A series of phenological events: Reconciling historic hypotheses regarding flower-leaf sequences in temperate forests

Daniel Buonaiuto, Nacho, Lizzie
June 17, 2019

To Do:

Abstract

As the the study of phenology has developed overtime, it has become increasing clear that it is not only individual phenological stages that affect organism fitness and ecological functioning, but also the relationship between stages. Of these phenological relationships, the flower-leaf phenological sequence (FLS) of deciduous woody plants have received particular treatment. The high incidence of a flowering-first species in the temperate zone has lead to the suggestion that FLS's are adaptive, and several hypotheses have been put forth to explain their evolution and function.

Classically, FLS's have been treated categorically and assigned at the species level. However, long-term phenological data suggest that not only are there substantial differences between species' FLS that are masked by categorization, there is also considerable intra-specific variation in FLS, at both the population and individual level. These observations suggest that the existing conceptual framework my be impeding our understanding of this phenological trait, and a new framework is needed.

In this paper, we first review the existing FLS hypotheses, and discuss how the existence of significant intra-specific variation would modify the hypotheses. We then present four case studies from temperature forest species which demonstrate that multiple hypotheses seem to be supported, and highlight that inference regarding the FLS hypotheses is highly sensitive to how FLS's are defined.

The review and case studies provide three major insights towards a new conceptual framework for the study of FLS. Fist, future research should allow for the coincidence of multiple hypotheses which can be achieved both by allowing for overlapping hypotheses in large community models and by testing individual hypotheses in smaller sub-groupings that control for other trait variation. Secondly, FLS's are better treated as continuous

variables and the classic categorization of FLS should be abandoned as continuous measures of FLS reduce the impact of observer bias, and allow for stronger inference regarding FLS variability both among and within species. Finally, researchers should move beyond hypothesis testing through trait associations and use intra-specific measures of FLS to test for fitness consequences in FLS variation.

Introduction

Phenology, the timing of seasonal life cycle events, allows organisms to synchronize important life history transitions with optimum environmental conditions (Forrest & Miller-Rushing, 2010), and is a critical component of ecosystem structure and function (Cleland *et al.*, 2007; Piao *et al.*, 2007). Recent work in woody plant phenology has begun to show that it is not only individual phenological stages that affect these processes, but also the relationship between them (Ettinger *et al.*, 2018).

One phenological relationship that has long received scientific interest (see Robertson (1895)), and recently, increased attention in the literature is the flower-leaf phenological sequence (FLS). In a typical model of plant life history, vegetative growth precedes reproduction. However, for many species in the deciduous forests of Eastern North America, it is not the green tips of new shoots that mark the commencement of the growing season, but the subtle, reds, and yellows of their flowers. This flowering-first FLS is common in these regions, and its prevalence has lead to the suggestion that this FLS has adaptive significance (Rathcke & Lacey, 1985). Recent advances have been made in characterizing the evolution and physiology of FLS (Gougherty & Gougherty, 2018; Savage, 2019), but many questions regarding the function of this phenological syndrome remain.

A deep inquiry into the nature of this phenological pattern is necessary and timely, because as can be seen in figure 1, anthropogenic climate change is altering FLS's, but the FLS response to climate change differs among species. In our comparison of three European tree species, all species increased the offset between flowering and leaf phenophases, but the rate of change varied between species. In fact, the mean FLS offset for one species, Fraxinus excelsior has already exceeded its historic range of variability, while Aesculus hippocastanum shows a more muted response. Depending on the function of FLS, this differential FLS sensitivity to climate change may have implications for community composition and population demography in the future.

As the study of phenology has matured as a discipline, it has become clear that measures of synchrony and variability are key components to understand the fitness benefits of distinct phenological syndromes (Ollerton & Lack, 1992). It would be expected this would hold true for phenological sequences as well. However, while some have found general correlation between flowering and leafing phenology (Lechowicz, 1995; Ettinger et al., 2018), fine scale FLS variability has never been evaluated. We suggest that characterizing FLS variation among individuals and populations will allow for a more biologically

relevant evaluation of the current FLS hypothesesm as well as reveal avenues for future, direct hypothesis testing.

Here we 1) Review the adaptive hypotheses of FLS and their respective predictions, 2) Evaluate variation in FLS, and explore how FLS variation within species, populations and individuals alters the predictions of the hypotheses, 3) Show how the incorporation of variation alters which hypotheses are supported using case studies from temperate forests, and 4) make recommendations for future study of FLS.

Defining FLS

Flower-leaf sequences have traditionally been classified into distinct qualitative categories that are almost always defined at the species level. The terms hysteranthy, protanthy, proteranthy or precocious flowering describe plants which produce flowers before leaves (Lamont & Downes, 2011; Heinig, 1899). A classic example of this FLS is *Acer rubrum*, which, as seen in figure 2, reaches peak flowering weeks before any sign of leave development. These species tend to exhibit a degree of physiological specialization, such as the separation of flower and leaf buds.

Seranthy, an FLS syndrome embodied in figure 2 by the species *Nyssa sylvatica*, describes species in which flowers begin to open after leaves are approach this full size. These species may still differentiate flower buds in the previous season, but may rely less on stored energy than flowering-first taxa.

But what about species whose FLS separation is less clear? It is possible to describe all species whose flowering period overlaps their leaf development as synanthous (Lamont & Downes, 2011), but this third category may obscure important inter-specific differences. Furthermore, both the flowering and leaf growth periods consist of several sub-stages making it difficult to fit FLS patterns neatly into these categories.

Take Betula alleghaniensis from figure 2 for example: One would be justified in classifying this species as hysteranthous because its flower buds tend to burst before its leaf buds, or as synanthous for the fact the its open flowers overlap the beginning of leaf growth. Can we really put this species in the same category as Acer rubrum, whose flowers open weeks before the leaves? Conversely, is this species truly similar to figure 2's Acer pensylvanicum whose flowers do not open until leaves are well along in their expansion? These kinds of decisions are best guided with a focus on their implications for the FLS hypotheses, which we will discuss in detail below.

