Temperature and photoperiod drive spring phenology across all species in a temperate forest community

Authors:

D. F. B. Flynn 1,2† & E. M. Wolkovich 1,2†*

Author affiliations:

¹Arnold Arboretum of Harvard University, 1300 Centre Street, Boston, Massachusetts, USA;

²Organismic & Evolutionary Biology, Harvard University, 26 Oxford Street, Cambridge, Mas-

sachusetts, USA

*Corresponding author: 617.396.3890; lizzie@oeb.harvard.edu

[†]Authors contributed equally

Keywords: climate change, forcing temperatures, chilling, temporal niche, daylength, winter temperatures, forest communities

Paper type: Full paper

Counts: Total word count for the main body of the text: 4502; Introduction: 1064; M & M: 914;

Results: 715; Discussion: 1809; 3 figures (all in color). Supporting Information of notes, 4 supple-

mental tables and 5 supplemental figures.

Summary:

- (1) Accurate predictions of spring plant phenology with climate change are critical for projections of growing seasons, plant communities and a number of ecosystem services, including carbon storage. Progress towards prediction, however, has been slow because the major cues known to drive phenology—temperature (including intensity of winter chilling and spring forcing) and photoperiod—generally covary in nature and may interact, which would make accurate predictions of plant responses to climate change complex and non-linear. Alternatively, recent work suggests many species may be dominated by one cue, which would make predictions much simpler.
- (2) We manipulated all three cues across 28 woody species from two North American forests.
- (3) All species responded to all cues examined. Chilling exerted a strong effect, especially on budburst, with responses to forcing and photoperiod greatest for leafout. Interactions between chilling and forcing suggest each cue may compensate somewhat for the other. Cues varied across species, leading to staggered leafout within each community and supporting the idea that phenology may be a critical aspect of species' temporal niches.
- (4) Our results suggest that predicting the spring phenology of communities will be difficult, as all species we studied could have complex, non-linear responses to future warming.

Plant phenology—the timing of recurring life history events, such as leafout and flowering—is critical to the structure and function of ecosystems (Cleland *et al.*, 2007). Spring plant phenology in particular drives local ecosystem properties, from the length of the growing season to energy balance between land and atmosphere, and scales up to impact global carbon cycles (Richardson *et al.*, 2009).

Phenology is also one of the major biological indicators of climate change, with plant phenology shifting earlier across the globe 4-6 days/°C with warming (IPCC, 2014). While this average response is strikingly consistent when considered across diverse datasets (Wolkovich et al., 2012), it masks considerable variation. Variation is extreme when examined across species (Wolkovich et al., 2014), but additional variation can be seen within species over space (Kramer et al., 2017; Vitasse et al., 2013) and time (Fu et al., 2015; Yu et al., 2010). Understanding this variation has been the goal of much recent work (Donnelly et al., 2017; Laube et al., 2015; Rutishauser et al., 2008; Zohner et al., 2017), with research focusing on two major linked aims: (1) identifying and quantifying the environmental cues that drive spring phenology (i.e., budburst and leafout), and (2) identifying what drives variation in cues between different species.

Decades of study on wild species spring phenology—mainly focused on temperate woody species— 16 show that three major cues underlie budburst and leafout: warm spring temperatures (forcing), in-17 creasing daylength (photoperiod), and length and intensity of winter temperature (chilling). Across 18 studies, increasing temperatures in the spring appear to be a dominant factor that controls spring phenology; yet many of these studies have been observational—making it nearly impossible to tease 20 out the co-varying effects of longer days and reduced cold temperatures, which generally reduce 21 chilling (Chuine, 2000; Cook et al., 2012). By contrast, experiments from controlled environments 22 (e.g., growth chambers) have highlighted the additional importance of photoperiod and chilling (Caffarra et al., 2011; Falusi & Calamassi, 1996; Foley et al., 2009; Ghelardini et al., 2010; Heide, 1993a), with longer days and increased chilling leading to more rapid budburst (Caffarra & Donnelly, 2011). Many of these cues are known to interact: photoperiod and chilling can together determine spring phenology through their complex impacts on dormancy release (Chuine, 2000),

insufficient chilling may be offset by additional forcing, and photoperiod and chilling often interact,
as a long photoperiod enhances cell growth, compensating for a lack of chilling during plants' winter
dormancy (Caffarra *et al.*, 2011; Heide, 1993a; Myking & Heide, 1995).

Yet, while such complexities have been identified in some species, a growing body of hypotheses 31 and experimental studies has suggested many species are dominated by one cue and may lack any response to other cues (Körner & Basler, 2010). If true, this would have critical implications for predicting responses to climate change. Species dominated by a forcing cue would be predicted to continue to advance their leafout timing with warming, while species with strong photoperiod 35 cues would instead stop advancing at some threshold point (Körner & Basler, 2010). This could lead to major separation in the phenology of communities, as some species shift earlier while others change little, with cascading consequences for species coexistence and invasion. Alternatively, if all three cues—forcing, photoperiod, and chilling—are present and interact then predictions would be far more complex (Chuine & Cour, 1999). A species experiencing a mild winter with insufficient chilling (as predicted with climate change) could still break bud, but it would require longer photoperiods and/or warmer temperatures (Heide, 1993a) than it has in the historical record—a trend increasingly seen in long-term observational records (e.g., Carter et al., 2017; Fu et al., 2015). If such complex cues are seen in all species within a community it could mean community phenology may shift more in step, with no dramatic separation between species.

Research to date shows cues clearly vary across species, and recent efforts have focused on understanding and predicting this variation. Studies have focused on attributes of species: native/exotic (Willis et al., 2010), the successional stage (i.e., pioneer or climax communities) to which species traditionally belong (Basler & Körner, 2012; Laube et al., 2014), and a variety of possibly related traits (Lechowicz, 1984; Polgar et al., 2014). Most of these studies hinge on an often implicit assumption that phenology—by helping define the temporal niche of a species—is a critical axis along which plant species assemble within communities (Gotelli & Graves, 1996; Loreau & de Mazancourt, 2008). Support for this hypothesis comes from work showing that phenology is often staggered within communities, and from the special case of plant invasions, where

research suggests that climate change has provided open temporal niche space for new species to occupy (Willis et al., 2010; Wolkovich et al., 2013). As the abiotic environment is not the sole contributor to plant performance, considering a suite of co-occurring species together is key for making progress in understanding the role phenology plays in shifts in community composition and ecosystem functioning (Cleland et al., 2007).

