Responses of sequential and hierarchical phenological events to warming and cooling in alpine meadows

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Organisms’ life cycles consist of hierarchical stages, from a single phenological stage (for example, flowering within a season), to vegetative and reproductive phases, to the total lifespan of the individual. Yet phenological events are typically studied in isolation, limiting our understanding of life history responses to climate change. Here, we reciprocally transfer plant communities along an elevation gradient to investigate plastic changes in the duration of sequential phenological events for six alpine species. We show that prolonged flowering leads to longer reproductive phases and activity periods when plants are moved to warmer locations. In contrast, shorter post-fruiting leaf and flowering stages led to shorter vegetative and reproductive phases, respectively, which resulted in shorter activity periods when plants were moved to cooler conditions. Therefore, phenological responses to warming and cooling do not simply mirror one another in the opposite direction, and low temperature may limit reproductive allocation in the alpine region.
solated phenological events (for example, green-up, flowering) have largely shaped our understanding of phenology as a bioindicator of climate change\(^1\)-\(^4\); yet in reality, phenological events are inter-related and hierarchical. For example, flower bud formation, flowering and fruit/seed maturation compose the reproductive phase of angiosperms; furthermore, vegetative and reproductive phases compose the annual total aboveground activity period of plants in seasonal habitats (hereafter activity period; Fig. 1). Certain phenological stages within the plant life cycle may limit later stages\(^5\)-\(^6\) and therefore affect the responsiveness of other phenological stages to environmental change. Thus, assessing an entire sequence of phenological events can reveal which individual stages have the largest effect on composite phenologies, and provide comprehensive insight into life history responses to climate change\(^6\)-\(^9\).

The degree to which plants respond to temperature changes by adjusting vegetative and reproductive phases should depend in part on the effect of temperature on resource allocation to growth versus reproduction. In addition, temperature-dependent physiological processes can regulate the duration of plant life history events. Plant growth and reproduction depend on the same internal resource pool\(^10\),\(^11\) and plants allocate a small percentage of their total resources to sexual reproduction in arctic and alpine regions\(^1\),\(^2\), presumably because they live in harsh environments with a short time window to complete their life cycles. Yet climate warming is resulting in a longer period of time that is conducive to growth and reproduction in many seasonal habitats\(^13\)-\(^16\).

Conversely, warmer environments may impose new constraints on the duration of phenological events if faster rates of development occur under warmer temperatures\(^8\),\(^17\). Given the harsh abiotic environment in alpine habitats, we expect that warmer temperatures will reduce constraints on allocation to sexual reproduction, in which case we expect longer reproductive phases and activity periods. Following this same logic, we expect reproductive phases to be shortened more so than vegetative phases under cooling, for a total reduction in activity periods.

In this study we reciprocally transferred intact plant communities in 1 m by 1 m blocks of soil at 3,200, 3,400, 3,600 and 3,800 m above sea level in the Qilian Mountains of China. Our experiment provides insight into how plants may adjust sequences of hierarchical phenological events within the life cycle (Fig. 1) to warmer temperatures expected under future climate change in addition to temperature anomalies, such as short-term cooling events\(^18\),\(^19\). In addition, by examining plastic responses in phenological duration to both warmer and cooler temperatures, our study improves our understanding of the basic role of temperature in shaping the relationships among phenological events in the plant life cycle, a need that has been highlighted by the widespread effects of climate change on phenological events\(^20\).

Intact soil blocks were transferred across all elevations in 2007. The phenology of six common perennial plants was monitored from 2008 to 2010 at all sites. These plants included two early-flowering sedge species, Kobresia humilis (Kh) and Carex scabrirostris (Cs); two mid-summer flowering forbs Potentilla anserine (Pa) and P. nivea (Pn); and two mid-summer flowering grasses, Poa pratensis (Pp) and Stipa aliena (Sa). We measured the durations of six phenological stages in tagged individual plants of each species: leaf out, flower bud, flowering, fruiting, post-flowering leaf and leaf colouring\(^21\) (Fig. 1) with 3–4 day intervals at each elevation. We report a metric of phenological sensitivity to temperature changes from the reciprocal transplants for each phenological event: change in duration divided by absolute change in temperature between each donor and receptor site (days °C\(^{-1}\)).

Here we find that warming significantly prolongs the duration of flowering, which is primarily responsible for longer reproductive phases and activity periods, but cooling significantly reduces both reproductive and vegetative phases and activity periods. Our results show that the temperature sensitivities of flowering duration to warming and post-fruiting leaf duration to cooling are greater compared with other phenological stages. Thus, the responses of phenological stages to warming and cooling do not simply mirror one another in the opposite direction, and low temperature may limit the reproductive allocation in the alpine region.

