

Spatial and temporal shifts in photoperiod with climate change

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Abstract

Climate change causes both temporal and geographic shifts in species; these shifts in turn affect the daylength (photoperiod) that species experience. As photoperiod is a common trigger of seasonal biological responses (e.g., affecting plant phenology in 84% of studies that manipulated photoperiod), such shifts in experienced photoperiod may have important implications for future distributions and fitness of many species. However, photoperiod has not been a focus of climate change forecasting to date, especially for early-season ('spring') events—which are often assumed to be driven by temperature. Here we show that impacts on experienced photoperiod due to temporal shifts could be quite large and may be orders of magnitude larger than impacts due to spatial shifts (e.g., 1.6 hours of change for expected temporal shifts versus only one minute for spatial shifts). Incorporating these effects into forecasts may be possible by leveraging existing experimental data; for example, growth chamber experiments on woody plant spring phenology often have data relevant for climate change impacts. We highlight how combining novel modeling approaches and empirical work on when, where, and how much photoperiod affects spring phenology, could rapidly advance our understanding and predictions of future spatial-temporal shifts due to climate change.

Introduction

Shifts in the timing of spring events—i.e., phenology, including flowering, bird arrival, egg hatching and myriad other biological activities—are some of the most widely documented signals of climate change. Across taxa, from plants and insects to mollusks and mammals, spring phenology is occurring earlier as temperatures warm, with average shifts of 1.2 to 5.1 days earlier per decade (Bradley et al., 1999; Parmesan and Yohe, 2003; Poloczanska et al., 2013; Root et al., 2003) or 1.3 to 5.6 days earlier per °C of warming (Polgar et al., 2013; Wolkovich et al., 2012). These changes are some of the largest climate change induced shifts observed, with early spring phenology shifting more rapidly than later season phenology in most cases (Bradley et al., 1999; Menzel et al., 2006), and suggest that temperature is a major driver of spring phenophases.

Spring phenology is not controlled solely by temperature, however. Photoperiod is also a critical cue for plants and animals, signaling changes in growth, mating, and reproduction across diverse species (e.g., Flynn and Wolkovich, 2018; Howe et al., 1996; Lagercrantz, 2009; Mcallan et al., 2006; Solbakken et al., 1994). Photoperiod is a useful cue to synchronize activities with seasonal climatic changes (e.g., Basler and Körner, 2012; Hsu et al., 2011; Singh et al., 2017) because it is consistent across years, especially compared to other seasonal cues such as temperature and precipitation (Saikkonen et al., 2012). For example, relying on a threshold photoperiod (see *Glossary*), rather than temperature alone, may prevent woody plants from leafing out during “false spring” events (unusually warm periods during winter that are followed by a return of cold temperatures, Gu et al., 2008). With current rapid warming photoperiod may also potentially slow the observed trend of advancing spring phenology.

Recent studies offer inconsistent views about whether photoperiod may eventually restrict advances in spring phenology in a warmer world. Some studies suggest that, with additional warming, photoperiod will limit phenological shifts of certain species such that they will not track rising temperatures (e.g., by leafing out earlier in the spring, Körner and Basler, 2010; Way and Montgomery, 2015). Instead, these species’ responses will increasingly become constrained by daylength and the trend of ever-earlier springs with warming may halt. Other studies, however, suggest that photoperiod will not constrain responses to warming for most species (Chuine et al., 2010; Zohner et al., 2016). The extent to which daylength constrains responses will depend in part on how rapidly photoperiod cues can acclimate or adapt to new environmental conditions, which remains poorly understood (Grevstad and Coop, 2015).

Perhaps because of these variable and uncertain responses, photoperiod is often not included in forecasts of biological responses to climate change, especially in the spring, even though it is known to be an important cue for biological activity (but see Caffarra et al., 2011a; Duputié et al., 2015; Grevstad and Coop, 2015). The exclusion of photoperiod may be problematic: although photoperiod itself is stable over time, the photoperiod that species *experience*, as they undergo climate change-induced shifts in space and time, is likely to be much less stable. In addition to shifting activity earlier with recent warming, many species have shifted their distributions poleward and upward in elevation (i.e., range shifts, Chen et al., 2011; Harsch et al., 2009; Parmesan, 2006; Peñuelas and Boada, 2003). These spatial and temporal shifts alter the photoperiod experienced by organisms (Fig. 1); altered photoperiods may have cascading effects on species’ performance, since daylength can affect the timing of development (Grevstad and Coop, 2015; Muir et al., 1994), migration (Dawbin, 1966), and other important responses.

The implications of potential climate change-induced shifts in experienced photoperiod are unclear, since the magnitude of potential shifts has not been described. Effects of photoperiod shifts may be relatively minor, especially because there can be substantial year-to-year variation in experienced photoperiod (Fig. 2). Alternatively, photoperiod may begin to constrain species’ responses to climate change (Körner and Basler, 2010).

Here, we ask:

1. How will climate change alter the photoperiod experienced by organisms?

2. What are the implications of altered photoperiods for biological responses to climate change?
3. Can research apply experiments that alter photoperiod to forecasting biological implications of climate change?

These questions are broadly relevant for diverse species. Here, we use a case study of spring woody plant phenology to illustrate our points (Box 1). We focus on spring events, as phenology during this time is one of the most widely observed and rapidly changing biological responses to climate change (Parmesan, 2006). Woody species are a useful focal group because they have been the subject of decades of growth chamber experiments, are at the center of an important and controversial debate on the relative effects of photoperiod versus temperature on their phenology, and because forecasting effects of climate change on their phenology (i.e., the length of the growing season) has critical implications for global carbon cycling and feedbacks to the climate system (Richardson et al., 2013). We use studies included in Observed Spring Phenology Responses in Experimental Environments (OSPREE), a new database of plant growth chamber studies that manipulate photoperiod and temperature to measure plant phenological responses, including budburst and flowering (Wolkovich et al., 2019).

How will climate change alter the photoperiod experienced by organisms?

Species experience different photoperiod regimes depending on their location on Earth (Fig. 1, 2), the seasonal timing of their activity, and inter-annual variation in climate. The daylength experienced by plants on the date that spring “green-up” occurs, for example, varies with latitude (Fig. 2a). This is in part because latitudinal variation in green-up date, which occurs earlier toward the equator and later toward the north pole, is strongly driven by climatic differences that affect phenology, and in part because of latitudinal variation in photoperiod (e.g., at the north pole, the daylength at the summer solstice is 24 hours). A general pattern of longer photoperiod at green-up toward the poles is consistent across years (Fig. 2b) and green-up does not appear to occur at daylengths less than 10 hours. There is strong spatiotemporal variation in experienced photoperiod across years (compare the photoperiod at green-up in “early” versus “late” years, Fig. 2): experienced photoperiod at green-up can vary two to three hours from one year to the next in the same location (Fig. 2c). Though green-up date corresponds to plant phenology, we expect that spatiotemporal patterns of variation in spring phenology would be similar for other organisms (Ovaskainen et al., 2013; Peñuelas et al., 2002).

Against this existing background variation, climate change will cause shifts in experienced photoperiod as species respond to warming temperatures. Spatial shifts in species’ ranges and temporal shifts in phenology will alter the photoperiods experienced by organisms with future climate change. The magnitude of these alterations will vary depending on the organism’s location and the type of shift(s) it undergoes. For example, poleward shifts in species’ ranges cause organisms to experience a wider range of daylength throughout the year (Fig. 1). Elevational shifts, in contrast, cause minimal changes in the range of daylength throughout the year.

