- Differences in flower and leaf bud responses to the environment
- drive shifts in spring phenological sequences of temperate woody

3 plants

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# 5 Introduction

6 One of the most widely documented biological effects of anthropogenic climate change are shifts in

phenology, the timing of life cycle events, in plants (Parmesan & Yohe, 2003; Menzel et al., 2006;

8 Cleland et al., 2007). While phenology is generally advancing with climate change, the strength

of these phenological shifts can vary substantially among specific phenological phases (Augspurger

½ Zaya, 2020). These differences alter the timing of phases relative to each other, changing the

the duration of inter-phase periods that make up phenological sequences (Ettinger et al., 2018).

12 Phenological sequences are a major driver of plant fitness that impact plant life history, resource al-

location, demography and ecosystem processes (Post et al., 2008). Shifts in phenological sequences

will likely alter many of these processes, but the effects these shifts depend both on the direction

15 (whether distinct phases are shifting closer together or farther apart) and magnitude (how much

they are shifting relative to each other).

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<sup>18</sup> Among deciduous woody plants, the relative timing of flower and leaf phenology, or flower-leaf se-

19 quences (FLSs), may be particularly consequential to fitness in temperate regions where flowering

prior to leaf development is common (Rathcke & Lacey, 1985; Gougherty & Gougherty, 2018).

Long-term phenological observations over the last several decades indicate that, like other phe-

nological sequences, FLSs are shifting due to anthropogenic climate change (Buonaiuto *et al.*, in review)—for several species, the time between flowering and leafing appears to be increasing, but the strength of this trend varies among species and the direction of FLS shifts are not consistent across populations. These changes could affect the important functions of FLSs, which may put some species at greater risk while benefiting others.

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Flowering before leaf development may be a critical adaptation for pollination efficiency in windpollinated taxa by eliminating pollen interception by the forest canopy (Whitehead, 1969). In
insect-pollinated taxa, flowering-first may increase the visibility of flowers to pollinators (Janzen,
1967; Savage, 2019). Species with decreasing FLS interphases with climate change may experience
increased pollen limitation as more wind pollen is intercepted by vegetative structures and flowers
are obscured by developing leaves. Conversely, pollination efficiency could improve for species with
lengthening FLS interphases (direction). A change in the FLS interphase of just a few days would
likely have little impact on these processes, but if shifts were on the order of weeks, the impact on
the pollination biology of a species could be highly significant (magnitude).

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Predicting the direction and magnitude of any FLS shifts requires identifying the underlying mechanisms that drive the different responses to climate change among these phenophases for a diversity of woody plant species. Decades of research suggests that for woody plants in temperate regions, cool winter temperatures (chilling), warm spring temperatures (forcing) and day-length (photoperiod) are the primary drivers of both reproductive and vegetative phenology (Forrest & Miller-Rushing, 2010; Flynn & Wolkovich, 2018). However, observed FLS shifts indicate that there must be differences in how these cues influence phenological activity in floral and leaf buds. Identifying these differences is a necessary step for predicting the direction and magnitude, and ultimately fitness impacts of FLS shifts with climate change.

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Studies that have attempted to identify the differences between reproductive and vegetative phe-

49 nology in woody plants have mostly focused on crop species and two common, yet competing,

50 explanations have emerged:

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One hypothesis suggests that reproductive and vegetative buds utilize the same underlying en-

vironmental cues, but have different threshold responses to forcing, with whichever bud type

bursts later—leaves or flowers—having a higher threshold (Guo et al., 2014; COSMULESCU &

55 CALUSARU, 2020; Cosmulescu & Ionescu, 2018). Under this hypothesis, which we call the pre-

cocity hierarchy hypothesis (PHH), leaf and flower buds share the same suite of cues and develop

57 similarly to non-forcing cues but they differ in the thermal units required for budburst.

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By contrast, an alternative hypothesis suggests that flower and leaf buds differ in the strength

of their phenological responses to the multiple environmental cues (Citadin et al., 2001; Gariglio

et al., 2006; Aslamarz et al., 2009; Mehlenbacher, 1991). Under this hypothesis, which we call the

differential sensitivity hypothesis (DSH), despite the fact that leaf and flower buds are exposed to

63 similar environmental conditions, each bud type may rely more or less on certain cues, producing

64 different and variable phenological patterns.

