Rethinking False Spring Risk

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Abstract

Temperate plants are at risk of being exposed to late spring freezes — often called false springs — which can
be damaging ecologically and economically. As climate change may alter the prevalence and severity of false
springs, our ability to accurately forecast such events has become more critical. Currently, many false spring
studies simplify the ecological and physiological information needed for accurate predictions of the level of
plant damage from late spring freezes. Here we review the complexity of factors driving a plant's false spring
risk. We highlight how species, life stage, and habitat differences contribute to the damage potential of false
springs. Integrating these complexities could rapidly advance forecasting of false spring events in climate
change and ecological studies.

24 Introduction

- Plants from temperate environments time their growth each spring to follow rising temperatures alongside increasing light and soil resource availability. While tracking spring resource availability, individuals that budburst before the last freeze date are at risk of leaf loss, damaged wood tissue, and slowed canopy development (Gu et al., 2008; Hufkens et al., 2012). These damaging late spring freezes are also known as false springs, and are widely documented to result in adverse ecological and economic consequences (Ault et al., 2013; Knudson, 2012).
- Climate change is expected to cause an increase in damage from false spring events due to earlier spring onset and potentially greater fluctuations in temperature in some regions (Inouye, 2008; Martin et al., 2010).

 Already, multiple studies have documented false springs in recent years (Augspurger, 2009, 2013; Gu et al., 2008; Menzel et al., 2015) and some have linked these events to climate change (Allstadt et al., 2015; Ault et al., 2013; Muffler et al., 2016; Vitra et al., 2017; Xin, 2016). This interest in false springs has led to a growing body of research investigating the effects on temperate forests. For this research to produce accurate predictions, however, researchers need methods that properly evaluate the effects of false springs across diverse species and climate regimes.

39 Measuring False Spring

- 40 Current metrics for estimating false springs events are generally simple, often requiring an estimate for the
- start of biological 'spring' (i.e. budburst) and whether temperatures below a particular threshold occurred in
- the following week. Such estimates inherently assume consistency of damage across species, functional group,
- 43 life stages, and other climatic regimes, ignoring that such factors can greatly impact plants' false spring risk.
- 44 As a result, such indices may lead to inaccurate estimates and predictions.
- 45 In this paper we highlight the complexity of factors driving a plant's false spring risk and provide a road
- map for improved metrics. We show how location within a forest or canopy (Augspurger, 2013), interspecific
- 47 variation in avoidance and tolerance strategies (Martin et al., 2010; Muffler et al., 2016), freeze temperature
- thresholds (Lenz et al., 2013), and regional effects (Muffler et al., 2016) unhinge simple metrics of false spring.
- We argue that a new approach that integrates these and other crucial factors would help accurately determine
- 50 current false spring damage and improve predictions of spring freeze risk under a changing climate while
- 51 potentially providing novel insights to how plants respond to and are shaped by spring frost.

Defining False Spring: An example in one temperate plant community

- Temperate forest plants experience elevated risk of frost damage during the spring due to the stochastic timing of frosts. Freezing temperatures following a warm spell can result in plant damage or even death
- ⁵⁶ (Ludlum, 1968; Mock et al., 2007). Many temperate species exhibit flexible spring phenologies, which help
- 57 them minimize spring freezing risk, but freeze damage can still occur. Once buds exit the dormancy phase,
- they are less freeze tolerant and less resistant to ice formation (Lenz et al., 2013; Taschler et al., 2004; Vitasse
- ₅₉ et al., 2014b). Intracellular ice formation from false spring events often results in severe leaf and stem damage
- 60 (Burke et al., 1976; Sakai & Larcher, 1987). Ice formation can also occur indirectly (i.e. extracellularly), which
- 61 results in freezing dehydration and mimics drought conditions (Beck et al., 2004; Hofmann & Bruelheide,
- ⁶² 2015; Pearce, 2001). Both forms of ice formation can cause defoliation and crown dieback (Gu et al., 2008).
- 63 An effective and consistent definition of false spring would accurately determine the amount and type of ice
- formation to properly evaluate the level of damage that could occur.
- 65 Currently there are several ways to define a false spring. A common definition describes a false spring as

having two phases: rapid vegetative growth prior to a freeze and a post freeze setback (Gu et al., 2008).

Other definitions instill more precise temporal parameters, specific to certain regions (e.g., in Augspurger, 2013, false spring for the Midwestern United States is defined as a warmer than average March, a freezing April, and enough growing degree days between budburst and the last freeze date). A widely used definition integrates a mathematical equation to quantify a false spring event. This equation, known as a False Spring Index (FSI), signifies the likelihood of damage to occur from a late spring freeze. Currently, FSI is evaluated annually by the day of budburst and the day of last spring freeze (often calculated at -2.2°C, Schwartz, 1993) through the simple equation (Marino et al., 2011):

$$FSI = \text{Day of Year}(LastSpringFreeze) - \text{Day of Year}(Budburst)$$
 (1)

Negative values indicate no risk situations, whereas a damaging FSI is currently defined to be seven or more days between budburst and the last freeze date (Equation 1) (Peterson & Abatzoglou, 2014). This seven day threshold captures the reality that leaf tissue is at high risk of damage from frost in the period after budburst, with later vegetative phases (e.g., after full leafout) being more resistant to such damage.