Improved understanding and predictions of phenology with climate change would benefit from 60 a fuller understanding of the interacting environmental cues that drive phenology within (and 61 eventually across) communities. To this aim we studied how forcing, photoperiod and chilling 62 cues vary in their impact on spring phenology across a community of 28 woody plant species from two temperate forest locations (Tables S1), separated by 4° latitude. We used clipped dormant branches, which have been shown to approximate whole plant responses (Vitasse & Basler, 2014), and forced them in controlled environments that varied forcing temperatures, photoperiod and chilling. We predicted that: (1) Cues would vary across species, driving staggered leafout across the spring, and (2) within-species cues would trade off, such that some species would be dominated by one or another cue, while others would show a mix of cues. To our knowledge this is the first multi-species study to assess all three cues in one experiment through a controlled environment approach; while several studies have done this for one species (Skuterud et al., 1994; Søgaard et al., 71 2008; Sønsteby & Heide, 2014; Worrall & Mergen, 1967), other studies of all three cues have used separate experiments (e.g., Caffarra & Donnelly, 2011) or relied on field sampling to assess one or more cues (e.g., Basler & Körner, 2012; Laube et al., 2014; Zohner et al., 2016).

75 Materials & Methods

76 Field sampling

Woody plant cuttings were made in January 2015 for 28 species at Harvard Forest (HF, 42.5°N, 72.2°W) and the Station de Biologie des Laurentides in St-Hippolyte, Québec (SH, 45.9°N, 74.0°W).

The typical late January temperatures are -3.4 and -22°C, respectively, with daylengths (across the year) ranging from 9 to 15.25 hours and 8.5 to 15.75 hours. Weather station data from each field

site were obtained for calculations of chilling units (see Table S7).

Species were chosen based on the dominant forest vegetation at each site, aiming to maximize
the number of shared species between the two sites. Of the 28 species, at least 19 occurred at both
sites. For each species, up to 15 representative healthy, mature individuals with branches accessible
by pole pruners from the ground were tagged in late summer and fall 2014. In winter 2015, six
individuals were located and 4-16 cuttings taken from each individual, depending on size of the
individual and number of treatments to be applied. Cuttings were kept cold and transported back
to the Arnold Arboretum in Boston, MA.

89 Growth Chamber Study

Cuttings were placed in growth chambers at the Arnold Arboretum in Erlenmeyer flasks with distilled water; water was changed every 7-10 days. The bases of cuttings were re-cut at each water change under water to prevent callusing. For 11 of the 28 species, sufficient cuttings were obtained from each individual tree to apply the full set of 12 experimental treatments: 2 temperature (20°C / 10°C warm vs. 15°C / 5°C cool) x 2 photoperiod (12 vs. 8 h) x 3 chilling (no additional chilling, additional 30 d at 4°C, or 30 d at 1.5°C) treatments. For the remaining 17 species, sufficient cuttings were obtained only to apply the temperature and photoperiod treatments, without the additional chilling levels. The total number of cuttings for a given species thus ranged from 24 to 144, depending on presence at each site and application of the chilling treatment (Fig. S1). Lighting was a combination of halogen incandescent bulbs and T5HO fluorescent lamps with the lamploft adjusted to provide roughly 400 $\mu mol/m^2/s$ as measured by Apogee QSO-A5E quantum PAR light 100 sensors in each chamber (sensor set to the height of the cuttings). Treatments were rotated across 101 chambers every two weeks, as was flask position within chamber, to remove any possible bias of 102 chamber or flask position. 103

Phenology of the cuttings was assessed using a BBCH scale, modified for use in trees (Finn et al., 2007), with observations on each of the 2,136 cuttings made every 2-3 days for the course of the 82-day experiment, a total of 48 observation days. The phenological stages assessed in the

present study are budburst, defined as beginning of sprouting or bud breaking or shoot emergence (Code 07 in Finn *et al.*, 2007) and leafout, defined as first leaves unfolded (Code 11 in Finn *et al.*, 2007). Additional stages up to flowering and stem elongation were also recorded and we provide a photographic guide to help visualize stages across species (Savas *et al.*, 2017). In total, we made over 19,320 phenological observations at the cutting level.

112 Statistical analysis

We analyzed our data using Bayesian hierarchical models because they allowed us to best-estimate 113 responses to our full experimental design at both the species and across-species levels. In particular 114 this approach takes into account the effect, variance and sample size for each species. For the two 115 phenological responses measured, days to budburst and leafout, we fit mixed-effects hierarchical 116 models using site, warming, photoperiod, and chilling treatments, and all two-way interactions as 117 predictors (fixed effects) and species as modeled groups (random effects). This approach allowed 118 us to calculate the posterior probabilities of the effects for each of the abiotic drivers individually 119 and interactively across all species sampled. The models were fit using the programming languages 120 Stan (Carpenter et al., 2016) (www.mc-stan.org), accessed via the rstan package (version 2.15.1) 121 in R (R Development Core Team, 2017), version 3.3.3. Stan provides efficient MCMC sampling via a No-U-Turn Hamiltonian Monte Carlo approach (more details can be found in Gelman et al. 123 (2014) and in Carpenter et al. (2016)). 124

The model was fit as follows:

$$y_{i} \sim N(\alpha_{sp[i]} + \beta_{site_{sp[i]}} + \beta_{forcing_{sp[i]}} + \beta_{photoperiod_{sp[i]}} + \beta_{chilling1_{sp[i]}} + \beta_{chilling2_{sp[i]}}$$

$$+ \beta_{forcing \times photoperiod_{sp[i]}} + \beta_{forcing \times site_{sp[i]}} + \beta_{photoperiod \times site_{sp[i]}}$$

$$+ \beta_{forcing \times chilling1_{sp[i]}} + \beta_{forcing \times chilling2_{sp[i]}}$$

$$+ \beta_{photoperiod \times chilling1_{sp[i]}} + \beta_{photoperiod \times chilling2_{sp[i]}}$$

$$+ \beta_{site \times chilling1_{sp[i]}} + \beta_{site \times chilling2_{sp[i]}})$$

The α and each of the 14 β coefficients were modeled at the species level, as follows:

1.
$$\beta_{site_{sp}} \sim N(\mu_{site}, \sigma^2_{site})$$

...

