Phenological sequences: how early-season events define those that

follow

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#### Abstract

#### Premise of the study

- Plant phenology is a critical trait, as the timings of phenophases such as budburst, leafout, flowering, and
- fruiting, are important to plant fitness. Despite much study about when individual phenophases occur and
- 14 how they may shift with climate change, little is known about how multiple phenophases relate to one another
- 15 across an entire growing season. We test the extent to which early phenological stages constrain later ones,
- throughout a growing season, across 25 angiosperm tree species.

#### 7 Methods

- We observed phenology (budburst, leafout, flowering, fruiting, and senescence) of 118 individual trees across
- 25 species, from April through December 2015.

#### 20 Key results

- 21 We found that early phenological events weakly constrain most later events, with the strongest constraints
- 22 seen between consecutive stages. In contrast, interphase duration was a much stronger predictor of phenology,
- 23 especially for reproductive events, suggesting that the development time of flowers and fruits may constrain
- the phenology of these events.

#### 5 Conclusions

- Much of the variation in later phenological events can be explained by the timing of earlier events and by
- 27 interphase durations. This highlights that a shift in one phenophase may often have cascading effects on later
- 28 phases. Accurate forecasts of climate change impacts should therefore include multiple phenophases within
- 29 and across years.

## 30 Key words

- angiosperm, arboretum, climate change, global warming, phenology curve, phenological mismatch, plant
- 32 phenology, tree

## 33 Introduction

- <sup>34</sup> Plant phenology, the timing of recurring life-events such as leafout and flowering, is a critical trait that
- affects individual fitness, population abundance, agricultural and natural productivity, and global climate,
- through its role in carbon sequestration (Chuine and Beaubien, 2001; Cleland et al., 2007; Willis et al., 2010;
- Miller-Rushing et al., 2010; Craine et al., 2012). Advancement of budburst, leafout, and other phenophases
- are some of the most widely documented biological impacts of anthropogenic climate change, and phenology
- 39 is likely to be further altered by future climate change (Parmesan, 2006). Because of its important role in
- 40 many ecosystem services and in the global climate cycle, improved understanding and forecasting of tree
- phenology would aid in planning and preparing for climate change impacts.
- Despite the observation that spring phenology generally shifts earlier with warmer temperatures, dramatic
- 43 variation exists in phenological responses to climate. Temperature is thought to be a major factor controlling
- 44 phenology of temperate tree species (Parmesan, 2006; Richardson et al., 2006; Morin et al., 2010; Schwartz

et al., 2013; Clark et al., 2014), but some populations and species have not shifted their phenology with recent warming (Wolkovich et al., 2012). In addition, different tree species vary widely in the timing of leafout and other phenological processes, even when exposed to the same environmental conditions (Lechowicz, 1984; Primack et al., 2009). Spring leafout, for instance, can span weeks among coexisting tree species (Lechowicz, 1984). It has been proposed that, in addition to external environmental drivers such as temperature, some drivers of plant phenology are "endogenous." That is, phenological sequences are affected by changes in 50 internal tree functions that may not be related to climate or other environmental factors (Borchert, 1992; 51 Marco and Páez, 2002). As an example, inflorescence architecture may affect the sequence of leafout to flowering in trees (Marco and Páez, 2002). 53 One important, but often neglected, feature of plant phenology is that events are sequential: leaf budburst 54 comes before leafout, flowering comes before fruiting, and so on. This ordering is an endogenous factor that may constrain phenological responses to climate change. For example, if flowering requires a minimum development time following leafout (e.g., to acquire sufficient carbon), then flowering time may be constrained 57 to shift no more than leafout timing has with recent climate change. This would be the case even if a greater advance in flowering may benefit the plant, for instance if warmer springs have caused pollinator activity to 59 shift earlier (Polgar et al., 2013; Thackeray et al., 2010) The importance of constraints between phenological events is unknown, however, because few studies have integrated across consecutive events throughout a growing season (Wolkovich and Ettinger, 2014). Instead, researchers generally focus on one or two phenophases per study. Early-season events (budburst and/or 63 leafout) have been extensively studied, often using climate-controlled growth chambers (e.g., Basler and Körner, 2012; Laube et al., 2014). A separate group of studies, comprised of long-term observational data, 65 focuses primarily on flowering (e.g., Fitter and Fitter, 2002; Miller-Rushing and Primack, 2008). Interest has recently surged in senescence, which had been less studied historically (Parmesan, 2006), but many of 67 these studies focus only on senescence (e.g., Taylor et al., 2008; Archetti et al., 2013; Jeong and Medvigy, 2014). A contemporary meta-analysis highlights the lack of data on multiple phenophases: only five out of 51 phenology studies (9.8%) included data on both leaf and flower phenology (Wolkovich et al., 2012). When research has looked across stages, important links have often been found (Li et al., 2016). For example, 71 later leafing in a given year may be associated with later flowering, and fall senescence has been associated with both fruit maturation and spring phenology, in different studies (Lechowicz, 1995; Keenan and Richardson, 2015; Liu et al., 2016). In contrast, other research that has quantified multiple phenological events over time has documented that some phenophases may shift asynchronously with climate change (e.g., spring events are occurring earlier as fall events have gotten later, first-flower dates have shifted earlier whereas 82