Hypotheses of FLS and their predictions

Wind pollination

The most prevalent FLS hypothesis associates hysteranthous flowering with pollination syndrome, suggesting that hysteranthy is an adaptation critical for effective wind pollination, with leafless flowering allowing for more efficient pollen dispersal and transfer (Whitehead, 1969; Stephen H. Spurr, 1980; Friedman & Barrett, 2009). This hypothesis hinges of the fact that the presence of leaves results in a substantial physical disruption to pollen transfer, a premise that we would not necessarily expect to be true for the early stages of leaf expansion, when tiny leaf primordia would have little impact on environmental structure. In this framework, trees that flower during the early stages of leave expansion would be expected to gain similar mechanical advantage to those who complete their flowering before any leaf activity. Therefore, this hypothesis predicts that wind pollinated species should flower before or with their leaves, while in animal pollinated species, FLS should be random or co-vary with pollinator activities.

Water dynamics

Another hysteranthous hypothesis, emerging from the dry deciduous tropics where flowering during the leafless season is also common (Janzen, 1967), suggest that flowering before leaf development is an adaptation to reduce water stress associated with maintaining floral hydration while leaves are transpiring (Franklin, 2016). This hypothesis suggests that there would be a significant cost to maintaining floral structures during any stage of leaf activity, and as such, it would be expected only species whose flowering occurred before any leaf expansion would gain this drought advantage. This hypothesis predicts that species that are drought tolerant should flower before leafing out, with minimal overlap between the floral and foliate phenophases. Species that are not drought tolerant get no real advantage from flowering first, so in these species FLS should be random.

Early flowering

A third possibility is flowering-first FLS is a physiological byproduct of selection for early flowering (Primack, 1987). Within this framework, there is no advantage to a species being hysteranthous vs. seranthous, as long as the absolute flowering time of the contrasting FLS's were the same. However, this equivalency may simply be a physiological impossibility. Recent work from Savage (2019) has demonstrated that flowers are primarily hydrated by the phloem and therefore are independent of xylem re-genesis, which is not the case for the xylem maintained leaf tissue. With physiological constraints on leaf phenology but not on flowering, selection for early flowering would drive hysteranthy. This would explain why hysteranthous species tend to be the earliest species to flower. Here, we would expect increased FLS offset (time between flowering and leaf out) to be

associated with generally earlier flowering phenology. We also would expect associations with other early flowering traits such as seed mass, dispersal season or cold tolerance to be more pronounced. However, this hypothesis does not require the selective driver of early flowering to be exclusively be one of these traits, and pollination syndrome or drought tolerance may still play a role in driving the early flowering (Savage, 2019).

This hypothesis predicts that species' flowering times should be strongly associate with flowering-first FLS. It also is likely there would be relationship between this FLS and other early flowering traits.

Phylogenetics

Finally, it is also possible that FLS's are highly conserved traits, and the preponderance of hysteranthy in the temperate zone is a product of phylogenetic representation of the region rather than an adaptive quality to the trait. There is no past biology to suggest differences between species that flower well in advance of the leaves or just prior to their emergence, but considering but extra categories may help see the patterning across the tree. This hypothesis predicts strong phylogenetic patterning in the FLS with no correlation with other traits to be expected.

Variation in FLS

All of the above hypotheses assume that FLS's are a species-level trait, however, this assumption has not been well examined in the literature (Gougherty & Gougherty, 2018). Intra-specific variation is the engine of natural selection, and if it is substantial in FLS patterns, there is much to be inferred from it regarding the origins and trajectory of this trait as the climate changes. We investigated individual FLS variation using a long term phenological data set collected at Harvard Forest in Petersham, Massachusetts (O'Keefe, 2015). There was substantial variation in FLS offset among years, with offset values varying by up to several weeks for most species. This variability can significantly blur FLS categorization. As seen in 3 Q. rubra, a species classically listed as flowering and leafing in synanthy, there are some years in which flower budburst is over a week before leaf budburst, and other years, in which leaf buds burst weeks prior to floral budburst. We also found there to significant population level variation in FLS, using the Pan European phenological database PEP725(Templ et al., 2018), with some populations differing in their mean FLS offset by a week or more.

Given the variability of FLS at the individual and population level, it is clear that considering FLS variability at only higher taxonomic levels may obscure important realities about the biology of this phenological trait. Below, we discuss how the observed variation below the species level may alter the existing FLS hypotheses.

How FLS variation alters predictions

Wind pollination

Intra-specific variability in FLS would complicate the wind pollination efficiency hypothesis considerably. It is well accepted that pollination syndrome is a species level trait, considered to be fairly immutable across ecological time and space. Under this hypothesis, one would not expect significant variation in FLS across population or individuals because one would not expect variation in pollination syndrome. However, as discussed above, a tree with no overlap between flowering and leafing phenology does not necessarily gain a significant pollen transfer advantage over an individual with some overlap. It is clear the the pollination efficiency advantage from flowering first diminishes as the canopy fills in, but the dynamics of this impact are not well characterized. We do not know at what point during leaf expansion pollination would become significantly encumbered, so it is possible, that interannual and population level variation in hysteranthous FLS could maintain a wind pollination advantage, as long as the overlap did not cross a certain unknown threshold. Therefore, based on the wind pollination efficiency hypothesis, would would not expect high levels of population or individual variation in FLS, but the detection of some FLS variability at these levels, does not inherently challenge the plausibility of the hypothesis.

Water dynamics

If FLS's are driven by water dynamics, we would expect there to be significant population level variation in FLS. Populations growing in drier habitats would be expected to show a stronger hysteranthous pattern than their counterparts growing in wetter habitats where there would be more relaxed selection for minimizing phenological overlap. Therefore, we would predict a correlation between FLS and average soil moisture. This hypothesis also suggests that water availability may drive interannual FLS variation, with drought years increasing hysteranthy, and wetter years permitting more FLS overlap. We might only expect to see a signal for the association between a drought tolerance and hysteranthy if the phenological observations for the species came from populations in drought prone regions.

