, I.H., Elmendorf, S.C., Normand, S., Rüger, N., Beck, P.S.A., et al. 2018. Plant functional trait change across a warming tundra biome. Nature, 562: 57–62. doi:10.1038/s41586-018-0563-7.
584 585 586 587	Borrell, J. S., Zohren, J., Nichols, R. A., and Buggs, R. J. 2020. Genomic assessment of local adaptation in dwarf birch to inform assisted gene flow. Evolutionary Applications, 13 (1): 161-175.
588 589 590 591 592	Box, J.E., Colgan, W.T., Christensen, T.R., Schmidt, N.M., Lund, M., Parmentier, FJ.W., et al. 2019. Key indicators of Arctic climate change: 1971–2017. Environ. Res. Lett. 14: 45010. doi:10.1088/1748-9326/aafc1b.
592 593 594 595 596	 Brown, R., Schuler, D. V., Bulygina, O., Derksen, C., Luojus, K., Mudryk, L., et al. 2017. Arctic terrestrial snow cover. In <i>Snow, Water, Ice and Permafrost in the Arctic (SWIPA) 2017</i> (pp. 25–64). Oslo, Norway: Arctic Monitoring and Assessment Programme (AMAP).
597 598 500	Buchanan, B. B., W. Gruissem, and R. L. Jones. 2015. Biochemistry and molecular biology of plants. 2nd ed. John Wiley & Sons, Hoboken, NJ
600 601 602 603	Busetto, L., Colombo, R., Migliavacca, M., Cremonese, E., Meroni, M., Galvagno, M., et al. (2010). Remote sensing of larch phenological cycle and analysis of relationships with climate in the Alpine region. Global Change Biology, 16(9): 2504–2517. doi: 10.1111/j.1365-2486.2010.02189.x

Arctic Science Downloaded from cdnsciencepub.com by 86.158.198.15 on 05/21/21 For personal use only. This Just-IN manuscript is the accepted manuscript prior to copy editing and page composition. It may differ from the final official version of record.