14.
$$\beta_{site \times chilling2_{sn}} \sim N(\mu_{site \times chilling2}, \sigma^2_{site \times chilling2})$$

For the μ and σ parameters, weakly informative priors were chosen (increasing the priors three-fold did not change the model results). We validated our model code could return valid parameter values using test data.

We ran four chains simultaneously, with 4 000 warm-up iterations followed by 3 997 sampling iterations, resulting in 15 998 posterior samples for each parameter. We used a non-centered parameterization on all interaction terms and assessed good model performance through \hat{R} close to 131 1 and high n_{eff} (15 998 for most parameters, but as low as 2440 for several parameters) as well as 132 visual consideration of chain convergence and posteriors (Gelman *et al.*, 2014).

In our figures we show means \pm 50% credible intervals from this model. We used 50% intervals because of our focus here is on the most likely value for each parameter (e.g., estimated response to forcing) and because they are computationally stable (Carpenter *et al.*, 2016; Gelman *et al.*, 2014). For those interested in a more traditional significance-testing approach (which is not our focus here), we provide 95% credible intervals in Table S2-S3. To assess relationships between responses to forcing, chilling and photoperiod across species we fit a simple linear relationship to each of the 139 1000 last iterations and report the mean and 50% credible intervals from the resulting distribution.

Results

141 Budburst success

Across all treatments, 9.8% of the cuttings did not break bud, while an additional 10.4% did not reach the leafout stage following budburst (see *Budburst and leafout success* in the Supporting Information). Variation was highest due to species identity and ranged from complete budburst and

leafout (e.g., Hamamaelis) to only 65% budburst (Quercus alba) or 50% leafout (Faqus grandifolia, Acer saccharum) across all treatments (Table S2). The percent of non-leafouts by site was similar, with 20.6% of Harvard Forest and 19.7% of St. Hippolyte samples failing to leafout. Additional 147 chilling decreased budburst and leafout success (Tables S3-S4; Fig. S2-S3), with 30 d at 1.5°C having 148 the largest effects (cuttings with 1.5°C were 14.2% less likely to reach budburst and 23.4% less likely 149 to reach leafout). Other effects were weaker and varied by budburst versus leafout (Tables S3-S4; 150 Fig. S2-S3): for example, forcing temperatures and photoperiod affected leafout only (increased 151 forcing increased leafout by 3.9% while longer days increased leafout by 4.2% though the total effect 152 of increased forcing and longer days together was only 3.0% due to a negative interaction of the 153 two effects, see Figure S3). 154

155 Days to budburst and leafout

Higher forcing temperatures, longer photoperiod (12 vs. 8 h), and additional chilling all caused 156 large advances in budburst and leafout (Fig. 1, Tables S2-S3). Forcing temperatures (20°C / 10°C 157 warm vs. 15°C / 5°C cool) and chilling (no additional chilling, additional 30 d at 4°C, or 30 d at 158 1.5°C) caused the largest advances in budburst and leafout, and these two effects offset one another, 159 as shown by their interactive delayed response (Fig. 1). Effects of chilling at 4°C were greater than 160 forcing for all species (for which chilling was assessed) for budburst and for most species for leafout 161 (effects of forcing were greater than chilling for Acer saccharum, Populus grandidentata, Quercus 162 rubra, Viburnum lantanoides). Effects of forcing were greater than photoperiod for all species for budburst for all but Prunus pensylvanica for leafout. We found similar effects of chilling across two 164 different base temperatures, with only minor differences: responses to the colder (1.5°C) chilling 165 treatment were similar or more muted compared to responses to the warmer (4°C) chilling treatment 166 (Fig. 1, S5-S6, Tables S2-S3). 167

Responses to cues were qualitatively similar across both budburst and leafout, but quantitatively varied greatly between the two phenophases (Fig. 1-2), with responses generally greater for leafout (Fig. 1). This change was dramatic for the response to forcing and photoperiod where the advance in days more than doubled from budburst to leafout (Fig. 1, Table S3-S4).

These three factors (forcing, chilling and photoperiod) did show some degree of substitutability, 172 meaning, for example, that a lack of chilling could be made up for by an increase in forcing. These 173 are indicated by the positive two-way interactions: for example, while forcing or chilling at 4°C 174 alone would advance leafout (-19 or -17 days, respectively), their combined effect would lead to an 175 advance of only 26 days because of their interaction (10 days, see Table S5). Chilling and forcing 176 temperature are more substitutable than chilling and photoperiod, for both budburst and leafout, 177 while forcing and photoperiod showed virtually no substitutability for budburst and only a small 178 amount for photoperiod (Fig. 1). 170

The two forest sites showed similar responses, with only a very minor possible delay in overall timing for the northern site, and a more pronounced effect of site through its interaction with chilling (Fig. 1, Tables S5-6). The full effect of chilling depended on site, with a larger impact of site on the lower chilling temperature (1.5°C). Effects of forcing temperatures and photoperiod were not clearly impacted by site.