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- last-flower dates have not, Menzel et al., 2006; CaraDonna et al., 2014). These insights and complications
  demonstrate the need to understand how phenological stages relate to one another across an entire growing
  season (Wolkovich and Ettinger, 2014).
- Here, we examine the extent to which early-season phenological events constrain later events across tree species planted in the same environment. Specifically, we test two hypotheses:
  - Hypothesis 1: Previous phenological events constrain later events; e.g., late-fruiting species set fruit late in the season, because they flower and leafout late (Figure 1). To be consistent with this hypothesis, we expected earlier events, such as flowering, would predict later events, such as fruiting. If constraints are strong, then, across all species, we expected that previous events should predict later events with a slope of one, indicating that the later event happens a set number of days (represented by the intercept) after the previous event (Figure 1). If constraints are weak, we expected to find relationships between later and earlier events, though the slope may be less than one. No relationship would suggest that earlier events do not constrain later events.
  - Hypothesis 2: Interphase duration constrains phenology; e.g., late-fruiting species set fruit late in the season because they require longer maturation time (Figure 1). To be consistent with this hypothesis, we expected that the interphase duration (the number of days between two phenological events) between earlier and later events would be a strong predictor (e.g., a high r<sup>2</sup>) of the later event, regardless of the timing of the earlier event (Figure 1).
- Testing these hypotheses addresses basic, critical questions about drivers of variation in temperate tree phenology. These questions remain unanswered despite decades of phenology research because previous field studies rarely examined multiple phenophases spanning the entire growing season across a large number of tree species.

#### $_{ iny 9}$ Materials and Methods

#### Study site and focal species

This study was conducted at the Arnold Arboretum of Harvard University (42.3074°N, 71.1208°W), a 281-acre park in Boston, Massachusetts, established in 1872. It contains a living collection of 3,825 woody plant taxa that are native to North America, Europe, and Asia. Arboreta are excellent resources for phenological studies

across many species (e.g., Primack and Miller-Rushing, 2009), particularly in temperate areas, because they
may contain a higher diversity of tree species growing in one location than nearby natural areas. In addition,
there is often great variation in phenology of species planted in arboreta, for public enjoyment of foliage and
flowers throughout the season. For this study, we selected 25 focal angiosperm species with diverse flowering
times, in order to maximize variation in phenology (Table 1). We selected three to five individuals of each
species for the study, yielding a total of 118 individuals.

