- 1 Timing of spring phenology and sensitivity to changing environmental drivers across a
- 2 community of temperate forest plants
- 3 Flynn, Wolkovich
- 4 The Arnold Arboretum of Harvard University
- 5 Accurate predictions of future spring plant phenology for with continued climate change are
- 6 critical for robust projections of future growing seasons, plant communities and a related
- 7 suite of critical ecosystem-level properties. Despite tremendous amounts of observational
- 8 data of plant phenology progress towards prediction has been hindered because the major
- 9 cues known to drive phenology—chilling temperatures in fall and winter, photoperiod, and
- spring forcing temperatures—generally co-vary in nature. Further, research to date using
- 11 controlled environments to separate these factors suggests that the cues are interactive,
- meaning accurate predictions of plant responses to climate change will be complex and
- 13 non-linear [1]. Recently, however, other research has suggested many species may be
- dominated by one of the three possible cues [2], with a tradeoff between photoperiod and
- 15 forcing temperature sensitivities, meaning some species' responses would be simple to
- 16 predict. To address this debate we present results of a full-factorial experiment manipulating
- 17 all three cues (spring forcing temperatures, photoperiod, and intensity of winter chilling)
- across 28 woody species and two latitudes (42.5°N and 46°N) in North American temperate
- 19 forests. In contrast to the predicted tradeoff between photoperiod and temperature cues we
- 20 find responses to these cues are largely coordinated across species; namely, species highly
- 21 sensitive to temperature were also highly sensitive to photoperiod. bud burst and leaf-out
- were more sensitive to temperature than to photoperiod.
- 23 Separated text for now: We need to report only some of the below once we narrow in on
- ₂₄ focus of the paper:
- 25 Winter chilling exerts a large role in driving advances in spring phenology, for both bud burst and leaf
- out stages, yet more intense chilling at 1.5°C resulted in less pronounced effects than at 4°C. Latitude
- of origin exerted surprisingly small effects on sensitivity to abiotic factors in driving spring phenology,
- indicating that local adaptation—at least across 4° of latitude—may not necessarily constrain woody

plant responses to climate change. Shrub and small tree species were less sensitive to changing 29 temperatures or photoperiod, but consistently earlier in their phenology. These results indicate that under warming conditions, communities could shift to a more canopy-tree dominated system with 31 generally later phenologies, counteracting advances in phenology at the ecosystem scale. Woody plant spring phenology 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 34 (add CITES). The crucial role that phenology plays in ecosystem processes, and the wealth of 35 observational data highlighting how rapidly plant and animal phenology are advancing [3] has led to increased interest in better understanding and prediction of how plant phenology will shift with 37 continued climate change. Decades of study on wild species spring phenology—mainly focused on temperate woody species—show that three major cues drive bud burst and leaf-out: spring temperatures (forcing). 40 length and intensity of winter temperature (chilling), and changing day length (photoperiod). Across 41 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 43 out the generally co-varying effects of longer days and reduced cold temperatures, which generally reduce chilling. In contrast studies from controlled environments (e.g., growth chambers or greenhouses) have highlighted the additional importance of photoperiod and chilling [4-8], with longer days and increased chilling leading to more rapid leaf-out (CITES). Further, photoperiod and chilling 47 often appear to interact, as long photoperiod enhances cell growth, compensating for a lack of chilling during plants' winter dormacy [4] [8] [9]. To an extent, all three factors—temperature, photoperiod, and chilling may be interchangeable in some species—such that a plant experiencing a mild winter with insufficient chilling can still break bud given sufficiently long photoperiods and warm temperatures [4]. A major challenge to build on this extensive study of how temperature and photoperiod drive spring 52 phenology is understanding—and possibly predicting—how the sensitivity to each cue and their 53 interactions varies across species. Temperate woody species are well known to have different sets of cues, depending on the species [2] (add CITES)—with some species showing stronger spring forcing or photoperiod cues, for example. A possible framework for understanding this variation in cues comes

from considering how adaptive pressures may drive temperate plant phenology (CITE - Wolkovich et