At the community level we found that all species were responsive to all cues (forcing tempera-185 tures, photoperiod and chilling, Fig. 3, S4-5), with each species having slightly different cues such that each species would budburst and leafout at a distinct time compared to other species (Fig. 187 3). Across species responses to forcing and photoperiod were related for budburst (mean slope of 188 0.31, CI of 0.15-0.48) and leafout (mean slope of 0.45, CI of 0.26-0.66), whereas responses between 189 forcing and chilling were very weak (budburst: mean slope of 0.12, CI of 0.04-0.20; leafout: mean slope of 0.11, CI of 0.04-0.20). Early species tended to show the smallest responses to all cues, 191 suggesting they bud burst and leaf out early because they require lower amounts of spring forcing, 192 longer days and chilling to start growth each season. In contrast, mid and late species relied on a 193 varying mix of cues to drive their spring phenology: for example, Populus grandidentata showed 194 a relatively strong response to forcing and chilling, but a milder response to photoperiod, while 195 Fagus grandifolia had a strong response to forcing and photoperiod and a much smaller response 196 to chilling. 197

98 Discussion

We found that all species responded to all three cues—spring forcing, winter chilling and photoperiod— 190 suggesting that future spring phenology with continued warming will most likely be complex and 200 non-linear. While some observational responses to date have suggested a potentially linear phe-201 nological response to warming (Ellwood et al., 2013) our results provide support for decades of 202 research that find spring phenology depends on a complex suite of multiple interactive cues (e.g., 203 Caffarra et al., 2011; Heide, 1993b). All three cues individually advanced budburst and leafout, 204 with the effects of chilling and forcing showing the largest interactive effect. The interactive ef-205 fects of forcing and chilling have been noted repeatedly before (e.g., Caffarra et al., 2011; Heide, 206 1993a) and highlight that insufficient chilling can be overcome by additional forcing—a hypothesis 207 suggested by recent studies that have found shifting temperature sensitivities in observational data 208 over time (Fu et al., 2015; Yu et al., 2010). Photoperiod and forcing also showed an interactive effect, but mainly for leafout and much smaller than either the singular effects of forcing or pho-210 toperiod (Figure 1). 211

212

213 Phenological cues: Multiple cues and interactive effects

In contrast to our expectations that within a species cues would trade off (i.e., a species could 214 be dominated by one cue), we found that species tended to show similar cues, especially between 215 forcing and photoperiod (Fig. 2a,c). Thus, a species with a strong response to forcing temperatures 216 generally also had a strong response to photoperiod and, similarly, a species with a comparatively 217 weak response to forcing also had a weaker response to photoperiod. This was also seen somewhat 218 with chilling (Fig. 2b,d), though we have fewer species with which to assess the relationship (see 219 Methods). There was substantial variation, however, such that if only a small subset of species 220 had been included in the study, it might have been concluded that a trade-off between photoperiod 221 sensitivity and warming sensitivity would exist. For example, Fagus grandifolia exhibited a relatively limited response to warming but substantial photoperiod sensitivity, while Rhamnus frangula showed a relatively limited response to photoperiod but substantial warming sensitivity.

These results also suggest that simple classification of some species as 'sensitive' or 'insensitive' 225 or similar bins (e.g., 'high', 'low', 'no' sensitivity) to any cue would be artificial, as species did not 226 form distinct clusters (Fig. 2). Previous studies have classified some of our studied species as non-227 or low-responsive to photoperiod (e.g., Alnus incana, Aronia melanocarpa in Zohner et al., 2016), 228 but we found these species were responsive to photoperiod and that any one species' cues were only 220 slightly different from the cues of several other species, yielding no clear way to define such binary 230 classifications. This is despite the fact that our species spanned a diversity of genera, including 231 canopy and understory species. We did find that shrubs tended to show smaller budburst responses 232 to photoperiod (Fig. 2a) than many trees, but this was not seen for leafout (Fig. 2c). 233

Our finding that all species responded strongly to all three cues is at odds with some recently 234 published work (Basler & Körner, 2012; Laube et al., 2014; Zohner et al., 2016), but is coherent 235 with many other studies (e.g., Heide, 1993a; Worrall & Mergen, 1967), with related process-based 236 models of woody plant phenology (Chuine et al., 2016, 2000) and with recent trends in long-term 237 data (e.g., Carter et al., 2017; Fu et al., 2015). The contrasting results may be due to varying methodologies and study aims. Our study used samples collected from the field in January—when 239 species in these locations had likely not fully met requirements for any cue—then used controlled 240 environments (growth chambers) to manipulate all three cues at once. In contrast, many other 241 studies have used multiple field sampling dates (i.e., sampling once every several weeks across the winter) to assess the effect of one cue, most often chilling (Laube et al., 2014; Weinberger, 1950; 243 Zohner et al., 2017), combined with controlled environments that manipulate the other cues (most 244 often forcing and photoperiod). This is done based on the assumption that chilling increases across 245 a winter season, yet forcing temperatures and photoperiod generally increase as well—meaning it 246 may be hard to assess fully any one cue using this method. Studies using this method may thus 247 underestimate the full suite of cues used to control spring phenology. 248

Study design plays an important role in all controlled environment phenology studies and can easily affect the findings and predictive utility of such studies in many ways. Studies that repetitively

sample throughout the winter may less accurately measure each cue, but have the advantage of 251 providing more realistic environmental conditions by capturing realistic shifts in all three cues across 252 the winter-spring season (Basler & Körner, 2012), and thus play an important role in predicting 253 near-term impacts of climate change. In contrast, many studies (e.g., Caffarra et al., 2011; Laube 254 et al., 2014) create more extreme changes in cues to assess better whether a cue is present. For 255 example, our study used a more extreme photoperiod difference between the two treatments, which 256 may better detect photoperiod responses. Our four-hour photoperiod difference corresponds to a 257 temporal change of 10-12 weeks in the spring at our two sites (see Supplemental Materials), which, 258 while extreme, is not an impossible change given projected warming and variation seen to date 250 (Stocker et al., 2013; Wolkovich et al., 2012). The drawback of this approach, however, is that 260 the design is much more artificial in its climate and, given the extreme treatments, may be less 261 relevant for near-term projections. Such designs may be more useful for longer-term predictions of 262 phenological responses with climate change and/or for use in parameterizing process-based models, 263 which often use a mix of results from observations and experiments. 264