Early flowering

This hypothesis predicts some variation on the population level based on local adaptation. Populations in which selection for earlier phenology is stronger, perhaps those in regions with shorter growing seasons, would be expected to show a higher degree of hysteranthy. At the individual level, FLS variability could be driven by interannual variability in spring conditions. Both flowering and leaf phenology are strongly cued to temperature and photoperiod (Flynn & Wolkovich, 2018; Rathcke & Lacey, 1985), but with leaf phenology constrained by xylem activity and flowering phenology relatively independent of

it, we would expect a stronger response in to environment in flowering time resulting in FLS variation. With the greater sensitivity to environment of flowering than leafing, we would expect FLS variation to be positively associated with flowering phenology variation. Below the species level, this hypothesis predicts that early flowering years or populations are associated with increased FLS offset for hysteranthous species.

Phylogenetics

With the lack of treatment of FLS variability in the literature below the species level, we have no strong basis for asserting whether the apparent variability in FLS is a product of genetic or environmental controls. If there is a strong genetic component to FLS as has been show in other phenophases (Wilczek *et al.*, 2010), some population level variation could be driven by reproductive isolation. With strong genetic control to FLS, we might also see consistent genotypic differences in FLS among individuals within a population, but would not predict high levels of interannual variation in FLS.

Available evidence for FLS hypotheses in temperate woody species

Despite a strong conceptual basis, direct tests of these hypotheses of hysteranthy in the literature are relatively rare, and support for them is mixed. Many studies only test a single hypothesis, making comparison between them difficult. For example, the primary evidence supporting the wind pollination hypotheses comes from pollen diffusion studies, e.g., particle movement through closed and open canopies (Niklas, 1985; Nathan & Katul, 2005; Milleron et al., 2012), which provide no framework for comparatively evaluating the other hysteranthy hypotheses. We are aware of no direct test that have tried and distinguish selection for hysteranthy from selection for early flowering, but Primack (1987) notes that hysteranthous, wind-pollinated species tend to also have large seed mass, and lack primary seed dormancy for germination, traits associated with early flowering in general. This raises the distinct possibility that hysteranthy may simply be one component of a larger suit of early flowering traits. We are also aware of no studies that have mechanistically evaluated the water dynamics hypothesis, though observations of flowering in the dry tropics by Borchert (1983); Reich & Borchert (1984) suggest that the timing of flowering in hysteranthous taxa is associated with a plant water status recovery due to leaf drop. Only recently has it even been suggested that this hypothesis might be relevant in the temperate zone as well, as it is not expected that water status would limit biological activity in the wet spring months of the temperate zone (Gougherty & Gougherty, 2018).

In contrast, studies testing multiple hypotheses have generally found support for more than one evolutionary driver of hysteranthy. One study by Bolmgren *et al.* (2003)

showed that wind pollinated species tend to also be earlier flowering than their biotocially pollinated sister taxa, suggesting a relationship between the early flowering and wind pollination hypotheses. A recent study by Gougherty & Gougherty (2018) tested multiple hypotheses by modeling associations between species' trait and FLS patterns in the Great Lakes regions. They found strong support for both the water dynamics and early flowering (flower timing and seed characteristics) hypotheses, and found strong phylogenetic clustering for FLS.

In all of these cases, variability in FLS below the species level was not addressed. Yet, there are datasets widely available that would allow for testing these several hysteranthy hypotheses concurrently, and at multiple taxonomic levels. To address this gap, we supplement our literature review by re-testing some previously-used datasets to examine all hypotheses, and we leverage several widely-available datasets to test how support for these hypotheses varies across the inter- to intra-specific levels.

We evaluated hysteranthy in four phenological datasets. Michigan Trees and its companion volume Michigan Shrubs and Vines (Barnes & Wagner, 1981,2004; Burton V. Barnes, Christopher W. Dick, 2016) (MTSV) contains categorical FLS information for 195 woody plant species. The USFS Silvics manual volume II (Burns et al., 1990) contains categorical FLS descriptions for 81 woody species. These data can be used to test inter-specific FLS variation. Within these datasets, we applied two alternative FLS classification schemes; physiological hysteranthy, which allowed for no overlap between floral and leaf phenophases, and functional hysteranthy, which allowed for a degree of overlap as predicted by the wind pollination hypotheses. The Harvard Forest dataset (HF) contains quantitative flowering and leaf phenology measurements for individuals of 24 woody species over a 15 year period, allowing for both inter- and intra-specific comparisons (O'Keefe, 2015). In this data set, we approximated the two hysteranthy classification schemes mentioned above by measuring the temporal offset between different floral and leaf phenophases. From the Pan European Phenological Database (PEP725) (Templ et al., 2018) we obtained spatially and temporally explicit, quantitative flowering and leaf phenology for four common European tree species. This allows for the evaluation of FLS only at the intra-specific level, but unlike the other datasets, it allows for population level variability to be assessed.

In considering each species-level data set separately and in tandem two clear trend emerge: One, in accordance with the recent literature, we found that multiple hypotheses were supported. There was generally a strong support for the early flowering and wind pollination hypotheses, poor support for the water dynamics hypothesis, and the phylogenetic signal was variable figure 7). The support for multiple hypotheses is not terribly surprising. We wouldn't expect the wind pollination hypothesis to explain hysteranthy in biotically pollinated taxa. Further, given the almost constant non-equilibrium state of temperate forest communities due to glacial cycles over the last 10,000 years (Stephen H. Spurr, 1980), it is not surprising our flora consists of species with radically different bio-geographic histories that may have evolved hysteranthous flower under very different

selection environments. The second clear signature from our analysis was that relative importance of each the predictors changed significantly depending on how hysteranthy was defined. Generally, this artifact was minimized when continuous measure of FLS were used over categorical.

