# False spring damage to temperate tree saplings is amplified with winter warming

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## Abstract

1. With warming temperatures, spring phenology (i.e., budburst and leafout) is advancing. Late spring freezes, however, may not advance at the same rate, leading to an increase in freezes that occur after trees initiate budburst—known as false springs—with continued warming. Understanding and predicting false springs with climate change requires understanding both how warming shifts budburst and the impacts of false springs on plant tissue. In particular, increasing research suggests warming winters may delay budburst through reduced chilling, which may also cause plants to leaf out slower or incompletely, subsequently decreasing spring freeze tolerance and potentially leading to higher damage from false

- spring events.
- 22 2. Here, we assessed the effects of over-winter chilling and false spring events on sapling phenology, growth
  23 and tissue traits, across eight temperate tree and shrub species in a lab experiment.
- 3. We found that false springs slowed subsequent budburst and leafout—extending the period of greatest risk for freeze damage, which increased damage to the shoot apical meristem, and decreased leaf toughness, leaf thickness and chlorophyll content. Longer chilling accelerated budburst and leafout, even under false spring conditions, thus chilling compensated for the adverse effects of false springs on phenology. Despite the effect of chilling on phenology, it did not cause any major re-ordering in the sequence of species leafout. We therefore expect climate change to reshape forest communities not through temporal reassembly but rather through impacts on growth and leaf traits from the coupled effects of false springs with decreases in over-winter chilling under future climate change scenarios.
- 4. **Synthesis:** With climate change and warming temperatures, over-winter chilling is anticipated to decrease and false springs are predicted to increase in certain regions. This combination could greatly impact plant performance, survival and shape species distributions, ultimately affecting important processes such as carbon uptake and nutrient cycling.
- Keywords: false spring, climate change, phenology, spring freeze, forest recruitment, temporal reassembly, budburst, temperate

# Introduction

The timing of spring in temperate deciduous forests shapes plant and animal communities and influences
ecosystem services from agriculture to forest management. With warming temperatures, spring phenology
(i.e., budburst and leafout, which are strongly cued by temperature) is advancing, causing longer growing
seasons (Chuine et al., 2001) and reshaping ecosystem dynamics. In one major example, advancing spring

phenology has led to increased carbon uptake across temperate forests, which are essential carbon sinks that
combat the negative effects of climate change (Keenan *et al.*, 2014). But climate change could diminish or
reverse these positive effects on carbon storage: specifically through cold snaps during the spring and warming
temperatures in the winter.

While climate change has warmed the Northern Hemisphere, cold snaps and freezing events are still occurring. These weather events can have big impacts on plant development each spring. One such event is known as a 'false spring', often defined as when temperatures drop below freezing (Schwartz et al., 2002, i.e., below -2.2°C) after budburst has initiated. Damage from false spring events can have cascading effects within a 50 forest, from changes to nutrient cycling and carbon uptake to shifts in seedling recruitment (Hufkens et al., 2012; Klosterman et al., 2018; Richardson et al., 2013). Furthermore, false springs can increase the chance of 52 additional freezes within a growing season by extending the period in which plants are most at risk—the time 53 between budburst and leafout (what we refer to as the 'duration of vegetative risk'), resulting in a positive feedback loop: observational studies suggest plants take longer to re-flush leaves after a false spring—up to 38 days (Augspurger, 2009, 2013; Gu et al., 2008; Menzel et al., 2015)—which could lead to additional false springs in a season (Augspurger, 2009). As false springs are predicted to increase in certain regions with increased climate change (Ault et al., 2015; Liu et al., 2018; Zohner et al., 2020) understanding their impacts is essential for robust forest management strategies and climate forecasting (Kral-O'Brien et al., 2019).

Warmer winters may also play a critical role in the future of forests as they directly impact one of the major cues plants use to time budburst: over-winter cold temperatures (chilling), in addition to warming spring temperatures (forcing) and longer daylengths (Chuine et al., 2016). Many temperate plants have evolved chilling requirements to avoid leafout during warm snaps in the middle of the winter, but with climate change, chilling requirements may not be met. If chilling is not met, plants may leaf out much slower or incompletely, which can in turn affect freeze tolerance. Thus more studies are needed in the face of climate change to understand the interplay of warming winters and false spring risk and the resulting impact to our forests.