**Results**

**Temperature sensitivity of hierarchical phenological events.** Inter-related phenological events within each hierarchy were analysed using a repeated-measures multivariate analysis of variance (MANOVA). The sensitivities of the activity period and vegetative and reproductive phases to temperature were significantly affected by temperature change direction (that is, warming and cooling) and/or species and their interaction (Tables 1 and 2). Warming prolonged the activity period and the reproductive phase, while cooling shortened the activity period and both vegetative and reproductive phases for all species (Fig. 2a–c). Warming significantly prolonged the activity period by an average of 15.7 days °C\(^{-1}\) across species (Fig. 2c), and the reproductive phase accounted for on average 83% of the increase in the duration of the activity period under warming (range 69–99% across species; Fig. 2d). In contrast, cooling shortened the activity period by an average of 14.8 days °C\(^{-1}\) across species (Fig. 2c), and the reproductive phase accounted for on average 44.8% of the decrease (range 20–68% across species) in the duration of the activity period under cooling (Fig. 2d).

**Temperature sensitivity of individual phenological event.** Plant species, temperature change direction and their interaction all significantly affected the sensitivity of duration to temperature in the six individual phenological stages (Table 3). Warming significantly prolonged the duration of the reproductive phase (11.1–15.1 days °C\(^{-1}\) across species) (Fig. 2b), which was mainly caused by the prolonged duration of flowering for all plant species (which ranged from 7.4 to 11.3 days °C\(^{-1}\) ) (Fig. 3a–f). Longer flowering also contributed to longer reproduction in four of the six species (0–5.7 days °C\(^{-1}\) ) (Fig. 3c–f) and longer flower budding in four of six species (Fig. 3a,c,e,f), but these stages always contributed less than the flowering stage. Warming also significantly prolonged the duration of the vegetative phase in three species (1.7–5.5 days °C\(^{-1}\) ) (Fig. 2a), which primarily derived from the prolonged duration of leaf colouring (1.5–4.4 days °C\(^{-1}\) ) (Fig. 3a,c–f; Cs is the exception (Fig. 3b)). In contrast, cooling significantly shortened the duration of the vegetative phase (4.6–9.6 days °C\(^{-1}\) ) (Fig. 2a), which resulted primarily from the...
Table 1 | Temperature sensitivity of duration change for total aboveground activity period.

|                     | Df | Sum Sq | Mean Sq | F value | Pr(> F) |
|---------------------|----|--------|---------|---------|---------|
| Year                | 1  | 134    | 134     | 0.861   | 0.354   |
| Species (S)         | 5  | 1,309  | 262     | 1.677   | 0.138   |
| TCD                 | 1  | 325,116| 325,116 | 2,083.0 | <0.001  |
| S * TCD             | 5  | 7,163  | 1,433   | 9.179   | <0.001  |

ANOVA, analysis of variance; TCD, temperature change direction (that is, warming and cooling).
Repeated-measures ANOVA is used to test the effects of year, species and temperature change direction on sensitivity of duration change to temperature for the total aboveground activity period.

Table 2 | Temperature sensitivity of duration change for vegetative and reproductive phases.

| df     | Pillai | approx F | num df | den df | Pr(> F) |
|--------|--------|----------|--------|--------|---------|
| Year   | 1      | 0.00248  | 0.65   | 2      | 0.521   |
| Species (S) | 5      | 0.04017  | 2.16   | 10     | 0.0181  |
| TCD    | 1      | 0.81048  | 1,124.7| 2      | <0.001  |
| S * TCD| 5      | 0.17802  | 10.30  | 10     | <0.001  |

MANOVA, multivariate analysis of variance; TCD, temperature change direction (that is, warming versus cooling).
Repeated-measures MANOVA analysis of sensitivity of duration to temperature for both vegetative and reproductive phases is used as the multivariate response and species, year and TCD as predictors.

Figure 2 | Temperature sensitivity of duration change of phenological events. The panels include the vegetative phase (a), reproductive phase (b), activity period (c) and their ratios of reproduction to activity period (d) under warming and cooling conditions. A t-test was used for each phenological event and species, and the α was adjusted following a Bonferroni correction. The mean value of phenological sensitivity in each plot in each year was the unit of replication (n = 45 for each t-test under warming and cooling conditions, respectively). Kh, K. humilis; Cs, C. scabrirostris; Pa, P. anserine; Pn, P. nivea; Pp, P. pratensis; and Sa, S. aliena. Δdays was the difference between receptor site (that is, away site) and donor site (that is, home site). Positive values indicate a prolonged phenological stage and negative values a shortened phenological stage when transferred compared with donor site. Means ± s.e. are shown in the figure. ‘*’, ‘**’ in the figure are the significant difference at 0.004 and 0.0008 levels between receptor site and donor site. All P values were <0.0001 except vegetative phase for Kh, Cs, Pn and Pp under warming, which were 0.031, 0.727, 0.002 and 0.101, respectively.
shortening of the post-fruiting leaf stage for all species (5.1–10.1 days °C\(^{-1}\)) (Fig. 3a–f). A shortened duration of fruit-set contributed to shorter reproductive phases (2.2–6.0 days °C\(^{-1}\)) under cooling for all plant species (Fig. 3a–f), although for Pa and Pn, shorter flowering stages contributed more than fruiting (Fig. 3c,d).