To date, where the scientific literature has addressed shifts in photoperiod with climate change, the focus has been on how spatial range shifts will affect photoperiod (e.g., Saikkonen et al., 2012; Way and Montgomery, 2015). However, shifting phenology—especially the large changes seen in spring phenology—will also alter experienced photoperiod, because of the seasonal patterns of daylength (Fig. 1).

Despite a focus on range shifts, current data suggest that temporal shifts will yield much larger changes in experienced photoperiod than spatial shifts (Fig. 1). For example, consider an insect that emerges from diapause or a tree that bursts its buds at latitude 45°, on average, around day of year 91 (April 2, when daylength is 12.8 hours). If the organism’s phenology shifts 30 days earlier over the next century (i.e., a rate

of 3 days per decade, as has been observed, Parmesan and Yohe, 2003), it will experience a daylength that is 1.6 hours shorter. This 1.6 hour decrease in daylength is equivalent to moving up 28.5° in latitude on this day of year. However, if the same species shifts its range up in latitude 0.5° (i.e., 60 km over the next century, comparable to observed rates, Parmesan and Yohe, 2003; Chen et al., 2011), it will experience a daylength that differs by less than a minute on the same day of year.

In many cases organisms may shift both their ranges and their phenology simultaneously (i.e., due to new climatic conditions, Duputié et al., 2015; Grevstad and Coop, 2015). In addition, photoperiod sensitivity (see *Glossary*) can vary with latitude, likely due to population-level differences in sensitivity (Caffarra et al., 2011b; Gauzere et al., 2017; Howe et al., 1996; Partanen et al., 2005; Saikkonen et al., 2012; Viherä-Aarnio et al., 2006). With future climate change, it is unclear how these complexities will affect the photoperiod experienced by organisms and if these shifts in photoperiod will have important implications for biological responses. This lack of clarity stems, in part, from the fact that phenology both affects and is affected by experienced photoperiod: climate change-induced shifts in phenology alter experienced photoperiod, which in turn affects phenology.

What are the implications of altered photoperiods for biological responses to climate change?

Daylength can play a role in controlling critical biological functions, including vegetative growth, cell elongation, budburst, and flowering in plants (Ashby et al., 1962; Erwin, 1998; Sidaway-Lee et al., 2010; Heide, 2011; Heide and Sønsteby, 2012; Hsu et al., 2011; Linkosalo and Lechowicz, 2006; Mimura and Aitken, 2007) and growth rate, maturation, migration, and diapause in animals (Bradshaw and Holzapfel, 2006; Dawbin, 1966; Muir et al., 1994; Saunders and Henderson, 1970; Tobin et al., 2008; Zydlewski et al., 2014). Climate change-induced shifts in photoperiod are therefore likely to alter these functions. Indeed, growth chamber studies demonstrate that the magnitude of daylength shifts we can expect with climate change (i.e., 1-2 hours of difference in daylength with temporal shifts over the next century) are substantial enough to affect spring phenology in trees (Table S1). The direction and magnitude of responses will vary, however, because of variation in photoperiod sensitivity, and because photoperiod often interacts with other environmental drivers, such as temperature, to affect phenology (Box 1).

The climate change-induced trend toward ever earlier springs means that experienced photoperiod may increasingly approach threshold photoperiod for many species, constraining their ability to respond to additional warming (Körner and Basler, 2010; Morin et al., 2010; Nienstaedt, 1966; Vittasse and Basler, 2013). Interactions between photoperiod and temperature may therefore result in muted phenological shifts, compared to what would be expected based on temperature change alone (Körner and Basler, 2010; Mimura and Aitken, 2007; Wareing, 1956). If photoperiod does become limiting, the average trend of earlier phenology with warming (Menzel, 2000; Ovaskainen et al., 2013; Peñuelas et al., 2002; Polgar et al., 2013) may stop.

A challenge in understanding the implications of altered photoperiods under climate change, and for forecasting whether and when the trend of earlier phenology with warming may slow or stop abruptly, is the wide range of observed photoperiod sensitivity across species (Flynn and Wolkovich, 2018; Sanz-Perez et al., 2009; Zohner et al., 2016), populations (Tanino et al., 2010), and ecotypes (Howe et al., 1995). How much genotype versus environment explain this variation is an active area of research (e.g., Franks et al., 2014; Gould et al., 2010; Mimura and Aitken, 2010; Fréjaville et al.). Environmental conditions clearly play a role, since different combinations of ambient temperature and photoperiod may explain some of this variation, because temperature cues can override photoperiod requirements under certain conditions (e.g., Tanino et al., 2010). In such cases, climate change-induced phenological shifts may occur at different rates than past shifts with warming. On the other hand, some of this variation may be due to underlying genetic differences, because photoperiod responses can be under strong genetic control (Bradshaw and Stettler, 1995; Keller et al., 2011; Weih, 2004, , see also Box 1). Teasing out the relative roles of genetics versus environmental conditions will

be critical to accurate forecasts of future phenology under climate change.

Species- and population-level variation in photoperiod sensitivity may result in altered communities as climate change progresses. For example, a species or population that is relatively insensitive to photoperiod can take advantage of warmer springs by having an earlier start to its growing season. Indeed, phenological tracking of temperature (e.g., earlier flowering, leafout, migration with warming) has been linked with higher performance in plants and animals (Cleland et al., 2012; Muir et al., 1994; Willis et al., 2010). Species or populations that are sensitive to temperature but relatively insensitive to photoperiod may therefore outcompete slower growing or later emerging ones that are limited by photoperiod and thus cannot take advantage of longer growing season conditions. To identify where, when, and how communities may be altered, methods for incorporating photoperiod into forecasting future phenology are critical.

Future directions: outstanding questions and incorporating photoperiod into forecasting

Incorporating photoperiod into forecasting is complex for a few major reasons. Future rates of phenological shifts are unlikely to be straightforward extrapolations from past and current rates. In addition, an organism's experienced photoperiod is both a driver and an effect of phenological shifts.

Approaches for forecasting can be grouped into two broad categories: statistical models and process-based models. These two modelling paradigms differ in at least two ways, in terms of relating phenology to climate change. First, statistical models generally assume linear relationships between species' responses and environmental variables (e.g., Flynn and Wolkovich, 2018; Van Belle et al., 2007; Ibáñez et al., 2010), instead process-based models often incorporate nonlinear threshold relationships as well (e.g. Chuine and Beaubien, 2001; Morin and Thuiller, 2009; Xie and Hsieh, 1989). Second, statistical models of phenology under climate change have typically ignored photoperiod, focusing instead on seasonal or annual temperature (e.g. Diez et al., 2012; Ibáñez et al., 2010; Van Belle et al., 2007, but see Richardson et al. (2013)). whereas process-based models of phenology more frequently incorporate photoperiod, along with temperature (Duputié et al., 2015; Morin and Thuiller, 2009; Xie and Hsieh, 1989; Zhao et al., 2013). A challenge of process-based models is that they require detailed data that is often not readily available (e.g., daily climate data, nonlinear biological responses to fine-scale changes in temperature). Perhaps because of this challenge, statistical models remain more commonly used in climate change forecasts of biological responses (e.g., Basler and Körner, 2012; Diez et al., 2012; García-Valdés and Morales-Castilla, 2016; Ibáñez et al., 2010; Van Belle et al., 2007; Zhu et al., 2012).