Understanding how winter chilling and false springs may affect future forests requires knowing whether
this interaction varies across species within a community. This is especially true if species have evolved
along a trade-off of risking spring freezes for early access to resources. While a single-species perspective
may predict all individuals to require high levels of chilling to delay budburst and ultimately diminish false
spring risk, competition between species for nutrients, water and light resources in the early spring likely
pushes some species—and some individuals within species—to leafout earlier (Augspurger, 2013). Young
trees and understory species generally initiate budburst before canopy trees to benefit from higher light levels
(Augspurger, 2008; Vitasse et al., 2013), which potentially puts these species and individuals at higher risk
of freeze damage (Vitasse et al., 2014). Thus, successful forest recruitment requires seedlings and saplings to
minimize false spring risk while maximizing growth.

The combination of species- and lifestage-level differences in responses to false springs, chilling and climate change could reshape the temporal assembly of forest communities. Species typically leafout in a similar sequence, with understory species leafing out earlier and higher canopy trees leafing out last but many studies are predicting substantial shifts in chronological order and reassembly of species' leafout with climate change (Roberts et al., 2015; Laube et al., 2013). As warming increases winter temperatures and false spring prevalence, phenological cues and their interactions are anticipated to also change, which could greatly alter competition and recruitment among forest species for early season resources and ultimately impact species diversity and carbon uptake in temperate forests.

Here, we assessed the effects of over-winter chilling length and false springs on sapling phenology and growth across eight temperate tree and shrub species. We exposed individuals to different levels of over-winter chilling crossed with a false spring event in growth chambers, and then following individuals for a growing season we asked: (1) How does over-winter chilling impact phenology, growth and physical leaf traits (2) how do false spring events impact phenology, growth and physical leaf traits and (3) how does the interaction between chilling and false springs impact community structure and phenological order?

### Materials and Methods

#### 93 Plant Selection and Material

We selected eight temperate woody plant tree and shrub species that span varying spring phenologies (e.g.,
early to later leafout), that were not used as crops or ornamental species: Acer saccharinum, Alnus incana
rugosa, Betula papyrifera, Betula populifolia, Cornus racemosa, Salix purpurea, Sorbus americana, and Viburnum dentatum (we originally included two additional species—Fagus grandifolia and Nyssa sylvatica, but the
plants were not delivered in a usable condition and thus we excluded them from the experiment). We used 48
dormant, one year old, bare root saplings—each measuring 6-12 inches—for each species from Cold Stream
Farm LLC (Freesoil, MI; 44°6' N -86°12' W) for a total of 384 individuals. Upon receipt, plants were potted
in 656 ml deepots with Fafard #3B Metro Mix soil and placed in growth chambers at the Weld Hill Research
Building of the Arnold Arboretum (Boston, MA; 42°17' N -71°8' W) at 4°C for different durations depending
on chilling treatment (explained below).

#### 104 Growth Chamber and Greenhouse Conditions

Individuals were randomly selected for one of six experimental treatments from a full factorial design of false spring (two levels: presence or absence of false spring) x chilling (three levels: four, six or eight weeks of chilling at 4°C with eight hour photoperiod, lighting was a combination of T5HO fluorescent lamps with halogen incandescent bulbs at roughly 250  $\mu mol/m^2/s$ ). Individuals were rotated within and among growth chambers every two weeks to minimize bias from possible growth chamber effects.

Once each chilling treatment was completed, we moved individuals to a greenhouse with mean daytime temperature of 15°C and a mean nighttime temperature of 10°C, and a photoperiod of 12 hour days throughout the spring until all individuals reached full leaf expansion. After all individuals of all species reached full leaf expansion, greenhouse temperatures and photoperiods were kept ambient, and all individuals were up-potted to 983 ml deepots and fertilized with SCOTTS 15-9-12 Osmocote Plus 5-6.