al. New Phyto 2014; and/or Wolkovich Cleland, Frontiers). Temperate woody plants should aim to maximize the carbon gain that comes from starting growth early in the season and gaining first access to critical resources such as light, soil nitrogen and water, while at once minimizing the risk of tissue loss to frost in early spring [2, 10, 11]. Across a community of co-occuring species we may expect a diversity of strategies to address these pressures—allowing each species to persist in the community. The ways that species manage such risks, and possible rewards, can vary in both when on average in 63 the spring they begin their growth, and also how flexible they are around that average timing. Related 64 to this, two tradeoffs have been described for spring phenology of woody plants in response to a variable spring environment: tolerance versus avoidance of freezing [12], and opportunistic versus conservative strategies [2]. The former describes how early or late in a season a species leafs out—with 67 tolerant species consistently leafing out early each year, while avoidance species would consistently leaf out late—while the latter describes how flexible a species or individual may be in response to unusually early warm-up events, with opportunistic species tending to have phenological cues that yield variable 70 leaf-out times across variable years and conservative species tending to leaf out consistently across 71 variable years. While related, these tradeoffs predict some differences in the characteristic of the cues and related traits of each species and may combine to produce contrasting changes in phenology. 73 Importantly, the combinations of these axes can result in non-intuitive responses at the community level. For example, under a relatively stationary long-term climate early tolerant but conservative species would enjoy the advantage of a long growing season. Yet, as that environment becomes nonstationary—as is the case with climate change—such advantages may quickly eroded. In contrast 77 under a warming environment, species that are relatively late in leafing out (avoidance) and flexible (opportunistic) close the gap in bud burst and leaf out times. Thus, understanding both the rank order 79 of phenology and the sensitivity of species to environmental cues across a whole community is 80 important in understanding how a changing climate will affect community dynamics. For the tolerance-avoidance axis, plants may either tolerate risk of spring freezing in two major ways: through phenological cues that allow them to leaf out well after all frost risk has passed (avoidance) or through investing in tissue which can withstand freezing (tolerant). For perennial plants, it has been found that leaf and wood tissues of species which have later phenologies can be more sensitive to frost damage [13, 14] (ADD CITE: Lechowicz 1984, AmNat), supporting the notion of a tradeoff between

tolerance and avoidance of freezing risk. Plants that leaf-out early but are tolerant would thus be expected to have a suite of tissue traits that allow them to withstand freezing: especially high tissue density, both of leaves and stems. In particular, wood density is positively related with leaf longevity 89 and resistance of xylem conduits to freezing-induced cavitation [15]. Thus high wood density would be advantageous for high-tolerance (early season) plant species. In contrast, avoidance-strategy plants would be expected to express lower tissue densities, with the shorter growing season being made up for by faster growth rates, less investment in structural elements of tissue, and relatively greater percent 93 nitrogen in leaves. 94 The axis of conservative versus opportunistic strategies makes specific predictions for the phenological 95 cues of differing species and may also predict the related traits of species. In opportunistic plant 96 strategies temperature is the dominant driver of spring phenology, while for species with conservative strategies photoperiod and chilling would be the major drivers. It has been found for several cases that short-lived, early successional species typically exhibit such opportunistic strategies, and 99 late-successional species are more typically chilling- and photoperiod-controlled in breaking of 100 dormancy [2] [16] [17]. Opportunistic species, showing greater ability to respond to early warm 101 temperatures in their timing of spring events, would be predicted to exhibit lower tissue densities, 102 greater leaf nitrogen concentrations (EXPLAIN a little MORE HERE). 103 Tests of how these two tradeoff axes drive phenology across a co-occurring community of species have 104 not previously been carried out. As the abiotic environment is not the sole contributor to plant 105 performance, considering a suite of co-occurring species together is key for making progress in 106 understanding the role phenology plays in shifts in community composition and ecosystem functioning. 107 To test the interactive effects of the three controlling drivers of spring phenology, temperature, 108 photoperiod, and chilling across latitudes, we carried out a study of 28 woody plants. We assessed 109 both bud burst and leaf-out to account for the potential different sensitives of these phenological 110 stages to abiotic drivers, and analyzed responses across all species to examine the support for how well 111 the tolerance-avoidance and opportunistic-conservative tradeoff axes represent temperature plant 112 spring phenology.

