- Phenological differences among species explain why early leafout
- extends the calendar but not thermal growing season
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5 1 Abstract

- 16 The extension of the growing season in the spring in temperate regions is one the most prominent biological
- 17 indicators of climate change. Most models of carbon storage assume that earlier growth will result in longer
- seasons and thereby enhance forest carbon storage, however current findings have challenged this assumption.

A recent hypothesis that has gained support in the literature is that plants dynamically adjust the end of their growing season based on their carbon-sink capacity, such that longer seasons do not increase total productivity.

If this is the case, variation in the calendar growing season (number of days) should be independent of variation in the thermal growing season (i.e., the period of favorable meteorological conditions for plant growth), which should remain relatively stable across years. We tested this prediction using rarely available plant-scale phenological measurements from a three year common garden experiment. The garden included 18 woody species, native to the Eastern United States from four populations of origin.

We found that at the community level, earlier leafout was correlated with earlier budset, resulting in a relatively stable calendar growing period over the three years. Because thermal conditions were lower at the start than end of the season, the relationship between earlier leafout and budset resulted in shorter thermal growing seasons when leafout was early—a relationship that was strongest in early-leafing species.

This relationship may explain some of the contrasting results of how climate change affects growing season length and productivity—if early leaf out results in earlier budset, as a whole plant are active for a proportionately large period of time where thermal conditions are less thermally optimal for photosynthesis.

This study shows linking phenological growing seasons to carbon storage requires integrating existing ecosystem measures with phenological variation at multiple scales, and that progress will require more efforts to understand and model species-level shifts in phenology.

36 2 Introduction

Terrestrial forests currently sequester 20% of greenhouse gas emissions annually (Shanley et al., 2024; Roe et al., 2021), providing a significant mitigation pathway for climate change. In mid and high latitudes, net carbon uptake is primarily determined by the length of the growing season (White et al., 1999). Most models of carbon storage assume that earlier spring leafout with climate change will drive longer seasons and increased carbon storage, in part offsetting future warming (Churkina et al., 2005; White et al., 1999;

- Keenan et al., 2014). Current findings, however, have called this major assumption into question. Recent research has suggested that plants adjust their end of season timing dynamically such that longer seasons do not increase total productivity, but the mechanism—and prevalence—of this effect is unclear (Zani et al., 2020; Norby, 2021a; Zohner et al., 2023).
- These contrasting results suggest fundamental gaps in our understanding of how early-season events, and the growing conditions of the vegetative period, shape end-of-season events. Some of this uncertainty is undoubtedly related to how the start and end of the season is measured, and the scale of observation. Many studies that have found growing season extension to be uncouple from increased prodictivity have used either largescale satellite measurements or small-scale single species 'pot' experiments, and contrast with findings from long-term large-scale CO_2 enrichment studies (Norby, 2021b). While results from satellite-derived metrics of 51 phenology (e.g., using NDVI) show a correlation between the start and end of the season, the signature that 52 is measured for 'end of season' is not clearly tied to a plant-scale event. Yet any connections would likely start at the individual plant level, where we rarely, if ever, have good measures of start and end of season events together in natural communities. While small scale pot experiments are better for capturing these plant-level measurements, results to date generally cannot differentiate between different species. Further, because end-of-season events are often more locally adapted than start-of-season events—with plants using unique photoperiods to cue important events such as budset (Bauerle et al., 2012; Soolanayakanahally et al., 2013)—these trends may vary across populations (Aitken & Bemmels, 2016).
- Additional uncertainty may stem from the fact that there are multiple ways to measure the growing season (Körner et al., 2023). The most common is calendar time (i.e., the number of days between the start and end of the growing season). Some studies have found that calendar growing seasons have lengthened with climate change (Menzel & Fabian, 1999; Liu et al., 2010), but several recent studies have found that earlier leafout is often correlated with earlier end-of-season events (Zani et al., 2020; Liu et al., 2016; Keenan & Richardson, 2015).
- The thermal growing season—defined here as the period of favorable meteorological conditions for plant