Future modelling can incorporate photoperiod by leveraging the large amount of experimental data on photoperiod responses (Fig. 4, Table S1), especially when process-based approaches are used. Researchers can use these data to first learn if the study species (or a phylogenetically closely related species) shows a photoperiod effect and, ideally, identify its threshold photoperiod and how it varies by population, ecotype, or other factors (Bradshaw and Holzapfel, 2006; Gwinner, 1996; Tobin et al., 2008). If there is evidence of a photoperiod response (e.g., *Fagus grandifolia*, or *Tilia americana* with low chilling in Fig. 3), daylength should be added to forecasting models, using the threshold photoperiod to define short-day and long-day conditions (Fig. 6). Given the large change in experienced photoperiod with temporal shifts (Fig. 1), this may be particularly important for phenological forecasting. Since spatial shifts are associated with smaller changes in experienced photoperiod, it may be less important for distribution forecasts. Many species, however, may shift in *both* space and time simultaneously. Thus, even though experienced photoperiod changes little as species distributions shift in space, phenology may be altered significantly.

For some species, experimental data can be immediately used in forecasting because experiments manipulate photoperiod at relevant scales (e.g., Basler and Körner, 2014; Heide and Sonsteby, 2015, Figs. 4, 5 A, Table S1). For example, photoperiod treatments from growth chamber experiments with *Fagus sylvatica*

span the variation in both current and expected future ranges (Fig. 5, Duputié et al., 2015), and may allow identification of threshold photoperiods (Fig. 6). In other cases, attempting to incorporate photoperiod into forecasts of future phenology will reveal gaps in our understanding of many aspects of photoperiod responses. For example, photoperiod treatments from existing experiments of *Quercus robur* do not accurately represent experienced photoperiods from current or future estimates, making fine scale projections difficult, even for this relatively well-studied species. This gap extends to many species, as most experiments manipulate photoperiod much more dramatically than will occur with climate change (Figs. 4, 5). Although these studies can be useful for understanding mechanistically how photoperiod responses work, extrapolating them to climate change models may not be reasonable.

Photoperiod is not fully integrated into most current forecasts of biological responses to climate change (but see Tobin et al., 2008), an omission that could affect the accuracy of forecasts. Forecasts from ecosystem models often incorporate photoperiod, along with other variables such as evaporative demand and temperature (e.g., ED Jolly et al., 2005; Medvigy et al., 2013), but photoperiod is rarely included in species distribution models (e.g., Morin and Thuiller, 2009; Zhu et al., 2012). The sensitivity of model outcomes to assumptions made about experienced photoperiod and threshold responses to photoperiod needs further study, including understanding how variation in photoperiod responses across ecosystems, species, populations, and life stages impacts forecasts.

As researchers more fully integrate photoperiod into forecasting, a critical area of further study is understanding *how* photoperiod acts as a cue. Photoperiod seems to interact with temperature to affect phenology (e.g., Zydlowski et al., 2014); this would explain the divergent effects of photoperiod observed across studies in woody plants (e.g., Fig. 3). However, exactly how it interacts with temperature is not well-defined for most species or populations (Boxes 1, S1). Understanding the drivers, as well as the consequences, of variations in photoperiod responses across species and populations will be particularly beneficial for forecasting. For example, what traits are associated with photoperiod sensitivity and does variation in photoperiod sensitivity or related traits have a strong genetic component? If so, are species or populations from some locations or lineages more likely than others to be constrained by photoperiod in their responses to climate change?

Conclusions

Organisms may undergo large changes to the photoperiod they experience with climate change, even if they do not shift their ranges spatially. Here we have shown that these altered photoperiods may result in stalled future advances of woody plant phenology with warming (Table S1, Fig. 4), with cascading effects on growth, fitness, and community composition due to the large variation in photoperiod responses across species and populations (Fig. 3). Shifts in photoperiod with climate change have implications for a variety of plant and animal responses, given that daylength affects critical activities for diverse species from insects (Bradshaw and Holzapfel, 2006; Linn et al., 1996) and salmon (Solbakken et al., 1994; Taranger et al., 2003) to birds (Dawson et al., 2001) and marsupials (McCallan et al., 2006; Solbakken et al., 1994). Given what we know, incorporating photoperiod into forecasting of climate change responses should improve model accuracy, and will illuminate additional experiments that could improve our mechanistic understanding of photoperiod as a critical cue for diverse biological responses.

Glossary

- budburst: when one or more leaf buds have visible green tips.
- chilling: the intensity and duration of winter temperature, often a certain sum of chilling that is required (e.g., some amount of hours or days of cold temperatures, defined by a specific critical temperature or

range of temperatures, such as between 0 and 7.2 °C, Richardson, 1974), that must be experienced for budburst to occur.

- daylength: the period of time during a 24-hour period during which an organism receives light.
- diapause: period of suspended development or growth, usually used to describe invertebrates during unfavorable environmental conditions such as winter
- dormancy: halted or reduced growth or activity, usually used to describe plants
- forcing: warm spring temperatures, often a certain sum of forcing that is required (e.g., some amount of hours or days above a specific temperature) for budburst or flowering can occur.
- green-up: The beginning of a new cycle of plant growth, usually evaluated at the landscape scale
- phenology: the timing of life cycle events in organisms
- photoperiod: the daily duration of light (daylength) and dark to which an organism is exposed; often used synonymously with daylength
- photoperiod sensitivity: the degree to which phenology is controlled by daylength; may be a nonlinear, or “threshold”, response in plants (Box S1) and animals (Tobin et al., 2008; Grevstad and Coop, 2015).
- photoperiodism: the ability to assess the length of day or night to regulate behavior, physiology, growth, development or reproduction.
- threshold photoperiod: length of day that causes an organism to switch from a short- to a long-day response (or vice versa). For example, in European larch (*Larix decidua*), budburst development may be constrained under short-day conditions, when daylengths are less than a threshold photoperiod of 10-11 hours (Migliavacca et al., 2008). Above this threshold photoperiod, the long-day response of unconstrained budburst development can occur.

Box 1. Are photoperiod effects widespread? A case study of woody plant spring phenology

Photoperiod responses are particularly well-studied in woody plant phenology. Decades of experimental growth chamber studies have shown that photoperiod is an important cue for spring budburst phenology in woody plants (e.g., Basler and Körner, 2014; Flynn and Wolkovich, 2018; Heide, 1993b). These experiments often manipulate photoperiod in combination with temperature to address basic questions about how these two environmental conditions act as biological cues. Temperature has a dual role in regulating woody plant phenology: chilling—the prolonged exposure to cold temperatures after growth cessation in the fall—is required to initiate budburst; and forcing—prolonged exposure to warm temperatures—is required for budburst to occur. Thus, chilling and forcing treatments are often altered in addition to photoperiod in growth chamber experiments (e.g., Campbell and Sugano, 1975; Falusi and Calamassi, 1990; Heide, 1977; Laube et al., 2014; Spann et al., 2004).

Woody plant growth chamber studies have been conducted for decades, but have only recently been synthesized (Wolkovich et al., 2019), revealing that photoperiod sensitivity is widespread, though with wide variation across studies and species. Growth chamber experiments in OSPREE suggest that the dominant photoperiod response in woody plant species is earlier and more rapid budburst with longer days (e.g., Caffarra and Donnelly, 2011). Thirty-one of the 85 studies in the OSPREE database included two or more different photoperiod treatments. Of these, 26 (84%) found significant photoperiod main effects or significant interactive effects with temperature (i.e., photoperiod x temperature effects), across 176 species (Table S1).