### Phenology and False Spring Treatment

We recorded phenology (using the BBCH scale (Meier, 2001)) every 2-3 days through full leaf expansion. 116 Budburst was denoted as BBCH stage 07, which is 'beginning of sprouting or bud breaking' and monitored 117 until full leaf expansion (BBCH stage 19) in order to evaluate the duration of vegetative risk (Chamberlain 118 et al., 2019) for each individual (Finn et al., 2007). Individuals in the 'false spring' treatment were placed 119 in a growth chamber set to mimic a false spring event during budburst, defined as once at least 50% of the 120 buds were at BBCH stage 07 but the individual had not yet reached BBCH stage 19 (that is, each sapling 121 was exposed to a false spring based on its individual phenological timing). False spring treatments lasted 122 approximately 14 hours, beginning at 18:00; temperatures were ramped down over 14 hours (Figure 1). At approximately 08:00 the following day, we placed false spring individuals back in the greenhouse. Once all individuals reached full leaf expansion (BBCH stage 19), we recorded phenology weekly until August 1st, then every 2-3 days again to monitor fall phenology. We monitored all individuals until complete budset.

## 27 Growth measurements

We measured height three times throughout the growing season (the day an individual reached full leaf 128 expansion, 60 days after full leaf out and when an individual reached complete budset). We measured the chlorophyll content of four leaves on each individual 60 days after full leaf out using an atLEAF CHL PLUS Chlorophyll meter, converting chlorophyll content to mg/cm<sup>2</sup> using the atLEAF CHL PLUS conversion tool. We measured leaf thickness using a Shars Digital Micrometer (accurate to 0.001mm) and leaf toughness 132 in Newtons using a Shimpo Digital Force Gauge on two leaves for each individual 60 days after full leaf 133 out. Additionally, we visually monitored damage to the shoot apical meristem, which consisted of complete 134 damage or disruption of growth in the main stem and resulted in early dormancy induction or reliance on 135 lateral shoot growth. Finally, we harvested each plant after it reached complete budset to dry, separate and 136 weigh belowground and aboveground biomass (including leaves). 137

#### 138 Data analysis

We used Bayesian hierarchical models (with the brms package, Bürkner, 2017, version 2.3.1) in R (R Devel-139 opment Core Team, 2017, version 3.3.1) to estimate the effects of chilling duration, false spring treatment 140 and all two-way interactions as predictors on: (1) duration of vegetative risk, (2) growing season length, (3) 141 shoot apical meristem damage, (4) total growth in centimeters, (5) total biomass, (6) chlorophyll content, (7) 142 leaf toughness and (8) leaf thickness. We modeled species hierarchically as grouping factors, which generated an estimate and posterior distribution for each species as well as an overall response across the eight species used in our experiment. We ran four chains, each with 4 000 iterations, of which 2 500 were warm-up iterations, for a total of 6 000 posterior samples for each predictor for each model using weakly informative priors (increasing priors three-fold did not impact our results). We evaluated our model performance based on  $\hat{R}$ values that were close to one, checked chain convergence and posterior predictive checks visually (Gelman et al., 2014), and made sure all models had 0 divergent transitions. Our models generally had high  $n_{eff}$  (6 000 for most parameters, but as low as 1 400 for a couple of parameters in the shoot apical meristem model). 150 All estimated values are reported in the text as means +/- standard errors relative to the no false spring 151 under four weeks of chilling treatment. 152

## Results

Chilling durations impacted individual phenology. As seen in many other studies, we found increases in chilling advanced day of budburst by  $-2.8 \pm 1.7$  days for six weeks of chilling and by  $-7.6 \pm 1.8$  days for eight weeks of chilling (as mentioned above, all values are given relative to the no false spring under four weeks of chilling treatment; Table S1). Longer chilling also accelerated leafout, reducing the duration of vegetative risk (8 weeks:  $-2.7 \pm 1.1$  days; Figure 2a, Table S2 and Table S3). Increased chilling shortened the growing season for individuals exposed to eight weeks of chilling ( $-9.7 \pm 5$  days; Figure 2b and Table S4), due to constraints on our experimental design: individuals in longer chilling treatments were put in the greenhouse two to four

weeks later than the four weeks of chilling group but all individuals experienced the same ambient conditions
when in the greenhouse, including shifts in temperature and light associated in the late summer/autumn that
generally trigger dormancy onset; thus shortening the growing season for the groups that experienced longer
chilling.