14 Results

Temperature and photoperiod individually and interactively determined timing of leaf-out, with the 115 strongest effects of temperature in short-day conditions. We found photoperiod sensitivity was 116 common and strong across all of the woody plants studies, consistently reducing time to phenological 117 responses for each species, across sites of origin. 118 For the 28 species studied, sensitivity to temperature and photoperiod cues for leaf-out times varied 119 substantially, and—in contrast to our hypotheses [that we set up in the intro]—co-varied overall. The coordinated response to warming temperatures and longer photoperiod was consistent with overall 121 pace of phenological events; earlier-leafing out species (namely the shrubs Spiraea alba, Viburnum 122 cassanoides, and Vaccinium myrtilloides) exhibited relatively limited advances to either warming or 123 longer days, while later leafing-out species showed ability to advance their phenology by in response to 124 both warming and longer days. Thus, no trade-off was observed between photoperiod-cued and 125 temperature-cued species, but rather species exhibit coordinated responses to both environmental 126 factors (Fig. 1). Of the other species, Fagus grandifolia exhibits relatively limited response to warming 127 but substantial photoperiod sensitivity, while Rhamnus frangula shows relatively limited response to 128 photoperiod but substantial warming sensitivity; if only a small subset of species including these two 129 had been included in the study, it might have been concluded that a tradeoff between photoperiod sensitivity and warming sensitivity would exist. 131 While both photoperiod and temperature cues were important for driving woody plant phenology, 132 responses to chilling were also substantial. bud burst day was accelerated most by the chilling 133 treatments. Tables 1 and 2 summarizes hierarchical mixed-effects model analysis of day of bud burst 134 and leaf-out, with negative values indicate earlier day of experiment for each event. Overall the 5°C 135 experimental warming resulted in 6.8 days earlier bud burst and 21.9 days earlier leaf out. Such 136 advance was delayed by the each chilling treatment, as indicated by the positive coefficient for the 137 temperature x chilling interactions. Latitude of origin (Site) overall had little direct effect on bud burst 138 or leaf-out, but populations from the northern site tended to exhibit slower bud burst and leaf-out, 139 with a more rapid bud burst and leaf out in response to the chilling treatments (indicated by negative 140 coefficients for site x chilling treatments).

Warming, photoperiod, and chilling individually and interactively acted to drive bud burst and leaf out
earlier across species. The strength of the acceleration in bud burst due to both warming and
photoperiod were similar, but the acceleration of leaf out due to warming exceeded that of
photoperiod for both phenological stages. Surprisingly, site of origin exerted limited effect on either
bud burst or leaf out across species.

147 Effect of chilling

The cuttings were harvested in late January 2015, and thus experienced substantial natural chilling by 148 the time they were harvested. Using weather station data from the Harvard Forest and St. Hippolyte 149 site, chilling hours (below 7.2°C), Utah Model chill portions (hours below 7.2°C and between 0°C and 150 7.2°C) and Dynamic Model [18] chill portions were calculated both for the natural chilling experienced 151 by harvest and the chilling experienced in the 4°C and 1.5°C treatments. The Utah Model and Dynamic Model of chill portions account for variation in the amount of chilling accumulated at 153 different temperatures, with the greatest chilling occurring approximately between 5-10°C, and fewer 154 chill portions accumulating at low temperatures and that higher temperatures can reduce accumulated 155 chilling effects. The two differ in the parameters used to determine the shape of the chilling 156 accumulation curve, with the Dynamic Model being shown to be the most successful in predicting 157 phenology for some woody species [19]. With both the Utah and Dynamic model, the more severe 158 chilling treatment resulted in fewer calculated chilling portions. Species varied widely in response to chilling treatments, with some exhibiting strong chilling 160

requirements (*Acer saccharum*, *Fagus grandifolia*), while others exhibited little change in phenological advancement under experimentally manipulated chilling. Overall, bud burst and leaf-out advanced by 22.1 or 26.4 days under additional 30 d of vernalization at 4°C, and advanced by a reduced amount of 19.7 or 26.1 days under 30 d of vernalization at 1.5°C. The reduced chilling effect at the lower temperature chilling is consistent with the Dynamic Model of chilling accumulation.