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While these mechanisms may produce similar phenological patterns under historic climate condi-

67 tions, they have different implications regarding the potential for FLS shifts with climate change.

The PHH suggests that FLS variation is largely a product of climate variation during the inter-

69 phase. If spring temperatures increase with climate change, the second phenophase of the FLS

with be accelerated relative to the first and the FLS interphases will decrease, but given the rel-

ative auto-correlation of spring temperatures (Di Cecco & Gouhier, 2018), these shifts should be

relatively muted.

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The DSH suggests that with significant cue use differences among bud types, there will be strongly

b localized effects of climate change on FLSs. Shifts in FLS variation will depend on the direction

and rate of change in cues at specific locations and the differential sensitivity of reproductive and vegetative phenology to cue combinations. This hypothesis allows not only for larger magnitude shift in FLSs, it also suggest that the magnitude of shifts may be highly divergent among populations of the same species.

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In this study we test these hypotheses by observing phenological responses to changing environmental conditions for both flower and leaf buds for a suite of temperate shrubs and trees. We subjected dormant twig cuttings of 10 species to multiple levels of forcing, chilling and photoperiod treatments in growth chambers and compared flower and leaf phenological responses to environmental change using a Bayesian hierarchical modeling approach. We then leveraged these data to to make generalized projections for how FLSs may shift with climate change and identify avenues for further research.

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### 89 Methods

# 90 Growth chamber study

We sampled all plant material used in this experiment from Harvard Forest in Petersham, MA.

On October 25, 2016, immediately after most plants in the area entered dormancy but before they

could accumulate any significant chilling in the field, we collected branch cuttings from 7-13 indi-

viduals of 12 woody plant species (4-12 cutting per individual for a total of 48-56 per species). The

95 species consisted of a mix of deciduous shrubs, understory and canopy trees commonly found in

mesic hardwood forests of the eastern United States (see tab. ?? for species list). We transported

all cuttings to the Arnold Arboretum in Boston, MA where they were re-cut in water to prevent

callousing and cavitation and placed in 500 ml Erlenmeyer flasks with distilled water.

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We randomly assigned cuttings to a full set of eight experimental treatments; two levels of chilling (4

vs 8 weeks at 4°C), two levels of temperature (24°C:18°C (day/night) warm vs 18°:12°C (day/night) 101 cool) and two levels of photoperiod (12 vs 8 hours). We alternated day/night temperature peri-102 odicity on a 12 hour schedule to reduce co-variation with photo-periodicity. We re-cut all twig 103 and changed the water every 7-10 days and rotated all treatments between growth chambers every 104 two weeks to minimize chamber effects. We made phenological observations every 2-3 days using a 105 modified BBCH scale for woody plants (Finn et al., 2007) for three month following release from 106 chilling conditions. In this period we assess three phenological phases: budbreak (BBCH phase 07), 107 leaf unfolding (BBCH phase 15) and first flower open (BBCH 60). At the conclusion of this period 108 we assessed all individuals that did not undergo budbreak and excluded any dead individuals for 109 analysis. 110

# Data analysis

To assess the sensitivity of each phase, we fit mixed-effect hierarchical models with chilling, forcing, photoperiod and all two-way interactions as the fixed effects and species as a grouping factor on both the slopes and the intercepts. We chose a Bayesian, hierarchical approach in order to identify systematic trends across species' responses while accounting for sample size, variance and the unique effect of each species. Two species *Betula allegheniensis* and *Acer saccharum* produced no flowers in our trial, so we excluded them from our analysis.

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We modeled the effects of environmental parameters on flower opening and leaf budburst separately. We also fit a model with FLS interphase (day of budburst- day of flowering) as a response variable to compare these estimates with field observations.