To demonstrate how the FSI definition works, we applied it to data from the Harvard Forest Long-term Ecological Research program in Massachusetts. We used three separate methodologies to calculate spring onset: long-term ground observational data (O'Keefe, 2014), PhenoCam data from Harvard Forest (Richardson, 2015), and USA National Phenology Network's (USA-NPN) Extended Spring Index (SI-x) data (USA-NPN, 2016). These spring onset values were then inputted into the FSI equation (Equation 1) to calculate FSI from 2008 to 2014 (Figure 1).

Each methodology rendered different FSI values, suggesting different false spring damage for the same site and same year. For most years, the observational FSI and PhenoCam FSI are about 10-15 days lower than the SI-x data. This is especially important for 2008, when the SI-x data indicates a false spring year, whereas the other two datasets do not. In 2012, the observational data and PhenoCam data diverge slightly and the PhenoCam FSI is over 30 days less than the SI-x value.

The reason for these discrepancies is that each method evaluates spring onset by integrating different attributes such as age, species or functional group. Spring phenology in temperate forests typically progresses by functional group: understory species and young trees tend to initiate budburst first, whereas larger canopy species start later in the season (Richardson & O'Keefe, 2009; Xin, 2016). The different FSI values determined in Figure 1 exemplify the differences in functional group spring onset dates and illustrate variations in forest demography and phenology. While the SI-x data (based on observations of early-active shrub species, including lilac, *Syringa vulgaris*) may best capture understory dynamics, the PhenoCam and observational FSI data integrate over larger canopy species. Such differences are visible each year, as the canopy-related metrics show lower risk, but are especially apparent in 2012. In 2012, a false spring event was reported through many regions of the US due to warm temperatures occurring in March (Ault *et al.*, 2015). These high temperatures would most likely have been too early for larger canopy species to initiate budburst but they would have affected smaller understory species, as is seen by the high risk of the SI-x FSI in Figure 1.

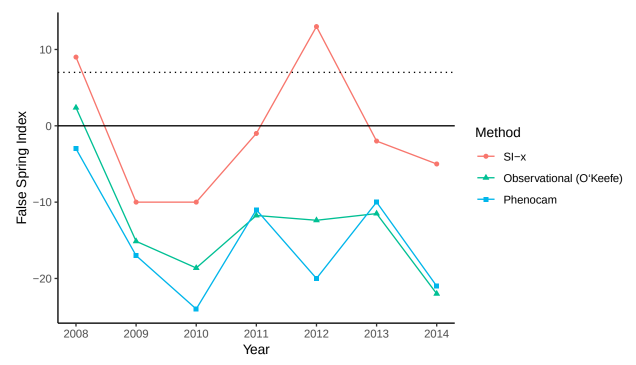


Figure 1: False Spring Index (FSI) values from 2008 to 2014 vary across methodologies. To calculate spring onset, we used the USA-NPN Extended Spring Index tool for the USA-NPN FSI values, which are in red (USA-NPN, 2016), long-term ground observational data for the observed FSI values, which are in green (O'Keefe, 2014), and near-surface remote-sensing canopy data for the PhenoCam FSI values, which are in blue (Richardson, 2015). The solid line at FSI=0 indicates a boundary between a likely false spring event or not, with positive numbers indicating a false spring likely occurred and negative numbers indicating a false spring most likely did not occur. The dotted line at FSI=7 indicates the seven day threshold frequently used in false spring definitions, which suggests years with FSI values greater than seven very likely had false spring events.

Yet, in contrast to our three metrics of spring onset for one site, most FSI work currently ignores variation across functional groups — instead using one metric of spring onset and assuming it applies to the whole community of plants (Allstadt et al., 2015; Marino et al., 2011; Mehdipoor & Zurita-Milla, 2017; Peterson

& Abatzoglou, 2014). The risk of a false spring varies across habitats and with species composition since spring onset is not consistent across functional groups (Martin *et al.*, 2010). Therefore, one spring onset date cannot be used as an effective proxy for all species. False spring studies should first assess the forest demographics and functional groups relevant to the study question in order to effectively estimate the date of spring onset. However, as we outline below, considering different functional groups is unlikely to be enough for robust predictions. It will also be important to integrate species differences within functional groups and to consider the various interspecific avoidance and tolerance strategies that species have evolved against false springs.