Species-specific responses to chilling demonstrate that chilling requirements are not uniform across
species, with of *Fagus grandifolia* to increasingly strong vernalization varies by latitude of origin and by
phenological stage; winter chilling reduced day to bud burst and leaf-out, but more strongly for

individuals from the northern site.

While nearly all species showed advances in spring phenology in response to the experimental chilling treatment, as indicated by fewer days to phenological events for the 4°C and 1.5°C treatments, the majority of species (e.g. *Populus grandidentata*) showed delays in both bud burst and leaf out at the more severe chilling treatment. Of the species exposed to the additional chilling, only *Fagus grandifolia* was consistently advanced by the more severe chilling.

75 Species-specific responses

Species traits partly explain variation in warming and photoperiod sensitivities of leaf out. Plants with
high nitrogen leaves, as well as high SLA (thinner, less dense) leaves, were significantly later in both
bud burst and leaf out. Thus early leaf out species tended to be tougher, less N-dense, and have
higher carbon investments than later species. Greater wood density had inconsistent effects as a driver,
with higher wood density driving later bud burst but tending to drive earlier leaf out.

Ring-porous species (*Fraxinus sp.*, *Lonicera*, *Myrica*, and *Quercus*; lower values of Pore Anatomy variable) exhibited significantly later bud burst and leaf out compared to diffuse-porous species, in line with previous work on wood anatomy and freezing risk [20].

Shrubs with low specific leaf area (thick/dense leaves) and high stem density were more likely to leaf out earlier. For trees, with an overall later leaf out pattern,

Rank order of leaf out and bud burst was stable across warming and photoperiod treatments. Chilling treatments shifted the order, for example *Fagus grandifolia* was the 23-28th species to burst bud with no additional chilling, but advanced to the 10-11th species to burst bud in with additional chilling.

Within chilling treatments, the consistency of the rank order was high, with standard deviation of the rank order ranging from 2.05 d (bud burst, no additional chilling) to 0.75 d (leaf out, additional chilling at 4°C). Compared to field observations, rank order of leaf out was generally most related in the cool, short-day treatment with no additional chilling (Fig. S10).

193 Nonleaf outs

Across all treatments, 20.2% of the cuttings did not break bud or leaf out. Across species, there was 194 no overall predictive effect of temperature, photoperiod, chilling, or site on the propensity to fail to 195 leaf out. Species ranged from complete leaf out (Hamamaelis) to only 50% leaf out (Fagus 196 grandifolia, Acer saccharum) across all treatments. The percent of nonleaf outs by site was similar, 197 with 20.6% of Harvard Forest and 19.7% of St. Hippolyte samples failing to leaf out. Examining 198 individual species, there was an interaction of temperature by day length for selected species, with 199 greater failure to leaf out in cool, short-day conditions for Acer pensylvanicum and Acer saccharum. 200 Site effects were inconsistent, with greater failure to leaf out for cuttings from St. Hippolyte in Acer 201 rubrum and Fagus grandifolia, and from Harvard Forest in Acer saccharum. 202