Main effects included responses such as growth (e.g., higher growth rates with longer days Ashby et al., 1962) and reproduction (e.g., increased flowering with longer days Heide and Sønsteby, 2012).

Growth chamber experiments highlight that responses to photoperiod vary depending on temperature conditions. For example, more rapid advancement of budburst was observed under long versus short days with low chilling, than with high chilling in *Betula paynifera* (Hawkins and Dhar, 2012) (Fig. 3). Frequently, long photoperiods can compensate for low amounts of chilling, resulting in enhanced cell growth (Heide, 1993a; Myking and Heide, 1995; Caffarra et al., 2011b).

Woody plant growth chamber experiments also demonstrate that, though photoperiod responses are common, they are variable (Fig. 3). Responses to photoperiod differ by species (e.g., Basler and Körner, 2012, 2014; Flynn and Wolkovich, 2018; Heide, 1993b; Howe et al., 1996; Zohner et al., 2016). For example, with longer chilling treatments some species seem insensitive to daylength (e.g., *Hammamelis* spp., *Prunus* spp., Zohner et al., 2016), whereas others (e.g. *Fagus* spp., Fig. 5A) seem to be highly sensitive to daylength, even with long chilling treatments (Zohner et al., 2016). In addition, some species demonstrated an opposing response to photoperiod than typically observed: *Tilia*, for example, showed delayed budburst with longer daylengths (Fig. 3, Ashby et al., 1962). Photoperiod sensitivity also varies by population and ecotype (e.g., Partanen et al., 2005) (Fig. 3). For example, photoperiod effects on budburst were more significant for lower latitude populations of *Betula pendula* and *B. pubescens* (Partanen et al., 2005).

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References

- Ashby, W., et al. 1962. Germination capacity in American Basswood *Tilia americana*. Transactions of the Illinois State Academy of Science 55:120–3.
- Basler, D., and C. Körner. 2012. Photoperiod sensitivity of bud burst in 14 temperate forest tree species. Agricultural and Forest Meteorology 165:73–81.
- . 2014. Photoperiod and temperature responses of bud swelling and bud burst in four temperate forest tree species. Tree Physiology 34:377–388.
- Bradley, N. L., A. C. Leopold, J. Ross, and W. Huffaker. 1999. Phenological changes reflect climate change in Wisconsin. Proceedings of the National Academy of Sciences 96:9701–9704.
- Bradshaw, H., and R. F. Stettler. 1995. Molecular genetics of growth and development in *Populus*. IV. Mapping QTLs with large effects on growth, form, and phenology traits in a forest tree. Genetics 139:963–973.
- Bradshaw, W. E., and C. M. Holzapfel. 2006. Evolutionary response to rapid climate change. Science 312:1477–1478.

- Caffarra, A., and A. Donnelly. 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.
- Caffarra, A., A. Donnelly, and I. Chuine. 2011a. Modelling the timing of *Betula pubescens* budburst. II. Integrating complex effects of photoperiod into process-based models. Climate Research 46:159–170.
- Caffarra, A., A. Donnelly, I. Chuine, and M. B. Jones. 2011b. Modelling the timing of *Betula pubescens* bud-burst. I. Temperature and photoperiod: A conceptual model. Climate Research 46:147.
- Campbell, R. K., and A. I. Sugano. 1975. Phenology of bud burst in Douglas-fir related to provenance, photoperiod, chilling, and flushing temperature. Botanical Gazette pages 290–298.
- Chen, I.-C., J. K. Hill, R. Ohlemüller, D. B. Roy, and C. D. Thomas. 2011. Rapid range shifts of species associated with high levels of climate warming. Science 333:1024–1026.
- Chuine, I., and E. G. Beaubien. 2001. Phenology is a major determinant of tree species range. Ecology Letters 4:500–510.
- Chuine, I., X. Morin, and H. Bugmann. 2010. Warming, photoperiods, and tree phenology. Science 329:277–278.
- Cleland, E. E., J. M. Allen, T. M. Crimmins, J. A. Dunne, S. Pau, S. E. Travers, E. S. Zavaleta, and E. M. Wolkovich. 2012. Phenological tracking enables positive species responses to climate change. Ecology 93:1765–1771.
- Dawbin, W. H. 1966. The seasonal migratory cycle of humpback whales. Whales, dolphins and porpoises pages 145–170.
- Dawson, A., V. M. King, G. E. Bentley, and G. F. Ball. 2001. Photoperiodic control of seasonality in birds. Journal of Biological Rhythms 16:365–380.
- Diez, J. M., I. Ibáñez, A. J. Miller-Rushing, S. J. Mazer, T. M. Crimmins, M. A. Crimmins, C. D. Bertelsen, and D. W. Inouye. 2012. Forecasting phenology: from species variability to community patterns. Ecology Letters 15:545–553.
- Duputié, A., A. Rutschmann, O. Ronce, and I. Chuine. 2015. Phenological plasticity will not help all species adapt to climate change. Global Change Biology 21:3062–3073.
- Erwin, J. E. 1998. Temperature and light effects on stem elongation (plant growth regulation by physical and mechanical stimuli, for further development of horticulture in East Asia). Journal of the Japanese Society for Horticultural Science 67:1113–1120.
- Falusi, M., and R. Calamassi. 1990. Bud dormancy in beech (*Fagus sylvatica* L.). Effect of chilling and photoperiod on dormancy release of beech seedlings. Tree Physiology 6:429–438.
- Flynn, D. F. B., and E. M. Wolkovich. 2018. Temperature and photoperiod drive spring phenology across all species in a temperate forest community. New Phytologist 219:1353–1362.
- Franks, S. J., J. J. Weber, and S. N. Aitken. 2014. Evolutionary and plastic responses to climate change in terrestrial plant populations. Evolutionary Applications 7:123–139.
- Fréjaville, T., B. Fady, A. Kremer, A. Ducouso, and M. Benito Garzón. ???? Inferring phenotypic plasticity and population responses to climate across tree species ranges using forest inventory data. Global Ecology and Biogeography .
- García-Valdés, R., and I. Morales-Castilla. 2016. Efectos del cambio climático en los ecosistemas forestales: integrando inventarios y modelos. Ecosistemas 25:51–59.