Plant Physiology and Diversity versus the Current False Spring Definition

Plants have evolved to minimize false spring damage through two strategies: avoidance and tolerance. Many temperate forest plants utilize various morphological strategies to be more frost tolerant: some have increased 115 'packability' of leaf primordia in winter buds, which may permit more rapid leafout (Edwards et al., 2017) and minimize the exposure time of less resistant tissues. Other species have young leaves with more trichomes to 117 act as a buffer against spring frosts (Agrawal et al., 2004; Prozherina et al., 2003). And many other individuals 118 are able to respond to abiotic cues such as consistently dry winters. Species living in habitats with drier 119 winters develop shoots and buds with decreased water content, which makes the buds more tolerant to drought 120 and also to false spring events (Beck et al., 2007; Kathke & Bruelheide, 2011; Hofmann & Bruelheide, 2015; 121 Morin et al., 2007; Muffler et al., 2016; Norgaard Nielsen & Rasmussen, 2009; Poirier et al., 2010). These strategies are probably only a few of the many ways plants work to morphologically avoid frost damage, and more studies are needed to investigate the interplay between morphological traits and false spring tolerance. 124 Rather than being more tolerant of spring freezing temperatures, some temperate forest species have evolved 125 to avoid frosts via their phenologies. Effective avoidance strategies require well-timed spring phenologies. 126 Most temperate deciduous tree species optimize growth and minimize spring freeze damage by using three cues 127 to initiate budburst: low winter temperatures (chilling), warm spring temperatures (forcing), and increasing 128 photoperiods (Chuine, 2010). The evolution of these three cues and their interactions has permitted temperate plant species to occupy more northern ecological niches (Kollas et al., 2014) and decrease the risk of false 130 spring damage (Charrier et al., 2011). One avoidance strategy, for example, is the interaction between overwinter chilling and spring forcing temperatures. Warm temperatures earlier in the winter will not result in
early budburst due to insufficient chilling (Basler & Körner, 2012). Likewise, photoperiod sensitivity is a
common false spring avoidance strategy: species that respond strongly to photoperiod cues in addition to
warm spring temperatures are unlikely to have large advances in budburst with warming, and thus may evade
false spring events as warming continues (Basler & Korner, 2014).

77 Defining Vegetative Risk

Phenology and frost tolerance are clearly intertwined — with important variation occurring across different phenological phases. Flowering and fruiting are generally more sensitive to false spring events than vegetative phases (Augspurger, 2009; Caradonna & Bain, 2016; Lenz et al., 2013), but false spring events that occur during the vegetative growth phenophases may impose the greatest freezing threat to deciduous plant species. Plants will suffer greater long-term effects from the loss of photosynthetic tissue, which could impact multiple years of growth, reproduction, and canopy development (Vitasse et al., 2014a; Xie et al., 2015). However, there is high variability in defining a damaging temperature threshold across species, including between agricultural and ecological studies (Figure 2).

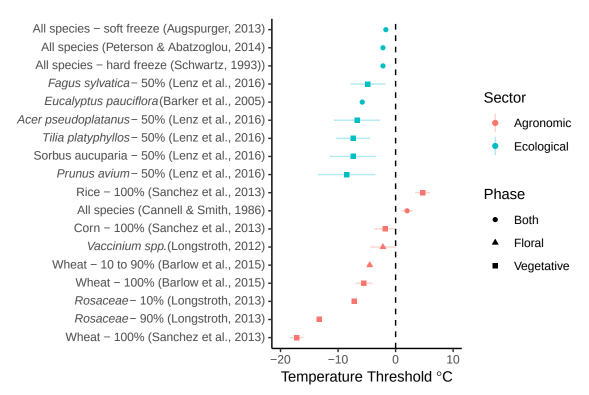


Figure 2: A comparison of damaging spring freezing temperature thresholds across ecological and agronomic studies. Each study is listed on the vertical axis along with the taxonomic group of focus. Next to the species name is the freezing definition used within that study (e.g. 100% is 100% lethality). Each point is the best estimate recorded for the temperature threshold with standard deviation if indicated in the study.

There is also important variation within certain phenological phases. Most notably, within the vegetative phases of spring leafout, plants that have initiated budburst but have not fully leafed out are more likely to sustain damage from a false spring than individuals past the leafout phase. This is because freezing tolerance is lowest after budburst begins until the leaf is fully unfolded (Lenz et al., 2016). Therefore, the rate of budburst and the length of time between budburst and leafout is essential for predicting the level of damage from a false spring event. We will refer to the timing between these phenophases — budburst to leafout — as the duration of vegetative risk (Figure 3).

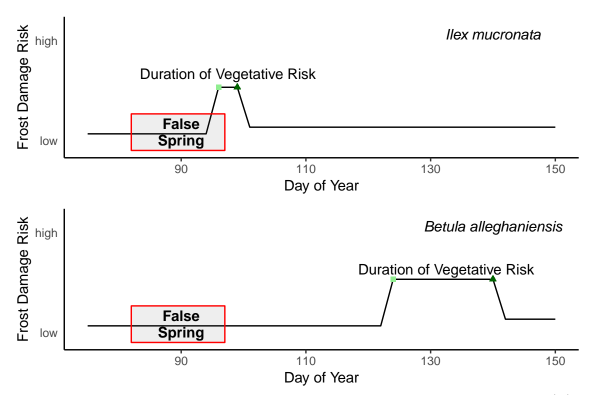


Figure 3: Differences in spring phenology and false spring risk across two species: *Ilex mucronata* (L.) and *Betula alleghaniensis* (Marsh.). We mapped a hypothetical false spring event based on historical weather data and long-term observational phenological data collected at Harvard Forest (O'Keefe, 2014). In this scenario, *Ilex mucronata*, which budbursts early and generally has a short period between budburst (light green squares) and leafout (dark green triangles), would be exposed to a false spring event during it's duration of vegetative risk (i.e. from budburst to leafout), whereas *Betula alleghaniensis* would avoid it entirely (even though it has a longer duration of vegetative risk), due to later budburst.