203 Discussion

Photoperiod sensitivity is common in northeastern woody plants, and greater photoperiod sensitivity is 204 related to, not instead of, temperature sensitivity. Taken together, this result shows that the 205 opportunism-conservatism tradeoff is not supported by the data for this suite of species. The most 206 sensitive species to both cues, namely the species which could advance their phenology in response to 207 both longer days and warmer temperatures, were the later-successional tree species, rather than the 208 shrubs. The trait data indicate partially that the species earliest to leaf out, namely the shrubs and 209 small trees, also had lower SLA and lower leaf %N, indicating greater investment in tissue structures. 210 These results support the tolerance-avoidance tradeoff, with the early phenology species being tolerant 211 to freezing but relatively less able to advance their phenology in a warming environment. These results 212 also indicate that the later-successional species have potentially the most to gain from a warming 213 world, as they can extend their growing seasons 214 While both photoperiod and temperature sensitives were common, chilling sensitivity greatly 215 outweighed both of these factors. It is important to note that the results from the chilling part of this 216 experiment are derived from 11, not 28 species, but the strength of this effect is notable. Strong 217 chilling requirements were detected both for bud burst and leaf out responses, and the most 218 substantial advance in spring phenology came from the more mild chilling treatment, at 4°C, with 219

220 reduced effectiveness of chilling at 1.5°C.

These three factors did show some degree of substituability, meaning for example that a lack of chilling could be made up for by an increase in temperature. These are indicated by the positive two-way interactions; chilling and forcing temperature are more substitutable than chilling and photoperiod, for both bud burst and leaf out.

We found only limited support for the northern populations showing more conservative

(photoperiod-cued) strategies in these 28 species was found, with small delays in both phenological

events for populations from the more northern site. The latitudinal range studied here is within the

range of the phenotypic flexibility of these species. Of these study species, we should not be overly

concerned about being photoperiod limited at the more northern sites; given sufficient pace of

dispersal, they will be able to track a changing climate.

bud burst is sensitive to the same environmental cues as leaf out, but species show idiosyncratic
orderings of their sensitivity to environmental cues at these two phenological stages; leaf out responses
can not necessarily be used to back-cast bud burst responses. bud burst showed a more limited total
response to environmental cues, and species were more tightly clustered in those responses.

Surprisingly, the smaller statured, earlier-leafing out shrubs and small trees exhibited reduced
sensitivity to all three factors of temperature, photoperiod, and chilling. They are relatively more fixed
in their timing of both bud burst and leaf out, perhaps indicating an alternative mechanism for timing
of spring phenology in these plants [21].

Given these results, the future of the northeastern forests may shift towards later-phenology, canopy
trees, as these species demonstrated a greater ability to lengthen their growing seasons
opportunistically in response to warmer temperatures.

Methods

Field sampling

Woody plant cuttings were made in January 2015 for 28 species which occurred in both Harvard Forest (HF, 42.5°N, 72.2°W) and the Station de Biologie des Laurentides in St-Hippolyte, Quebec

length between these two sites differs by a maximum of 45 minutes. Weather station data from each field site was obtained for calculations of chilling units. 248 Species were chosen based on the dominant forest vegetation at each site, aiming to maximize the 249 number of shared species between the two sites. Of the 28 species, 19 occurred at both sites. 250 Comparing only shared species, the mean days to bud burst and leaf out across all treatments for 251 Harvard Forest and St. Hippolyte was 25.6/36.8 and 24.8/36.1 days, respectively (Table S1). For each 252 species, up to 15 representative healthy, mature individuals with branches accessible by pole pruners 253 from the ground were tagged in late summer and fall 2014. In winter 2015, six individuals were located 254 and 4-16 cuttings taken from each individual, depending on size of the individual and number of 255 treatments to be applied. Cuttings were kept cold and transported back to the Arnold Arboretum in 256 Boston, MA. 257

Cuttings were placed in growth chambers at the Arnold Arboretum in Erlenmeyer flasks distilled water,

(SH, 45.9°N, 74.0°W). The typical late January temperatures are -3.4 and -22°C, respectively; day