#### 110 Phenology data collection

We visited each individual once every 6-10 days throughout the growing season, and, when necessary, used 111 binoculars to assist with phenology observations. Phenology observations in the spring began on 6 April 2015 112 (prior to any budburst occurring) and fall phenology observations ended on 2 December 2015 (at which point 113 more than 95% of leaves on all trees had changed color and dropped). We observed five phenological stages, which were quantified following the National Phenology Network protocols (for a full description see Denny 115 et al., 2014). The budburst phase was characterized by green leaf tips being visible at the tips of buds, and the leafout phase was characterized by visible fully unfolded leaves and petioles that had completely emerged 117 from the buds. The flowering phase was when open flowers were visible, and the fruiting phase was defined by ripe fruit being visible. Leaf senescence was characterized by leaves changing from green to fall colors. 119 On each observation day, we noted whether or not each phenophase was present. If a particular phase, such 120 as budburst, was present, we estimated the abundance of the phenophase on the individual tree, following 121 Denny et al. (2014). For budburst and fruiting phenophases, abundance categories were, <3, 11-100, 101-122 1000, 1001-10000, or >10,000 buds bursting or fruits present. For leafout, flowering, and leaf senescence 123 phenophases, abundance categories were by percentage: <5%, 5-24%, 25-49%, 50-74%, 75-94%, and 95% or 124 more. For leafout, we estimated the potential canopy space full with leaves; for flowering, we estimated the percentage of flower buds that were open; for senescence, we estimated the percentage of potential canopy 126 space full with non-green leaves. In all these estimations, we ignored dead branches.

From the field observation data, we extracted the day-of-year (DOY) of the first observed occurrence of a given phenological phase. Budburst and fruiting DOY were defined as the first day when three or more burst leaf buds or ripe fruits, respectively, were observed on the individual. Leafout, flowering, and leaf senescence DOY were defined as the first day when 5% or more of the individual was leafing out, had open flowers, or showed fall colors, respectively (Denny et al., 2014). From these individual tree phenology observations, we calculated species-level mean start dates for all phenophases, for use in our statistical analyses. We used these

mean start dates to calculate interphase durations between phenophases. We calculated interphase duration
as the number of days between the start of the later phenophase and the start of the earlier phase (Figure
1).

#### 137 Statistical analyses

To understand the extent to which previous phenological events constrain later events across species (Hypoth-138 esis 1, Figure 1), we fit linear models in which the response variable was phenological stage (i.e., the species' 139 mean DOY of leafout, flowering, fruiting, or senescence), and the predictor was previous phenological stage. 140 Thus, budburst was excluded as a response variable, because it was the earliest stage we quantified, and 141 senescence was excluded as a predictor variable because it was the latest stage we quantified. We therefore 142 fit 10 separate regression models, estimating the intercept of the relationship between later and previous 143 phenological phases, and forcing the slope to be one (Hypothesis 1, Figure 1). In addition, we fit 10 models, with the same predictor and response variables, in which we estimated the best-fit slope and intercept (via 145 least-squares, e.g., a standard regression model). Under Hypothesis 1, with strong constraints, we expected that the models with forced slopes should provide similar fit to the data as the standard regression models 147 that estimate both slopes and intercepts. We compared fit of these two model structures using r-squared values, as well as Akaike's Information Criterion (AIC). 149

To understand the extent to which interphase duration constrains later phenological events (Hypothesis 2, Figure 1), we fit linear models in which the response variable was phenological stage (DOY), and the predictor 151 was the number of days between phenological stages. Thus, as above, budburst was excluded as a response variable. We therefore fit 10 different models, each with one of four phenological stages as the response variable 153 and one of the interphase durations preceding the focal phenological stage as a predictor. To investigate the effect of interphase duration, we fit models in which the interphase durations were randomized with respect 155 to the timing of the earlier phenophase across species. We did this resampling of interphase duration 999 156 times for each model structure. If our expectation of later events being constrained by interphase duration 157 was supported, then the best-fit slope should fall within the resampled models' slope estimates. Furthermore, 158 if the constraints imposed by interphase duration were strong, then the randomized models should include a 159 narrow range of possible slopes. To examine these possibilities, we compared the range of slopes of all the 160 resampled models to the slope of the fitted model.

All analyses were conducted in R version 3.2.4 (R Core Team, 2017), and code is available with the dataset online at Knowledge Network for Biocomplexity (Gee et al., 2017).