3 How Species Phenological Cues Shape Vegetative Risk

Predictions of false spring critically depend on understanding what controls the duration of vegetative risk across species. For temperate species, the three major cues (winter chilling temperatures, spring warming temperatures and photoperiod) that control budburst (Chuine, 2010) play a dominant role. Most phenological studies currently focus on one phenophase (i.e. budburst or leafout) but, to examine false spring risk, it is important to examine the effects of the three phenological cues and their interactions on the duration of vegetative risk—that is, researchers must collect data on both budburst and leafout timing.

Such cues may provide a starting point for predicting how climate change will alter the duration of vegetative risk. Robust predictions will require more information, especially the emissions scenario realized over coming

decades (IPCC, 2015), but some outcomes with warming are more expected than others. For example, higher 162 temperatures are generally expected to increase forcing and decrease chilling in many locations, as well as 163 to trigger budburst at times of the year when daylength is shorter. Using data from a recent study that manipulated all three cues and measured budburst and leafout (Flynn & Wolkovich, 2018) shows that any 165 one of these effects alone can have a large impact on the duration of vegetative risk (Figure 4): more forcing 166 shortens it substantially (-15 to -8 days), while shorter photoperiods and less chilling increase it to a lesser 167 extent (+3 to 9 days). Together, however, the expected shifts generally shorten the duration of vegetative 168 risk by 4-13 days, both due to the large effect of forcing and the combined effects of multiple cues. How shortened the risk period is, however, varies strongly by species and highlights how climate change may speed 170 some species through this high risk period, but not others. Additionally, as our results are for a small set of species we expect other species may have more diverse responses, as has already been seen in shifts in 172 phenology with warming (Cleland et al., 2006; Fu et al., 2015; Xin, 2016).

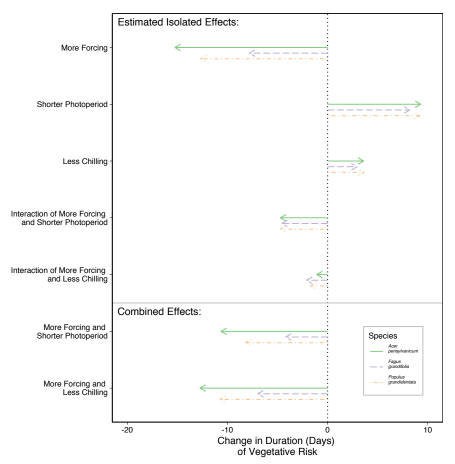


Figure 4: We examine the effects of phenological cues on the duration of vegetative risk across three species: Acer pensylvanicum, Fagus grandifolia, and Populus grandidentata. 'More Forcing' is a 5°C increase in spring warming temperatures, 'Shorter Photoperiod' is a 4 hour decrease in photoperiod and 'Less Chilling' is a 30 day decrease in over-winter chilling. Along with the estimated isolated effects, we the show the combined predicted shifts in phenological cues with potential climate change (i.e. more forcing with shorter photoperiod and more forcing with less chilling) and the subsequent shifts in duration of vegetative risk across species. To calculate the combined effects, we added the estimated isolated effects of each cue alone with the interaction effects for the relevant cues for each species.

These findings highlight the need for further studies on the interplay between chilling, forcing, and photoperiod cues and the duration of vegetative risk across species. This is especially true for species occupying ecological niches more susceptible to false spring events; even if warming causes a shortened duration of vegetative risk for such species, the related earlier budburst dates could still lead to greater risk of false spring exposure.

Predictable Regional Differences in Climate, Species Responses and False Spring Risk

Robust predictions must consider the interplay of species cues with a specific location's climate. Climate and 180 thus false spring risk vary across regions. Some regions experience harsher winters and greater temperature variability throughout the year (Figure 5 e.g. Maine, USA), and these more variable regions often have a 182 much higher risk of false spring than others (Figure 5 e.g. Lyon, France). Understanding and integrating 183 spatiotemporal effects and regional differences when investigating false spring risk and duration of vegetative 184 risk would help improve predictions as climate change progresses. Such differences depend both on the local 185 climate, the local species and the cues for each species at that location, as a single species may have varying 186 cues across space. Therefore, based on cues alone, different regions may have different durations of vegetative 187 risk for the same species (Caffarra & Donnelly, 2011; Partanen, 2004; Vihera-aarnio et al., 2006). Studies also show that different species within the same location can exhibit different sensitivities to the three cues 189 (Basler & Körner, 2012; Laube et al., 2013), further amplifying the myriad of climatic and phenological shifts that determine false spring risk in a region. 191