When considering our intra-specific datasets, contrary to what we predicted given the water dynamics hypothesis, intra-annual variation in water available (drought vs non drought years) does not increase hysteranthy. Rather, it seems that drought years correlate with a decrease in FLS offset in hysteranthous species, largely due to delayed flowering. This observation limits our expectation for the water dynamics hypothesis, but does not eliminate the possibility that intra-specific differences in FLS offset are adaptive for populations growing in regions with sustained differences in water availability. If fact, when we examined the relationship between 30 year soil moisture records and population level variation in FLS offset across Germany, we did find a weak association between lower average soil moisture levels and increased hysteranthous offset as predicted by the water dynamics hypothesis. However, when we incorporate other predictors such as flowering time into our analysis, the negative association between soil moisture and increasing hysteranthous offset disappeared (see figure 6). Our intraspecific models suggested that variability in hysteranthous offset is much more tightly correlated with variability in flowering time vs. leafing time (figure 8). However, this contrast was far less stark in seranthous Aesculus hippocastum. These patterns match expectations from recent physiological work by Savage (2019) who found that flower phenology was more less constrained than leaf phenology in for a suite of hysteranthous and synathous woody plants. Though our intraspecific data set is species limited, we see that plasticity in the first phenophase of the season (flowering for hysteranthous species and leafing for seranthous species) seems to drive variability in in FLS offset, but this observation should be tested more rigorously and explicitly in future work. While the inter- and intra-specific case studies are not perfectly comparable (ie pollination hypothesis can be evaluated intra-specifically), the inference from the intra-specific study supports the relationships found in the inter-specific case studies, and provided novel insights of its own. But perhaps more important that the results of all of these specific case studies themselves, is that through considering them together, we are provided a more comprehensive picture of where our understanding of this phenological trait is currently, and where it needs to go.

Future

Our interrogation of the nature of flower-leaf phenological sequences leads to realization that it is instructive to test questions of hysteranthy at many scales. Because trait modeling in large community level datasets seem to support multiple hypotheses and are confounded by species' identities and observer bias, the utility of these data has its limits. While there is certainly value to broad taxonomic studies, and future large-scale analyses should continue, it is possible the evolutionary dynamics of hysteranthy may be better

explored with a more mechanistic approach, which may mean utilizing a more taxonomically restricted focus.

One option is to look within the hypotheses to address sub-grouping of taxa in which overlap between hypotheses could be controlled. For example, what drives hysteranthy among biotically pollinated taxa? It certain isn't wind pollination efficiency. Or, what factors accounts for variability in hysteranthy among wind pollinated taxa? Incorporating a more explicit phylo-biogeographic approach would be instructive at this level, for example: are their phylogeopraphic commonalities between biotically pollinated hysteranthous species in Eastern flora?

But even with drilling down to sub-groupings, interspecific trait association models can only can take us so far. One reality of these kind of studies is that we never know we are picking the right traits. For example we used minimum precipitation across a species' range, one of the only available quantitative drought metrics at the scale of large interspecific models, to represent the water dynamics hypothesis. Is this really a good proxy for drought tolerance? Further, species evolve a suit of traits for any function, and unmeasured traits might bias our results (Davies et al., 2019). For example, wind pollinated species could compensate for a pollen intercepted by a synanthous or seranthous FLS by over producing pollen or through self-pollination. To really understand FLS across large taxonomic space, one would have to compare species across an unfeasibly large, N-dimensional trait space.

Considering variation in hysteranthy at the intra-specific level overcomes many of these limitations, and is the next frontier in testing the evolutionary and ecological significance of FLS. Evolutionary theory predicts that intra-specific variation should follow the same trends as interspecific variation. The agreement between our intra- and inter-specific models supports this, and may suggests that we are narrowing in on certain hypotheses. Further, though our datasets were taxonomically and geographically limited, they demonstrate that FLS variability is significant over time and space. Looking within species holds most other traits relatively equal, avoiding the problem of latent tradeoffs with unmeasured traits.

There are also clear advantages of treating hysteranthy as a continuous trait. As mentioned above, continuous data minimizes the observer bias that comes with categorization. It also reveals important inter-specific differences that are masked by categorization. For example, two categorically hysteranthous species may have dramatically different FLS offsets. Through working with continuous measures of hysteranthy, substantial intra-specific differences in FLS emerge, and as will be discussed more below, these will be valuable for hypothesis testing. All and all, our work shows categorizing hysteranthy into groups is biased and biologically problematic; future studies about phenological sequences should avoid these categories and treat FLS as continuous traits whenever possible.

With this equalizing nature of intra-specific comparisons and continuous measurements of FLS we can move beyond trait associations and actually begin to look at fitness consequences of FLS variation through experimental manipulations and observations.

This next step is intuitive because fitness actually drives trait evolution, and the hysteranthy hypotheses themselves make fitness predictions. It is tough to tease these apart at the inter-specific level because of the latent traits mentioned above, but the hypotheses predict that variability in hysteranthy would lead to variability into fitness outcome at the intra-specific level. For example, the wind pollination hypothesis predicts that years with increased hysteranthy should correlate with more pollination success. The water dynamics hypothesis suggests populations with increased hysteranthous offset should better tolerate drought. These predictions could be directly assessed through well designed experiments.

Looking at fitness consequences will not only help clarify basic scientific hypotheses, but is essential for understanding how global change induced alterations to FLS's will impact species demographics. For example, if hysteranthy is driven by pollination efficiency, increased hysteranthy with climate change might favor hysteranthous species. Or, if climate changes reduces FLS offset, hysteranthous species may be at greater risk for reproductive failure. A better understanding of consequences of variation in hysteranthy is essential both for understanding the evolutionary origins of this trait, and for predicting the fate of species with this phenological syndrome as global climate continues to change.