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The models we fit appear below:

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$$y_{[i]} \sim N(\alpha_{sp_{[i]}} + \beta_{forcing_{sp_{[i]}}} + \beta_{chilling_{sp_{[i]}}} + \beta_{photoperiod_{sp_{[i]}}} + \beta_{forcingxchilling_{sp_{[i]}}} + \beta_{forcingxphotoperiod_{sp_{[i]}}} + \beta_{forcingxphoto$$

127

Where  $y_{[i]}$  is either the day of the experiment leaf budburst, day of first flower opening or FLS interphase length. We modeled the  $\alpha$  and each  $\beta$  parameter at the species level using the formula:

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$$\alpha_{x_{sp}}$$
 or  $\beta_{x_{sp}} \sim N(\mu_x, \sigma_x^2)$ 

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We fit all models using the R package "brms" (Bürkner, 2018). We ran each model on four chains with 4000 iterations with a 3000 iteration warm up for a total of 4000 sampling iterations. In both models we used weakly informative priors and increasing the priors 5-fold did not affect the model results.

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#### 38 Climate change predictions

To apply our model results to general climate change projections we chose our environmental treat-139 ments in this experiment to broadly reflect historic and future conditions at our sampling site. Our 140 low forcing treatment approximated average spring temperature (March/April) at the site while 141 our high temperature treatment reflects a 5 °C increase. Average field chilling (calculated from 15 Oct - 15 April, measured in Utah units) at Harvard Forest is 979.64, approximately 60% of the 143 difference between our low and high chilling treatment (Fig. ??). Thus, our low chilling treatment 144 represents a feasible estimate for a decrease in chilling with climate change and our high chilling 145 treatment approximate reasonable increase. We should note that our low photoperiod treatment (8 hours of daylight) is well below the photoperiod experienced at Harvard Forest, but given that 147 the photoperiod effects are expected to be small, we chose more extreme values in order to robustly 148 estimate an effect (i.e., increasing statistical power). For this reason, our climate change projections 149 for FLS variation are based on our high photoperiod treatment alone. 150

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We used our flower and budburst models to project for each species in our study:

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- 1. FLSs under average environmental conditions (treatments: low forcing, 6.5 weeks of chilling treatment)
- 2. FLS shifts with spring warming only (high forcing, 6.5 weeks of chilling treatment)
- 3. FLS shifts with warming and increased chilling ((high forcing, 8 weeks of chilling treatment)
- 4. FLS shifts with warming and decreased chilling ((high forcing, 4 weeks of chilling treatment)

To validate our predictions, we compared our FLS interphase model estimates of "average" condition FLS interphases to long term phenological records from Harvard Forest (O'Keefe, 2015) for five species common to both datasets (Fig. ??), and found them to be comparable. Given the variable dynamics of shifts in environmental forcing and chilling with climate change over time and space, these projections should not be treated as absolute predictions of the magnitude of FLS shifts with climate change. Instead, we provide these projections to identify general trends in how FLSs could shift with warming and demonstrate the range of possibilities vary based on individual characteristics of plant species and the specific climate dynamics.

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# 168 Results

#### 169 Growth chamber study

Both flower and leaf buds advanced with higher forcing and longer chilling duration (flowers: 21 day advance/ $\delta$ chillingduration, 18dayadvance/ $\delta$ forcingtemperature, leaves: 30dayadvance/ $\delta$ chilling, 17dayadvance

While leaf and flower bud phenological responses to environmental cues were qualitatively similar, the strength of their responses to each cue differed substantially. Leaf buds responded more strongly to chilling than flower buds (1.4x), and had a stronger response to all cue interactions (forcing:chilling 2x, photoperiod:chilling 7.1x, photoperiod:forcing 2.4x) (Fig. 1Tab. ??).

Across all species both bud types displayed a relatively proportionate response to forcing (18 and 175 17 day advance for flower and leaves respectively) (Fig. 1, Tab. ??). While there was significant variation among species in their strength of their response to forcing between bud types, no species displayed the characteristic sensitivity pattern of the PHH in which the sensitivity to forcing of the second phase twice as strong as the sensitivity of the first phase (simulations for supp?). Rather, the differences in the strength of the responses of each bud type to each environmental cue combination is signature of the DSH.