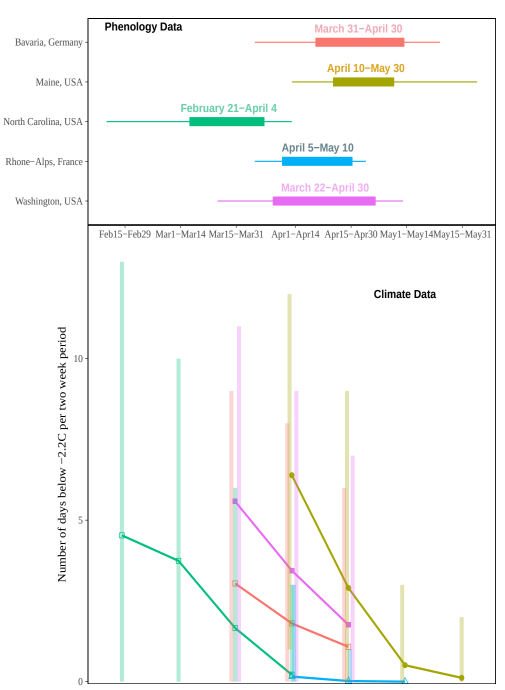


Figure 5: False spring risk can vary dramatically across regions. Here we show the period when plants are most at risk to tissue loss – between budburst and leafout (upper, lines represent the range with the thicker line representing the interquartile range) and the variation in the number of freeze days (-2.2°C) (Schwartz, 1993) that occurred on average over the past 50 years for five different sites (lower, bars represent the range, points represent the mean). Data come from USA-NPN SI-x tool (1981-2016) and observational studies (1950-2016) for phenology (Schaber & Badeck, 2005; Soudani et al., 2012; USA-NPN, 2016; White et al., 2009) and NOAA Climate Data Online tool for climate (from 1950-2016).

How a single species phenological cues varies across space is not yet well predicted. Some studies have investigated how phenological cues for budburst vary across space, including variation across populations, by 193 using latitudinal gradients (Gauzere et al., 2017; Søgaard, Gunnhild and Johnsen, Øystein and Nilsen, Jarle and Junttila, Olavi, 2008; Way & Montgomery, 2015; Zohner et al., 2016). Fewer, however, have integrated 195 distance from the coast (but see Aitken & Bemmels, 2015; Harrington & Gould, 2015; Myking & Skroppa, 196 2007) or regional effects. Some studies assert that the distance from the coast is a stronger indicator of 197 budburst timing than latitude (Myking & Skroppa, 2007), with populations further inland initiating budburst 198 first, whereas those closer to the coast budburst later in the season. Therefore, to better understand the interplay between duration of vegetative risk and climatic variation it is important to recognize how climate 200 regime extremes (e.g. seasonal trends, annual minima and annual maxima) vary across regions and how they will shift in the future: as climatic regimes are altered by climate change false spring risk could vary 202 in intensity across regions and time (i.e. regions currently at high-risk of false spring damage could become low-risk regions in the future and vice versa). 204

205 Conclusion

Temperate forest trees are most at risk to frost damage in the spring due to the stochasticity of spring freezes. 206 With warm temperatures advancing in the spring but last spring freeze dates advancing at a slower rate, there could be more damaging false spring events in the future, especially in high-risk regions (Gu et al., 2008; 208 Inouye, 2008; Liu et al., 2018). Current equations for evaluating false spring damage (e.g. Equation 1) largely simplify the myriad complexities involved in assessing false spring damage and risks. More studies aimed 210 at understanding relationships between species avoidance and tolerance strategies, climatic regimes, and physiological cue interactions with the duration of vegetative risk would improve predictions. Additionally, 212 research to establish temperature thresholds for damage across functional types and phenophases will help 213 effectively predict false spring risk in the future. An integrated approach to assessing past and future spring 214 freeze damage would provide novel insights into plant strategies, and offer more robust predictions as climate 215 change progresses, which is essential for mitigating the adverse ecological and economic effects of false springs.