False springs also impacted individual phenology. Individuals exposed to the false spring treatment had longer durations of vegetative risk given four weeks of chilling (3  $\pm$  0.8 days) and slightly longer durations for given six weeks of chilling (1.5  $\pm$  1.1 days; Figure 2a and Table S3). Effects on the duration of vegetative risk from false spring treatments (which led to longer durations) and from increased chilling (which led to shorter durations) were generally additive, resulting in no major changes in the durations of vegetative risk for individuals exposed to a false spring that received eight weeks of chilling (0.9  $\pm$  1.1 days, Figure 2a and Table S3).

False springs impacted growth habit and shoot growth but not total biomass. Across all chilling treatments, 172 especially for the four and eight week chilling treatments, false springs led to more damage to the shoot apical 173 meristem (4 weeks: 52.5% increase in probability of damage under false spring treatment or  $2.1 \pm 1$ ; 6 weeks: 174 32.5% or a  $1.3 \pm 1.4$ ; and 8 weeks: 55% or a  $2.2 \pm 1.3$ ; Figure 3a and Table S5). Shoot growth over the 175 growing season increased with eight weeks of chilling  $(11 \pm 4 \text{ cm})$  but growth did not increase for eight weeks 176 of chilling under false spring conditions (4.7  $\pm$  5.5 cm; Figure 3b and Table S6). False springs led to slightly 177 lower total biomasses when saplings were exposed to only four weeks of chilling (-3.4  $\pm$  2.8 g) but there was very little change in total biomass under false spring conditions (compared to the no false spring) for both 179 the six  $(-3.6 \pm 4 \text{ g})$  and the eight weeks of chilling treatments  $(2.9 \pm 3 \text{ g}; \text{Figure S1})$  and Table S7).

False springs also affected physical leaf traits. False spring treatments decreased leaf toughness across all chilling treatments (4 weeks:  $-0.1 \pm 0$  N; 6 weeks:  $-0.1 \pm 0$  N; and 8 weeks:  $-0.1 \pm 0$  N; Figure 4a and Table S8) and decreased chlorophyll content in leaves with six and eight weeks of chilling (6 weeks:  $-1.5 \pm 1.2$  mg/cm<sup>2</sup>; and especially for 8 weeks:  $-2 \pm 1.1$  mg/cm<sup>2</sup>; Figure S2 and Table S9). Additionally, false springs led to decreased leaf thickness across four and eight weeks of chilling treatments, but there was little change

for the six weeks of chilling treatment (4 weeks: -8.9  $\pm$  3.7  $\mu$ m; 6 weeks: -3.5  $\pm$  5.3  $\mu$ m; and 8 weeks: -15.8  $\pm$  5.3  $\mu$ m for eight weeks of chilling; Figure 4b and Table S10).

False springs and chilling treatments were generally consistent across species, though not always. Duration of vegetative risk decreased for most species with longer chilling (i.e., the eight weeks), except for Salix 189 purpurea, which experienced longer durations of vegetative risk with longer chilling (Figure 2a). False springs 190 led to meristem damage agross all species except for Betula populifolia and Sorbus americana. Additionally, 191 Viburnum dentatum experienced consistent meristem damage under all treatments (Figure 3a). Effects on 192 leaf thickness were especially variable across species under the longer chilling treatments, specifically with 193 Sorbus americana and Viburnum dentatum having thicker leaves with increased chilling (Figure 4b). 194 Despite large treatment effects on phenology, we found no major effects on phenological rank within the 195 community. Order of leafout timing was consistent across all treatments, with Salix purpurea always being 196 first to leafout, followed by Betula papyrifera, B. populifolia and Cornus racemosa, followed by Alnus rugosa, Sorbus americana, Viburnum dentatum and Acer saccharinum (Figure 5). Viburnum dentatum was the

first to leafout, followed by Betula papyrifera, B. populifolia and Cornus racemosa, followed by Alnus rugosa,

Sorbus americana, Viburnum dentatum and Acer saccharinum (Figure 5). Viburnum dentatum was the

only species to change rank across treatments, though it was consistently grouped with the later-leafout

species. Order of budset timing was also consistent across all treatments, with Cornus racemosa and Sorbus

americana being first to set bud, followed by Betula papyrifera, Acer saccharinum and Viburnum dentatum,

followed by B. populifolia, Salix purpurea and Alnus rugosa (Figure S3). Acer saccharinum was the only

species to change budset rank across treatments, though it was grouped consistently with Betula papyrifera

and Viburnum dentatum.