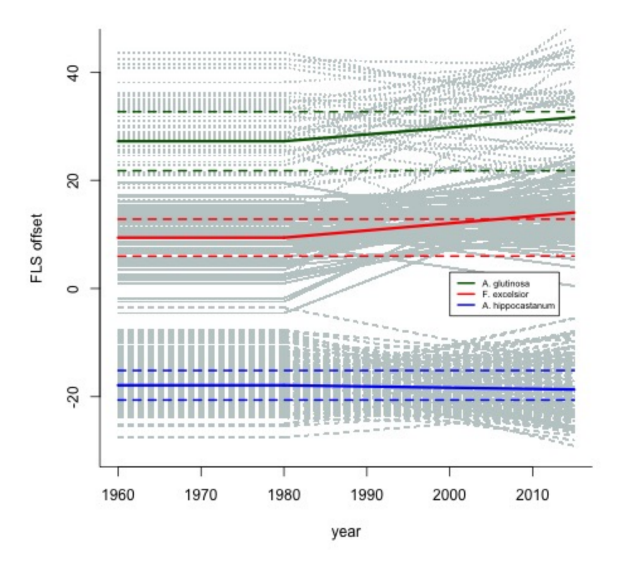


Figure 1: Trends in average FLS offset across Europe for 3 tree species from 1960 to 2015. Dashed lines indicate historic range of FLS variability. All species are increasing their offset, but the rate of change differs between species and and sites

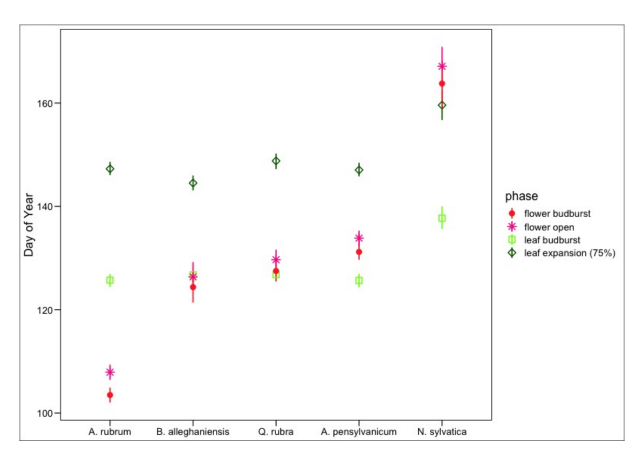


Figure 2: Average day of phenological events for highlighted woody plant species at Harvard Forest in Petersham, MA from 1990-2015

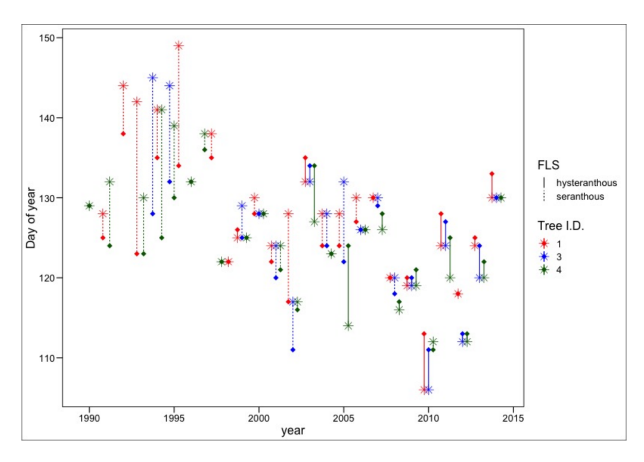


Figure 3: FLS variability among years and within a population of $Quercus\ rubra$ at Harvard forest.

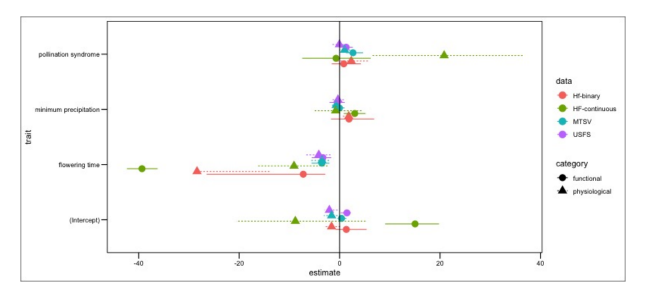


Figure 4: All case study model estimates, phyloglm with 95% bootstrap intervals

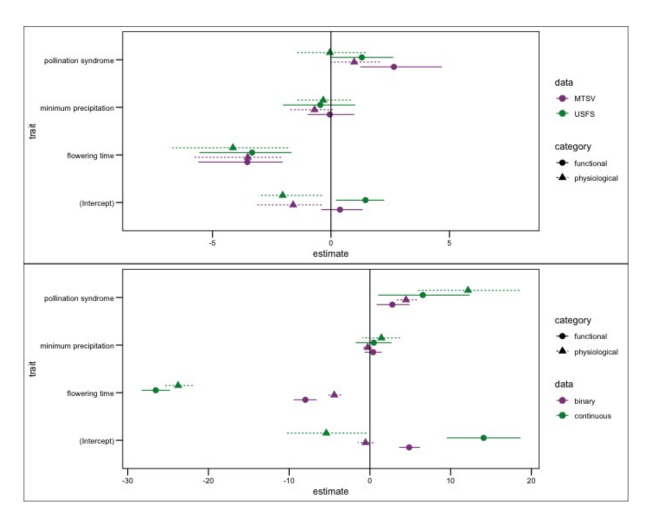


Figure 5: alternatue to figure 4: 4a is phylog glm with 95% bootstrap intervals, 4b is brms estiamtes with 80% CIs