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References

- Agrawal, A.A., Conner, J.K. & Stinchcombe, J.R. (2004) Evolution of plant resistance and tolerance to frost damage. *Ecology Letters* 7, 1199–1208.
- Aitken, S.N. & Bemmels, J.B. (2015) Time to get moving: assisted gene flow of forest trees. *Evolutionary*Applications 9, 271–290.
- Allstadt, A.J., Vavrus, S.J., Heglund, P.J., Pidgeon, A.M., Wayne, E. & Radeloff, V.C. (2015) Spring plant
- phenology and false springs in the conterminous U. S. during the 21st century. Environmental Research
- 227 Letters (submitted) **10**, 104008.
- Augspurger, C.K. (2009) Spring 2007 warmth and frost: Phenology, damage and refoliation in a temperate deciduous forest. Functional Ecology 23, 1031–1039.
- Augspurger, C.K. (2013) Reconstructing patterns of temperature, phenology, and frost damage over 124 years: Spring damage risk is increasing. *Ecology* **94**, 41–50.
- ²³² Ault, T.R., Henebry, G.M., de Beurs, K.M., Schwartz, M.D., Betancourt, J.L. & Moore, D. (2013) The false
- spring of 2012, earliest in North American record. Eos, Transactions American Geophysical Union 94,
- 181–182.
- ²³⁵ Ault, T.R., Schwartz, M.D., Zurita-Milla, R., Weltzin, J.F. & Betancourt, J.L. (2015) Trends and natural
- variability of spring onset in the coterminous United States as evaluated by a new gridded dataset of spring
- indices. *Journal of Climate* **28**, 8363–8378.
- Barker, D., Loveys, B., Egerton, J., Gorton, H., Williams, W. & Ball, M. (2005) CO₂ Enrichment predisposes
- foliage of a eucalypt to freezing injury and reduces spring growth. Plant, Cell and Environment 28, 1506–
- 240 1515.
- Barlow, K., Christy, B., O'Leary, G., Riffkin, P. & Nuttall, J. (2015) Simulating the impact of extreme heat
- 242 and frost events on wheat crop production: A review. Field Crops Research 171, 109–119.

- Basler, D. & Körner, C. (2012) Photoperiod sensitivity of bud burst in 14 temperate forest tree species.
- Agricultural and Forest Meteorology 165, 73–81.
- Basler, D. & Korner, C. (2014) Photoperiod and temperature responses of bud swelling and bud burst in
- four temperate forest tree species. Tree Physiology 34, 377–388.
- Beck, E.H., Fettig, S., Knake, C., Hartig, K. & Bhattarai, T. (2007) Specific and unspecific responses of
- plants to cold and drought stress. Journal of Biosciences 32, 501–510.
- Beck, E.H., Heim, R. & Hansen, J. (2004) Plant resistance to cold stress: Mechanisms and environmental
- signals triggering frost hardening and dehardening. Journal of Biosciences 29, 449–459.
- ²⁵¹ Burke, M., Gusta, L., Quamme, H., Weiser, C. & Li, P. (1976) Freezing and injury in plants. Annual Review
- of Plant Physiology **27**, 507–528.
- ²⁵³ Caffarra, A. & Donnelly, A. (2011) The ecological significance of phenology in four different tree species:
- Effects of light and temperature on bud burst. International Journal of Biometeorology 55, 711–721.
- ²⁵⁵ Caradonna, P.J. & Bain, J.A. (2016) Frost sensitivity of leaves and flowers of subalpine plants is related to
- tissue type and phenology. Journal of Ecology 104, 55–64.
- ²⁵⁷ Charrier, G., Bonhomme, M., Lacointe, A. & Améglio, T. (2011) Are budburst dates, dormancy and cold ac-
- climation in walnut trees (Juglans regia L.) under mainly genotypic or environmental control? International
- Journal of Biometeorology 55, 763-774.
- 260 Chuine, I. (2010) Why does phenology drive species distribution? Philosophical Transactions of the Royal
- Society B: Biological Sciences **365**, 3149–3160.
- ²⁶² Cleland, E., Chiariello, N., Loarie, S., Mooney, H. & Field, C. (2006) Diverse responses of phenology to global
- changes in a grassland ecosystem. PNAS 103, 13740–13744.
- ²⁶⁴ Edwards, E.J., Chatelet, D.S., Spriggs, E.L., Johnson, E.S., Schlutius, C. & Donoghue, M.J. (2017) Correla-
- tion, causation, and the evolution of leaf teeth: A reply to Givnish and Kriebel. Am J Bot 104, 509-515.
- Flynn, D.F.B. & Wolkovich, E.M. (2018) Temperature and photoperiod drive spring phenology across all
- species in a temperate forest community. New Phytologist 219.
- Fu, Y.H., Zhao, H., Piao, S., Peaucelle, M., Peng, S., Zhou, G., Ciais, P., Huang, M., Menzel, A., Peñuelas,
- J. & et al. (2015) Declining global warming effects on the phenology of spring leaf unfolding. Nature 526,
- 270 104-107.