#### 164 Results

We monitored five phenophases, which varied in duration. First budburst date occurred over 32 days and 165 first leafout date occurred over 30 days in the spring, across all focal individuals (Appendix S1) and species 166 (Figure 2). Flowering phenology occurred over a longer period than budburst and leafout, spanning 131 167 days from late April to September. The first observation of ripe fruit spanned 175 days, and the start of leaf 168 senescence occurred over 56 days across all individuals and species. Most species (20/25) spent the majority of 169 the growing season in the reproductive phenological phases (i.e., flowering and fruit development), and most 170 species (23/25) began leaf budburst prior to flowering, though leaf development overlapped with flowering in 171 some species (Figure 2). The majority of species (15/25) produced ripe fruit prior to beginning senescence 172 (Figure 2). 173

We found that the timing of early phenological stages predicted the timing of later stages in many cases (Figure 174 3, Appendix S2). The strongest relationships (i.e., with the most variation explained) occurred between 175 adjacent stages (those along the diagonal in Figure 3, such as leafout and budburst, fruiting and flowering). 176 Even for adjacent phases, however, neither the forced slope models nor the regression models explained the 177 majority of the variation in phenology (r<sup>2</sup><0.30, in most cases). For three relationships (budburst versus 178 flowering, leafout versus flowering, and flowering versus fruiting), the model fits were consistent with the strong 179 constraint hypothesis—the forced slope model provided similar fit to the regression models and both models explained a significant amount of variation in phenology (Figure 3, Appendix S2). For four relationships, 181 standard regression models explained much more variation than the forced slope models. For two of these relationships (budburst versus leafout, and fruiting versus senescence), the regression models had slopes that 183 were less than one, and for the other two relationships (budburst versus fruiting and leafout versus fruiting), the regression models had slopes that were greater than one (Figure 3, Appendix S2). 185

We observed strong relationships ( $r^2>0.7$ ) between phenology and interphase duration for the two reproductive phenophases (flowering and fruiting time, Figure 4, Appendix S3). Flowering DOY was strongly predicted by days between flowering and leafout ( $r^2=0.93$ ), as well as by days between flowering and budburst ( $r^2=0.87$ ). Fruiting DOY was strongly predicted by days between fruiting and flowering stages ( $r^2=0.74$ ), by days between fruiting and leafout ( $r^2=0.98$ ), and by days between fruiting and budburst ( $r^2=0.97$ ). Senescence was predicted by days between senescence and budburst ( $r^2=0.74$ ), days between senescence and leafout ( $r^2=0.82$ ), and days between senescence and flowering ( $r^2=0.17$ ); senescence was not affected by days between senescence and fruiting. Leafout was not predicted by interphase duration (i.e., time since budburst). In all cases where later events were well predicted ( $r^2>0.7$ ) by interphase duration, the best fit model was predicted by our null model (i.e., the randomizations). However, the relationship between senescence and days between senescence and flowering was weaker than expected by our null model (see resampling estimates in Figure 4,Appendix S3).

#### 198 Discussion

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The ordering inherent in phenology means most phenological stages are dependent—in some way—on preceding stages: dormancy must be broken before buds can burst and flowers must be produced before fruit.

Here we show how the major phenological stages of temperate trees are constrained by one or more earlier phenological stages.

All phenological stages we observed support Hypothesis 1: timing appears to be constrained by at least one 203 previous phenological stage. Our findings are consistent with recent work suggesting that senescence time can be affected by earlier phenophases (Keenan and Richardson, 2015; Liu et al., 2016; Li et al., 2016), and suggest that this one relationship is part of a larger suite of correlated phenophases. Consecutive events were 206 correlated across both growth and reproductive phenophases (i.e., flowering and leafout were correlated to 207 a similar degree as fruiting and flowering, Figure 3). These associations may occur because of endogenous 208 dependencies between the two phases, because of a shared external driver such as growing degree days, or a combination of endogenous and external factors (Lechowicz, 1995). Thus, environmental conditions in the 210 winter or spring that may directly affect only early phenological stages, such as budburst, are likely to have cascading effects on later stages such as leafout, flowering, and fruiting. Our data suggest that, for most 212 events, these effects are more apparent for consecutive stages (i.e., those along the diagonal in Figure 3), and 213 are well-approximated by the forced slope model in some cases (Figure 3).