### Discussion

Our experiment examined the consequences of two major interactive effects of climate change across eight deciduous forest tree species—false springs and chilling. Our results confirmed the major features of false springs (plant damage) and chilling (advancing spring phenology) then highlighted how these two effects

altered multiple aspects of plant phenology, plant growth and leaf traits. Importantly, we found false springs 209 and chilling have opposing additive effects on the duration of vegetative risk. This suggests that the combi-210 nation of increased false springs and warmer winters could be especially detrimental to forest communities. 211

#### False springs and chilling interactively determined risk and damage

Chilling length greatly influences spring phenology during the critical budburst to leafout phases, and thus may compensate for the detrimental effects of false springs on phenology. With false springs increasing the 214 duration of vegetative risk, the risk of multiple false springs occurring in one season also increases. But our 215 experiment found that chilling can compensate for this increase in duration of vegetative risk: with increased 216 chilling, the duration of vegetative risk does not increase under false spring conditions. This suggests chilling 217 is more important for saplings in terms of exposure to multiple false springs. With climate change and 218 warming temperatures, over-winter chilling is anticipated to decrease (Laube et al., 2013) and false springs 219 are predicted to increase in certain regions (Ault et al., 2015; Liu et al., 2018). This combination could greatly 220 impact plant performance, survival and shape species distributions, ultimately affecting ecosystem processes, 221 such as carbon uptake and nutrient cycling. 222 We found that false springs impacted sapling growth, regardless of chilling duration. Past studies suggest 223 early-budburst species can withstand lower temperature thresholds (Lenz et al., 2013; Muffler et al., 2016; 224 Zohner et al., 2020), but our results suggest false springs consistently impair shoot apical meristem growth, 225 regardless of a species phenological order. Damage to the shoot apical meristem can lead to reliance on lateral shoot growth, rendering inefficient growth patterns and—if damage is significant within a forest stand—it can lead to declines in recruitment (Rhodes & Clair, 2018). Though overall height increased with more overwinter chilling, we found that false springs impacted all individuals similarly across all chilling treatments, suggesting an interplay between false spring damage to the meristem and overall plant height and thus further emphasizing the detrimental effects of meristem damage.

False springs greatly impacted physical leaf characteristics in our experiment: with chlorophyll content, leaf

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toughness and leaf thickness decreasing, the quality of the leaf decreased. This reduction in quality could subsequently lead to an increase in herbivory risk (Onoda et al., 2011). We found that increased chilling levels decreased leaf toughness and decreased chlorophyll content under false spring conditions. Further studies that assess the secondary compounds and total phenolic content (Ayres, 1993; Webber & Mason, 2016) as well as photosynthetic rate of the leaves would help to assess the interactive effects of chilling and false springs on leaf level traits and better predict herbivory risk and the overall leaf quality under climate change.

#### False springs and chilling did not reshape temporal assembly

Climate model projections and experimental studies with very low chilling predict substantial shifts in species 240 leafout order under future climate change conditions (Roberts et al., 2015; Laube et al., 2013); other studies us-241 ing long-term phenology observations suggest leafout phenology order is consistent across years (Wesołowski, 242 Tomasz and Rowiński, Patryk, 2006). We did not find major shifts in species leafout order—thus consistent 243 with observational studies (Wesołowski, Tomasz and Rowiński, Patryk, 2006)—except for Viburnum denta-244 tum, though it still leafed out within the late-leafout group of species across all treatments. Therefore, we do not predict major reassembly of forest communities due to winter warming or false spring incidence. Our results are also in line with some experimental studies, for example, in one full factorial growth chamber experiment, most treatments did not lead to substantial phenological reordering, except when individuals experienced little to no field chilling (Laube et al., 2013). Future climate projections suggest major shifts in chilling due to warmer winters, which our design may not fully capture, but chilling is also not well understood (Nanninga, Claudia and Buyarski, Chris R and Pretorius, Andrew M and Montgomery, Rebecca A, 2017). Some regions may experience more chilling rather than less with warming (Fraga, Helder and Pinto, Joaquim G. and Santos, João A., 2019). Our findings that chilling has cascading effects on phenology, growth and leaf traits, suggests we need to better understand over-winter chilling in order to make accurate predictions. Phenological rank remained consistent across all of our false spring and chilling treatments—where all species 255 were affected equally. In nature, however, not all of our study species are at equal risk of false springs, with