growth (Körner et al., 2023)—offers an alternative to the calendar growing season. It is also widely used, 67 most commonly in the metric of growing degree days, a temperature derived measure of time that accumulates when temperatures are above a certain minimal threshold (Chuine, 2000; Moore & Remais, 2014; Yang et al., 69 1995). Because plant photosynthetic rates increase with temperature (Farguhar et al., 1980) the thermal growing season is mechanistically related to primary productivity, and thus may be a better proxy than calendar time for relating growing season length to carbon gain. Importantly, depending on the temperature over the course of a growing season, years with substantially different calendar growing seasons can have very similar thermal growing seasons, or vice versa. Recent work has suggested early, increased productivity in the calendar growing season drives early senescence (Zani et al., 2020), and proposed that plants adjust their phenological trajectory mid-season based on a combination of growing season temperature and daylength (Zohner et al., 2023), thus maintaining a consistent thermal growing season. Here, we examine how start-of-season events (i.e., leafout) may affect end-of-season events (i.e., budset) to determine the length of the growing season. Using rarely available plant-scale data—phenological observations over three years from a multi-species common garden study—we test how correlated leafout and budset are in different species from four populations across a 4.5 latitudinal cline and examine which phenological event more strongly influences variation in growing season length. From this, we can understand how start- and end-of-season events together impact the calendar growing season, and how this relates to the thermal growing season, which connects to potential productivity. This study offers insights into physiology that will allow us

3 Results & Discussion

87 3.1 Variation in leafout and budset

Our multi-species common garden captured high variation in leafout, budset and leaf coloration, allowing us to
examine how these phases correlate, but also providing important insights into how both vary across species,

to scale from ecosystem level observations to individual mechanisms too, and improve forecasting.

populations and years. Consistent with the high number of studies that have found species and year-to-year environmental variation drive leafout variation (Delpierre et al., 2024; Donnelly et al., 2017; Polgar & Primack, 2011), we found high variance in leafout timing among species and years ($\sigma_{species}$:8.12 UI_{95} [4.96,11.65], σ_{year} :10.61 UI_{95} [3.37,21.19], Figure 1a,b). Population level variation was low ($\sigma_{population}$:0.64, UI_{95} [0,1.76], Figure 1c). Sambucus racemosa was typically the first species to leafout in the spring, leafing out approximately two weeks before Sorbus americana, the last species to leaf out (Figure 1a). There were no differences in leafout timing among the four populations included in our study (Figure 1b). Leafout was the earliest in 2019 and the latest in 2020 (Figure 1c).

Spring phenological phases are reported to be more plastic than autumn ones (McKown et al., 2014; Aitken & Bemmels, 2016; Vico et al., 2021), but we found that, relative to leafout, variance in budset timing was higher for species, years and populations ($\sigma_{species}$:9.77 UI_{95} [6.25,13.97], σ_{year} :15.25 UI_{95} [4.38,32.2], $\sigma_{population}$:2.47, UI_{95} [0,6.56], Figure 1), but followed similar relative contributions (highest variance in year, lowest in population). Budset was earliest for Amelanchier canadensis and latest for Alnus incana and Betula paperifera with more than three weeks between them (Figure 1b). Following trends in leafout and our finding that earlier leafout correlates with earlier budset, 2019 had the earliest budset and 2020 the latest (Figure 1a).

These results are somewhat surprising as autumn phenological phases like budset and leaf coloration are commonly thought to be strongly dependent on population, with different populations requiring different critical photoperiods to trigger these phases and leading to relatively stable phase dates across years (Soolanayakanahally et al., 2013), (and find some review paper). Supporting this—and in contrast to leafout—we found that populations did vary in their budset and their leaf coloration. Populations from the Second College Grant (the site with the fewest frost free days) set buds approximately two days before those from the White Mountains, but these differences were weak (Figure 1b) and small. Leaf coloration for populations from the Second College Grant occurred about three days before populations from the White Mountains and four days before populations from St. Hippolyte and Harvard Forest, qualitatively following an expected cline of frost free days

(ADD SUPPLIMENTAL FIGURE), though as with budset, these trends were weak and small. Our results suggest we need much more work on additional species, as results to date have focused mainly on one genus (*Populus*) and more efforts to understand how environmental factors beyond photoperiod may affect budset and senescence. Even for *Populus balsamifera*, the species suggested to be mainly photoperiod-controlled, recent work suggests temperature may also play a major role (Michelson *et al.*, 2018).

Budset and leaf coloration phenology were generally correlated (put coefficient in here), but we observed 120 notable species-level variation. For example species like Sambucus racemosa, Spirea tomentosa were about 121 the lastest to set but had earlier leaf coloration relative the trest of the community. Other species like Betula populifolia and Alnus incana had both later budset and leaf coloration relative to the community. These 123 differences may reflect High variation across species in their leafout and budsetlead to species driving the 124 most variation in primary growing season length ($\sigma_{species}$:14.37, UI_{95} [9.33,20.48], with less variation among 125 years (σ_{year} :4.89, UI_{95} [0.38,13.3], Figure 1b) and little variation explained by population ($\sigma_{population}$:2.48, 126 $UI_{95}[0,6.18]$, Figure 1). Due to it's early leafout and late budset S. racemosa had the longest calendar growing 127 season of the species in our study. A. canadensis and S. americana had the shortest growing seasons, though 128 for A. canadensis this was due to early budset and for S. americana late leafout (Figure 1a). Population 129 level differences in calendar growing season were determined by differences in budset, and followed the same pattern with the Second College Grant populations marginally earliest and White Mountains latest, with high uncertainty (Figure 1b). (Figure 1b).