**Correlations among phenological events.** There were significant negative correlations between the sensitivity of the vegetative (VP) and reproductive phases (RP) to warming and cooling regardless of whether the analysis was pooled across all species (Table 4) or conducted separately for each species (Supplementary Tables 2–13; except for Pp (Supplementary Table 3). Temperature sensitivity of duration change for six individual phenological stages.

|                | df | Pillai | approx F | num df | den df | Pr(>F) |
|----------------|----|--------|----------|--------|--------|---------|
| Year           | 1  | 0.01675| 1.48     | 6      | 522    | 0.1821  |
| Species (S)    | 5  | 0.33889| 6.37     | 30     | 2,630  | <0.001  |
| TCD            | 1  | 0.82313| 404.89   | 6      | 522    | <0.001  |
| S * TCD        | 5  | 0.53236| 10.45    | 30     | 2,630  | <0.001  |

MANOVA, multivariate analysis of variance; TCD, temperature change direction (that is, warming versus cooling). Repeated-measures MANOVA analysis of sensitivity of duration to temperature for the six individual phenological stages is used as the multivariate response and species, year and TCD as predictors.

Repeated-measures MANOVA analysis of sensitivity of duration to temperature for the six individual phenological stages is used as the multivariate response and species, year and TCD as predictors.
phenology14,22. Instead, sensitivity of duration to temperature found no evidence of lag or build-up effects of warming on plant experimental studies in high-elevation or high-latitude sites, we common response to the alpine environment. Unlike other consistent despite the inclusion of three different plant life forms both vegetative and reproductive phases, for shorter annual moving plants up in elevation to cooler temperatures shortened longer reproductive phases; in particular flowering. Conversely, activity periods in individuals of all six species. These patterns are in composite phenophases. In all three years of the study, moving and allowed us to identify the individual stages that drive changes in the duration of phenological events in six alpine plant species.

Discussion
Our reciprocal transfer experiment revealed substantial plasticity in the duration of phenological events in six alpine plant species and allowed us to identify the individual stages that drive changes in composite phenophases. In all three years of the study, moving plants down in elevation to warmer temperatures prolonged activity periods of individuals in all six species, mainly because of longer reproductive phases; in particular flowering. Conversely, moving plants up in elevation to cooler temperatures shortened both vegetative and reproductive phases, for shorter annual activity periods in individuals of all six species. These patterns are consistent despite the inclusion of three different plant life forms in our study: sedges, grasses and forbs, which is suggestive of a common response to the alpine environment. Unlike other experimental studies in high-elevation or high-latitude sites, we found no evidence of lag or build-up effects of warming on plant phenology14,22. Instead, sensitivity of duration to temperature was consistent across years in all species, despite interannual variation in temperature (for example, 2008 was cooler than 2009 and 2010 by 0.6–0.7 °C on average)3.

Most studies understandably focus on the effects of warming on plant phenophases in response to climate change, but in our study we also address a need for more basic research on relationships among inter-related phenological events23 by asking whether responses to cooling are simply the opposite of responses to warming or whether events respond differentially. The outcome depends on hierarchy level, which highlights the utility of considering phenological sequences within a hierarchical framework. For the upper hierarchies of the activity period, reproductive phase and vegetative phase, it appears that opposite responses are indeed observed under warming versus cooling (longer versus shorter). However, cooler temperatures do not simply have the opposite effect of warmer temperatures on the duration of individual life history events, especially within the vegetative phase. Therefore, it should not be assumed that effects of cooler temperatures on the duration of phenological events will simply mirror responses to warmer temperatures in the opposite direction.

The onset of reproduction signals a change in resource allocation from growth and survival to fecundity within the plant life cycle24,25. Longer reproductive phases under warming and shorter phases under cooling suggest that allocation to reproduction is probably increased and reduced according to temperature in these alpine plants. Our results are consistent with the observation that alpine plants allocate a small proportion of their resources to sexual reproduction under harsh environmental conditions12. Thus we find no evidence of a faster rate of development of reproductive structures under warmer temperatures as seen in other studies5,17. Instead, our results support the hypothesis that reproductive allocation in these plants is generally constrained by cold-temperature conditions in the alpine environment.

Longer reproduction and activity periods under warmer temperatures were primarily driven by longer flowering stages in all species (Fig. 2d). The flowering stage accounted for 70% of the extension to the reproductive phase under warming and for 58% of the extension to the activity period on average across species. Advanced first flowering appears to be the main contribution to prolonged flowering duration under warming.