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#### Climate change predictions

Our model predicted that both flower and leaf phenology will advance in most of our generalized 183 scenarios for most species, but shifts in FLS depended strongly on how forcing levels change relative 184 to chilling duration (Fig. 3). Following the significant differences in sensitivity to chilling between flowering and leafing phenology we found in our model, FLS interphases were more strongly influ-186 enced by changes in chilling exposure than increased forcing alone. The direction and magnitude of 187 shifts in FLS interphases depended on species and the specifics of FLS phase order, with flowering-188 first and flowering-concurrently species tending to show more profound alterations to FLS patterns than leafing-first taxa. Under some warming scenarios, our model predicted that FLS interphases 190 for some species may effectively disappear or the order of phenophases in the FLS may switch (Fig. 191 3). 192

#### Discussion Discussion

In our study, variation in FLS patterns of deciduous woody plants was dictated by differences in the strength of the response of flower and leaf buds to the primary environmental cues of spring phenology, with differences in the chilling response among bud types being the strongest driver of FLS variation. These result suggest that climate change has potential to significantly disrupt FLSs

as global warming alters historic chilling patterns across the temperate zone. Across all of our treatment combinations, there were species level differences in the magnitude of differential sensitivity 199 to environmental cues, but with high chilling duration treatment, which approximated the historic 200 chilling levels at our collection site, the FLSs for most species followed the predicted pattern of the 201 PHH, with the sensitivity of the second phase of the FLS to forcing approximately twice as strong 202 as that of the first phase 2. This may explain why the two FLS hypotheses have been difficult to 203 distinguish under current field conditions where in most locations chilling requirements for both bud type were frequently met under historic climate conditions (). In conjunction with site-specific 205 FLS shifts and species-specific FLS functions, the difficulty of assessing differential sensitivity in 206 contemporary field conditions there is a need for generalizing principles from experimental studies 207 such as this one, to more fully anticipate the implications of FLS shifts and focus research efforts 208 to the species that may be most affected by FLS shifts with climate change. 209

# 210 Reconciling the DSH and the PHH

The strong differential sensitivity to chilling between flower and leaf buds we found in our study 211 reveals a possible mechanistic link between the DSH and PHH, and offers insight into why these 212 hypotheses have been difficult to differentiate in the past. Our data show that the PHH can be 213 considered to be a special case of the DSH- when the chilling requirement for both flower and leaf 214 buds is met, an an individual appear to follow the predicted pattern of PHH, with temperature 215 during the FLS interphase dictating the inter-annual variability in FLSs. Long term studies sug-216 gest that under historic climate conditions, chilling requirements were generally met (), which may 217 explain why support for the PHH most often associated with observational studies (e.g.). This is 218 consistent with findings in other phenological studies that suggest simple growing degree models 219 (which underlie the PHH) accurately predict phenology under current climate, but under-perform 220 under climate change scenarios when shifts in chilling accumulation become more pronounced (??). 221

By contrast, experimental studies which manipulate chilling levels beyond which was historically

observed in the field tend to support the DSH (e.g. ). The results of our study in wild species are consistent with experimental manipulations of tree-crop phenology which also found a higher sensitivity to chilling for leaf buds (Gariglio et al., 2006; Citadin et al., 2001). Our findings suggest that as climate continues to change, differential sensitivity to the environment between flower and leaf phenology should become more apparent in field observations. However, the adherence of FLSs to the precocity hierarchy patterning under historic conditions suggest that historical range of FLS variation does not constrain the potential for FLS shifts with climate change.