3.2 Comparing calendar to thermal growing seasons: why longer calendar seasons may not result in increased primary productivity

Across all species, populations and years, a later start of spring (i.e., leafout) in calendar time was associated with shorter growing seasons (mean estimated effect: -0.44, $UI_{95}[-0.73, -0.16]$). This trade-off in earlier leafout and shorter calendar-time growing season was apparent in all species in our common garden, but to varying degrees (Fig. 2a,b). When we controlled for species and population level differences, however, we

found earlier leafout led to earlier budset (Pearson's correlation coefficient of 0.32 $CI_{95}[0.25, 0.39]$), resulting in a relatively stable calendar growing season 1c).

In contrast, we found an apparently fundamental—and unexpected trade-off—between early, longer calender growing seasons and thermal growing seasons. Across species, populations and years, later leafout in calendar time resulted in a longer thermal growing season (mean estimated effect: $5.49, UI_{95}[1.54, 9.23]$; Figure 2c).

These contrasting results—of a relatively stable growing season measured in calendar days at the community scale, but one that is 'shorter' in thermal time with earlier leafout—may explain some of the contrasting results of how climate change affects end-of-season events and productivity (Zani et al., 2020).

Our results suggest that earlier leafout may have little effect on the thermal growing season because of 147 unfavorably low temperatures combined with the observed correlation between leafout and budset (Figure 148 1). Given that photosynthesis is limited by cold temperatures in early spring, earlier leafout appear to provide 149 limited opportunity for substantial growth yet may deprive plants of fully using late-season warmth. This 150 may explain why multiple studies have failed to find correlations between longer seasons and increased plant 151 growth (Cufar et al., 2015; Camarero et al., 2022; Dow et al., 2022; Silvestro et al., 2023). This relationship, 152 however, was strongly species-dependent. Later leafout in calendar time led to longer thermal seasons in 153 species that typically leaf out earlier in the spring relative to others. This included shrubs such as Sambucus 154 racemosa, Viburnum cassiodes, Spirea alba, Diervella lonicera, Aronia melanocarpa Spirea tomentosa and the 155 tree species Betula populifolia. Later-leafout species showed a weaker relationship (Betula papyrifera, Betula 156 allegheniensis and Alnus incana, Figure 2d). 157

We can see these dynamics play out by tracking the phenology of four individuals plants as an example.

The earlier individual of *Aronia melanocarpa* (Figure 3, dark green bars) starts growing 24 days before a

later individual (light green bars), but only ceases 13 days before it (i.e., it has a 14 day longer calendar

growing season). However, because the 24 day growth advantage it has occurs when thermal conditions are

less favorable, it ends up having a shorter thermal growing season (i.e., less change for carbon assimilation)

than its later conspecific (Figure 3a,b). This is not the case for the later leafing species *Myrica gale* where

the both the earlier (dark blue bars) and later leafing individual (light blue bars) start growing under more optimal thermal conditions, so the 20 day "head start" the earlier individual incurs results in a both a longer calendar and thermal growing season (Figure 3a,c).

₆₇ 3.3 Ecological and forecasting implications

Our multi-species common garden study showed that for early-leafing species, earlier leafout does not extend their thermal growing season—a proxy for potential carbon uptake period—despite extending the calendar 169 growing season. For early-leafout species, years with delayed leafout resulted in a longer thermal growing 170 season. This relationship was in part explained by positive correlations between leafout and budset where 171 later leafing individuals also set buds later, extending their growth into that later part of the season when 172 thermal conditions were more favorable. For later leafing species, earlier leafout did not substantially reduce 173 their thermal growing season. This is likely because for them, an earlier leafout still occurred in thermally 174 favorable times of the season, and a relatively small advance in calendar time resulting in a proportionately 175 larger gain in thermal sums. 176

Our results show that there is little advantage from a carbon or primary productive perspective for leafing out
too early in the season, as thermal conditions are not favorable for photosynthesis and assimilation. Many of
the species that are most phenologically sensitive to climate change are already among the earliest species to
leaf out in temperate plant communities (Shen et al., 2014; Geng et al., 2020), implying there may be little
to gain from a carbon perspective.