Although some of the variation in reproductive phenology (flowering and fruiting) was explained by previous phenology (Hypothesis 1), much more variation was explained by interphase duration (Hypothesis 2). Later flowering species generally required more time between flowering and leafout. Similarly, late-fruiting species had longer interphase durations between the first observation of ripe fruit and first flowering date. It may be that late-fruiting species require longer fruit development times to produce larger fruits or more highly-provisioned seeds. This would be consistent with previous theories that trees investing more resources into their offspring (i.e., having larger seeds) require more time to build resources (Bolmgren and Cowan, 2008; Sun and Frelich, 2011). There were notable exceptions to this general relationship, however. Some species set fruit later than expected, given their interphase duration between flowering and fruiting (Catalpa speciosa,

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Tilia americana, T. japonica, Figure 4). These species also flowered later than expected, given their leafout DOY (Figure 3). External factors related to their ecology may be the cause; for example, these species 225 are all insect-pollinated, so the timing of their pollinator activity may have affected their floral phenology (Elzinga et al., 2007). Other external factors may also shape the time of fruiting: for example several species 227 ripened fruit after leaf senescence, which was most dramatic for two species (Liquidambar styraciflua, Platanus occidentalis) that are wind-dispersed, suggesting they may ripen after leaf senescence to benefit from fewer 229 leaves, higher winds, and thus further dispersal of their seeds. The phenology classifications we used did not 230 allow us to separate out exactly when fruit development and ripening begins and ends; detailed measurements 231 of resource allocation and translocation, as well as observations of individual flower buds, will be required to 232 understand the physiology underlying the phenological patterns we observed. 233

Despite our expectation of strong relationships between later phenophases and interphase durations, our results show two relationships that are weaker than expected (Figure 4). Leafout was not predicted by the interphase duration between budburst and leafout, and senescence was not predicted by the interphase duration between fruiting and senescence (Figure 4). We had expected that these two sets of phases would demonstrate stronger constraints of interphase duration because they occur at the beginning and end of a bounded growing season (Letten et al., 2013). The weak ability of interphase duration to predict leafout may be due to the distinct weather patterns in 2015 and how they interacted with species' cues for spring phenology. Trees have species-specific chilling and forcing requirements that must be met prior to leafing out, and are generally understood to be related to accumulations of cold and warm temperatures (e.g., Schwartz and Hanes, 2010; Chuine, 2010; Clark et al., 2014). The pattern of how quickly cooler and warmer temperatures accumulate across a growing season can therefore impact how variable leafout is across species. In contrast to some years that have high variation in leafout date across species (Lechowicz, 1995), in our study year (2015) many species leafed out close to DOY 130 (10 May), regardless of leafout-budburst interphase duration (which ranged from 0 to 20 days, Figure 4). Temperature conditions particular to 2015 may have caused this: temperatures were colder than average in January through March, and then switched to aboveaverage in late April and early May (www.bluehill.org). Such long periods of cold followed by rapid warming may have meant that chilling requirements were met for all species well before warm temperatures began, and then forcing requirements were rapidly met for many species (even if they had diverse requirements) leading to a flush of leafout in early May, across diverse species. Variation in the environment each year may alter how later phenophases are constrained by earlier events. This highlights the need to understand how such constraints vary across years and what underlies the remaining variation not explained by earlier phenophases.

Our results indicate that both Hypothesis 1 and Hypothesis 2 are operating and can explain a substantial amount of the variation observed among species in a single year. For example, although later fruiting is generally associated with later flowering (Figure 3), some species, such as Quercus alba and Quercus grandifolia, flower relatively early and fruit late; later fruiting for these species is instead associated with longer interphase duration between fruiting and flowering (Figure 4). Understanding drivers of variation in phenological responses among species requires disentangling the ways that earlier phenology and interphase duration interact with one another, and with environmental conditions. Experimental manipulations will be beneficial for discerning the physiological and genetic bases for the relationships we observed (Flint, 1974), as will long-term studies with enough years of data to encompass large amounts of natural variation in temperature (e.g., Inouye, 2008).