### Table 4 | Correlations among temperature sensitivities of phenological duration.

| Leaf-out | Flower bud | Flower | Fruit | Post-fruit leaf | Leaf colour | VP | RP |
|----------|------------|--------|-------|-------------|-------------|----|----|
| Warming  |            |        |       |             |             |    |    |
| Flower bud | 0.12       |        |       |             |             |    |    |
| Flower    | -0.60**    | 0.28** |       |             |             |    |    |
| Fruit     | -0.14      | 0.55** | 0.03  |             |             |    |    |
| Post-fruit leaf | -0.41**  | -0.25** | -0.08 | -0.10      |             |    |    |
| Leaf colour | -0.18     | -0.07 | -0.04 | 0.05        | 0.05        |    |    |
| WP        | 0.37**     | -0.31** | -0.55** | -0.14     | 0.37**      |    |    |
| RP        | -0.48**    | 0.72** | 0.70** | 0.70**      | -0.17       | -0.01 | -0.50** |
| AP        | -0.31**    | 0.62** | 0.44** | 0.71**      | 0.05        | 0.40** | 0.09 | 0.82** |
| Cooling   |            |        |       |             |             |    |    |
| Flower bud | 0.18       |        |       |             |             |    |    |
| Flower    | -0.38**    | 0.15   |       |             |             |    |    |
| Fruit     | 0.06       | 0.28** | 0.19* |             |             |    |    |
| Post-fruit leaf | -0.31**   | -0.26** | 0.13  | 0.04        |             |    |    |
| Leaf colour | -0.29**  | -0.20 | -0.11 | -0.47**     | -0.29**     |    |    |
| WP        | 0.04       | 0.33** | 0.16  | -0.26**     | 0.73**      | 0.21* |    |
| RP        | -0.10      | 0.54** | 0.66** | 0.81**      | 0.01        | -0.41** | -0.34** |
| AP        | -0.06      | 0.25** | 0.49** | 0.56**      | 0.65**      | -0.21* | 0.47** | 0.67** |

AP, activity period; RP, reproductive phase; VP, vegetative phase.

* and ** indicate significant correlations at P < 0.0014 and 0.0003 levels, respectively.

The temperature sensitivity of each phenological event is included under warming and cooling, across all three years of the study (2008–2010). All species were pooled for ease of interpretation, because results are largely consistent if each species is analysed separately (Supplementary Tables 1–12). The sample unit is the mean sensitivity value for each species in each plot and year (n = 270 for warming and for cooling, respectively).
as opposed to later last flowering. Some experiments have found either longer flowering or maintenance of flowering duration under warmer temperatures, depending on the species, whereas other experiments have found a shorter duration of flowering and other individual reproductive stages under warming. This variation across species in how the duration of flowering responds to warming may in part reflect different warming methods (open-top chambers versus infrared heaters versus transplants) in addition to ecosystem type—specifically the timing of abiotic extremes that may constrain individual stages within a growing season. For example, divergence in entire reproductive phases and flowering stages at the species level has been detected in two habitats with extreme abiotic conditions in the middle of the growing season—high temperatures and drought—so that a mid-season gap in reproduction appears at the community level. In sum, plastic responses to climate change are likely to include phenological shifts away from parts of the season where abiotic conditions are becoming more extreme and toward parts of the season where abiotic extremes are being attenuated.

The response of flowering to warming may be attributed to genetic and physiological mechanisms, and the genetic basis of flowering is the most well understood plant life history phase. The onset of flowering occurs earlier under warming and later under cooling in these six species, which agrees with laboratory research on genetic pathways regulating how the timing of flowering responds to temperature in Arabidopsis thaliana. Thus, the responses of flowering duration in this study are probably mediated through similar pathways at the genetic level that determine the switch to allocation to reproduction. However, in contrast to what we find here, a recent gene regulatory model predicts shorter flowering durations under climate warming in Anemone halleri, a perennial relative of A. thaliana. We suspect this discrepancy is related to effects of warmer temperatures on vernalization requirements, which appear not to play a role over the range of temperature variation in our study but could be important under continued climatic warming.

The population and community-level consequences of extended phenological stages under climate change will depend on biotic interactions with antagonists and mutualists, in addition to which type of interaction has a stronger effect on plant fitness. Longer flowering durations in individual plants may reduce the likelihood of phenological mismatch with animal pollinators and may also compensate or buffer against potentially declining pollinator populations under climate change. Longer flowering durations of individuals may also alter patterns of assortative mating, with potential consequences for additive genetic variation within the population, depending on whether synchrony among individuals is maintained or increased. Longer activity periods could increase exposure of plants to herbivory and disease in both vegetative and floral tissue. Indeed, over longer time scales that span multiple plant generations, biotic interactions like the ones described above are likely to drive adaptive responses in phenological duration, which could differ from the short-term plastic responses reported in our study.