- Gauzere, J., Delzon, S., Davi, H., Bonhomme, M., Garcia de Cortazar-Atauri, I. & Chuine, I. (2017) Inte-
- grating interactive effects of chilling and photoperiod in phenological process-based models. A case study
- with two European tree species: Fagus sylvatica and Quercus petraea. Agricultural and Forest Meteorology
- 274 **244-255**, 9-20.
- ²⁷⁵ Gu, L., Hanson, P.J., Post, W.M., Kaiser, D.P., Yang, B., Nemani, R., Pallardy, S.G. & Meyers, T. (2008)
- The 2007 Eastern US spring freeze: Increased cold damage in a warming world. BioScience 58, 253.
- Harrington, C.A. & Gould, P.J. (2015) Tradeoffs between chilling and forcing in satisfying dormancy require-
- ments for Pacific Northwest tree species. Frontiers in Plant Science 6.
- 279 Hofmann, M. & Bruelheide, H. (2015) Frost hardiness of tree species is independent of phenology and macro-
- climatic niche. Journal of Biosciences 40, 147–157.
- Hufkens, K., Friedl, M.A., Keenan, T.F., Sonnentag, O., Bailey, A., O'Keefe, J. & Richardson, A.D. (2012)
- Ecological impacts of a widespread frost event following early spring leaf-out. Global Change Biology 18,
- 2365-2377.
- Inouye, D.W. (2008) Effects of climate change on phenology, frost damage, and floral abundance of montane
- wildflowers. *Ecology* **89**, 353–362.
- ²⁸⁶ IPCC (2015) Climate change 2014: mitigation of climate change, vol. 3. Cambridge University Press.
- ²⁸⁷ Kathke, S. & Bruelheide, H. (2011) Differences in frost hardiness of two Norway spruce morphotypes growing
- at Mt. Brocken, Germany. Flora Morphology, Distribution, Functional Ecology of Plants 206, 120–126.
- Knudson, W. (2012) The economic impact of the spring's weather on the fruit and vegetable sectors. The
- 290 Strategic Marketing Institute Working Paper 0.
- Kollas, C., Körner, C. & Randin, C.F. (2014) Spring frost and growing season length co-control the cold
- range limits of broad-leaved trees. Journal of Biogeography 41, 773–783.
- Laube, J., Sparks, T.H., Estrella, N., Höfler, J., Ankerst, D.P. & Menzel, A. (2013) Chilling outweighs
- photoperiod in preventing precocious spring development. Global Change Biology 20, 170–182.
- Lenz, A., Hoch, G., Körner, C. & Vitasse, Y. (2016) Convergence of leaf-out towards minimum risk of freezing
- damage in temperate trees. Functional Ecology 30, 1–11.

- Lenz, A., Hoch, G., Vitasse, Y. & Körner, C. (2013) European deciduous trees exhibit similar safety margins
 against damage by spring freeze events along elevational gradients. New Phytologist 200, 1166–1175.
- Liu, Q., Piao, S., Janssens, I.A., Fu, Y., Peng, S., Lian, X., Ciais, P., Myneni, R.B., Peñuelas, J. & Wang, T.
- (2018) Extension of the growing season increases vegetation exposure to frost. Nature Communications 9.
- Longstroth, M. (2012) Protect blueberries from spring freezes by using sprinklers. url.
- Longstroth, M. (2013) Assessing frost and freeze damage to flowers and buds of fruit trees. url.
- Ludlum, D.M. (1968) Early American Winters: 1604-1820. 3, Boston: American Meteorological Society.
- Marino, G.P., Kaiser, D.P., Gu, L. & Ricciuto, D.M. (2011) Reconstruction of false spring occurrences over
- the southeastern United States, 1901–2007: an increasing risk of spring freeze damage? Environmental
- Research Letters 6, 24015.
- Martin, M., Gavazov, K., Körner, C., Hattenschwiler, S. & Rixen, C. (2010) Reduced early growing season
- freezing resistance in alpine treeline plants under elevated atmospheric CO_2 . Global Change Biology 16,
- 309 1057–1070.
- Mehdipoor, H. & Zurita-Milla, E.I.V.R. (2017) Continental-scale monitoring and mapping of false spring: A
- cloud computing solution. University of Leeds.
- Menzel, A., Helm, R. & Zang, C. (2015) Patterns of late spring frost leaf damage and recovery in a European
- beech (Fagus sylvatica L.) stand in south-eastern Germany based on repeated digital photographs. Frontiers
- in Plant Science **6**, 110.
- Mock, C.J., Mojzisek, J., McWaters, M., Chenoweth, M. & Stahle, D.W. (2007) The winter of 1827–1828 over
- eastern North America: a season of extraordinary climatic anomalies, societal impacts, and false spring.
- 317 Climatic Change **83**, 87–115.
- Morin, X., Ameglio, T., Ahas, R., Kurz-Besson, C., Lanta, V., Lebourgeois, F., Miglietta, F. & Chuine, I.
- 319 (2007) Variation in cold hardiness and carbohydrate concentration from dormancy induction to bud burst
- among provenances of three European oak species. Tree Physiology 27, 817–825.
- Muffler, L., Beierkuhnlein, C., Aas, G., Jentsch, A., Schweiger, A.H., Zohner, C. & Kreyling, J. (2016) Dis-
- tribution ranges and spring phenology explain late frost sensitivity in 170 woody plants from the Northern
- Hemisphere. Global Ecology and Biogeography 25, 1061–1071.