258 Growth Chamber Study

246

259

with water changed every 7-10 days. The base of cuttings was re-cut at each water change under 260 water to prevent callusing. For 11 of the 28 species, sufficient cuttings were obtained from each 261 individual tree to apply the full set of 12 experimental treatments: 2 temperature (20°C / 10°C warm 262 vs. 15°C / 5°C cool) x 2 photoperiod (12 vs. 8 h) x 3 chilling (no additional chilling, additional 33 d at 263 4°C, or 33 d at 1.5°C) treatments. For the remaining 17 species, only sufficient cuttings were obtained 264 to apply the temperature and photoperiod treatments, without the additional chilling levels. The total 265 number of cuttings for a given species thus ranged from 24 to 144, depending on presence at each site 266 and application of the chilling treatment. 267 Phenology of the cuttings was assessed using a modified BBCH scale [22], with observations on each 268 of the 2,136 cuttings made every 2-3 days for the course of the 82-day experiment, a total of 48 269 observation days. The phenological stages assessed in the present study are bud burst, defined as 270 beginning of sprouting or bud breaking or shoot emergence (Code 07 in [22]) and leaf out, defined as 271 first leaves unfolded (Code 11 in [22]). Additional stages up to flowering and stem elongation were 272 also recorded. In total, we made 19,318 phenological observations at the cutting level.

Functional trait collection In summer 2015, the same individuals previously tagged in the field were 274 revisited as part of an additional study. Six individuals of each species were sampled for several plant functional traits, following standard protocols [23]. In some cases, the individual used in the growth 276 chamber study was missing, in poor condition, or had no remaining branches to sample, and was 277 replaced by a nearby representative individual. For each individual, height and diameter at breast 278 height (DBH) were recorded, and leaf and stem material were sampled from the middle of the canopy 279 or the greatest height reachable with pole pruners. Leaf material was kept cool and moist, and within 280 several hours was scanned for leaf area and weighed fresh. Stem volume was measured using a 281 water-displacement method. Samples were oven dried at 70°C and weighed within several days of 282 sampling, and specific leaf area (SLA) were calculated stem density. Leaf tissue was further processed 283 for carbon:nitrogen ratio using an elemental analyzer (Perkin-Elmer Elemental Analyzer) at Harvard 284 Forest. Since in not all cases the same individual used for the growth chamber experiments was the individual sampled for functional traits. 286

287 Statistical analysis

- For the two phenology responses measured, we fit mixed effect models separately for day of year, using site, warming, photoperiod, and chilling treatments as predictors and species as a modeled groups (random effects). For each model, two-way interactions for effects of site, warming, and each of the chilling treatments were included. Simplified versions of models were initially fit using the *lme4* package in the statistical programming environment R, then full versions of the model were fit using a Markov Chain Monte Carlo sampling approach in the programming language

 Stan [24](www.mc-stan.org).
- The model was fit using a Baeysian approach with weak priors with five main effects and each of their two-way interactions. The model was fit as follows:

$$y_{i} \sim N(\alpha_{sp[i]} + \beta_{site_{sp[i]}} + \beta_{temperature_{sp[i]}} + \beta_{photoperiod_{sp[i]}} + \beta_{chilling1_{sp[i]}} + \beta_{chilling2_{sp[i]}} + \beta_{temperature \times photoperiod_{sp[i]}} + \beta_{temperature \times site_{sp[i]}} + \beta_{photoperiod \times site_{sp[i]}} + \beta_{temperature \times chilling1_{sp[i]}} + \beta_{temperature \times chilling2_{sp[i]}} + \beta_{photoperiod \times chilling1_{sp[i]}} + \beta_{photoperiod \times chilling2_{sp[i]}} + \beta_{site \times chilling1_{sp[i]}} + \beta_{site \times chilling2_{sp[i]}})$$

Each of the 14 β coefficients was modeled at the species level, as follows

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

14. $\beta_{site \times chilling_{2sp}} \sim N(\mu_{site \times chilling_2}, \sigma^2_{site \times chilling_2})$

For the mu and sigma parameters, weakly informative priors were chosen.

298 Phylogenetic methods

We tested the influenced of phylogenetic relatedness on the relationship between functional traits and 299 sensitivities to warming, photoperiod, and chilling treatment. Sensitivities were extracted as the slopes 300 of the species-level responses of leaf out day to each of the experimental factors; more negative values 301 indicate greater advance in leaf out in response to that factor. Using a phylogenetic tree resolved at 302 the genus level from Phylomatic (www.phylodiversity.net), and the caper package in R, we fit 303 phylogenetic generalized linear models between the sensitivities at the species level to the functional 304 traits of stem density, SLA, and percent leaf nitrogen (%N). In this type of model, the parameter λ 305 represents the strength of the phylogenetic symbol, with values close to 1 indicating that closely 306 related species have more similar responses to the abiotic drivers.