In contrast to the individual reproductive stages that largely changed in concert with one another under warming and cooling, different individual vegetative stages accounted for the effects of warming and cooling on the duration of the vegetative phase (Table 1). In general, leaf senescence was extended under warming, whereas the post-fruit leaf stage was shortened under cooling (Fig. 3). Shorter post-fruit leaf stages are consistent with a strategy of maintaining allocation to growth and reproduction under cooling, instead of allocating to tissue maintenance following reproduction, although in two species the flowering or flowering stages were reduced in a similar magnitude to the post-fruit leaf stage. Leaf senescence is an integrated response of leaf cells to endogenous developmental and external environmental conditions, and our results are consistent with another study that found longer leaf coloration periods in deciduous trees under warmer fall temperatures. Longer growing seasons often lead to increased primary production and carbon sequestration. However, longer activity periods may not translate into increased carbon sequestration in our study because senescence was the main driver of longer vegetative phases and longer reproduction was the main driver of longer activity periods under warming. Furthermore, the leaf-out stage was only longer under warming in one species. The balance between respiration and photosynthesis under elevated temperatures will ultimately determine whether longer activity periods result in increased or decreased primary production.

Temperature is not the only factor that varied over the elevation gradient in our study and is only one of several factors that may affect plant phenology under climate change. Although higher elevation sites tended to be drier, the 3,600 m site had the highest soil moisture. Thus, our conclusions of the effects of temperature are unlikely to be confounded by concurrent changes in soil moisture because these two abiotic factors do not precisely covary across our four sites. Although photoperiod can play a role in the regulation of alpine plant life history events, photoperiod does not vary across our sites that are separated by a maximum distance of 9 km. Increasing atmospheric levels of CO₂ can also extend the duration of life history events under warming, and will most likely affect vegetative stages directly. Additionally, the timing of spring snowmelt often predicts the timing of flowering in high-elevation and high-latitude environments, but in our study area the ground is usually snow-free unless there is a large snow event. Thus, the timing of snowmelt is unlikely to be an important phenological cue in our study. Our experimental protocol controlled for plant community context and below-ground soil communities, but other biotic factors may vary across the sites. For example, the activity and effectiveness of animal pollinators may have varied across the sites, but pollinators would only affect the reproduction of the two forb species, and tradeoffs between pollinator visitation rates and pollinator effectiveness across elevation gradients can compensate for such variation.

Finally, we focused on the duration of events at the individual plant-level in this study, but the hierarchical nature of phenological events can be categorized in various ways. For example, measuring the duration of individual leaves and flowers would provide further insight into the consequences of phenological responses to temperature change.

A main goal of ecological research is to understand what drives the abundance and distribution of species in space and time, and climate change presents ecologists with a unique opportunity to advance basic research in temporal ecology. Rapidly shifting species’ phenologies under climate change have brought to light that we still need an improved understanding of how various abiotic and biotic factors affect the duration of life cycle events in order to predict species’ responses to climate change. Extended phenological durations may or may not be adaptive under climate change, but in our study species, the duration of flowering, post-fruiting leaf, and senescence should be particularly sensitive to climate change in the near future.

Methods
Experimental site and data collection. The experimental site and design are described elsewhere in detail. In brief, the experiment was conducted at Haibei Alpine Meadow Ecosystem Research Station of the Chinese Academy of Sciences, located at latitude 37°57′N, and longitude 101′12′E along a 3,200–3,800 m elevational gradient (3,200, 3,400, 3,600 and 3,800 m) on the south slope of the
Qilian Mountains in Qinghai, China. The four sites included four different plant communities within 9 km of one another (maximum distance between lowest and highest elevations). At the centre of each site, HOBO weather stations (Onset Computer Corporation, Cape Cod, Massachusetts, USA) were used to monitor soil temperature at 5 cm depth and soil moisture at 20 cm depth. Data were sampled at 1 min intervals, and 30 min averages were stored in the data logger. Annual average soil temperatures at 5 cm depth were 3.9, 2.5, 2.0 and 0.4 °C, and average annual soil moisture at 20 cm depth was 11.8, 11.3, 12.7 and 10.2% at 3,200, 3,400, 3,600 and 3,800 m during the experimental period, respectively. Photoperiod is consistent across the four sites because of their close proximity.