This result raises question about why some species leafout early during these unfavorable conditions, and why species tracking spring warming due to climate change have increased performance relative to non-trackers (Cleland et al., 2012). In our study we only evaluated the thermal conditions that may affect photosynthesis, rather than photosynthesis itself, which also depends strongly on light availability. In forest systems, light availability is strongly dependent on canopy conditions, and is highly dependent on biotic interactions. In our study, the species that leafed out the earliest are understory shrubs, for whom access to light becomes

severely limited later in the growing season as canopy trees leaf out. It may be that for these species, the light availability early in the season necessitates leafing out in sub-optimal thermal conditions. In fact, some studies suggest that understory shrubs get all/most of their carbon before canopy closure (AUGSPURGER et al., 2005).

This study shows linking phenological growing season to primary productivity requires accounting for phe-192 nological variation at multiple scales (individual, species level, multiple phases). These results suggest that 193 progress will require more efforts to understand and model species-level shifts in phenology. While satellite 194 observations can document intriguing trends (e.g., Zohner et al., 2023), observations that include far more 195 species at a finer scale are likely critical for mechanistic understanding. In particular our results suggest 196 budset may be far more variable year to year than often suggested (Michelson et al., 2018) but see(McKown 197 et al., 2014; Vander Mijnsbrugge et al., 2016). Recent work supports this, by highlighting an important role 198 for temperature alongside photoperiod, in driving budset (Olsen et al., 2014; Rohde et al., 2011). Further, 199 most work on budset in broadleaf species comes from one genus (*Populus spp.*) highlighting a critical gap, 200 as a focus on single species may not always predict broad scale trends (Morales-Castilla et al., 2024). 201

Of course, budset is just one of many ways that plants begin to transition from growth to dormancy each year

generally at different times for different events (Michelson et al., 2018). Understanding which metrics of

end of season events correlate best with growth and carbon gain are well-established critical needs (Gallinat

et al., 2015). Our results suggest we also need to study how these events shift with earlier and warmer years.

Studies of leaf longevity have begun to examine this, but more work across different metrics of end of season

and across many more species is critical.

$_{208}$ 4 Methods

209 4.0.1 The Common Garden

In 2014-2015, we collected seeds of 18 species woody plants from multiple parent plants at four field sites in eastern Northern America spanning approximately a 3.5 degree latitudinal gradient. The four field sites included Harvard Forest (42.55°N, 72.20°W), the White Mountains (44.11°N, 52.14°W), Second College Grant of Dartmouth College (44.79°N, 50.66°W), and St. Hippolyte, QC, CAN (45.98°N, 74.01°W). We transported all seeds back the the Weld Hill Research Building at the Arnold Arboretum in Boston MA 214 (42.30°N, 71.13°W) where we germinated seeds following standard germination protocols, and grew them to 215 seedling stages in the research greenhouse. In the spring of 2017 we out-planted seedlings to establish the 216 Common garden at Weld Hill. Plantings were randomized between 16 plot blocks. Individuals that were too 217 small to survive outside were maintained in the growth facilities for an additional year and out-planted in 218 the early spring of 2018. Plots were divided between tree plots which included species Acer pensylvanicum, 219 Amelanchier canadensis, Alnus incana, Betula papyrifera, Betula populifolia, Beluta alleghaniensis, Quercus 220 alba, and Quercus rubra and shrub plots which included the remaining species and shade cloth (Table 1). 221 Plots were regularly weeded and watered throughout the duration of the study and were pruned in the fall 222 of 2020. Survival of Acer spicatum, Acer pensylvanicum, Vaccinium myrtilloides Quercus alba, and Quercus 223 rubra was limited and these species are therefore not included in the following analyses. Based on survivorship 224 in the common garden, our subsequent analyses are based on offspring from 1-10 individual parent plants 225 per field site.

227 4.0.2 Phenological monitoring:

For the years of 2018-2019, we made phenological observations of all individuals in the common garden twice
per week from February to December. In 2020 due to the COVID 19 pandemic, we monitored once per week
from March to November. We describe phenological stages using a modified BBCH scale (Finn et al., 2007)

a common metrics for quantify woody plant phenological progression. We observed all major vegetative stages (budburst BBCH 07, leafout BBCH 15, end of leaf expansion bbch 19, leaf coloration/drop BBCH 97, reproductive phases flowering BBCH 60-65, fruiting BBCH 72-79 and fruit/cones fully ripe BBCH 89). We added additional phases for budset and labelled full budset as BBCH 102.