- Gauzere, J., S. Delzon, H. Davi, M. Bonhomme, I. G. de Cortazar-Atauri, and I. Chuine. 2017. Integrating 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 244:9–20.
- Gould, P. J., C. A. Harrington, and J. B. St. Clair. 2010. Incorporating genetic variation into a model of budburst phenology of coast douglas-fir (*pseudotsuga menziesii* var. *menziesii*). Canadian journal of forest research 41:139–150.
- Grevstad, F. S., and L. B. Coop. 2015. The consequences of photoperiodism for organisms in new climates. Ecological Applications 25:1506–1517.
- Gu, L., P. J. Hanson, W. M. Post, D. P. Kaiser, B. Yang, R. Nemani, S. G. Pallardy, and T. Meyers. 2008. The 2007 Eastern US spring freeze: Increased cold damage in a warming world. BioScience 58:253.
- Gwinner, E. 1996. Circadian and circannual programmes in avian migration. Journal of Experimental Biology 199:39–48.
- Harsch, M. A., P. E. Hulme, M. S. McGlone, and R. P. Duncan. 2009. Are treelines advancing? A global meta-analysis of treeline response to climate warming. Ecology Letters 12:1040–1049.
- Hawkins, C. D., and A. Dhar. 2012. Spring bud phenology of 18 *Betula papyrifera* populations in British Columbia. Scandinavian Journal of Forest Research 27:507–519.
- Heide, O. 1993a. Daylength and thermal time responses of budburst during dormancy release in some northern deciduous trees. Physiologia Plantarum 88:531–540.
- . 1993b. Dormancy release in beech buds (*Fagus sylvatica*) requires both chilling and long days. Physiologia Plantarum 89:187–191.
- Heide, O. M. 1977. Photoperiod and temperature interactions in growth and flowering of strawberry. Physiologia Plantarum 40:21–26.
- . 2011. Temperature rather than photoperiod controls growth cessation and dormancy in *Sorbus* species. Journal of Experimental Botany 62:5397–5404.
- Heide, O. M., and A. Sønsteby. 2012. Floral initiation in black currant cultivars (*Ribes nigrum* L.): Effects of plant size, photoperiod, temperature, and duration of short day exposure. Scientia Horticulturae 138:64–72.
- Heide, O. M., and A. Sonsteby. 2015. Simultaneous dormancy induction interferes with short day floral induction in black currant (*Ribes nigrum* L.). Scientia Horticulturae 185:228–232.
- Howe, G. T., G. Gardner, W. P. Hackett, and G. R. Furnier. 1996. Phytochrome control of short-day-induced bud set in black cottonwood. Physiologia Plantarum 97:95–103.
- Howe, G. T., W. P. Hackett, G. R. Furnier, and R. E. Klevorn. 1995. Photoperiodic responses of a northern and southern ecotype of black cottonwood. Physiologia Plantarum 93:695–708.
- Hsu, C.-Y., J. P. Adams, H. Kim, K. No, C. Ma, S. H. Strauss, J. Drnevich, L. Vandervelde, J. D. Ellis, B. M. Rice, et al. 2011. FLOWERING LOCUS T duplication coordinates reproductive and vegetative growth in perennial poplar. Proceedings of the National Academy of Sciences 108:10756–10761.
- Ibáñez, I., R. B. Primack, A. J. Miller-Rushing, E. Ellwood, H. Higuchi, S. D. Lee, H. Kobori, and J. A. Silander. 2010. Forecasting phenology under global warming. Philosophical Transactions of the Royal Society B-Biological Sciences 365.
- Jolly, W. M., R. Nemani, and S. W. Running. 2005. A generalized, bioclimatic index to predict foliar phenology in response to climate. Global Change Biology 11:619–632.

- Keller, S. R., R. Y. Soolanayakanahally, R. D. Guy, S. N. Silim, M. S. Olson, and P. Tiffin. 2011. Climate-driven local adaptation of ecophysiology and phenology in balsam poplar, *Populus balsamifera* L. (Salicaceae). American Journal of Botany 98:99–108.
- Körner, C., and D. Basler. 2010. Warming, photoperiods, and tree phenology response. Science 329:278–278.
- Lagercrantz, U. 2009. At the end of the day: a common molecular mechanism for photoperiod responses in plants? Journal of Experimental Botany 60:2501–2515.
- Laube, J., T. H. Sparks, N. Estrella, J. Höfler, D. P. Ankerst, and A. Menzel. 2014. Chilling outweighs photoperiod in preventing precocious spring development. Global Change Biology 20:170–182.
- Linkosalo, T., and M. J. Lechowicz. 2006. Twilight far-red treatment advances leaf bud burst of silver birch (*Betula pendula*). Tree Physiology 26:1249–1256.
- Linn, C. E., M. G. Campbell, K. R. Poole, W.-Q. Wu, and W. L. Roelofs. 1996. Effects of photoperiod on the circadian timing of pheromone response in male *Trichoplusia ni*: relationship to the modulatory action of octopamine. Journal of Insect Physiology 42:881–891.
- McCallan, B. M., C. R. Dickman, and M. S. Crowther. 2006. Photoperiod as a reproductive cue in the marsupial genus *Antechinus*: ecological and evolutionary consequences. Biological Journal of the Linnean Society 87:365–379.
- Medvigy, D., S.-J. Jeong, K. L. Clark, N. S. Skowronski, and K. V. Schäfer. 2013. Effects of seasonal variation of photosynthetic capacity on the carbon fluxes of a temperate deciduous forest. Journal of Geophysical Research: Biogeosciences 118:1703–1714.
- Menzel, A. 2000. Trends in phenological phases in Europe between 1951 and 1996. International Journal of Biometeorology 44:76–81.
- Menzel, A., T. H. Sparks, N. Estrella, E. Koch, A. Aasa, R. Ahas, K. Alm-Kuebler, et al. 2006. European phenological response to climate change matches the warming pattern. Global Change Biology 12:1969–1976.
- Migliavacca, M., E. Cremonese, R. Colombo, L. Busetto, M. Galvagno, L. Ganis, M. Meroni, E. Pari, M. Rossini, C. Siniscalco, et al. 2008. European larch phenology in the alps: can we grasp the role of ecological factors by combining field observations and inverse modelling? International journal of biometeorology 52:587–605.
- Mimura, M., and S. Aitken. 2007. Adaptive gradients and isolation-by-distance with postglacial migration in *Picea sitchensis*. Heredity 99:224.
- . 2010. Local adaptation at the range peripheries of sitka spruce. Journal of evolutionary biology 23:249–258.
- Morin, X., J. Roy, L. Sonié, and I. Chuine. 2010. Changes in leaf phenology of three European oak species in response to experimental climate change. New Phytologist 186:900–910.
- Morin, X., and W. Thuiller. 2009. Comparing niche-and process-based models to reduce prediction uncertainty in species range shifts under climate change. Ecology 90:1301–1313.
- Muir, W. D., W. S. Zaugg, A. E. Giorgi, and S. McCutcheon. 1994. Accelerating smolt development and downstream movement in yearling chinook salmon with advanced photoperiod and increased temperature. Aquaculture 123:387–399.
- Myking, T., and O. Heide. 1995. Dormancy release and chilling requirement of buds of latitudinal ecotypes of *Betula pendula* and *B. pubescens*. Tree physiology 15:697–704.