early-budburst species (e.g., Salix americana or Betula papyrifera) generally more at risk than later-budburst 257 species (e.g., Acer saccharinum or Viburnum dentatum). This suggests that climate change could reshape 258 forest communities, though not directly through temporal assembly. Instead, our results suggest that change 259 may come from physical damage and leaf trait impacts of false springs and decreases in chilling. Some 260 temperate tree and shrub species utilize various leaf characteristics to lower their risk of false spring damage: 261 increased 'packability' of leaf primordia in winter buds, which allows for more rapid leafout (Edwards et al., 262 2017), increased trichome density on young leaves to protect leaf tissue against freezes (Agrawal et al., 2004; 263 Prozherina et al., 2003) and buds with decreased water content to increase freeze tolerance (Beck et al., 2007; Hofmann & Bruelheide, 2015; Kathke & Bruelheide, 2011; Morin et al., 2007; Muffler et al., 2016; Norgaard Nielsen & Rasmussen, 2009; Poirier et al., 2010). Our study did not directly examine these traits, but our results suggest that the complex interplay of a changing climate and species-specific mosaics of traits, including their freeze-tolerance physiologies, could influence phenological rank and forest community structure.

Our results suggest that predicting future forest recruitment will require integrating the effects of shifting 270 false springs on inter- and intraspecific competition. With over-winter chilling decreasing with climate change, 271 saplings—which generally leafout earlier than later lifestages to gain access to light (Augspurger, 2009)—are likely more at risk of damage from false spring events. This could lead to dieback of saplings, most especially of early-budbursting species, in temperate forests with climate change. Thus, climate change could greatly impact early-budburst species, which will likely see increases in durations of vegetative risk from the dual effects of lower chilling and heightened false spring risk. Understanding how false springs are changing and how equally—or not—these effects impact different species, especially their seedlings and saplings, is crucial for future projections. By integrating the additive and adverse effects of decreasing over-winter chilling and 278 increasing false spring risk—and how false springs are changing across various species, we can better predict 279 shifts in forest communities and recruitment under climate change. 280

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- 286 and manuscript.

## 287 Author Contribution

- 288 C.J.C. and E.M.W. conceived of the study, identified species to use in the study and determined which
- phenological and growth measurements to observe. C.J.C. performed the analyses and produced all figures
- 290 and tables. C.J.C. wrote the paper, and both authors edited it.

# Data Availability:

- Data and code from the analyses will be available via KNB upon publication and are available to all reviewers
- upon request. Raw data, Stan model code and output are available on github at https://github.com/cchambe12/chillfreeze
- 294 and provided upon request.

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# 7 Tables and Figures

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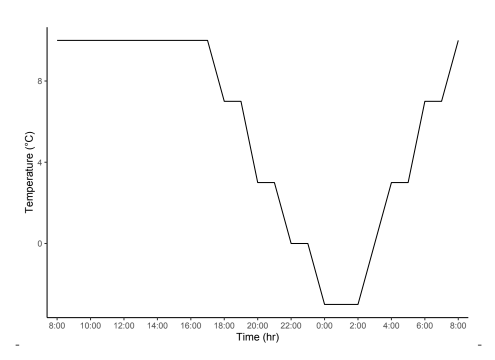


Figure 1: False spring treatment temperature regime in the growth chamber

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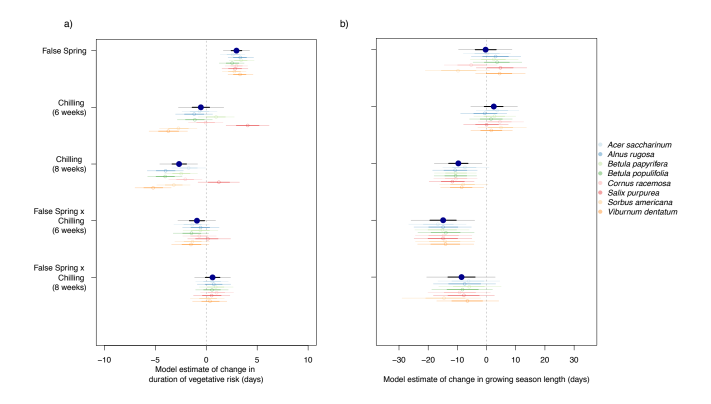