235 4.0.3 Data analysis

To better understand the role that variation in leafout and budset phenology play in determining calendar growing season length among species populations and years we fit a Bayesian hierarchical model with a normal (Gaussian) probability distribution. We calculated growing season duration by subtracting the day of leafout from the day of budset. We fit an intercept only model with phenophase timing (leafout, budset or growing season duration) as the response variable and partial pooling across species, populations and years. We only included observations with both budset and leafout observed on the same plant in this analysis (n= 595).

To assess the relationship between variation in leafout timing and calendar and thermal growing seasons we
fit two additional regression models with thermal or calendar growing season length as the response variable
and day of leafout as the main prediction. To account for species-level differences we included partially
pooling on the slope and intercept of species.

We define the thermal growing season as the cumulative growing degree day heat sums between the day of leafout and the day of budset for each species. We calculated daily heat sums using the R package "pollen" (Jakub Nowosad, 2019) using a 10 C base temperature with minimum and maximum daily temperature data from the weld hill weather station.

All models were fit in the R package "brms" (Bürkner, 2018) on 4 chains with a 4000 iteration warm-up and 1000 sampling iterations on each chain for a total of 4,000 sampling iterations across all chains. We evaluated model fit, with no divergent transitions, rhats less than 1.01 and high effective sample sizes. We performed all analyses in R version 4.1.2 (R Core Team, 2021).

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²⁵⁸ 6 Figures

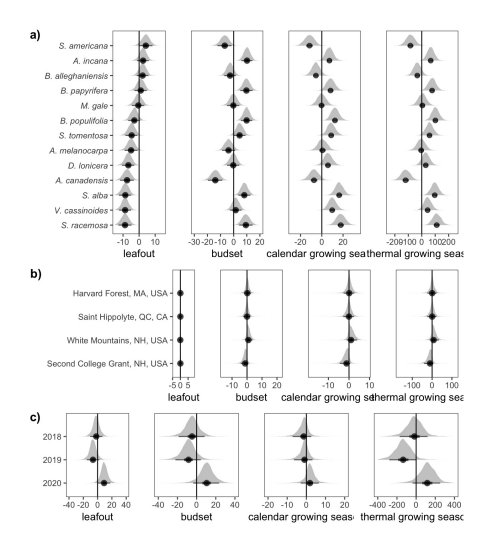


Figure 1: Difference in leafout, budset and calendar growing season length partitioned between species (a) populations (b) and years (c). Point represent the median effect size estimate, and bars the 50% uncertainty intervals. The grey distribution depict the full uncertainty estimate around the estimate.

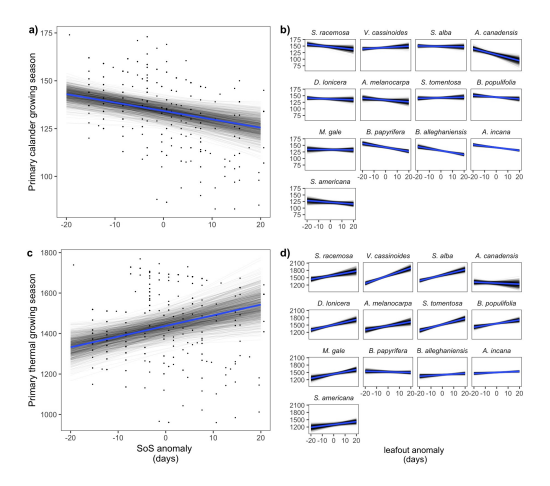


Figure 2: The relationship between Start of Spring (SoS; calendar day of leafout) and growing season length differs between the calendar growing season and the thermal growing season. Later leafout resulted in a shorter calendar growing season (a) and this pattern was consistent across species in our study (b). In contrast, an increasing later SoS resulted in a longer thermal growing season (c) though this effect was stronger for species that typically leafout earlier in the season—panels in c) display in the typical order of leafout among species. The blue trend lines represent the mean effect of SoS timing on growing season length and black lines represent 1000 draws from the posterior distribution as a measure of uncertainty. Points in a), and c) represent the raw data.