Further research could also illuminate at what other biological levels these constraints occur, and how they are shaped by genotypic and environmental variation. We exploited the diverse species plantings of an arboretum 267 to examine phenological constraints across species, but such constraints may also operate within species. Phe-268 nology frequently varies across populations within a species, because of environmental differences (i.e., plants leaf out earlier in warmer locations) and/or local adaptation (Rathcke and Lacey, 1985). Local adaptation 270 to diverse environments could potentially drive stronger constraints at the intraspecific level: for example, 271 environments with short growing seasons may select for shorter interphase durations, whereas longer growing 272 seasons may select for the opposite, and studies across such environments may shower stronger correlations among phenophases. Addressing this, however, requires teasing apart the influence of environment (even to 274 the microclimatic level, see Schwartz et al., 2014), versus genotype and thus may be best accomplished in a 275 common garden setting with multiple individuals of the same genotype. Especially beneficial for predictions 276 would be multiple common gardens across and beyond species' ranges. In addition, given that trees can 277 respond to environmental factors at even finer scales (e.g., the branch level, Nakamura et al., 2010), the phenological constraints we observed may affect phenology within an individual tree, as well. 279

Our findings have important implications for improved forecasting of climate change-induced shifts in phenology. A shift in one phase may have cascading effects on later phases, since each phase is linked to phases
that occur before and after it (Wolkovich and Cleland, 2014). Phenology models covering the whole annual
cycle are rare for wild plants (but see Chuine and Beaubien, 2001; Li et al., 2016) and have not been used to
forecast phenology at the community level. Our results highlight a clear need to conduct future multi-species
phenology studies across entire growing seasons (Wolkovich and Ettinger, 2014), and begs the question of
how phenophases may be linked across years, as well (e.g., Elmendorf et al., 2016). For example, the timing
of spring budburst in one year may be related to the timing of budset the previous fall (Mimura and Aitken,

2010). Although relationships between phenophases have not been widely studied, there is a growing ecological literature on the concept of "ecological memory," or the capacity of past states to influence present or
future responses (Ogle et al., 2015). The ecological memory of phenology has not been quantified, but may
be critical for accurate forecasting, particularly for species like *Quercus rubra*, which require more than one
year for fruit maturation, as well species that preform buds multiple years in advance of budburst (Diggle,
1997; Klimes et al., 2007). For example, one study of desert shrubs found that one of the best predictors
of leaf phenology in a given year was the phenology of the of the same individual in prior years (Cody and
Prigge, 2003). Accurate forecasts of community-wide phenological shifts are likely to require species-specific
(or even individual-specific) information, in addition to climate data (Diez et al., 2012).

### Conclusions

We have shown that early and late phenological stages are strongly linked across the growing season, providing a new approach to explain some of the dramatic variation in phenological responses observed to date. Many 299 studies have sought to identify the particular environmental drivers of phenology (e.g., Morin et al., 2010; 300 Schwartz et al., 2013). Our findings here suggest that timing and duration of previous phenological states 301 should also be examined. In addition, identifying the appropriate temporal window for both environmental 302 and endogenous drivers is essential (Teller et al., 2016). Because earlier phenophases define those that follow, 303 the relevant time period for these drivers may extend further back in time than the single growing season 304 we evaluated here. Multi-year studies will be critical to evaluate the extent to which phenological patterns are consistent among years that may vary in climate, as well as biotic conditions (i.e., pollinator or pest populations, Lechowicz, 1995). A fuller understanding of phenological constraints and drivers of phenological variation offers the potential for improved forecasts of phenological shifts with climate change to help predict 308 how ecosystem functions will be altered in the future.

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## Data Accessibility

The data set and R code used in this study are available online at Knowledge Network for Biocomplexity

(Gee et al., 2017).