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Figures and Tables

Table 1: Chill units in field and field and growth chamber conditions.

Site	Treatment	Chilling Hours	Utah Model	Chill portions
Harvard Forest	Field chilling	892	814.50	56.62
	$4.0~^{\circ}\text{C} \times 30~\text{d}$	2140	2062.50	94.06
	$1.5~^{\circ}\text{C} \times 30~\text{d}$	2140	1702.50	91.17
St. Hippolyte	Field chilling	682	599.50	44.63
	$4.0~^{\circ}\text{C} \times 30~\text{d}$	1930	1847.50	82.06
	$1.5~^{\circ}\text{C} \times 30~\text{d}$	1930	1487.50	79.18

Figure 1: Modeled effects plots, bud burst and leaf out

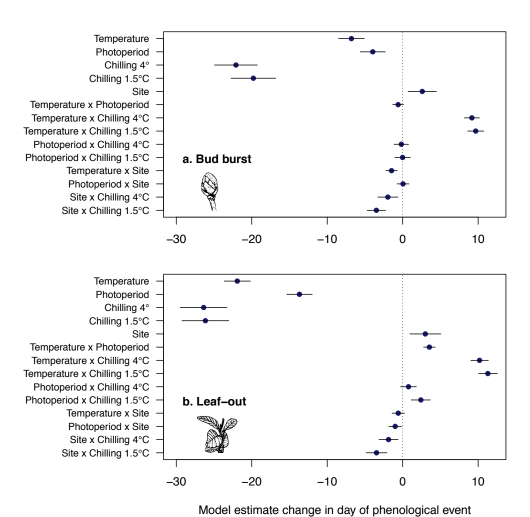


Figure 2: Sensitivity of bud burst and leaf out to warming, leaf out, and chilling.

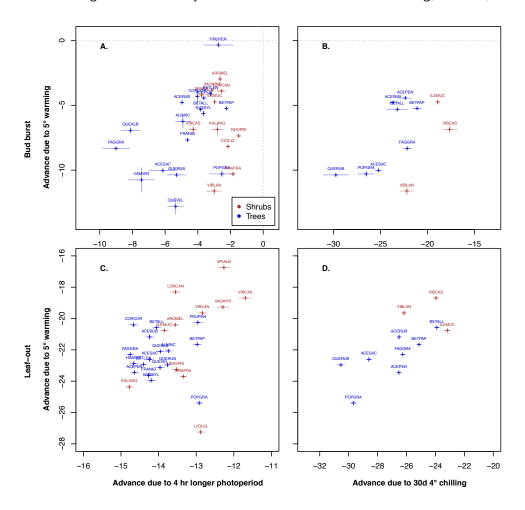


Table 2: Phylogenetic signal in timing of bud burst and leaf out and species specific traits, as estimated in the caper package with simultaneous fitting of lambda. Pore anatomy (ring- versus diffuse-porous species) was highly clustered phylogenetically, but no other trait examined demonstrated significant phylogenetic signal

Relationship	Lambda
SLA - Temperature	0.000
SLA - Photoperiod	0.000
SLA - Chilling 4 °C	0.000
SLA - Chilling 1.5 °C	0.000
Wood Density - Temperature	0.000
Wood Density - Photoperiod	0.000
Wood Density - Chilling 4 °C	0.000
Wood Density - Chilling 1.5 °C	0.000
% N - Temperature	0.285
% N - Photoperiod	0.203
% N - Chilling 4 °C	0.127
% N - Chilling 1.5 °C	0.130
Pore anatomy - Temperature	1.000
Pore anatomy - Photoperiod	1.000
Pore anatomy - Chilling 4 °C	1.000
Pore anatomy - Chilling $1.5~^\circ\text{C}$	1.000