# 231 Implications of the DSH for flower leaf sequences with climate change

The strong differential sensitivity to chilling the between flower and leaf buds we found in our study 232 suggests complex FLS dynamics with climate change. Predicted shifts in chilling are highly variable 233 across both time and space-because chilling only accumulates at intermediately low temperatures 234 warming may increase chilling at some locations while decreasing it in others (). This suggests that the direction and magnitude of FLS shifts is likely to vary substantially among populations based 236 on the specific cue combinations at a given locality. Long-term phenology records show there was 237 already substantial intra-specific variation in FLSs at the population level (Buonaiuto et al., in 238 review) and our findings suggest that these populations level differences may be further amplified by climate change. In this way, all the three generic FLS climate change scenarios depicted in Fig. 240 3 should not be considered alternatives to each other, but rather, could occur contemporaneously 241 across a species' range. 242

Increased population level heterogeneity of in FLSs has potential to alter patterns of pollen dispersal and gene flow across the landscape (). For example, advancing canopy closure relative to
flowering impedes long-distance pollen transport (?). With divergent FLS shifts at the population level, sires from populations in which climate dynamics are extending FLS interphases may
increase their contribution landscape patterns of gene flow relative to populations in which FLSs
are reduced. Depending on the spatial arrangement of these populations and other factors such as

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pollinator movement or prevailing wind directions, this could either facilitate or impeded genetic rescue of climate stressed populations (). Despite these important implications, there is currently little scholarship regarding how inter-population variation in FLS patterns may impact landscape processes like gene flow and recruitment and this should remain an active area of research inquiry.

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The implications of our study's observed differential sensitivity to photoperiod to FLS shifts with 255 climate change are more difficult to characterize. Climate change does not directly impact photoperiod, but warming does shift the time of year when plants become phenologically active, changing 257 the photoperiod they experience. However, depending on the latitude, phenology would have to 258 shifts by at minimum several weeks before the experience photoperiod would change substantially 259 (?). For this reason we modeled climate change scenarios with a constant photoperiod in our FLS 260 projections with climate change, but at high latitudes where photoperiod changes more rapidly 261 over the season, the experienced photoperiod may mute the FLS shifts captured in our projections. 262 This may be particularly important as species shift shift their distribution pole ward with climate 263 change and begin to encounter novel photoperiod regimes (WAY & MONTGOMERY, 2015).

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Our study also highlights that the direction and magnitude of FLS shifts with climate change 266 are species specific. Not only is it likely that the function of FLS variation differs among species 267 (Buonaiuto et al., in review), but we found that FLSs of some species were very sensitive to changing climate conditions while other remain fairly robust (Fig. 3). These differences suggest 269 that some FLS shifts will impact some more than others, and researchers should focus their efforts 270 towards species or populations that are likely to be most vulnerable. However, identify vulnerable 271 species is challenging. At present, observational studies cannot capture the magnitude of FLS 272 shifts with climate, and using artificial environments to manipulate FLSs for all species of interest 273 is impractical and would be prohibitively labor intensive. Therefore, there is a strong need for 274 generalizing principles to aid in identify species with potential for consequential FLS shifts with 275 climate change. While one study cannot hope to represent the taxonomic diversity of a temperate 276

forest, we identified several patterns in the FLS responses of our multi-species experiment that may serve as starting point for further inquiry.

# 279 0.1 Generalizing principles for species-specific FLS shifts

In our study several species, Acer rubrum, Ilex verticillata, Prunus pensylvanicum, Prunus virginiana, and Viburnum acerifolium, had FLSs that were relatively robust to changing environments.
For other species, Acer pensylvanicum, Vaccinium corymbosum and Ilex mucronata, which typically
begin to produce leaves shortly before flowers open, projected FLS shifts were moderate, with the
combination of increased forcing coupled with a reduction in chilling driving the strongest shifts
in FLSs. The two species with the most significant FLS shifts across treatment combinations and
climate change projections were Comptonia peregrina and Corylus cornuta (Fig. 3). In all of our
climate change scenarios, the FLS interphase was dramatically reduced in these taxa.