Figure 2: Effects of false spring treatment, six weeks of chilling and eight weeks of chilling relative to four weeks of chilling with no false spring on a) duration of vegetative risk (days) and b) growing season length (days). Larger, blue dots represent overall estimates across all species, while smaller dots are estimates for each species. Dots and thin lines show means and 90% uncertainty intervals and thicker lines show 50% uncertainty intervals.

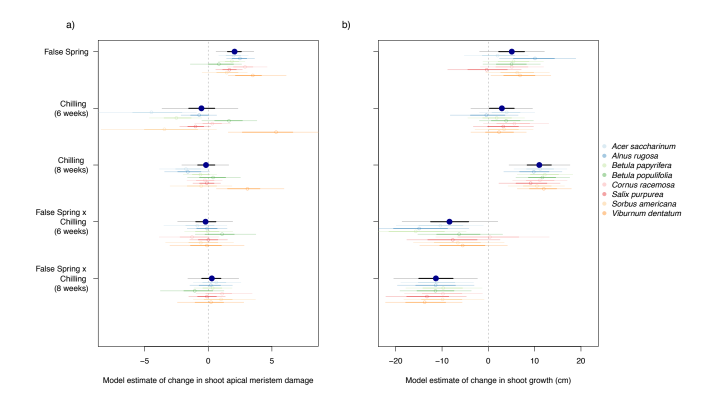


Figure 3: Effects of false spring treatment, six weeks of chilling and eight weeks of chilling on a) shoot apical meristem damage and b) total shoot growth (cm). Larger, blue dots represent overall estimates across all species, while smaller dots are estimates for each species. Dots and thin lines show means and 90% uncertainty intervals and thicker lines show 50% uncertainty intervals.

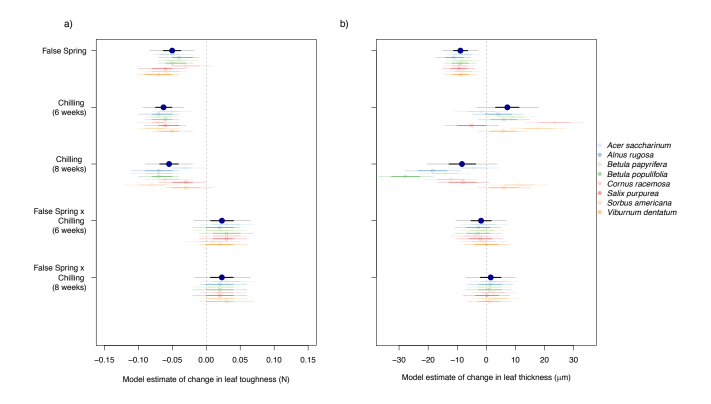


Figure 4: Effects of false spring treatment, six weeks of chilling and eight weeks of chilling on a) leaf toughness (N) and b) leaf thickness ( $\mu$ m). Larger, blue dots represent overall estimates across all species, while smaller dots are estimates for each species. Dots and thin lines show means and 90% uncertainty intervals and thicker lines show 50% uncertainty intervals.

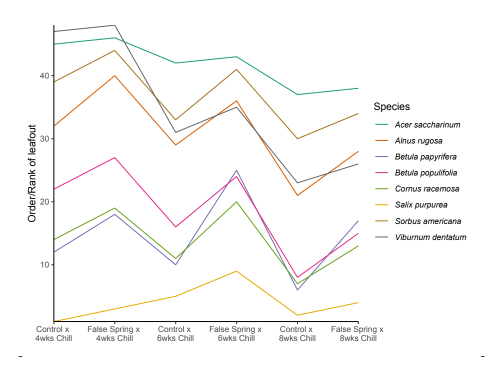


Figure 5: Rank order of leafout across all species using mean trends.