- Nienstaedt, H. 1966. Dormancy and dormancy release in white spruce. *Forest Science* 12:374–384.
- Ovaskainen, O., S. Skorokhodova, M. Yakovleva, A. Sukhov, A. Kutenkov, N. Kutenkova, A. Shcherbakov, E. Meyke, and M. del Mar Delgado. 2013. Community-level phenological response to climate change. *Proceedings of the National Academy of Sciences* 110:13434–13439.
- Parmesan, C. 2006. Ecological and evolutionary responses to recent climate change. *Annual Review of Ecology Evolution and Systematics* 37:637–669.
- Parmesan, C., and G. Yohe. 2003. A globally coherent fingerprint of climate change impacts across natural systems. *Nature* 421:37.
- Partanen, J., H. Hänninen, and R. Häkkinen. 2005. Bud burst in Norway spruce (*Picea abies*): preliminary evidence for age-specific rest patterns. *Trees* 19:66–72.
- Peñuelas, J., and M. Boada. 2003. A global change-induced biome shift in the Montseny mountains (NE Spain). *Global Change Biology* 9:131–140.
- Peñuelas, J., I. Filella, and P. Comas. 2002. Changed plant and animal life cycles from 1952 to 2000 in the mediterranean region. *Global Change Biology* 8:531–544.
- Polgar, C. A., R. B. Primack, E. H. Williams, S. Stichter, and C. Hitchcock. 2013. Climate effects on the flight period of lycaenid butterflies in Massachusetts. *Biological Conservation* 160:25–31.
- Poloczanska, E. S., C. J. Brown, W. J. Sydeman, W. Kiessling, D. S. Schoeman, P. J. Moore, K. Brander, J. F. Bruno, L. B. Buckley, M. T. Burrows, et al. 2013. Global imprint of climate change on marine life. *Nature Climate Change* 3:919.
- Richardson, A. D., T. F. Keenan, M. Migliavacca, Y. Ryu, O. Sonnentag, and M. Toomey. 2013. Climate change, phenology, and phenological control of vegetation feedbacks to the climate system. *Agricultural and Forest Meteorology* 169:156–173.
- Richardson, E. 1974. A model for estimating the completion of rest for 'redhaven' and 'elberta' peach trees. *HortScience* 9:331–332.
- Root, T. L., J. T. Price, K. R. Hall, S. H. Schneider, C. Rosenzweig, and J. A. Pounds. 2003. Fingerprints of global warming on wild animals and plants. *Nature* 421:57–60.
- Saikkonen, K., K. Taulavuori, T. Hyvönen, P. E. Gundel, C. E. Hamilton, I. Vännen, A. Nissinen, and M. Helander. 2012. Climate change-driven species' range shifts filtered by photoperiodism. *Nature Climate Change* 2:239.
- Sanz-Perez, V., P. Castro-Diez, and F. Valladares. 2009. Differential and interactive effects of temperature and photoperiod on budburst and carbon reserves in two co-occurring Mediterranean oaks. *Plant Biology* 11:142–51.
- Saunders, R. L., and E. B. Henderson. 1970. Influence of photoperiod on smolt development and growth of Atlantic salmon (*Salmo salar*). *Journal of the Fisheries Board of Canada* 27:1295–1311.
- Sidaway-Lee, K., E.-M. Josse, A. Brown, Y. Gan, K. J. Halliday, I. A. Graham, and S. Penfield. 2010. SPATULA links daytime temperature and plant growth rate. *Current Biology* 20:1493–1497.
- Singh, R. K., T. Svystun, B. AlDahmash, A. M. Jönsson, and R. P. Bhalerao. 2017. Photoperiod-and temperature-mediated control of phenology in trees—a molecular perspective. *New Phytologist* 213:511–524.
- Solbakken, V. A., T. Hansen, and S. O. Stefansson. 1994. Effects of photoperiod and temperature on growth and parr-smolt transformation in Atlantic salmon (*Salmo salar* L.) and subsequent performance in seawater. *Aquaculture* 121:13–27.

- Spann, T. M., J. G. Williamson, and R. L. Darnell. 2004. Photoperiod and temperature effects on growth and carbohydrate storage in southern highbush blueberry interspecific hybrid. *Journal of the American Society for Horticultural Science* 129:294–298.
- Tanino, K. K., L. Kalcsits, S. Silim, E. Kendall, and G. R. Gray. 2010. Temperature-driven plasticity in growth cessation and dormancy development in deciduous woody plants: a working hypothesis suggesting how molecular and cellular function is affected by temperature during dormancy induction. *Plant Molecular Biology* 73:49–65.
- Taranger, G., E. Vikingstad, U. Klenke, I. Mayer, S. Stefansson, B. Norberg, T. Hansen, Y. Zohar, and E. Andersson. 2003. Effects of photoperiod, temperature and GnRH α treatment on the reproductive physiology of Atlantic salmon (*Salmo salar* L.) broodstock. *Fish Physiology and Biochemistry* 28:403–406.
- Tobin, P. C., S. Nagarkatti, G. Loeb, and M. C. Saunders. 2008. Historical and projected interactions between climate change and insect voltinism in a multivoltine species. *Global Change Biology* 14:951–957.
- Van Belle, J., J. Shamoun-Baranes, E. Van Loon, and W. Bouten. 2007. An operational model predicting autumn bird migration intensities for flight safety. *Journal of Applied Ecology* 44:864–874.
- Viherä-Aarnio, A., R. Häkkinen, and O. Junntila. 2006. Critical night length for bud set and its variation in two photoperiodic ecotypes of *Betula pendula*. *Tree Physiology* 26:1013–1018.
- Vitasse, Y., and D. Basler. 2013. What role for photoperiod in the bud burst phenology of European beech. *European Journal of Forest Research* 132:1–8.
- Wareing, P. 1956. Photoperiodism in woody plants. *Annual Review of Plant Physiology* 7:191–214.
- Way, D. A., and R. A. Montgomery. 2015. Photoperiod constraints on tree phenology, performance and migration in a warming world. *Plant, Cell & Environment* 38:1725–1736.
- Weih, M. 2004. Intensive short rotation forestry in boreal climates: present and future perspectives. *Canadian Journal of Forest Research* 34:1369–1378.
- Willis, C. G., B. R. Ruhfel, R. B. Primack, A. J. Miller-Rushing, J. B. Losos, and C. C. Davis. 2010. Favorable climate change response explains non-native species' success in Thoreau's woods. *Plos One* 5:e8878.
- Wolkovich, E. M., B. I. Cook, J. M. Allen, T. M. Crimmins, J. L. Betancourt, S. E. Travers, S. Pau, J. Regetz, T. J. Davies, N. J. B. Kraft, T. R. Ault, K. Bolmgren, S. J. Mazer, G. J. McCabe, B. J. McGill, C. Parmesan, N. Salamin, M. D. Schwartz, and E. E. Cleland. 2012. Warming experiments underpredict plant phenological responses to climate change. *Nature* 485:494–497.
- Wolkovich, E. M., A. K. Ettinger, D. Flynn, T. Savas, C. Chamberlain, D. Buonaiuto, and J. Samaha. 2019. Observed Spring Phenology Responses in Experimental Environments (OSPREE). doi:10.5063/F1QV3JQR.
- Xie, L., and W. W. Hsieh. 1989. Predicting the return migration routes of the Fraser River sockeye salmon (*Oncorhynchus nerka*). *Canadian Journal of Fisheries and Aquatic Sciences* 46:1287–1292.
- Zhao, M., C. Peng, W. Xiang, X. Deng, D. Tian, X. Zhou, G. Yu, H. He, and Z. Zhao. 2013. Plant phenological modeling and its application in global climate change research: overview and future challenges. *Environmental Reviews* 21:1–14.
- Zhu, K., C. W. Woodall, and J. S. Clark. 2012. Failure to migrate: lack of tree range expansion in response to climate change. *Global Change Biology* 18:1042–1052.
- Zohner, C. M., B. M. Benito, J. C. Svenning, and S. S. Renner. 2016. Day length unlikely to constrain climate-driven shifts in leaf-out times of northern woody plants. *Nature Climate Change* 6:1120–1123.
- Zydlowski, G. B., D. S. Stich, and S. D. McCormick. 2014. Photoperiod control of downstream movements of Atlantic salmon *Salmo salar* smolts. *Journal of Fish Biology* 85:1023–1041.