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It is likely that these three difference response patterns we observed correlate to broader anatom-289 ical, physiological and phenological differences among species. The species that maintained FLS 290 structure across climate change scenarios generally shared a strongly leafing-first FLS, with a fairly 291 long FLS interphase. These species tended to have mixed buds so there may be strong physical 292 constraints on their FLSs. By contrast, the species that were most sensitive to FLS shifts were 293 monoecious, flowering-first, wind-pollinated shrubs. This result may reflect other evidence that 294 wind-pollinated species appear to be more sensitive to climate change than biotically pollinated 295 taxa (Ziello et al., 2012). Given the hypothesized function of FLS in wind-pollinated species, the direction and magnitude of FLS shifts we observed could suggest that that these species, and flowering-first, wind-pollinated taxa in general, may face particular risk for reproductive perfor-298 mance reductions. 290

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Much of the conversation around phenology and pollination in the context of global change has
centered around trophic mismatches between pollinator and floral phenology (Memmott *et al.*,

2007), which is of little relevance to abiotically pollinated taxa. By contrast, the possibility that
the effect of FLS shifts with climate change may be particularly important for abiotically pollinated
woody plants and the scope and impact of FLS shifts in these taxa suggest they should be explored
in greater detail in the future.

# Conclusion:

Our experiment provides strong evidence that while flower and leaf buds respond to the same suite of environmental cues to initiate spring phenological activity, the different bud types rely on 309 each cue with differing strength. This differential sensitivity to cues drives variation in flower-leaf 310 sequences and will dictate the magnitude and direction of FLS shifts with climate change. Shifts in 311 FLSs with climate change are likely to vary across forest communities and depend on the specific 312 combinations of cue levels at a given locality and the species represented there. More research is 313 needed to identify species' traits that may correlate with the potential for FLS shifts, but flowering-314 first, wind-pollinated species may be particular sensitive to FLS shifts and merit continued research 315 focus.

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# Tables Tables

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	Estimate	Est.Error	Q25	Q75	phase
Intercept	77.54	10.01	70.91	84.01	flower
	70.30	8.93	64.56	76.01	budburst
Chill	-21.31	7.54	-26.14	-16.78	flower
	-30.35	5.20	-33.66	-27.06	budburst
Light	-5.99	5.83	-9.73	-2.12	flower
	5.95	5.12	2.68	9.29	budburst
Force	-18.87	6.36	-22.85	-14.85	flower
	-17.39	5.16	-20.70	-14.01	budburst
Chill:Light	-0.70	6.17	-4.60	3.44	flower
	-5.04	4.16	-7.73	-2.26	budburst
Chill:Force	6.75	6.62	2.73	10.96	flower
	12.31	4.77	9.28	15.42	budburst
Light:Force	-5.42	6.22	-9.39	-1.30	flower
	-12.90	4.12	-15.54	-10.17	budburst

Table 1: Model estimates of the effect of variation in chilling, forcing and photoperiod on the flower and leaf phenology of 10 temperate woody plant species suggest that the strength of phenological responses to environmental change is phase specific.

# Figures

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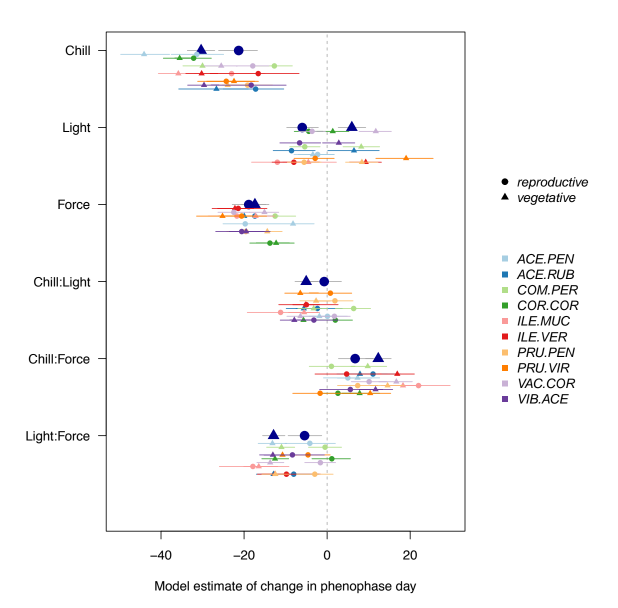


Figure 1: Experimental results suggest differential sensitivity to environmental cues between flower and leaf buds. Vegetative buds (circles) as more sensitive to chilling and interaction between chilling and forcing. Flower buds (triangles) advance with photoperiod increases but leaf buds appear to delay. These differential sensitivities dictate how FLS patterns vary with changing environmental conditions.