604	
605	Chapin III, F.S., and Shaver, G.R. 1985. Individualistic growth response of tundra plant species
606	to environmental manipulations in the field. Ecology. 66: 564–576.
607	http://www.esziournals.org/doi/abs/10.2307/19/0/05
6007	10.2307/1940403.
608	
609	Chapin III, F. S., Shaver, G. R., and Kedrowski, R. A. 1986. Environmental controls over
610	carbon nitrogen and phosphorus fractions in <i>Eriophorum vaginatum</i> in Alaskan tussock
611	tundra Jaural of Foology 74(1): 167, 105, http://www.istor.org/stable/0260257
011	tundra. Journal of Ecology $74(1)$: 107–195. <u>http://www.jstor.org/stable/2200557</u>
612	
613	Chapin III, F.S., Van Cleve, K., and Chapin, M.C. 1979. Soil temperature and nutrient cycling in
614	the tussock growth form of <i>Eriophorum vaginatum</i> Journal of Ecology 67. 169–189
615	
015	
616	Cohen, J., Screen, J. A., Furtado, J. C., Barlow, M., Whittleston, D., Coumou, D., et al. 2014.
617	Recent Arctic amplification and extreme mid-latitude weather. Nature Geoscience, 7(9):
618	627-637 doi: 101038/ngeo2234
010	02/ 05/. 401. 10.1050/1Be0225 1
610	Crowley M. I. (2007) The P. Pook Wiley, Chickester UK
019	Clawley, IVI. J. (2007). The K DOOK. Whey. Chichester, OK.
(0)	
620	Curasi, S. R., Parker, T. C., Rocha, A. V., Moody, M. L., Tang, J., and Fetcher, N. 2019.
621	Differential responses of ecotypes to climate in a ubiquitous Arctic sedge: implications for
622	future ecosystem C cycling New Phytologist 223 (1) [•] 180–192 doi:10.1111/nph.15790
022	
623	Elmandarf S.C. Hanry G.H.P. Hallister P.D. Björk P.G. Bjorkman A.D. Callaghan T
025	Linichuori, S.C., Henry, O.H.K., Homster, K.D., Djork, K.O., Djorkman, A.D., Canagnan, T.
624	V,, et al. 2012a. Global assessment of experimental climate warming on fundra vegetation:
625	heterogeneity over space and time. Ecol. Lett. 15: http://dx.doi.org/10.1111/j.1461-
626	0248.2011.01716.x.
627	Elmendorf S.C. Henry G.H.R. Hollister R.D. Biork R.G. Boulanger-Lanointe N. Cooper
(20)	Elinendon, S.C., Henry, C.H.K., Hollister, K.D., Djork, K.C., Doulanger-Lapoline, N., Cooper,
628	E.J., et al. 2012b. Plot-scale evidence of tundra vegetation change and links to recent
629	summer warming. Nat. Clim. Chang. 2: 453–457. doi:10.1038/nclimate1465.
630	Epstein, H. E., Raynolds, M. K., Walker, D. A., Bhatt, U. S., Tucker, C. J., and Pinzon, J. E.
631	2012 Dynamics of above ground phytomass of the circumpolar Arctic tundra during the past
(22)	three deep log. Environmental Descent Letters 7(1): 01550(
632	three decades. Environmental Research Letters, 7(1): 015506.
633	Fetcher, N., and Shaver, G.R. 1982. Growth and tillering patterns within tussocks of <i>Eriophorum</i>
634	<i>vaginatum</i> . Ecography, 5 : 180–186. doi:10.1111/j.1600-0587.1982.tb01034.x.
635	Fetcher N and Shaver G R 1990 Environmental sensitivity of ecotypes as a notential influence
626	an primary productivity. Am Not 126:126 121 doi:10.1096/205005
030	on primary productivity. Ani. Nat. 130 . 120–151. doi.10.1080/283083.
<i>.</i>	
637	Hamilton, T. D. 2003. Glacial Geology of the Toolik Lake and Upper Kuparuk River Regions.
638	Retrieved from University of Alaska. Institute of Arctic Biology website:
639	https://scholarworks.alaska.edu/bitstream/11122/1502/1/BiologicalPaners_no26.pdf
057	hups,//sonolai //orks.aluska.edu/onstream/11122/1002/1/Diologican upers_no20.put
640	Hereford I 2009 A quantitative survey of local adaptation and fitness trade-offs American
040	110101010, J. 2007. A quantitative survey of local adaptation and numers naue-ons. American
641	Naturalist $1/3(5)$: $5/9-588$. doi: 10.1086/59/611
642	Høye, T. T., Post, E., Meltofte, H., Schmidt, N. M., and Forchhammer, M. C. 2007. Rapid
643	advancement of spring in the High Arctic Current Biology 17(12). R449–R451 doi:
6/1	10 1016/j cub 2007 0/ 0/7
044	10.1010/J.0u0.2007.0T.0T/

- Junttila O., and Kaurin Å. 1985. Climatic control of apical growth cessation in latitudinal
 ecotypes of *Salix pentandra* L. Plant production in the North: proceedings from Plant
 Adaptation Workshop, Tromso, Norway, September 4-9, 1983/edited by Ase Kaurin, Olavi
 Juntilla and Jarle Nilsen, pp. 85—91.
- Kaufman, D. S., and Manley, W. F. 2004. Pleistocene maximum and Late Wisconsinan glacier
 extents across Alaska, USA. Developments in Quaternary Sciences, 2: 9-27.

Kaufman, D. S., Young, N. E., Briner, J. P., and Manley, W. F. 2011. Alaska Paleo-Glacier Atlas
(Version 2). In J. Ehlers, P. L. Gibbard, & P. Hughes (Eds.), Quaternary Glaciations Extent
and Chronology, Part IV: A Closer Look. Developments in Quaternary Science, 15: 427–
445.