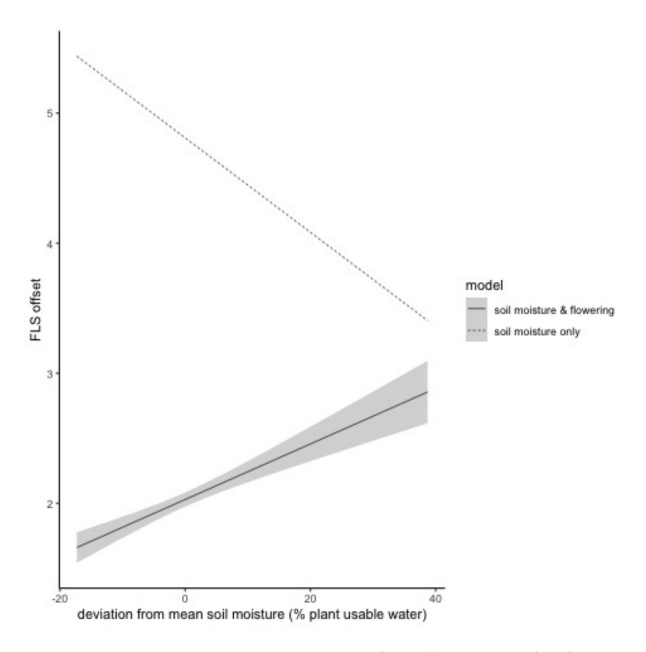


Figure 6: effect of soil moisture on FLS offset with (dashed) and without (solid) considering flowering time

References

(2019) Dwd climate data center (cdc): Multi-annual grids of soil moisture in 5cm depth under grass and sandy loam, version 0.x,. digital.

Barnes, B.V. & Wagner, W.H.J. (1981,2004) Michigan Trees: A guide to the Trees of the Great Lakes Region. University of Michigan Press.

Bolmgren, K., Eriksson, O. & Linder, H.P. (2003) Contrasting flowering phenology and

- species richness in abiotically and biotically pollinated angiosperms. *Evolution* **57**, 2001–2011.
- Borchert, R. (1983) PHENOLOGY AND CONTROL OF FLOWERING IN TROPICAL TREES. *BIOTROPICA* **15**, 81–89.
- Bürkner, P.C. (2018) Advanced bayesian multilevel modeling with the r package brms. R Journal 10, 395–411.
- Burns, R.M., Honkala, B.H. & coordinators], T. (1990) Silvics of north america: Volume 2. hardwoods. Tech. rep., United States Department of Agriculture (USDA), Forest Service.
- Burton V. Barnes, Christopher W. Dick, M.E.G. (2016) Michgan Shrubs Vines: A guide to species of the Great Lakes Region. University of Michigan Press.
- Cleland, E.E., Chuine, I., Menzel, A., Mooney, H.A. & Schwartz, M.D. (2007) Shifting plant phenology in response to global change. *Trends in Ecology Evolution* **22**, 357 365.
- Davies, T.J., Regetz, J., Wolkovich, E.M. & McGill, B.J. (2019) Phylogenetically weighted regression: A method for modelling non-stationarity on evolutionary trees. *Global Ecology and Biogeography* **28**, 275–285.
- Ettinger, A., Gee, S. & M. Wolkovich, E. (2018) Phenological sequences: how early season events define those that follow. *American Journal of Botany* **105**.
- Flynn, D.F.B. & Wolkovich, E.M. (2018) Temperature and photoperiod drive spring phenology across all species in a temperate forest community. *New Phytologist* **219**, 1353–1362.
- Forrest, J. & Miller-Rushing, A.J. (2010) Toward a synthetic understanding of the role of phenology in ecology and evolution. *Philosophical Transactions of the Royal Society B: Biological Sciences* **365**, 3101–3112.
- Franklin, D.C. (2016) Flowering while lea ess in the seasonal tropics need not be cued by leaf drop: evidence from the woody genus brachychiton (malvaceae). *Plant Ecology and Evolution* **149**, 272–279.
- Friedman, J. & Barrett, S.C.H. (2009) Wind of change: new insights on the ecology and evolution of pollination and mating in wind-pollinated plants. *Annals of Botany* **103**, 1515–1527.
- Gellman, A. & Hill, J. (2997) Data Analysis Using Regression and Multilevel/Hierarchical Models. Cambridge University Press.

- Gougherty, A.V. & Gougherty, S.W. (2018) Sequence of flower and leaf emergence in deciduous trees is linked to ecological traits, phylogenetics, and climate. *New Phytologist* **220**, 121–131.
- Heinig, R. (1899) Glossary of the botanic terms used in describing flowering plants. Calcutta, India.
- Ho, L.S.T. & Ane, C. (2014) A linear-time algorithm for gaussian and non-gaussian trait evolution models. *Systematic Biology* **63**, 397–408.
- Ives, A.R. & Garland, Jr., T. (2010) Phylogenetic logistic regression for binary dependent variables. Systematic Biology **59**, 9–26.
- Ivits, E., Horion, S., Fensholt, R. & Cherlet, M. (2013) Drought footprint on european ecosystems between 1999 and 2010 assessed by remotely sensed vegetation phenology and productivity. *Global Change Biology* **20**, 581–593.
- Janzen, D.H. (1967) Synchronization of sexual reproduction of trees within the dry season in central america 21, 620–637.
- Lamont, B.B. & Downes, K.S. (2011) Fire-stimulated flowering among resprouters and geophytes in australia and south africa. *Plant Ecology* **212**, 2111–2125.
- Lechowicz, M.J. (1995) Seasonality of flowering and fruiting in temperate forest trees. Canadian Journal of Botany 73, 175–182.
- Milleron, M., Lopez de Heredia, U., Lorenzo, Z., Perea, R., Dounavi, A., Alonso, J., Gil, L. & Nanos, N. (2012) Effect of canopy closure on pollen dispersal in a wind-pollinated species (Fagus sylvatica L.). PLANT ECOLOGY 213, 1715–1728.
- Nathan, R. & Katul, G. (2005) Foliage shedding in deciduous forests lifts up long-distance seed dispersal by wind. *PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF THE UNITED STATES OF AMERICA* **102**, 8251–8256.
- Niklas, K.J. (1985) The aerodynamics of wind pollination 51, 328–386.
- O'Keefe, J. (2015) Phenology of woody species at harvard forest since 1990.
- Ollerton, J. & Lack, A. (1992) Flowering phenology: An example of relaxation of natural selection? *Trends in Ecology / Evolution* **7**, 274 276.
- Orme, D., Freckleton, R., Thomas, G., Petzoldt, T., Fritz, S., Isaac, N. & Pearse, W. (2013) caper: Comparative Analyses of Phylogenetics and Evolution in R.
- Piao, S., Friedlingstein, P., Ciais, P., Viovy, N. & Demarty, J. (2007) Growing season extension and its impact on terrestrial carbon cycle in the northern hemisphere over the past 2 decades. *Global Biogeochemical Cycles* **21**.