Further, most phenological studies face limitations on how fully they can assess cues because of limited understanding of dormancy and its release preceding budburst (Chuine et al., 2016; Cooke 266 et al., 2012). Because dormancy release cannot be easily assessed (Chuine et al., 2016) most studies 267 to date using individuals sampled from the field do not fully know at what stage in endo- or eco-268 dormancy an individual is before the experiment or exactly how much chilling or forcing has been received. At our sampling date in January all individuals would have received some degree already 270 of all three cues, but still responded significantly to all treatments. In particular our finding that 271 all species responded to chilling suggests our sampled individuals must have all still been in en-272 dodormancy, since major responses to chilling are not expected after plants have moved from endo 273 to ecodormancy (Chuine et al., 2016). Our sampling date may also have affected our findings with regard to site effects. Chilling was the only factor to show noticeable differences due to site effects, 275 which could indicate that chilling requirements vary across populations due to local adaptation, or 276 it could be due to the field chilling experienced before we took cuttings for our experiments (see 277

Table S4). Additionally, our finding of no major difference in the two different chilling temperatures could indicate that plants cannot assess chilling temperatures below some threshold (Coville, 1920; Guy, 2014; Harrington et al., 2010) or that most met their chilling requirements at the higher 280 chilling treatment. If the latter is true, then studies that sample much earlier in the season should 281 find an elevated response to lower chilling temperatures. Finally, our limited knowledge of what 282 controls dormancy release also makes determining which temperatures are 'forcing' and which are 283 'chilling' difficult. Chilling is often assumed to happen below 5°C, but may occur also between 5-284 10°C (or higher) depending on the species and study (Harrington & Gould, 2015; Luedeling et al., 285 2013). This may make our low forcing treatment (15/5°C) a possible nighttime chilling treatment, 286 depending on the species and exact conditions. This design, however, had the benefit of holding 287 the diurnal temperature range—which has been suggested to alter budburst timing (Rossi & Isabel, 2017)—constant across treatments. 289

290

291 Phenological cues at the community scale

At the community level we found that each species had a unique suite of cues, leading to a generally staggered leafout (Fig. 3). This provides support for the idea that spring phenology is an important component of the temporal niche (Gotelli & Graves, 1996; Loreau & de Mazancourt, 2008). Species cues varied depending on the phenophase considered, meaning species' responses also shuffled between the two stages (Fig. 2). This fundamentally means that the species that bursts bud first will not necessarily leaf out first.

These quantitatively diverging findings for each phenophase suggest complex dynamics in the
early season within a community of woody plant species. Increasing evidence suggests the period
between budburst and leafout is when plants are at greatest risk of tissue loss from frost (Lenz
et al., 2013), and these new insights have come at the same time that research suggests risk of frost
damage may increase with climate change (Augspurger, 2009; Dai et al., 2013). For early-season
species in particular, this period may be critical to their current and future performance. Our results
suggest that the cues for each stage are not identical and support other work suggesting cues on

bud swelling and budburst may be distinct from the cues governing the development afterwards

(Basler & Körner, 2014). Understanding budburst is particularly difficult as it is the first observable

event after a series of unobservable (but see Rinne et al., 2011), yet important physiological events

required for budburst (Caffarra et al., 2011; Vitasse et al., 2014) and our results echo calls for

increased research in this topic (Chuine et al., 2016), which spans molecular, cellular and whole

plant areas of study (Morin et al., 2009; Rinne et al., 2011; Singh et al., 2017).

Our community-level findings may provide an additional way to build on our understanding 311 of what factors ultimately shape each species' mix of cues for budburst and leafout. Recent work 312 has critically begun to address this issue using by examining how attributes such native/invasive 313 status, climatic range, or climatic history predict cues (e.g., Laube et al., 2015; Zohner et al., 2017). 314 Building on these insights will require improved understanding of phenology's role in defining a 315 species niche and controlling its inclusion in a community. For example, assembly theory suggests 316 early-active species could out-compete later-active species through priority effects, which would 317 produce communities where all species leafout early. When this is not the case (as in our data 318 and many other systems) trade-offs may explain variation in phenology at the community-level (?). In temperate forests one dominant hypothesis for this trade-off is that early-active species 320 should also have traits that allow them to survive or avoid tissue loss to frost (?) while later-active 321 species would need traits that allow them to be competitive for resources when other species have 322 already had access to resource pools (e.g., soil nutrients or light). Testing these hypotheses requires matched trait and phenology data with a focus on careful measures of frost sensitivity (e.g., the 324 minimum temperatures tissues can experience without damage) and traits related to competition 325 (e.g., resource uptake metrics or growth rates under varied nutrient and competitive environment 326 regimes). 327

328 Conclusions

Across the two communities we studied, our results suggest species within a community have staggered budburst and leafout due to a mix of all three major environmental cues: forcing temperatures, photoperiod and chilling. In contrast to our hypothesis (and others', e.g., Körner & Basler,

2010), we found no evidence of any species being dominated by one or another cue; instead, species tended to show similar cues, especially between forcing and photoperiod cues. Thus, accurately predicting the phenology of any one of our studied species under diverse environmental conditions 334 would require considering how all three cues will change in concert. Shifting climate has already 335 clearly altered forcing and potentially chilling across the globe (IPCC, 2014; Stocker et al., 2013) 336 with trends expected to continue and possibly accelerate. In contrast, photoperiod has not, and 337 will not, shift; however plants that advance their leafout will experience large shifts in photope-338 riod. These trends combined with our results mean that all 28 species we studied could potentially 330 show complex, non-linear responses in the future, with cascading community and ecosystem con-340 sequences.

342 Acknowledgements

We thank T. Savas for technical and field assistance as well as countless observing hours, J. Samaha,
H. Eyster for help with field collections, E. Borjigin-Wang, J. Samaha, N. Farrant, and T. Chen
for recording chamber observations, the *Stanleyi* group for model help, and B. Cook, A. Ettinger,
S. Joly, and Y. Vitasse for comments that improved the manuscript.

347 Author Contributions:

DFBF and EMW conceived of the study design, performed analyses and wrote the paper. DFBF also carried out the experiment.

350 Data, Code & Model Output:

Raw data will be available via the Harvard Forest Data Archive upon publication and are available to all reviewers upon request. Stan model code and output provided upon request.