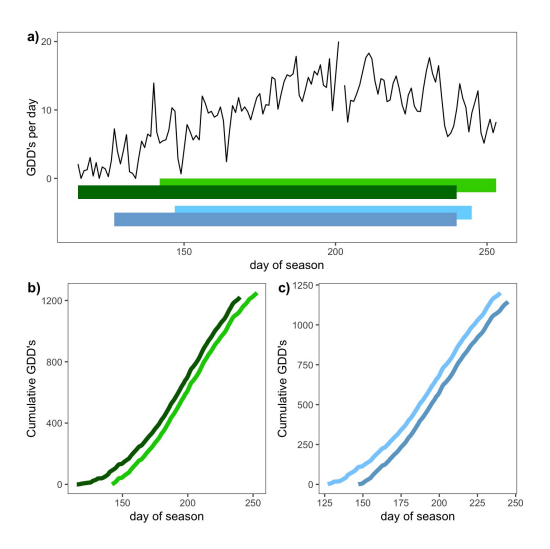


Figure 3: Thermal conditions vary across the calendar growing season, which can generate a complex relationship between the calendar and thermal growing seasons. Panel a) depicts the daily heat sums at the Weld Hill Research Building in 2019 and the calendar growth season of early and late leafing individuals of Aronia melanocarpa (green bars) and Myrica gale (blue bars). Despite the fact the the early individual of A. melanocarpa leafs out 24 days before it's later con-generic and only sets bud 13 days before it (i.e., it has a 14 day longer calendar growing season) it's thermal growing season is shorter (panel b) because most of its growth advantage (explain this better) takes place in the unfavorable early spring. In contrast for M. gale where both the early and late individual leaf out later in the spring, the 20 day head start and 5 day earlier finish of the earlier individual (15 day longer calendar growing season) results in a longer thermal growing season for it as well (panel c)

Table 1: Species list

Species	functional group	n
Acer pensylvanicum	tree	2
Acer spicatum	tree	NA
$Alnus\ incana$	shrub	31
$A melanchier\ canadensis$	shrub	6
$Aronia\ melanocarpa$	shrub	12
$Betula\ alleghaniens is$	tree	24
Betula papyrifera	tree	13
Betula populifolia	tree	24
Diervilla lonicera	shrub	16
Myrica gale	shrub	15
Quercus alba	tree	NA
Quercus rubra	tree	NA
$Sambucus\ racemosa$	shrub	11
Sorbus americana	shrub	5
Spiraea alba	shrub	19
$Spiraea\ tomentosa$	shrub	21
Vaccinium myrtilloides	shrub	NA
$Viburnum\ cassinoides$	shrub	25

References

- Aitken, S. N. & Bemmels, J. B. (2016). Time to get moving: assisted gene flow of forest trees. *Evol Appl*, 9, 271–90.
- AUGSPURGER, C. K., CHEESEMAN, J. M. & SALK, C. F. (2005). Light gains and physiological capacity of understorey woody plants during phenological avoidance of canopy shade. *Functional Ecology*, 19, 537–546.
- Bauerle, W. L., Oren, R., Way, D. A., Qian, S. S., Stoy, P. C., Thornton, P. E., Bowden, J. D., Hoffman, F. M.
 & Reynolds, R. F. (2012). Photoperiodic regulation of the seasonal pattern of photosynthetic capacity and
 the implications for carbon cycling. Proceedings of the National Academy of Sciences, 109, 8612–8617.
- Bürkner, P.-C. (2018). Advanced bayesian multilevel modeling with the r package brms. *R Journal*, 10, 395–411.
- Camarero, J. J., Campelo, F., Colangelo, M., Valeriano, C., Knorre, A., Solé, G. & Rubio-Cuadrado, Á.
 (2022). Decoupled leaf-wood phenology in two pine species from contrasting climates: Longer growing
 seasons do not mean more radial growth. Agricultural and Forest Meteorology, 327, 109223.
- ²⁷² Chuine, I. (2000). A unified model for budburst of trees. Journal of Theoretical Biology, 207, 337–347.
- Churkina, G., Schimel, D., Braswell, B. H. & Xiao, X. (2005). Spatial analysis of growing season length control over net ecosystem exchange. *Global Change Biology*, 11, 1777–1787.
- Cleland, E. E., Allen, J. M., Crimmins, T. M., Dunne, J. A., Pau, S., Travers, S. E., Zavaleta, E. S. & Wolkovich, E. M. (2012). Phenological tracking enables positive species responses to climate change.

 Ecology, 93, 1765–1771.
- Čufar, K., De Luis, M., Prislan, P., Gričar, J., Črepinšek, Z., Merela, M. & Kajfež-Bogataj, L. (2015). Do variations in leaf phenology affect radial growth variations in Fagus sylvatica? International journal of biometeorology, 59, 1127–1132.
- Delpierre, N., Garnier, S., Treuil-Dussouet, H., Hufkens, K., Lin, J., Beier, C., Bell, M., Berveiller, D.,
 Cuntz, M., Curioni, G., Dahlin, K., Denham, S. O., Desai, A. R., Domec, J.-C., Hart, K. M., Ibrom, A.,
 Joetzjer, E., King, J., Klosterhalfen, A., Koebsch, F., McHale, P., Morfin, A., Munger, J. W., Noormets, A.,
 Pilegaard, K., Pohl, F., Rebmann, C., Richardson, A. D., Rothstein, D., Schwartz, M. D., Wilkinson, M.
 Soudani, K. (2024). Phenology across scales: An intercontinental analysis of leaf-out dates in temperate
 deciduous tree communities. Global Ecology and Biogeography, 33, e13910.
- Donnelly, A., Yu, R., Caffarra, A., Hanes, J., Liang, L., Desai, A. R., Liu, L. & Schwartz, M. D. (2017).
 Interspecific and interannual variation in the duration of spring phenophases in a northern mixed forest.