Khorsand Rosa, R., Oberbauer, S. F., Starr, G., Parker La Puma, I., Pop, E., Ahlquist, L., and
Baldwin, T. 2015. Plant phenological responses to a long-term experimental extension of
growing season and soil warming in the tussock tundra of Alaska. Global Change Biology,
21(12): 4520–4532. doi: 10.1111/gcb.13040

Leimu, R., and Fischer, M. 2008. A meta-analysis of local adaptation in plants. PLoS ONE
3(12): 1–8. doi: 10.1371/journal.pone.0004010

- Linhart, Y. B., and Grant, M. C. 1996. Evolutionary significance of local genetic differentiation
 in plants. Annual Review of Ecology and Systematics, 27: 237–277. doi:
 10.1146/annurev.ecolsys.27.1.237
- Mark, A.F., Fetcher, N., Shaver, G.R., and Chapin III, F.S. 1985. Estimated ages of mature
 tussocks of *Eriophorum vaginatum* along a latitudinal gradient in central Alaska, U.S.A.
 Arct. Alp. Res. 17: 1–5. doi:10.2307/1550957.

May, J.L., Healey, N.C., Ahrends, H.E., Hollister, R.D., Tweedie, C.E., Welker, J.M., Gould, W.A., and Oberbauer, S.F. 2017. Short-Term Impacts of the Air Temperature on Greening and Senescence in Alaskan Arctic Plant Tundra Habitats. Remote Sensing, 9(12): 1338. doi:10.3390/rs9121338.

- May, J.L., Hollister, R.D., Betway, K.R., Harris, J.A., Tweedie, C.E., Welker, J.M., et al. 2020.
 NDVI changes show warming increases the length of the green season at tundra
 communities in northern Alaska: A fine-scale analysis. Frontiers in Plant Science, 11. doi:
 10.3389/fpls.2020.01174
- McGraw, J. B., Chester A. L., and Stuart L. 1983. A note on July senescence in tundra plants at
 Eagle Creek, Alaska, USA. Arctic and Alpine Research 15:267–269.
- McGraw, J. B., Turner, J. B., Souther, S., Bennington, C. C., Vavrek, M. C., Shaver, G. R., and
 Fetcher, N. 2015. Northward displacement of optimal climate conditions for ecotypes of *Eriophorum vaginatum* L. across a latitudinal gradient in Alaska. Global Change Biology,
 21(10): 3827–3835. doi: 10.1111/gcb.12991
- Mohl, J.E., Fetcher, N., Stunz, E., Tang, J., and Moody, M.L. 2020. Comparative transcriptomics
 of an arctic foundation species, tussock cottongrass (*Eriophorum vaginatum*), during an
 extreme heat event. Sci. Rep. 10: 1–14. doi:10.1038/s41598-020-65693-8.

667

668 669

Arctic Science Downloaded from cdnsciencepub.com by 86.158.198.15 on 05/21/21 For personal use only. This Just-IN manuscript is the accepted manuscript prior to copy editing and page composition. It may differ from the final official version of record.