353 References

- Augspurger CK. 2009. Spring 2007 warmth and frost: phenology, damage and refoliation in a temperate deciduous forest. Functional Ecology, 23: 1031–1039.
- 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, Körner C. 2014. Photoperiod and temperature responses of bud swelling and bud burst in four temperate forest tree species. *Tree Physiology*, **34**: 377–388.
- Caffarra A, Donnelly A. 2011. The ecological significance of phenology in four different tree species: effects of light and temperature on bud burst. *Int J Biometeorol*, **55**: 711–21.
- Caffarra A, Donnelly A, Chuine I, Jones MB. 2011. Modelling the timing of Betula
 pubescens bud-burst. I. Temperature and photoperiod: A conceptual model. Climate Research,
 46: 147.
- Carpenter B, Gelman A, Hoffman M, Lee D, Goodrich B, Betancourt M, Brubaker

 MA, Guo J, Li P, Allen R. 2016. Stan: A probabilistic programming language. Journal of

 Statistical Software, (in press).
- Carter JM, Olive ME, Gerhart LM, Stern JH, Marchin RM, Nagel J, Ward JK. 2017.
- Warmest extreme year in U.S. history alters thermal requirements for tree phenology. *Oecologia*,
- **183**: 1197–1210.
- Chuine I. 2000. A unified model for budburst of trees. Journal of Theoretical Biology, 207: 337–347.
- Chuine I, Bonhomme M, Legave JM, de Cortazar-Atauri IG, Charrier G, Lacointe A

 Ameglio T. 2016. Can phenological models predict tree phenology accurately in the future?

 the unrevealed hurdle of endodormancy break. Global Change Biology, 22: 3444–3460.

- Chuine I, Cambon G, Comtois P. 2000. Scaling phenology from the local to the regional level: advances from species-specific phenological models. *Global Change Biology*, **6**: 943–952.
- Chuine I, Cour P. 1999. Climatic determinants of budburst seasonality in four temperate-zone tree species. New Phytologist, 143: 339–349.
- Cleland EE, Chuine I, Menzel A, Mooney HA, Schwartz MD. 2007. Shifting plant phenology in response to global change. Trends in Ecology & Evolution, 22: 357–365.
- Cook BI, Wolkovich EM, Parmesan C. 2012. Divergent responses to spring and winter
 warming drive community level flowering trends. Proceedings of the National Academy of Sciences
 of the United States of America, 109: 9000–9005.
- Cooke JEK, Eriksson ME, Junttila O. 2012. The dynamic nature of bud dormancy in trees:
 environmental control and molecular mechanisms. *Plant Cell and Environment*, **35**: 1707–1728.
- Coville FV. 1920. The influence of cold in stimulating the growth of plants. Proceedings of the

 National Academy of Sciences of the United States of America, 6: 434–435.
- Dai J, Wang H, Ge Q. 2013. The decreasing spring frost risks during the flowering period for woody plants in temperate area of eastern China over past 50 years. *Journal of Geographical* Sciences, 23: 641–652.
- Donnelly A, Yub R, Caffarra A, Hanesa J, Liang L, Desai AR, Liu L, Schwartz MD.
- 2017. Interspecific and interannual variation in the duration of spring phenophases in a northern mixed forest. Agricultural and Forest Meteorology, 243: 55–67.
- Ellwood ER, Temple SA, Primack RB, Bradley NL, Davis CC. 2013. Record-breaking early flowering in the eastern united states. *Plos One*, 8. Ellwood, Elizabeth R. Temple, Stanley A. Primack, Richard B. Bradley, Nina L. Davis, Charles C.
- Falusi M , Calamassi R. 1996. Geographic variation and bud dormancy in beech seedlings

 (Fagus sylvatica L). Annales des Sciences Forestières, 53: 967–979.

- Finn GA, Straszewski AE, Peterson V. 2007. A general growth stage key for describing
 trees and woody plants. Annals of Applied Biology, 151: 127–131.
- Foley ME, Anderson JV, Horvath DP. 2009. The effects of temperature, photoperiod, and
- vernalization on regrowth and flowering competence in Euphorbia esula (Euphorbiaceae) crown
- buds. Botany, 87: 986–992.
- 405 Fu YSH, Zhao HF, Piao SL, Peaucelle M, Peng SS, Zhou GY, Ciais P, Huang MT,
- Menzel A, Uelas JP et al. 2015. Declining global warming effects on the phenology of spring
- leaf unfolding. *Nature*, **526**: 104–107.
- Gelman A, Carlin JB, Stern HS, Dunson DB, Vehtari A, Rubin DB. 2014. Bayesian
- 409 Data Analysis. 3rd edn. CRC Press, New York.
- Ghelardini L, Santini A, Black-Samuelsson S, Myking T, Falusi M. 2010. Bud dormancy
- release in elm (*Ulmus spp.*) clones—a case study of photoperiod and temperature responses. *Tree*
- physiology, **30**: 264–274.
- 413 Gotelli NJ, Graves GR. 1996. The temporal niche. In: Null Models in Ecology (eds. Gotelli
- NJ. & Graves GR.). Smithsonian Institution, Washington, DC, pp. 95–111.
- 415 Guy RD. 2014. The early bud gets to warm. New Phytologist, 202: 7-9.
- 416 Harrington CA, Gould PJ. 2015. Tradeoffs between chilling and forcing in satisfying dormancy
- requirements for pacific northwest tree species. Frontiers in Plant Science, 6.
- Harrington CA, Gould PJ, St Clair JB. 2010. Modeling the effects of winter environment
- on dormancy release of Douglas-fir. Forest Ecology and Management, 259: 798–808.
- 420 Heide O. 1993a. Daylength and thermal time responses of budburst during dormancy release in
- some northern deciduous trees. *Physiologia Plantarum*, **88**: 531–540.
- 422 **Heide O. 1993b**. Dormancy release in beech buds (Fagus sylvatica) requires both chilling and
- long days. Physiologia Plantarum, 89: 187–191.