 Agricultural and Forest Meteorology, 243, 55–67.
- Dow, C., Kim, A. Y., D'Orangeville, L., Gonzalez-Akre, E. B., Helcoski, R., Herrmann, V., Harley, G. L.,
 Maxwell, J. T., McGregor, I. R., McShea, W. J. et al. (2022). Warm springs alter timing but not total
 growth of temperate deciduous trees. Nature, 608, 552–557.
- Farquhar, G. D., von Caemmerer, S. & Berry, J. A. (1980). A biochemical model of photosynthetic co2 assimilation in leaves of c3 species. *Planta*, 149, 78–90.
- Finn, G., Straszewski, A. & Peterson, V. (2007). A general growth stage key for describing trees and woody plants. *Annals of Applied Biology*, 151, 127–131.
- Gallinat, A. S., Primack, R. B. & Wagner, D. L. (2015). Autumn, the neglected season in climate change research. *Trends in Ecology Evolution*, 30, 169–176.

- Geng, X., Fu, Y. H., Hao, F., Zhou, X., Zhang, X., Yin, G., Vitasse, Y., Piao, S., Niu, K., De Boeck,
 H. J., Menzel, A. & Peñuelas, J. (2020). Climate warming increases spring phenological differences among
 temperate trees. *Global Change Biology*, 26, 5979–5987.
- Jakub Nowosad (2019). pollen: Analysis of Aerobiological Data.
- Keenan, T. F., Gray, J., Friedl, M. A., Toomey, M., Bohrer, G., Hollinger, D. Y., Munger, J. W., O'Keefe, J., Schmid, H. P., Wing, I. S., Yang, B. & Richardson, A. D. (2014). Net carbon uptake has increased through warming-induced changes in temperate forest phenology. *Nature Climate Change*, 4, 598–604.
- Keenan, T. F. & Richardson, A. D. (2015). The timing of autumn senescence is affected by the timing of spring phenology: implications for predictive models. *Global Change Biology*, 21, 2634–2641.
- Körner, C., Möhl, P. & Hiltbrunner, E. (2023). Four ways to define the growing season. *Ecology Letters*, 26, 1277–1292.
- Liu, B., Henderson, M., Zhang, Y. & Xu, M. (2010). Spatiotemporal change in china's climatic growing season: 1955–2000. *Climatic Change*, 99, 93–118.
- Liu, Q., Fu, Y. H., Zhu, Z., Liu, Y., Liu, Z., Huang, M., Janssens, I. A. & Piao, S. (2016). Delayed autumn phenology in the northern hemisphere is related to change in both climate and spring phenology. *Global Change Biology*, 22, 3702–3711.
- McKown, A. D., Guy, R. D., Klápště, J., Geraldes, A., Friedmann, M., Cronk, Q. C. B., El-Kassaby, Y. A.,
 Mansfield, S. D. & Douglas, C. J. (2014). Geographical and environmental gradients shape phenotypic
 trait variation and genetic structure in opulus trichocarpa. New Phytologist, 201, 1263–1276.
- Menzel, A. & Fabian, P. (1999). Growing season extended in europe. Nature, 397, 659-659.
- Michelson, I. H., Ingvarsson, P. K., Robinson, K. M., Edlund, E., Eriksson, M. E., Nilsson, O. & Jansson, S. (2018). Autumn senescence in aspen is not triggered by day length. *Physiologia Plantarum*, 162, 123–134.
- Moore, J. L. & Remais, J. V. (2014). Developmental models for estimating ecological responses to environmental variability: Structural, parametric, and experimental issues. *Acta Biotheoretica*, 62, 69–90.
- Morales-Castilla, I., Davies, T. J., Legault, G., Buonaiuto, D. M., Chamberlain, C. J., Ettinger, A. K.,
 Garner, M., Jones, F. A. M., Loughnan, D., Pearse, W. D., Sodhi, D. S. & Wolkovich, E. M. (2024).
 Phylogenetic estimates of species-level phenology improve ecological forecasting. *Nature Climate Change*,
 14, 989–995.
- Norby, R. J. (2021a). Comment on "increased growing-season productivity drives earlier autumn leaf senescence in temperate trees". *Science*, 371, eabg1438.
- Norby, R. J. (2021b). Comment on "increased growing-season productivity drives earlier autumn leaf senescence in temperate trees". *Science*, 371, eabg1438.
- Olsen, J. E., Lee, Y. & Junttila, O. (2014). Effect of alternating day and night temperature on short dayinduced bud set and subsequent bud burst in long days in norway spruce. Frontiers in Plant Science, 5.
- Polgar, C. A. & Primack, R. B. (2011). Leaf-out phenology of temperate woody plants: from trees to ecosystems. *New Phytologist*, 191, 926–941.
- R Core Team (2021). R: A Language and Environment for Statistical Computing. R Foundation for Statistical Computing, Vienna, Austria.