684 685	Mudryk, L., Brown, R., Luojus, K., Decharme, B., and Helfrich, S. 2019. Terrestrial Snow Cover [In Arctic Report Card 2019]. http://www.arctic.noaa.gov/Report-Card.
686	
687 688 689	Myers-Smith, I.H., Forbes, B.C., Wilmking, M., Hallinger, M., Lantz, T., Blok, D., et al. 2011. Shrub expansion in tundra ecosystems: dynamics, impacts and research priorities. Environ. Res. Lett. 6: 045509. doi:10.1088/1748-9326/6/4/045509.
690	
691 692	Myers-Smith, I.H., Thomas, H.J.D., and Bjorkman, A.D. 2019. Plant traits inform predictions of tundra responses to global change. New Phytol. 221 : 1742–1748. doi:10.1111/nph.15592.
693	
694 695 696	Nilsen J. 1985. Light climate in northern areas. Plant production in the North: proceedings from Plant Adaptation Workshop, Tromso, Norway, September 4-9, 1983/edited by Ase Kaurin, Olavi Juntilla and Jarle Nilsen, pp. 62-72.
09/	Observer S. F. Elmandarf, S. C. Tarrelan, T. C. Hallisten, D. D. Dasha, A. V. Dret Harts, M.
698 699 700 701 702	S., et al. 2013. Phenological response of tundra plants to background climate variation tested using the International Tundra Experiment. Philosophical Transactions of the Royal Society of London. Series B, Biological Sciences, 368 (1624): 20120481. doi: 10.1098/rstb.2012.0481
/02	
703 704 705 706	Park, T., Ganguly, S., Tømmervik, H., Euskirchen, E.S., Høgda, KA., Karlsen, S.R., et al. 2016. Changes in growing season duration and productivity of northern vegetation inferred from long-term remote sensing data. Environ. Res. Lett. 11: 084001. doi:10.1088/1748- 9326/11/8/084001.
707	Park, T., Chen, C., Macias-Fauria, M., Tømmervik, H., Choi, S., Winkler, A., et al. 2019.
708 709	Changes in timing of seasonal peak photosynthetic activity in northern ecosystems. Global Change Biology, 25 : 2382–2395. doi: 10.1111/gcb.14638
710	
711	Park, K., Kang, S. M., Kim, D., Stuecker, M. F., and Jin, FF. 2018. Contrasting local and
712 713	31(8): 3155–3166. doi: 10.1175/JCLI-D-17-0600.1
714	Parker T.C. Tang, I. Clark, M.B. Moody, M.M. and Fetcher, N. 2017. Ecotypic differences
715	in the phenology of the tundra species <i>Erionhorum vaginatum</i> reflect sites of origin. Ecology
716	and Evolution, 7(22): 9775–9786. doi: 10.1002/ece3.3445
717	Pinheiro J Bates D DebRoy S Sarkar D & R Core Team 2017 nlme: Linear and
718	nonlinear mixed effects models. https://cran.r-project.org/package=nlme. R-Project.
719	Post E Forchhammer M.C. Bret-Harte M.S. Callaghan T.V. Christensen T.R. Elberling
720	B et al 2009 Ecological dynamics across the Arctic associated with recent climate change
721	Science, 325 : 1355–1358. doi:10.1126/science.1173113.
700	
722 723 724 725	 Prevey, J., Vellend, M., Rüger, N., Hollister, R. D., Bjorkman, A. D., Myers-Smith, I. Het al. 2017. Greater temperature sensitivity of plant phenology at colder sites: Implications for convergence across northern latitudes. Global Change Biology, 23: 2660–2671. doi: 10.1111/gcb.13619