Twelve intact soil blocks (100 × 100 cm wide × 30–40 cm deep) (30 cm depth at 3,800 m due to shallower soil layer) with attached vegetation from each elevation were transferred across the elevation gradient after the soils started to thaw in early May, 2007 (12 soil blocks per elevation × 4 elevations, for a total of 48 soil blocks or plots). There were three replicate transfers from each elevation, and these intact soil blocks were monitored for the entire growing season. All plots were censused at an interval of 3 or 4 days from early April to the end of October in each year, and the same individuals were monitored from 1 year to the next. Observations were made at an interval of 3 or 4 days from early April to the end of October in each year, and recordings were made of the onset and end dates of six phenological events, listed here in chronological order: leaf emergence, flower budding, flowering, fruit-set for forbs or seed-set for graminoids, post-fruit/seeding vegetative stage and leaf colouring (number of days from first yellowing or browning of leaves to complete discoloration). Because phenology was assessed at 3–4 day intervals, onset and end dates of each phenological event were estimated; for onset, we used the average of the first date on which the event was recorded and the date of the prior census in which the event was not recorded (vice versa for end date). For example, if flowering was observed on 5 June for the first time, and the census before that on 1 June had no flowering, the estimated onset date was 3 June. The duration of each phenological event was the number of days between the estimated starting and ending dates for each event, for each individual plant or stem. We calculated a phenological event as the number of days between the estimated starting and ending dates for each event, for each individual plant or stem. We calculated a phenological duration to temperature changes from the reciprocal transplants: change in duration per change in °C (Adams’ C°C). For example, if the difference in mean flowering duration between a receptor and donor site is 5 days, and the temperature difference between the two sites is 5°C, for a sensitivity of 1 day °C, each species-level response was an average across 10 individuals per plot, and this calculation was performed for each phenological event for each species in every plot.

Phenological responses to warming and cooling within each hierarchy level were analysed with the method repeated-measures ANOVA (Tables 2 and 3), except total activity period, which was analysed with a repeated-measures ANOVA (Table 1) because it is a single event. MANOVA investigates whether there are statistically significant mean differences among groups in a linear combination of several variables. Several phenological responses within each hierarchy level are correlated with one another (Table 4 and Supplementary Tables 2–13), indicating that a MANOVA is necessary to test for an overall effect of predictors on the multiple phenological responses. Pillai’s statistic (the default) was applied to the MANOVA output. Only minor differences in results were observed with the use of other statistics (Wilks’ Lambda). Repeated-measures ANOVA was used to test the contributions of the factors year, species and temperature change direction (two levels, that is, warming and cooling) to the sensitivity of the duration of the activity period to temperature (Table 1). The base package R-3.2.3 (ref. 61) was used to conduct statistical analyses.

The effects of cooling and warming on sensitivity of duration for different phenological events were analysed by comparison to the controls (the plots transferred within the same elevations). A t-test was used for each phenological event and species, and the z was adjusted following a Bonferroni correction (Fig 2 and 3). Significant differences were reported at P < 0.004 and 0.0008 levels in the text. The mean value of phenological sensitivity in each plot in each year was the unit of replication (n = 45 for each t-test under warming and cooling, respectively).

Correlation analysis was performed among the sensitivities of duration change to temperature for all phenological events to determine their relationships in direction and magnitude for each species and across all species during the experimental periods. Since 36 tests were presented in Table 1, the significance level for each individual test was set to 1/36, a conservative correction for multiple comparisons. We found that significant correlations were reported at P < 0.0014 and 0.0003 levels in the text.

Data availability. The data that support the findings of this study are available from the corresponding author upon request.

References

1. Wolkovich, E. M. et al. Warming experiments underpredict plant phenological responses to climate change. Nature 485, 494–497 (2012).

2. Iler, A. M., Høye, T. T., Inouye, D. W. & Schmidt, N. M. Nonlinear flowering responses to climate: Are species approaching their limits of phenological change? Philos. Trans. R. Soc. Lond. B Biol. Sci. 368, 20120489 (2013).

3. Wang, S. P. et al. Asymmetric sensitivity of first flowering date to warming and cooling in alpine plants. Ecology 95, 3387–3398 (2014).

4. Jiang, L. L. et al. Relatively stable response of fruiting stage to warming and cooling relative to other phenological events. Ecology 97, 1961–1969 (2016).

5. Sola, A. J. & Ehrlén, J. Vegetative phenology constrains the onset of flowering in the perennial herb Lathyrus vernus. J. Ecol. 95, 208–216 (2007).

6. Lost, E. S., Pedersen, C., Wilmers, C. C. & Forchhammer, M. C. Phenological sequences reveal aggregate life history response to climatic warming. Ecology 89, 363–370 (2008).

7. Donohue, K. Germination timing influences natural selection on life history characters in Arabidopsis thaliana. Ecology 83, 1006–1016 (2002).

8. Hagerty, B. P. & Galloway, L. F. Response of individual components of reproductive phenology to growing season length in a monocarpic herb. J. Ecol. 99, 242–253 (2011).

9. Sherry, R. A. et al. Changes in duration of reproductive phases and lagged phenological response to experimental climate warming. Plant Ecol. Divers. 4, 23–35 (2011).

10. Guggisberg, S., Kesselring, H., Stöcklin, J. & Hamann, E. Lower plasticity exhibited in mean flowering duration between a receptor and donor site is 5 days, this indicates that a MANOVA is necessary to test for an overall effect of predictors on and cooling relative to other phenological events. Ecology 97, 1961–1969 (2016).

11. Molau, U. Relationships between flowering phenology and life-history strategies in tundra plants. Arct. Alp. Res. 25, 391–402 (1993).