- Myking, T. & Skroppa, T. (2007) Variation in phenology and height increment of northern *Ulmus glabra*populations: Implications for conservation. *Scandinavian Journal of Forest Research* **22**, 369–374.
- Norgaard Nielsen, C.C. & Rasmussen, H.N. (2009) Frost hardening and dehardening in *Abies procera* and other conifers under differing temperature regimes and warm-spell treatments. *Forestry* **82**, 43–59.
- ³²⁸ O'Keefe, J. (2014) Phenology of woody species at Harvard Forest since 1990. Tech. rep.
- Partanen, J. (2004) Dependence of photoperiodic response of growth cessation on the stage of development in *Picea abies* and *Betula pendula* seedlings. *Forest Ecology and Management* **188**, 137–148.
- Pearce, R. (2001) Plant freezing and damage. Annals of Botany 87, 417–424.
- Peterson, A.G. & Abatzoglou, J.T. (2014) Observed changes in false springs over the contiguous United

 States. Geophysical Research Letters 41, 2156–2162.
- Poirier, M., Lacointe, A. & Ameglio, T. (2010) A semi-physiological model of cold hardening and dehardening in walnut stem. *Tree Physiology* **30**, 1555–1569.
- Prozherina, N., Freiwald, V., Rousi, M. & Oksanen, E. (2003) Interactive effect of springtime frost and elevated ozone on early growth, foliar injuries and leaf structure of birch (*Betula pendula*). New Phytologist 159, 623–636.
- Richardson, A. & O'Keefe, J. (2009) Phenological differences between understory and overstory: a case study using the long-term harvard forest records, pp. 87–117. A. Noormets (Ed.), Phenology of Ecosystem Processes, Springer, New York.
- 342 Richardson, A.D. (2015) PhenoCam images and canopy phenology at Harvard Forest since 2008.
- Sakai, A. & Larcher, W. (1987) Frost Survival of Plants. Springer-Verlag.
- Sánchez, B., Rasmussen, A. & Porter, J.R. (2013) Temperatures and the growth and development of maize and rice: a review. *Global Change Biology* **20**, 408–417.
- Schaber, J. & Badeck, F.W. (2005) Plant phenology in germany over the 20th century. Regional Environmental

 Change 5, 37–46.
- Schwartz, M.D. (1993) Assessing the onset of spring: A climatological perspective. *Physical Geography* **14(6)**, 536–550.

- Søgaard, Gunnhild and Johnsen, Øystein and Nilsen, Jarle and Junttila, Olavi (2008) Climatic control of bud burst in young seedlings of nine provenances of Norway spruce. *Tree Physiology* **28**, 311–320.
- Soudani, K., Hmimina, G., Delpierre, N., Pontailler, J.Y., Aubinet, M., Bonal, D., Caquet, B., de Grandcourt,
- A., Burban, B., Flechard, C. & et al. (2012) Ground-based Network of NDVI measurements for tracking
- temporal dynamics of canopy structure and vegetation phenology in different biomes. Remote Sensing of
- Environment **123**, 234–245.
- Taschler, D., Beikircher, B. & Neuner, G. (2004) Frost resistance and ice nucleation in leaves of five woody timberline species measured in situ during shoot expansion. *Tree Physiology* **24**, 331–337.
- USA-NPN (2016) USA National Phenology Network Extended Spring Indices.
- Vihera-aarnio, A., Hakkinen, R. & Junttila, O. (2006) Critical night length for bud set and its variation in two photoperiodic ecotypes of *Betula pendula*. Tree Physiology **26**, 1013–1018.
- Vitasse, Y., Lenz, A., Hoch, G. & Körner, C. (2014a) Earlier leaf-out rather than difference in freezing resistance puts juvenile trees at greater risk of damage than adult trees. *Journal of Ecology* **102**, 981–988.
- Vitasse, Y., Lenz, A. & Körner, C. (2014b) The interaction between freezing tolerance and phenology in temperate deciduous trees. Frontiers in Plant Science 5, 541.
- Vitra, A., Lenz, A. & Vitasse, Y. (2017) Frost hardening and dehardening potential in temperate trees from
 winter to budburst. New Phytologist 216, 113–123.
- Way, D.A. & Montgomery, R.A. (2015) Photoperiod constraints on tree phenology, performance and migration in a warming world. *Plant, Cell & Environment* 38, 1725–1736.
- White, M.A., De Beurs, K.M., Didan, K., Inouye, D.W., Richardson, A.D., Jensen, O.P., O'Keefe, J., Zhang, G., Nemani, R.R., Van Leeuwen, W.J.D. & Al., E. (2009) Intercomparison, interpretation, and assessment of spring phenology in North America estimated from remote sensing for 1982-2006. *Global Change Biology* 15, 2335–2359.
- Xie, Y., Wang, X. & Silander, J.A. (2015) Deciduous forest responses to temperature, precipitation, and drought imply complex climate change impacts. *Proceedings of the National Academy of Sciences* 112, 13585–13590.

- Xin, Q. (2016) A risk-benefit model to simulate vegetation spring onset in response to multi-decadal climate variability: Theoretical basis and applications from the field to the Northern Hemisphere. Agriculture and
- ³⁷⁸ Forest Meteorology **228-229**, 139–163.
- Zohner, C.M., Benito, B.M., Svenning, J.C. & Renner, S.S. (2016) Day length unlikely to constrain climatedriven shifts in leaf-out times of northern woody plants. *Nature Climate Change* **6**, 1120–1123.