726	Prevéy, J. S., Rixen, C., Rüger, N., Høye, T. T., Bjorkman, A. D., Myers-Smith, I. H. et al. 2019.
727	Warming shortens flowering seasons of tundra plant communities. Nature Ecology and
728	Evolution, 3(1): 45–52. doi: 10.1038/s41559-018-0745-6
729	R Development Core Team. 2016. R: A Language and Environment for Statistical Computing.
730	Savolainen, O., Lascoux, M., and Merilä, J. 2013. Ecological genomics of local adaptation.
731	Nature Reviews Genetics, 14(11): 807–820. doi: 10.1038/nrg3522
732	Schedlbauer, J. L., Fetcher, N., Hood, K., Moody, M. L., and Tang, J. 2018. Effect of growth
733	temperature on photosynthetic capacity and respiration in three ecotypes of <i>Eriophorum</i>
734	<i>vaginatum</i> . Ecology and Evolution, 8(7): 3711–3725. doi: 10.1002/ece3.3939
735 736 737 738	 Semenchuk, P. R., Gillespie, M. A. K., Rumpf, S. B., Baggesen, N., Elberling, B., and Cooper, E. J. 2016. High Arctic plant phenology is determined by snowmelt patterns but duration of phenological periods is fixed: An example of periodicity. Environmental Research Letters, 11(12): 125006. doi: 10.1088/1748-9326/11/12/125006
739	Shaver, G. R., and Laundre, J. 1997. Exsertion, elongation, and senescence of leaves of
740	<i>Eriophorum vaginatum</i> and <i>Carex bigelowii</i> in Northern Alaska. Global Change Biology, 3 :
741	146–157. doi: 10.1111/j.1365-2486.1997.gcb141.x
742 743	Shaver, G. R., Fetcher, N., and Chapin III, F. S. 1986. Growth and flowering in <i>Eriophorum vaginatum</i> : annual and latitudinal variation. Ecology, 67 (6): 1524–1535.
744	Shi, M., Parazoo, N., Jeong, S., Birch, L., Lawrence, P., Euskirchen, E. S., and Miller, C. E.
745	2019. Exposure to cold temperature affects the spring phenology of Alaskan deciduous
746	vegetation types. Environmental Research Letters, 15(2): 025006.
747	http://iopscience.iop.org/10.1088/1748-9326/ab6502
748	Souther, S., Fetcher, N., Fowler, Z., Shaver, R. G., and McGraw, J. B. 2014. Ecotypic
749	differentiation in photosynthesis and growth of <i>Eriophorum vaginatum</i> along a latitudinal
750	gradient in the Arctic tundra. Botany, 92 (8): 551–561. doi: 10.1139/cjb-2013-0320
751	Stone, R. S., Dutton, E. G., Harris, J. M., and Longenecker, D. 2002. Earlier spring snowmelt in
752	northern Alaska as an indicator of climate change. Journal of Geophysical Research:
753	Atmospheres, 107 (9–10): 10–11. doi: 10.1029/2000jd000286
754	Sullivan, P.F., and Welker, J.M. 2005. Warming chambers stimulate early season growth of an
755	arctic sedge: results of a minirhizotron field study. Oecologia, 142: 616–26.
756	doi:10.1007/s00442-004-1764-3.
757	Wagner, I., and Simons, A. M. 2009. Divergence among arctic and alpine populations of the
758	annual, <i>Koenigia islandica</i> : Morphology, life-history, and phenology. Ecography, 32 (1):
759	114–122. doi: 10.1111/j.1600-0587.2008.05497.x
760 761	Wein, R.W. 1973. Biological flora of British-Isles - <i>Eriophorum vaginatum</i> L. J. Ecol. 61 : 601–615. doi:10.2307/2259047.
762 763	Walker, D.A., Raynolds, M.K., Daniëls, F.J.A., Einarsson, E., Elvebakk, A., Gould, W.A., et al. 2005. The circumpolar Arctic vegetation map. J. Veg. Sci. 16: 267–282.

http://dx.doi.org/10.1111/j.1654-1103.2005.tb02365.x.

Arctic Science Downloaded from cdnsciencepub.com by 86.158.198.15 on 05/21/21 For personal use only. This Just-IN manuscript is the accepted manuscript prior to copy editing and page composition. It may differ from the final official version of record.

ion of record.	765 766 767 768	Walker, D. A., Daniëls, F. J. A., Alsos, I., Bhatt, U. S., Breen, A. L., Buchhorn, M., et al. 2016. Circumpolar Arctic vegetation: A hierarchic review and roadmap toward an internationally consistent approach to survey, archive and classify tundra plot data. Environmental Research Letters, 11 (5): 055005. doi:10.1088/1748-9326/11/5/055005
final official vers	769 770 771	Walker, T.W.N., Weckwerth, W., Bragazza, L., Fragner, L., Forde, B.G., Ostle, N.J., et al. 2018. Plastic and genetic responses of a common sedge to warming have contrasting effects on carbon cycle processes. Ecol. Lett. 22: 159–169. doi:10.1111/ele.13178.
m the fins	772 773	Woods, C., and Caballero, R. 2016. The role of moist intrusions in winter Arctic warming and sea ice decline. Journal of Climate, 29 (12): 4473–4485.
05/21/21 It may differ fro	774 775 776 777	Wookey, P.A., Aerts, R., Bardgett, R.D., Baptist, F., Bråthen, K., Cornelissen, J.H.C., et al. 2009. Ecosystem feedbacks and cascade processes: Understanding their role in the responses of Arctic and alpine ecosystems to environmental change. Glob. Chang. Biol. 15: 1153– 1172. doi:10.1111/j.1365-2486.2008.01801.x.
98.15 on position. l	778	
/ 86.158.1 page comj		
ub.com by iting and j		
nsciencep o copy ed		
d from cd ipt prior t		
ownloade d manusci		
science D		
Arctic S script is th		
-IN manus		
This Just-		
use only.		
· personal		
For		