- Primack, R.B. (1987) Relationships among flowers, fruits, and seeds. *Annual Review of Ecology and Systematics* **18**, 409–430.
- Rathcke, B. & Lacey, E.P. (1985) Phenological patterns of terrestrial plants. *Annual Review of Ecology and Systematics* **16**, 179–214.
- Reich, P. & Borchert, R. (1984) WATER-STRESS AND TREE PHENOLOGY IN A TROPICAL DRY FOREST IN THE LOWLANDS OF COSTA-RICA. *JOURNAL OF ECOLOGY* **72**, 61–74.
- Robertson, C. (1895) The philosophy of flower seasons, and the phaenological relations of the entomorphilous flora and the anthophilous insect fauna 29, 97–117.
- Savage, J.A. (2019) A temporal shift in resource allocation facilitates flowering before leaf out and spring vessel maturation in precocious species. *AMERICAN JOURNAL OF BOTANY* **106**, 113–122.
- Stephen H. Spurr, B.V.B. (1980) Forest Ecology. John Wiley Sons, 3rd edn.
- Templ, B., Koch, E., K.Bolmgren, Ungersböck, M., Paul, A., Scheifinger, H. & et al. (2018) Pan european phenological database (pep725): a single point of access for european data. *Int. J. Biometeorology*.
- Whitehead, D.R. (1969) Wind pollination in the angiosperms: Evolutionary and environmental considerations 23, 28–35.
- Wilczek, A.M., Burghardt, L.T., Cobb, A.R., Cooper, M.D., Welch, S.M. & Schmitt, J. (2010) Genetic and physiological bases for phenological responses to current and predicted climates. *Philosophical transactions of the Royal Society of London. Series* B, Biological sciences 365, 3129–3147.
- Zanne, A.E., Tank, D.C., Cornwell, W.K., Eastman, J.M., Smith, S.A., FitzJohn, R.G., McGlinn, D.J., O'Meara, B.C., Moles, A.T., Reich, P.B. & et al. (2013) Three keys to the radiation of angiosperms into freezing environments. *Nature* **506**, 89–92.

Supplement

Methods

Climate Change and FLS:

To evaluate how FLS patterns have overtime in association with climate change we obtained phenological data for three European woody plant species with long term records of both flower (BBCH 60) and leafout phenology (BBCH 11) from the Pan European

Phenological Database (Templ *et al.*, 2018). We restricted the data set to include only stations with great than 50 years worth of data. For each species, we modeled FLS offset (day of year leafing- day of year flowering) as a function of time, using a hinge model with 1980 as break point in accordance with climate change models of ?). For each species, we displayed the pre-1980 mean and standard devation of FLS offset and the post-1980 change in mean FLS offset that can be associated with climate change.

Case studies

MTSV and USFS: For these two, categorical, species level case studies, we converted verbal descriptions of flower-leaf sequences into a binary response variable. For our more inclusive "functional" definition of hysteranthy, we included species entries with descriptions "flowers before the leaves", "flowers before or with leaves" and textit flowers with leaves" as hysteranthous. Our more restrictive "physiological" hysteranthy definition only included species described as "flowers before the leaves" as hysteranthous.

For modeling trait associates we chose three predictors to represent the three major FLS hypothesis; pollination syndrome, average flowering time and minimum precipitation levels across the species range. Pollination syndrome and average flowering time we obtained directly from the data sources, and estimates of minimum precipitation came from the USDA/NRCS Conservation plants characteristics (). We coded pollination syndrome as binary, biotic or wind pollinated, with known ambophilous species in the genus *Salix* assigned to the ancestral, biotic pollinated, state of angiosperms. Flowering time was the average of the range of months reported in each data source.

HF: For each species in the HF data set, we calculated a continuous mean off FLS "offset" defined here as the average foliate day of the year - average floral day of the year. We approximated our "physiological" FLS characterization be defining offset as (day of leaf budburst-day of flower bud burst) and our "functional" FLS categorization by defining offset as (day of leaf expansion to 75% of final size- day of first flower open). We also recoded the HF continuous offset variables as continuous with positive offset values coded as hysteranthous and negative values as seranthous.

For all species-level case studies (USFS MTSV and HF), associations between hysteranthy and the trait predictors were modeled with logistical regressions in phylogenetic generalized linear modeling framework (Ives & Garland, 2010) using the R package phylolm (Ho & Ane, 2014). Our models incorporated a published angiosperm phylogenetic tree (Zanne et al., 2013) pruned to match the species list for each case study. Species found in the trait data set but not in the original phylogenetic tree were added to the pruned tree at the genus level root. In total 32 species were added to the generic roots for the MTSV data set . We The models were run with 599 bootstrapped re-sampling iterations for each data set (?). Continuous predictors were re-scaled by subtracting the mean and dividing by two standard deviations to allow for a reasonable comparison of effect sizes between the binary and continuous predictors in this model (Gellman & Hill,

2997). To assess the phylogenetic structure of hysteranthous flowering, we used the Caper package (Orme *et al.*, 2013) to calculate a phylogenetic D statistic.