- Roe, S., Streck, C., Beach, R., Busch, J., Chapman, M., Daioglou, V., Deppermann, A., Doelman, J., Emmet-Booth, J., Engelmann, J., Fricko, O., Frischmann, C., Funk, J., Grassi, G., Griscom, B., Havlik,
- P., Hanssen, S., Humpenöder, F., Landholm, D., Lomax, G., Lehmann, J., Mesnildrey, L., Nabuurs, G.-J.,
- Popp, A., Rivard, C., Sanderman, J., Sohngen, B., Smith, P., Stehfest, E., Woolf, D. & Lawrence, D.
- (2021). Land-based measures to mitigate climate change: Potential and feasibility by country. Global
- Change Biology, 27, 6025–6058.
- Rohde, A., Bastien, C., Boerjan, W. & Thomas, S. (2011). Temperature signals contribute to the timing of photoperiodic growth cessation and bud set in poplar. *Tree Physiology*, 31, 472–482.
- Shanley, C. S., Graves, R. A., Drever, C. R., Schindel, M., Robertson, J. C., Case, M. J. & Biswas, T. (2024).

 Mapping forest-based natural climate solutions. *Communications Earth & Environment*, 5, 502.
- Shen, M., Tang, Y., Chen, J., Yang, X., Wang, C., Cui, X., Yang, Y., Han, L., Li, L., Du, J., Zhang, G. & Cong, N. (2014). Earlier-season vegetation has greater temperature sensitivity of spring phenology in northern hemisphere. *PLOS ONE*, 9, 1–11.
- Silvestro, R., Zeng, Q., Buttò, V., Sylvain, J.-D., Drolet, G., Mencuccini, M., Thiffault, N., Yuan, S. & Rossi, S. (2023). A longer wood growing season does not lead to higher carbon sequestration. *Scientific reports*, 13, 4059.
- Soolanayakanahally, R. Y., Guy, R. D., Silim, S. N. & Song, M. (2013). Timing of photoperiodic competency causes phenological mismatch in balsam poplar (populus balsamifera l.). *Plant, cell & environment*, 36, 116–127.
- Vander Mijnsbrugge, K., Depypere, L., Michiels, B. & De Cuyper, B. (2016). Genetic and temporal plastic variation in bud burst, bud set and flower opening responses of local versus non-local provenances of prunus spinosa in a provenance trial. *Basic and Applied Ecology*, 17, 262–272.
- Vico, G., Karacic, A., Adler, A., Richards, T. & Weih, M. (2021). Consistent poplar clone ranking based on leaf phenology and temperature along a latitudinal and climatic gradient in northern europe. *BioEnergy Research*, 14, 445–459.
- White, M. A., Running, S. W. & Thornton, P. E. (1999). The impact of growing-season length variability on carbon assimilation and evapotranspiration over 88 years in the eastern us deciduous forest. *International Journal of Biometeorology*, 42, 139–145.
- Yang, S., Logan, J. & Coffey, D. L. (1995). Mathematical formulae for calculating the base temperature for growing degree days. *Agricultural and Forest Meteorology*, 74, 61–74.
- Zani, D., Crowther, T. W., Mo, L., Renner, S. S. & Zohner, C. M. (2020). Increased growing-season productivity drives earlier autumn leaf senescence in temperate trees. *Science*, 370, 1066–1071.
- Zohner, C. M., Mirzagholi, L., Renner, S. S., Mo, L., Rebindaine, D., Bucher, R., Palouš, D., Vitasse, Y., Fu, Y. H., Stocker, B. D. & Crowther, T. W. (2023). Effect of climate warming on the timing of autumn leaf senescence reverses after the summer solstice. *Science*, 381, eadf5098.