- 424 IPCC. 2014. Climate Change 2014: Impacts, Adaptation, and Vulnerability. Cambridge University
- Press, Cambridge, United Kingdom and New York, NY, USA.
- 426 Körner C, Basler D. 2010. Phenology under global warming. Science, 327: 1461–1462.
- 427 Kramer K, Ducousso A, Gomory D, Hansen JK, Ionita L, Liesebach M, Lorent A,
- Schueler S, Sulkowska M, de Vries S et al. 2017. Chilling and forcing requirements for
- foliage bud burst of European beech (Fagus sylvatica L.) differ between provenances and are
- phenotypically plastic. Agricultural and Forest Meteorology, 234: 172–181.
- Laube J, Sparks TH, Baessler C, Menzel A. 2015. Small differences in seasonal and thermal
- niches influence elevational limits of native and invasive balsams. Biological Conservation, 191:
- 433 682-691.
- 434 Laube J, Sparks TH, Estrella N, Hofler J, Ankerst DP, Menzel A. 2014. Chilling
- outweighs photoperiod in preventing precocious spring development. Global Change Biology, 20:
- 436 170–182.
- Lechowicz MJ. 1984. Why do temperate deciduous trees leaf out at different times? Adaptation
- and ecology of forest communities. The American Naturalist, 124: 821–842.
- 439 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:
- 441 1166-1175.
- 442 Loreau M, de Mazancourt C. 2008. Species synchrony and its drivers: Neutral and nonneutral
- community dynamics in fluctuating environments. American Naturalist, 172: E48–E66.
- Luedeling E, Guo L, Dai JH, Leslie C, Blanke MM. 2013. Differential responses of trees to
- temperature variation during the chilling and forcing phases. Agricultural and Forest Meteorology,
- 446 **181**: 33–42.

- 447 Morin X, Lechowicz MJ, Augspurger C, O'Keefe J, Viner D, Chuine I. 2009. Leaf
- phenology in 22 north american tree species during the 21st century. Global Change Biology, 15:
- 449 961–975.
- 450 Myking T, Heide O. 1995. Dormancy release and chilling requirement of buds of latitudinal
- ecotypes of Betula pendula and B. pubescens. Tree Physiology, 15: 697–704.
- Polgar C, Gallinat A, Primack RB. 2014. Drivers of leaf-out phenology and their implications
- for species invasions: insights from thoreau's concord. New phytologist, 202: 106–115.
- 454 R Development Core Team. 2017. R: A Language and Environment for Statistical Computing.
- R Foundation for Statistical Computing, Vienna, Austria.
- Richardson AD, Hollinger DY, Dail DB, Lee JT, Munger JW, O'keefe J. 2009. Influence
- of spring phenology on seasonal and annual carbon balance in two contrasting new england forests.
- 458 Tree Physiology, **29**: 321–331.
- Rinne PLH, Welling A, Vahala J, Ripel L, Ruonala R, Kangasjarvi J, van der Schoot
- 460 C. 2011. Chilling of dormant buds hyperinduces FLOWERING LOCUS T and recruits GA-
- Inducible 1,3-beta-Glucanases to reopen signal conduits and release dormancy in *Populus*. *Plant*
- 462 *Cell*, **23**: 130–146.
- Rossi S, Isabel N. 2017. Bud break responds more strongly to daytime than night-time tem-
- perature under asymmetric experimental warming. Global Change Biology, 23: 446–454.
- 465 Rutishauser T, Luterbacher J, Defila C, Frank D, Wanner H. 2008. Swiss spring plant
- phenology 2007: Extremes, a multi-century perspective, and changes in temperature sensitivity.
- 467 Geophysical Research Letters, 35.
- 468 Savas T, Flynn DFB, Wolkovich EM. 2017. A standardized photographic guide to woody
- plant spring phenology. doi:10.5063/F1M906MP.

- 470 Singh RK, Svystun T, AlDahmash B, Jonsson AM, Bhalerao RP. 2017. Photoperiod- and
- temperature-mediated control of phenology in trees a molecular perspective. New Phytologist,
- **213**: 511–524.
- Skuterud R, Dietrichson J et al. 1994. Budburst in detached birch shoots (betula pendula)
- of different varieties winter-stored in darkness at three different temperatures. Silva Fennica.
- Søgaard G, Johnsen Ø, Nilsen J, Junttila O. 2008. Climatic control of bud burst in young
- seedlings of nine provenances of norway spruce. Tree Physiology, 28: 311–320.
- Sønsteby A, Heide OM. 2014. Chilling requirements of contrasting black current (Ribes
- nigrum L.) cultivars and the induction of secondary bud dormancy. Scientia Horticulturae, 179:
- 479 256-265.
- Stocker T, Qin D, Platner G. 2013. Climate change 2013: The physical science basis. Working
- 481 Group I Contribution to the Fifth Assessment Report of the Intergovernmental Panel on Climate
- Change. Summary for Policymakers (IPCC, 2013).
- Vitasse Y, Basler D. 2014. Is the use of cuttings a good proxy to explore phenological responses
- of temperate forests in warming and photoperiod experiments? Tree Physiology, **34**: 174–183.
- Vitasse Y, Hoch G, Randin CF, Lenz A, Kollas C, Scheepens JF, Koerner C. 2013.
- Elevational adaptation and plasticity in seedling phenology of temperate deciduous tree species.
- 487 Oecologia, **171**: 663–678.
- Vitasse Y, Lenz A, Korner C. 2014. The interaction between freezing tolerance and phenology
- in temperate deciduous trees. Frontiers in Plant Science, 5.
- Weinberger JH. 1950. Chilling requirements of peach varieties. Proceedings of the American
- Society for Horticultural Science, **56**: 122–128.