#### 323 Author contributions

- 324 All authors conceived of and designed the study and edited the manuscript; S.G. conducted the fieldwork;
- S.G. and A.E. analyzed the data, and A.E. wrote the manuscript.

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# Tables

Table 1: **Study species.** Twenty-five angiosperm species were selected based on their flowering phenology in long-term records of the Arnold Arboretum. The flowering patterns we observed during our one year of data collection did not always perfectly match these long-term patterns. The number of individuals of each species observed at the Arnold Arboretum from spring through fall 2015 is in parentheses.

Early-season flowering	Mid-season flowering	Late-season flowering
Aesculus flava (5)	Carya glabra (5)	Catalpa speciosa (5)
Betula alleghaniensis (5)	Carya ovata (5)	Kalopanax septemlobus (3)
Betula nigra (5)	Crataegus crus-galli (5)	Styphnolobium japonicum (5)
Gleditsia triancanthos (5)	Fagus engleriana (4)	Tilia americana (5)
Liriodendron tulipifera (5)	Fagus grandifolia (5)	Tilia japonica (5)
Phellodendron amurense var. lavallei (4)	Fraxinus chinensis (5)	
Populus deltoides ssp. deltoides (5)	Liquidambar styraciflua (5)	
Pyrus calleryana var. dimorphophylla (3)	Platanus occidentalis (5)	
Pyrus ussuriensis var. hondoensis (5)	Quercus glandulifera (4)	
Quercus alba (5)	Quercus rubra (5)	

## Figure legends

Figure 1: **Hypotheses.** We show flowering and fruiting as examples of consecutive phenological events. We expected the same patterns for other consecutive events, such as leaf budburst and leafout. Interphase duration is the time between phenological events, e.g., the number of days between the first day of flowering and the first day that ripe fruits were observed.

Figure 2: Species' phenology during the 2015 growing season, ordered by mean first-flower dates. Growth phenology is shown for budburst (from its mean start day-of-year to the mean start day-of-year for leafout, across all individuals within a species), leafout (from the mean day-of-year when fully-expanded leaves were first observed through the start of senescence), and senescence (from the mean day-of-year when leaves first began changing color through the mean day-of-year when more than 95% of leaves on the tree had changed color). Reproductive phenology is shown for flowering (from the mean day-of-year when flowers first appeared to the mean day-of-year when fruits first appeared to the mean day-of-year when fruits first appeared to the mean day-of-year when more than 95% of fruits were first observed as ripe).

Figure 3: Relationships among phenological stages across the 25 focal species. Linear models were fit with the species-level mean day-of-year (DOY) of the later phenological stages as the response variable, and mean day-of-year of earlier stage as the explanatory variable. Models with a forced slope of 1 are shown by dashed red lines, and  $\rm r^2$  is given when  $\rm r^2{>}0.10$ . ("fs", in red).  $\rm r^2$  for standard regression ("reg," in black) and lines for these models are shown when  $\rm r^2{>}0.10$  (solid black lines). Asterisks highlight when Hypothesis 1 was supported; full model statistics are summarized in Appendix S2 in the Supplemental Materials. Species in the legend are ordered from early to late first-flower dates.

Figure 4: Relationships among phenological stages and interphase duration across the 25 focal species. Interphase duration (x-axis) is the time between the start of the earlier phenological event and the start of the later phenological event (e.g., the number of days between the species' mean start of flowering and its mean start of fruiting). Linear models were fit with the species-level mean day-of-year (DOY) of the later phenological stages as the response variable, and interphase duration as the explanatory variable. Solid lines (representing model fit) and  $\rm r^2$  are shown when  $\rm r^2 > 0.10$ . Gray lines represent model fits when interphase was randomized with respect to the timing of the earlier phenophase (in some cases the range of possible relationships is quite narrow, given the constraints of ordering inherent in phenological events). When our null expectation of later events being constrained by interphase duration was supported, the best-fit slope (black line) falls within the randomized lines (in gray). Asterisks highlight when Hypothesis 2 was supported; full model statistics are summarized in Appendix S3 in the Supplemental Materials. Species in the legend are ordered from early to late first-flower dates.