12. Mylénis, R. B., Koeleg, C. D., Tucker, C. J., Asrar, G. & Nemani, R. I. Increased plant growth in the northern high latitudes from 1981 to 1991. Nature 386, 698–702 (1997).

13. Arft, A. M. et al. Responses of tundra plants to experimental warming: meta-analysis of the international tundra experiment. Ecol. Monogr. 69, 491–511 (1999).

14. Oberbauer, S. F. et al. Phenological response of tundra plants to background climate variation tested using the International Tundra Experiment. Philos. Trans. R. Soc. Lond. B Biol. Sci. 368, 20120481 (2013).

15. Tara, P. J., Iler, A. M. & Inouye, D. W. Shifts in flowering phenology reshape a subalpine plant community. Proc. Natl Acad. Sci. USA 111, 4916–4921 (2014).

16. Guggisberg, S., Kesselring, H., Stöcklin, J. & Hamann, E. Lower plasticity exhibited by high- versus mid-elevation species in their phenological responses to manipulated temperature and drought. Ann. Bot. 116, 953–962 (2015).

17. Maye, F. E. & Cosyns, L. C. Impact of the Younger Dryas cooling event upon lowland vegetation of Maritime Canada. Ecol. Monogr. 65, 130–134 (1995).

18. Sirocko, F., Brückn, H. & Pflädl, S. Solar influence on winter severity in central Europe. Geophys. Res. Lett. 39, L116704 (2012).

19. Wolkovich, E. M., Cook, B. I. & Davies, T. J. Progress towards an interdisciplinary science of plant phenology: building prediction across space, time and species diversity. New Phytol. 201, 1156–1162 (2014).

NATURE COMMUNICATIONS | DOI: 10.1038/ncomms12489 | www.nature.com/naturecommunications 7
21. Wang, S. P. et al. Timing and duration of phenological sequences of alpine plants along an elevation gradient on the Tibetan plateau. *Agric. For. Meteorol.* **139-140**, 220–228 (2014).

22. Hoffmann, A. A. et al. Phenological changes in six Australian subalpine plants in response to experimental warming and year-to-year variation. *J. Ecol.* **98**, 927–937 (2010).

23. Bolmgren, K. & Eriksson, O. Are mismatches the norm? Timing of flowering, fruiting, dispersal and germination and their fitness effects in *Frangula alnus* (Rhamnaceae). *Oikos* **124**, 639–645 (2015).

24. Bazzaz, F. A. & Grace, J. *Plant Resource Allocation* (Academic Press, 1997).

25. Roff, D. A. *Life History Evolution* (Sinauer Associates, 2002).

26. Price, M. V. & Waser, N. M. Effects of experimental warming on plant reproductive phenology in a subalpine meadow. *Ecology* **79**, 1261–1271 (1998).

27. Dunne, J. A., Harte, J. & Taylor, K. J. Subalpine meadow flowering phenology respon-A to climate change: integrating experimental and gradient methods. *Ecol. Monogr.* **73**, 69–86 (2003).

28. Reyes-Fox, M. et al. Elevated CO2 further lengthens growing season under warming conditions. *Nature* **510**, 529–526 (2014).

29. Sherry, R. A. et al. Divergence of reproductive phenology under climate warming. *Proc. Natl Acad. Sci. USA* **104**, 198–202 (2007).

30. Aldridge, G., Inouye, D. W., Forrest, J. E. R., Barr, W. A. & Miller-Rushing, A. J. Emergence of amid-season period of low floral resources in a montane meadow ecosystem associated with climate change. *J. Ecol.* **99**, 905–913 (2011).

31. Andrés, F. & Coupland, G. The genetic basis of flowering responses to seasonal cues. *Nat. Rev.* **13**, 627–639 (2012).

32. Obeso, J. R. The costs of reproduction in plants. *New Phytol.* **155**, 321–348 (2002).

33. Blázquez, M. A., Ahn, J. H. & Weigel, D. A thermosensory pathway controlling flowering time in *Arabidopsis thaliana*. *Nat. Genet.* **36**, 168–171 (2003).

34. Kumar, S. V. et al. Transcription factor PIF4 controls the thermosensory activation of flowering. *Nature* **484**, 242–245 (2012).

35. Lee, J. H. et al. Role of SVP in the control of flowering time by ambient temperature in *Arabidopsis*. *Genes Dev.* **21**, 397–402 (2007).

36. Liu, J. Z., Feng, L. L., Li, J. M. & He, Z. H. Genotypic and epigenetic control of plant heat responses. *Front. Plant Sci.* **6**, 267 (2015).

37. Wigge, P. A. et al. Integration of spatial and temporal information during floral induction in *Arabidopsis*. *Science* **309**, 1056–1059 (2005).

38. Satoh, A. et al. Forecasting flowering phenology under climate warming by modelling the regulatory dynamics of flowering-time genes. *Nat. Commun.* **4**, 2303 (2013).