- 492 Willis CG, Ruhfel BR, Primack RB, Miller-Rushing AJ, Losos JB, Davis CC. 2010.
- Favorable climate change response explains non-native species' success in Thoreau's woods. *PLoS*
- 494 *ONE*, **5**: e8878.
- 495 Wolkovich EM, Cook BI, Allen JM, Crimmins TM, Betancourt JL, Travers SE, Pau
- S, Regetz J, Davies TJ, Kraft NJ et al. 2012. Warming experiments underpredict plant
- phenological responses to climate change. *Nature*, **485**: 494–7.
- 498 Wolkovich EM, Cook BI, Davies TJ. 2014. Progress towards an interdisciplinary science of
- plant phenology: building predictions across space, time and species diversity. New Phytologist,
- **201**: 1156–1162.
- Wolkovich EM, Davies TJ, Schaefer H, Cleland EE, Cook BI, Travers SE, Willis CG,
- Davis C. 2013. Temperature-dependent shifts in phenology contribute to the success of exotic
- species with climate change. American Journal of Botany, 100: 1407–1421.
- Worrall J, Mergen F. 1967. Environmental and genetic control of dormancy in picea abies.
- Physiologia Plantarum, 20: 733–745.
- 506 Yu HY, Luedeling E, Xu JC. 2010. Winter and spring warming result in delayed spring
- phenology on the Tibetan Plateau. Proceedings of the National Academy of Sciences of the
- United States of America, **107**: 22151–22156.
- Zohner CM, Benito BM, Fridley JD, Svenning JC, Renner SS. 2017. Spring predictability
- explains different leaf-out strategies in the woody floras of north america, europe and east asia.
- 511 Ecology Letters, **20**: 452–460.
- ⁵¹² Zohner CM, Benito BM, Svenning JC, Renner SS. 2016. Day length unlikely to constrain
- climate-driven shifts in leaf-out times of northern woody plants. Nature Climate Change, 6:
- 514 1120-1123.

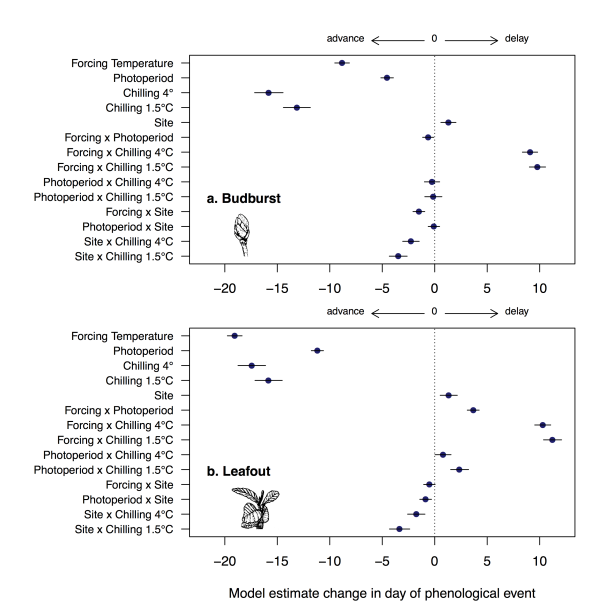


Figure 1: Effects of forcing temperature, photoperiod, chilling and site on budburst (a) and leafout (b) days across 28 species. Dots and bars show mean and 50% credible intervals from a Bayesian hierarchical model that also incorporated species-level variations (see Tables S2-S3; Figs. 1, S2-S3). Advances in phenology are shown by negative numbers; delays are shown as positive. Forcing temperatures and photoperiods were two levels each (see Methods), and chilling treatments were applied for 30 days. Budburst and leafout images from Finn et al. (2007).

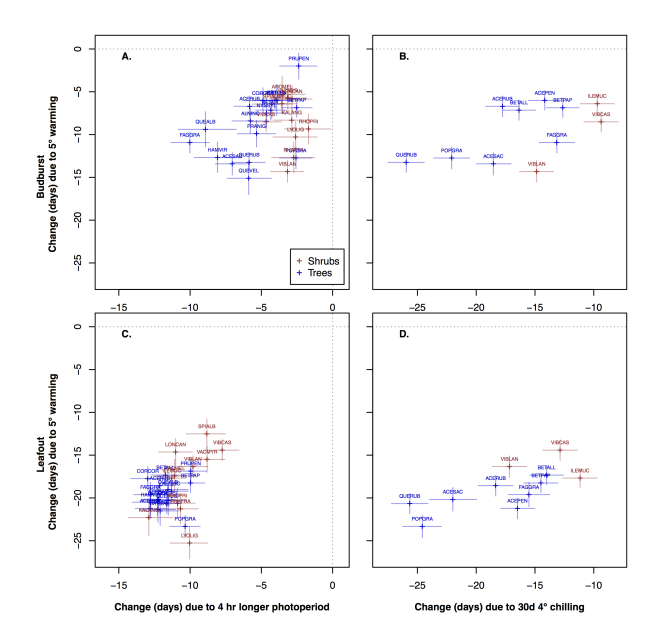


Figure 2: Effects of photoperiod, temperature and chilling across species (shrub species shown in red, tree species in blue): Crosses and bars show mean and 50% credible intervals from a Bayesian hierarchical model (see Tables S2-S3; Figs. 1, S2-S3). For visualization purposes, species names are represented by the first three letters of the genus and first three letters of the species epithet (see Table S1 for full species names and Fig. S5-S6 for additional versions of figure).

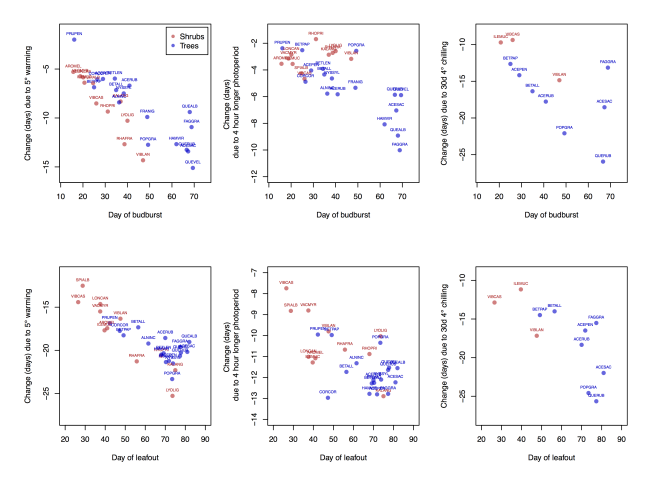


Figure 3: Effects of photoperiod, temperature and chilling across species compared to day of budburst (upper panels) or leafout (lower panels): we show mean estimates of sensitivity to warming, photoperiod, and chilling from a Bayesian hierarchical model (see Tables S2-S3; Figs. 1, S2-S3). For visualization purposes, species names are represented by the first three letters of the genus and first three letters of the species epithet (see Table S1 for full species names).