Figures

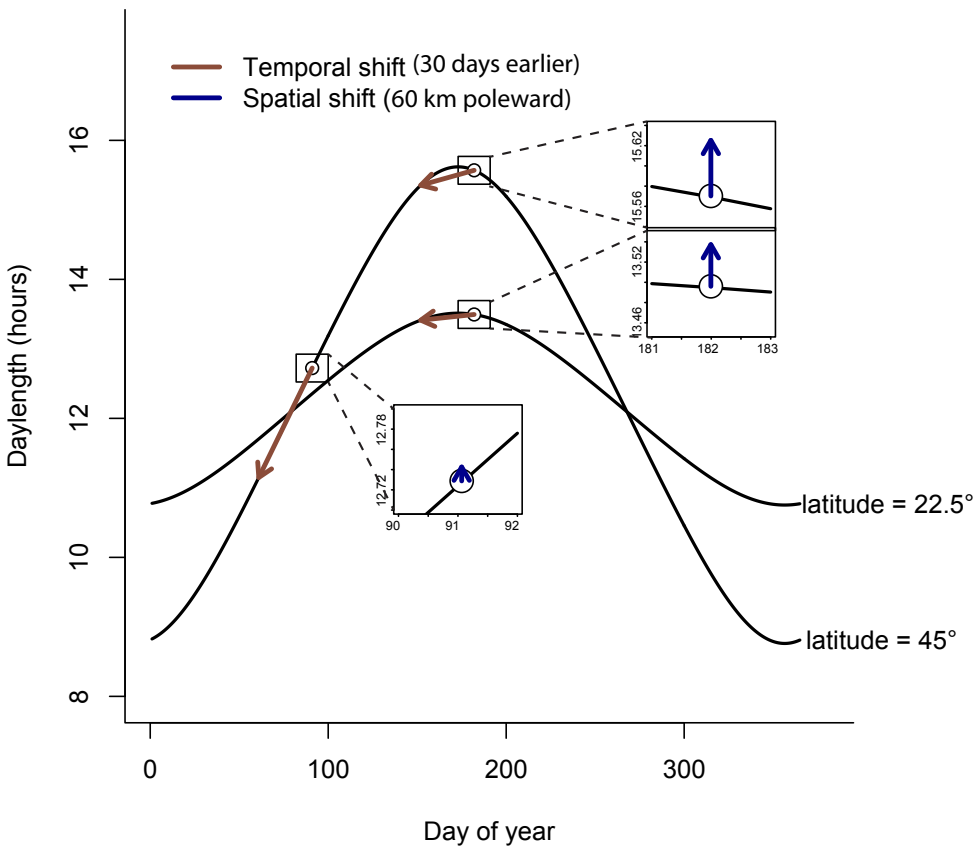


Figure 1: **Photoperiod varies with latitude and by day of year**, such that temporal shifts in activity yield larger changes in experienced photoperiod compared with spatial shifts. Here, we show this variation at two latitudes (22.5° , 45°), using hypothetical spatial and temporal shifts. These shifts, based on observed average rates with recent global warming—16.9 kilometers per decade, or approximately 1.5 degrees in 100 years, for spatial shifts (Parmesan, 2006), and 2.3 days per decade, or 23 days in 100 years, for temporal shifts (Chen et al., 2011)—highlight the greater magnitude in daylength changes close to the equinox (e.g., day of year 91), versus close to the summer solstice (e.g., day of year 182).

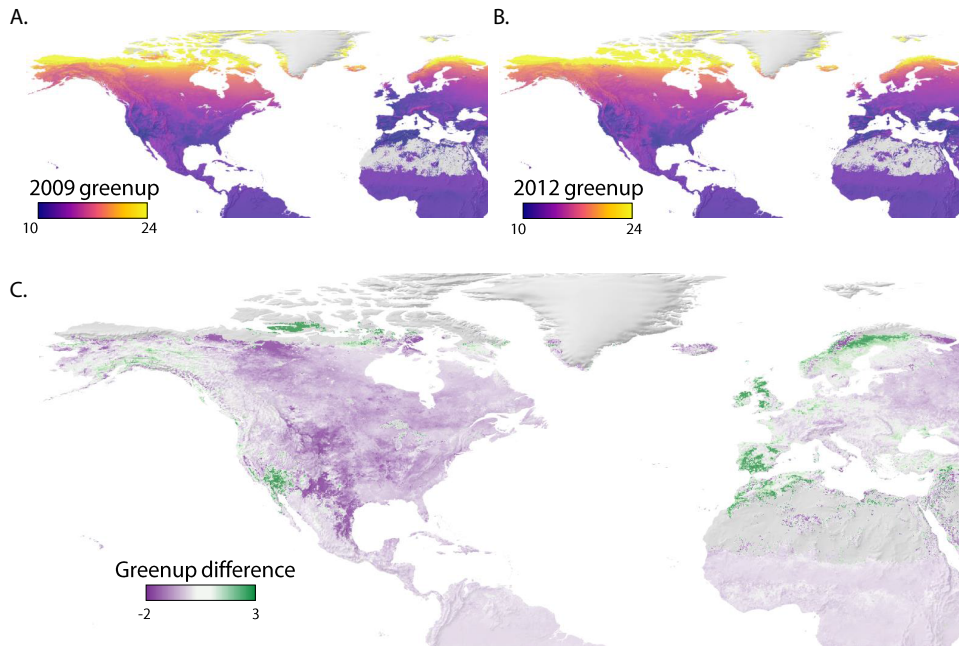


Figure 2: Photoperiod on the “green-up” date varies over space and between years “Green-up” is the beginning of seasonal greening, identified by satellite remote sensing measurements taken regularly throughout the year of the concentrations of green leaf vegetation. Hours of daylight on the date of spring green-up (here from MODIS satellite data) across North America and Europe for an average (2009, A) and early (2012, B) North American start of spring. The differences between the years (in hours of daylength) are shown in (C). A negative difference signifies earlier green-up in 2012 versus 2009; a positive difference is the result of later green-up in 2012 compared with 2009. See ‘Quantifying and mapping differences in green-up across the United States and Europe’ in the Supplemental Materials for more details.

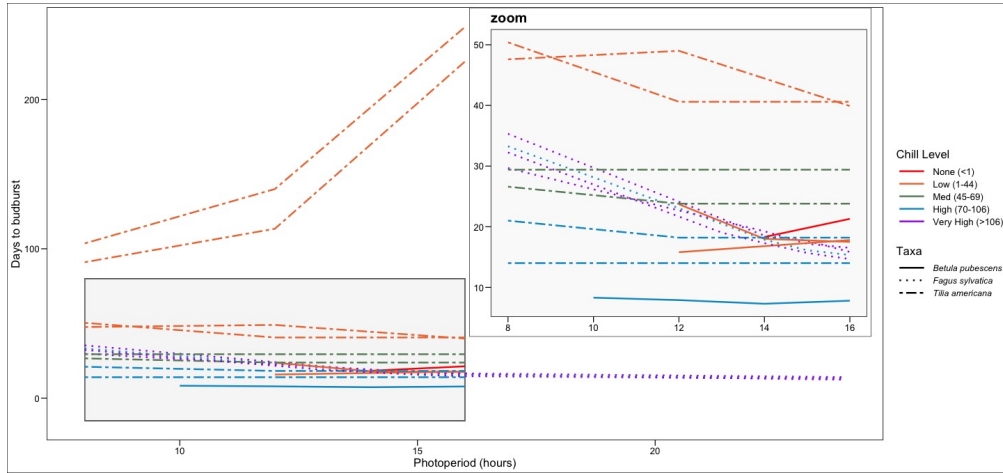


Figure 3: **Nonlinearities in phenological responses to daylength** are apparent in spring woody plant phenology experiments (from the OSPREE database) in which three or more photoperiod treatment levels were applied. The shape of the response curves for *Betula pubescens* (Caffarra et al., 2011b), *Fagus sylvatica* (Heide, 1993b) and *Tilia americana* (Ashby et al., 1962) differ depending on the amount of winter chilling received (measured in Chill portions). Species and chilling levels with multiple lines represent plant material from different populations.