39. Memmott, J., Craze, P. G., Waser, N. M. & Price, M. V. Global warming and the disruption of plant-pollinator interactions. *Ecol. Lett.* **10**, 710–717 (2007).

40. Hegland, S. J., Nielsen, A., Lazaro, A., Bjerknes, A. L. & Totland, O. How does climate warming affect plant-pollinator interactions? *Ecol. Lett.* **12**, 184–195 (2009).

41. Thomson, J. D. Flowering phenology, fruiting success and progressive deterioration of pollination in an early-flowering geophyte. *Philos. Trans. R. Soc. Lond. B Biol. Sci.* **365**, 3187–3199 (2010).

42. McKinney, A. M. et al. Asynchronous changes in phenology of migrating broad-tailed hummingbirds and their early-season nectar resources. *Ecology* **93**, 1987–1993 (2012).

43. Iler, A. M. et al. Maintenance of temporal synchrony between syrphid flies and floral resources despite differential phenological responses to climate. *Global Change Biol.* **19**, 2348–2359 (2013).

44. Bunkle, L. A., Marlin, J. C. & Knight, T. M. Plant-pollinator interactions over 120 years: loss of species, co-occurrence, and function. *Science* **339**, 1611–1615 (2013).

45. Haye, T. T. et al. Phenology of high-arctic butterflies and their floral resources: species-specific responses to change. *Curr. Zool.* **60**, 243–251 (2014).

46. Fox, G. Assortative mating and plant phenology: evolutionary and practical consequences. *Evol. Ecol. Res.* **5**, 1–18 (2003).

47. Franks, S. J. The unique and multifaceted importance of the timing of flowering. *Annu. J. Bot.* **102**, 1401–1402 (2015).

48. Elzinga, J. A. et al. Time after time: flowering phenology and biotic interactions. *Trends Ecol. Evol.* **22**, 432–439 (2007).

49. Lim, P. O., Woo, H. R. & Nam, H. G. Molecular genetics of leaf senescence in *Arabidopsis*. *Trends Plant Sci.* **8**, 272–278 (2003).

50. Archetti, M., Richardson, A. D., O’Keeffe, J. & Delpierre, N. Predicting climate change impacts on the amount and duration of autumn colors in a New England forest. *PLoS ONE* **8**, e57373 (2013).

51. Nemani, R. R. et al. Climate-driven increases in global terrestrial net primary production from 1982 to 1999. *Science* **300**, 1560–1563 (2003).

52. Piao, S. L. et al. Net carbon dioxide losses of northern ecosystems in response to autumn warming. *Nature* **451**, 49–52 (2008).

53. Richardson, A. D. et al. Influence of spring and autumn phenological transitions on forest ecosystem productivity. *Philos. Trans. R. Soc. Lond. B Biol. Sci.* **365**, 3227–3246 (2010).

54. Cleland, E. E., Chiariello, N. R., Loarie, S. R., Mooney, H. A. & Field, C. B. Diverse responses of phenology to global changes in a grassland ecosystem. *Proc. Natl Acad. Sci. USA* **103**, 13740–13743 (2006).

55. Keller, F. & Körner, C. The role of photoperiodism in alpine plant development. *Arct. Antarct. Alp. Res.* **35**, 361–368 (2003).

56. Dörr, T. et al. Plant functional traits mediate reproductive phenology and success in response to experimental warming and snow addition in Tibet. *Glob. Chang. Biol.* **19**, 459–472 (2013).

57. Kearns, C. A. Anthophilous fly distribution across an elevation gradient. *Am. Midl. Nat.* **127**, 172–182 (1992).

58. Bingham, R. A. & Orthner, A. R. Efficient pollination of alpine plants. *Nature* **391**, 238–239 (1998).

59. Forrest, J. R. K. & Thomson, J. D. An examination of synchrony between insect emergence and flowering in Rocky Mountain meadows. *Ecol. Monogr.* **81**, 469–491 (2011).

60. Scheiner, S. M. & Gurevitch, J. (eds) *Design and Analysis of Ecological Experiments* (Oxford University Press, 2001).

61. R Development Core Team. *A Language and Environment for Statistical Computing* (R Foundation for Statistical Computing, 2015).

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X.E.L., L.L.J., F.D.M., S.P.W., H.S.N. and A.M.I. contributed equally to the paper. S.P.W. and X.Q.Z. designed the research; J.C.D., Z.H.Z., C.Y.L., Y.N.L., M.Y.D. and L.Z. performed the research; X.E.L., F.D.M., H.S.N., L.R.Z., Y.M.L. and X.Y.B. analysed the data; and X.E.L., L.L.J., F.D.M., S.P.W., H.S.N., A.M.I., J.P. and G.J.W. wrote the paper.

### Additional information

**Supplementary Information** accompanies this paper at http://www.nature.com/naturecommunications

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