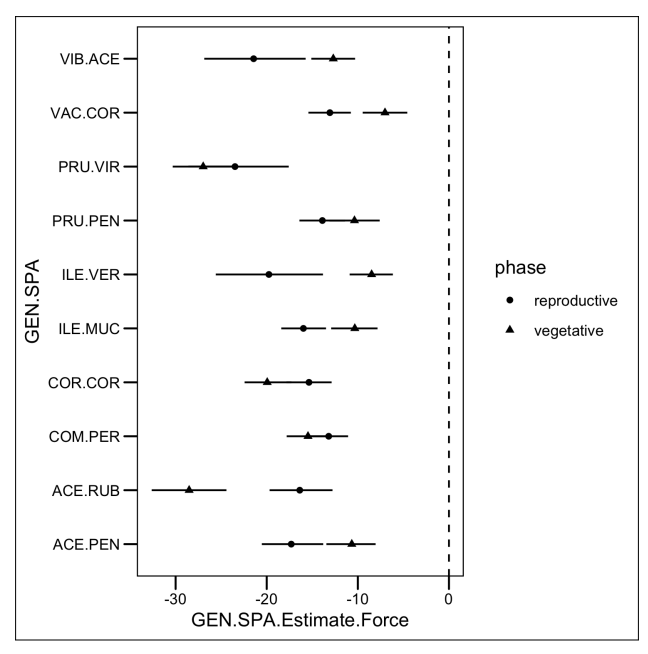


Figure 2: With high chilling and photoperiod, the comparative response to forcing among leaf and flower buds resemble patterns predicted by the precocity hierarchy hypothesis (PHH) Say a bit more about this.

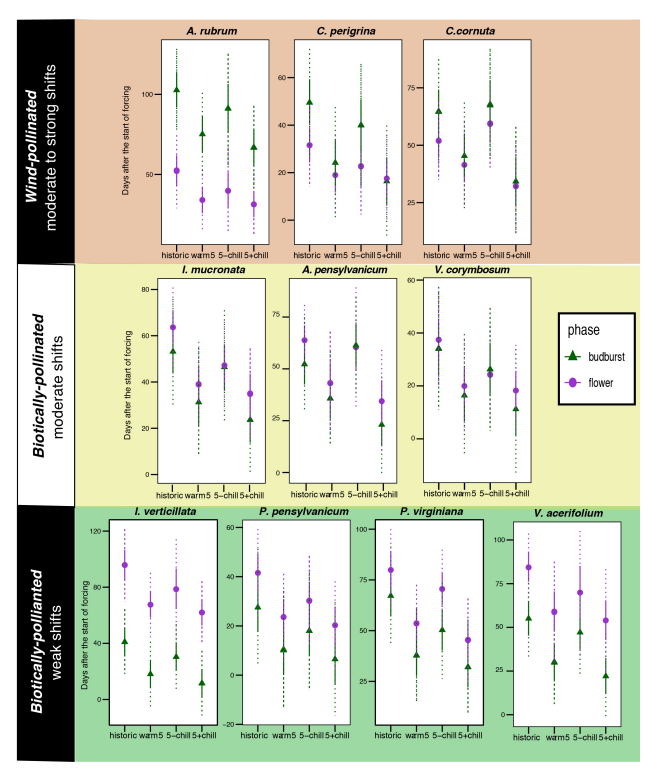


Figure 3: Flower-leaf sequences (FLSs) of temperate, woody species will shift with climate change, but the magnitudes of these shifts vary by species and depend on the specific dynamics of temperature at a given location. We used Bayesian, hierarchical models comparing flower and leaf bud responses to variable temperature combinations to predict FLSs patterns under current climate conditions and three climate change scenarios; an increase in spring warming alone (warm 5), increase in spring warming and increase in winter chilling (warm 5 +chill) and an increase in spring warming and decrease in winter chill (warm 5 -chill). Projected FLS shifts are most pronounced in wind-pollinated, flowering-first species but FLS shifts for all species depend on the relationship between forcing and chilling changes which is likely to vary by location with climate change.