PEP 725: For intra-specific analysis, we utilized phenological records from PEP725 stations in Germany with more than 10 years worth of flowering and leafout records (Templ et al., 2018) for species Alnus glutinosa, Fraxinus excellsior and Betula pendula. To test for associations between FLS variability and inter-annual water availability we modeled the association between FLS and drought years from 2003-2010 using a linear mixed modeling framework with the R package lme4 (?). Drought years were determined based on Ivits et al. (2013). To test associations for population level variation in FLS and long term soil moisture, we obtained average August soil moisture raster grids 1991-2010 for Germany from the German Weather Service (DWD, 2019), and extracted soil moisture values at every cell. We then tested associations between average soil moisture at each PEP725 phenological station and average FLS for species Aesculus glabra, Alnus glutinosa, Fraxinus excellsior and Betula pendula using a Bayesian linear mixed model framework with the brms package in R (Bürkner, 2018). We also repeated the analysis with average April soil moisture data from the same time period and results were robust.

Using same PEP725 species records as above, we used linear models to test the relationship between flowering and leaf timing and FLS offset.

Data set	Physiological	Functional
MTSV	0.22	0.07
USFS	0.13	0.65
HF	0.01	0.27

Figure 7: Phylo.D estimates of phylogentic signal for hysteranthous flowering in three case studies

Species	Effect size flowering (sd)	\mathbb{R}^2	Effect size leafing (sd)	\mathbb{R}^2
Alnus glutinosa	-0.67 (0.002314)	0.58	0.24(0.005)	0.03
Fraxinus excellsior	-0.53 (0.002)	0.41	0.24 (0.004)	0.05
Aesculus hippocastum	-0.28 (0.002)	0.10	0.411 (0.0019)	0.26

Figure 8: Flowering vs leafing influence on hysteranthy

Phylogenies

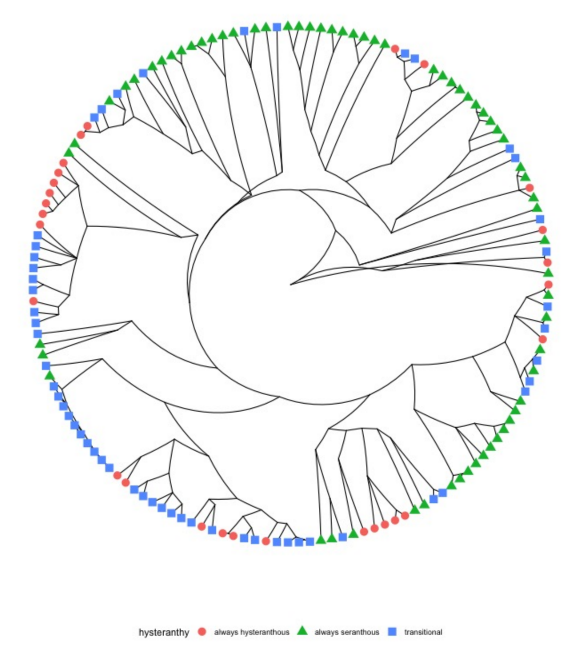


Figure 9: MTSV phylogeny

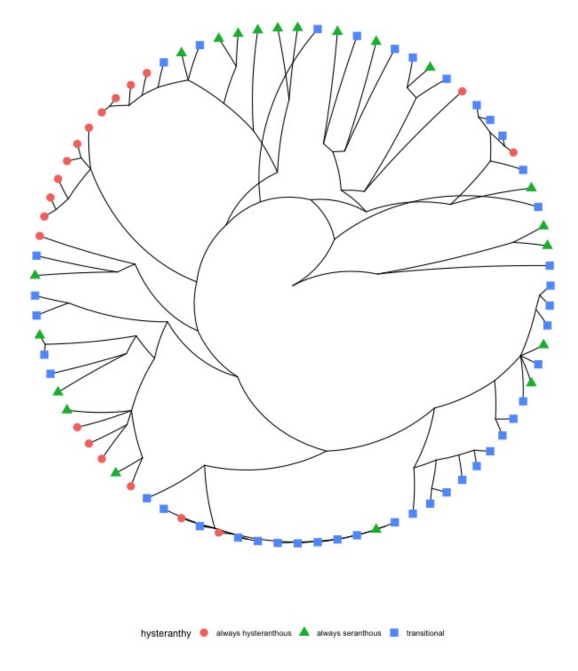


Figure 10: USFS Phylogeny

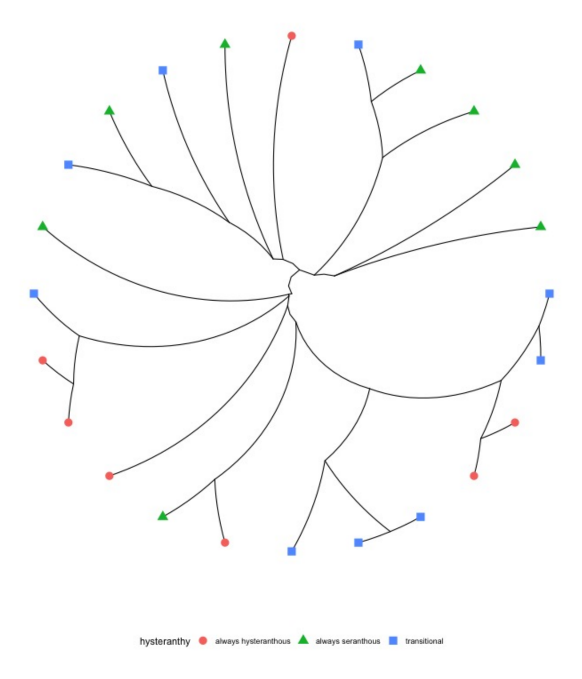


Figure 11: HF Phylogeny