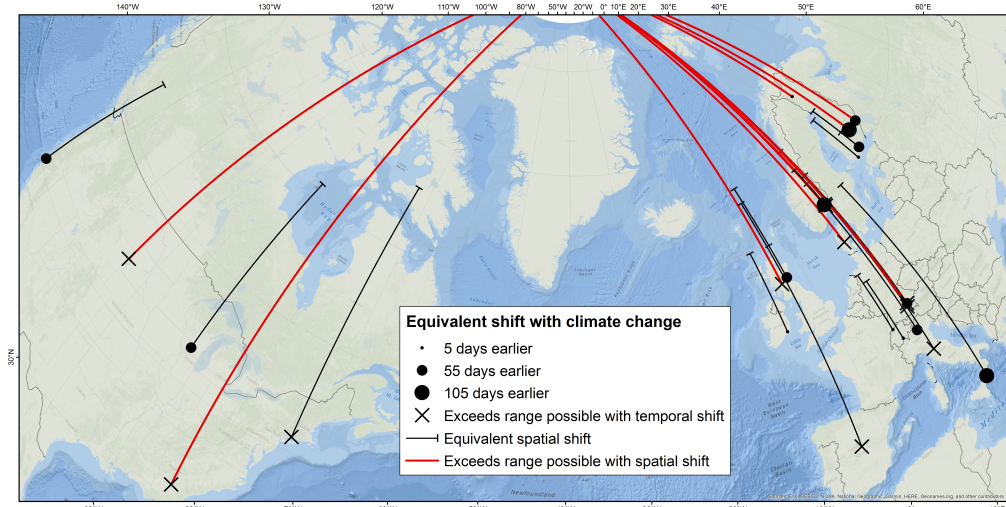


Figure 4: **Experimental photoperiod treatments and their equivalent spatial and temporal shifts** for experiments in the OSPREE database that manipulated photoperiod. See ‘Mapping temporal and spatial shifts in space and time’ in the Supplemental Materials for details on how we calculated the required spatial (lines) or temporal (circles and Xes) shifts to be equivalent to photoperiod treatments in each experiment.

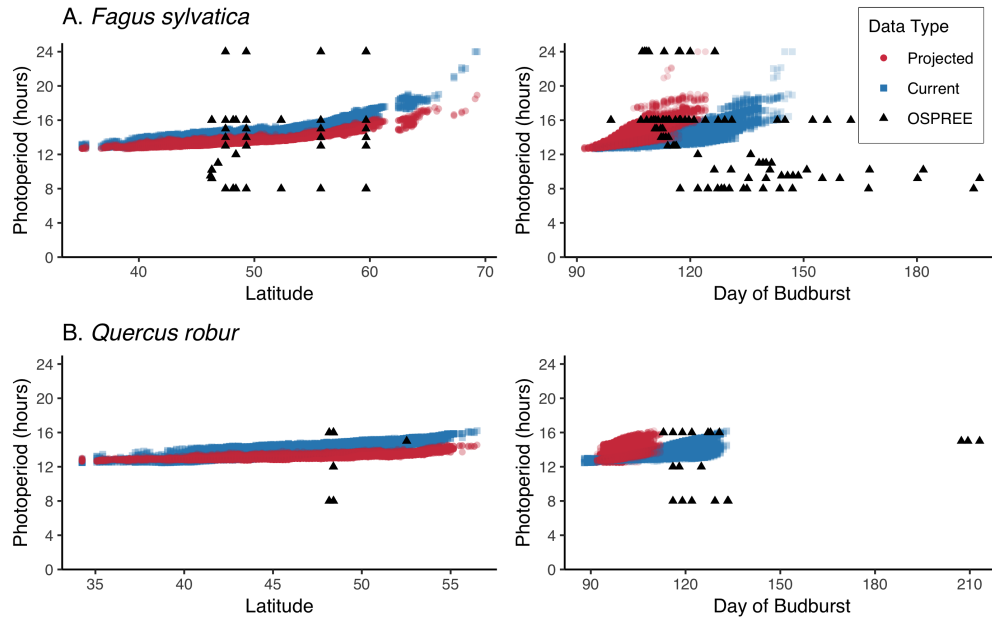


Figure 5: Experienced photoperiods in experiments differ from those in the natural world, shown here by latitude (left panels) and by day of budburst (right panels) for *Fagus sylvatica* (A, upper panels) and *Quercus robur* (B, lower panels). Triangles show experimental treatments of photoperiod in the OSPREE database. To illuminate potential gaps between experiments and the natural world, we show the photoperiod when budburst occurs in its current (1981-2000) and projected ranges (2081-2100, using the A1Fi Phenofit scenario, see Duputié et al., 2015). We scaled the days to budburst for all OSPREE data points by adding the day of budburst from the first Phenofit observation. See Supplemental Materials and Duputié et al. (2015) for additional details.

Accounting for photoperiod to improve forecasts of biological responses to climate change

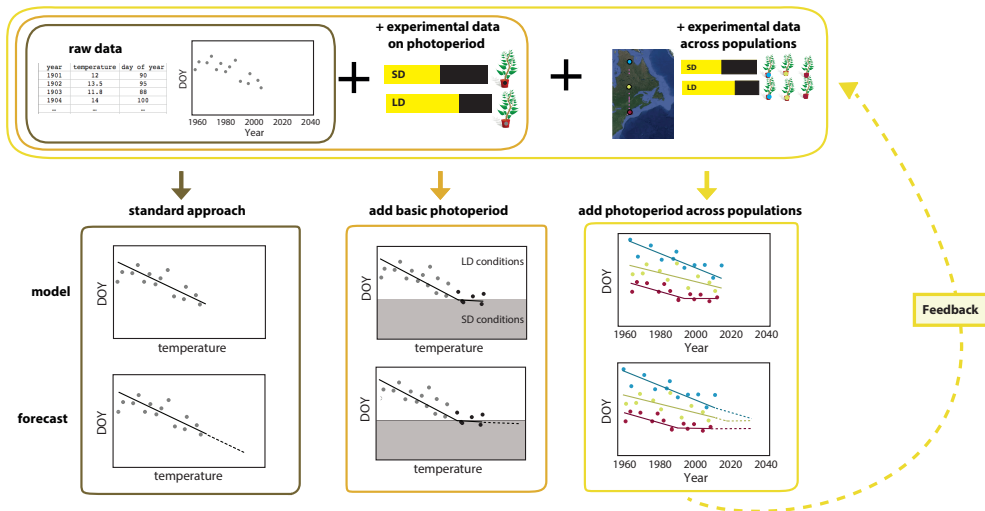


Figure 6: **Conceptual diagram of how to include photoperiod in forecasting biological responses to climate change.** Current approaches for forecasting spring phenology with climate change frequently rely on linear relationships between historical temperature data and observed dates of spring phenology (left panels). Adding responses to photoperiod, which commonly operate as threshold responses to short days (SD) versus long days (LD, see “photoperiod sensitivity” in the *Glossary*), will alter these forecasts (center panel) in ways that differ across species with divergent threshold photoperiods. Other factors that interact with photoperiod, such as population-level variation in photoperiod responses, can be incorporated into forecasts to further improve their accuracy (right panel).