Thermo- vs. Photo-periodicity

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March 2019

1. Introduction

- (a) Phenology is important
- (b) Experiments show phenology cued by temperature and photoperiod.
- (c) Interactions are important and need to be pursued.
- (d) Testing interactions is complex; we will consider photoperiod and forcing
- (e) Must be orthogonal

2. Experimental Designs for interactions

- (a) Each cue has possible 2 axes, intensity and period.
 - i. Photoperiod: usually concerned with period only
 - ii. Forcing, both period and intensity (temperature)
- (b) Three major experiment designs could be utilized.
 - i. Temp intensity and photo period
 - A. Pro: Simple, easy to make truly orthogonal
 - B. Con: Some literature says this is bad for temperature. (find it). As such is rarely done.
 - ii. Covary thermo- and photo-period (most common).
 - A. Pro: more accurate representation of forcing. Simulates nature so good for comparison to field.
 - B. Con: Non-orthogonal for GDH. Can determine between photo-period and thermo-period effects.
 - iii. Decouple thermo- and photo-period (never really done).
 - A. Pro: Orthogonal for GDH.
 - B. Con: New Non-orthogonal issue. Dawn T. (Find citations)
- (c) Design II will probably continue to be most popular. And we can and should use math to more accurately interpret our results.

3. Mathematical Predictions

(a) Explain Data

- (b) Assumptions
 - i. Forcing is most important
 - ii. Others
- (c) Show the math
- (d) make predictions
- 4. Experimental Validation
 - (a) Explain experiment two.
 - (b) Present comparison results

Introduction

Phenology, the timing of life cycle transitions in organisms, is important component of ecosystem structure and function (Chuine & Beaubien, 2001; Piao et al., 2007; Cleland et al., 2007). While phenology has been of interest to biologists for over a century, it has begun to receive increased attention in the recent decades as observed phenological shifts across a broad taxonomic spectrum have emerged as one of the most widely observed effects of anthropogenic climate change to date (Parmesan & Yohe, 2003; Root et al., 2003; Menzel et al., 2006; Wolkovich & Ettinger, 2014).

It has long been recognized that experimental manipulations in artificial environments are ideal for mechanistically characterizing phenological response to environmental fluctuations (Ettinger et al.; Primack et al., 2015). This body of experimental work has demonstrated that temperature (winter chilling and spring forcing) and photoperiod are the primary cues of phenology for plants in the temperate/boreal zones (Rathcke & Lacey, 1985; Visser et al., 2010; Forrest & Miller-Rushing, 2010).

Recent advances have demonstrated that plant phenological responses are nonlinear, due largely to interactions between cues(Flynn & Wolkovich, 2018; Laube et al., 2014). This highlights the need for experiments to be designed to evaluate the strength of these interactions. To truly test interactions, treatments must be orthogonal: that is there must be more than one level of each treatment, and each level of each treatment must be crossed with every level of the other. This adds complexity to experimental setups, and experimental artifacts that interfere with interpretation of results are readily introduced (Wolkovich et al., 2012). In this paper, we discuss a particular challenge that arises when investigation interaction between forcing temperate and photoperiod.

Considering light and temperature in phenology experiments

For both light and temperature cues, there are two axes that can be manipulated in experiments. Then intensity of the cue (ie temperature of forcing or light intensity/wavelength) and the period (duration of temperature exposure or light periods). While light intensity has

been to shown to affect many aspects of organism's biological activity including phenology (Brelsford & Robson, 2018; Cober et al., 1996), the period of light exposure is generally considered to be the dominate light cue, and phenological research interests tends to focus on photoperiod rather than intensity (FindCitation). For temperature, though intensity is considered a major driver of phenology, the very common and successful model framework for quantifying temperature effects on phenology, the growing degree day or growing degree hour, integrates temperature with its period. Researchers must be thoughtful about how to incorporate these three main treatment axes(photo-period, theromo-period and thermo-intensity) in their experiments, and be wary of how these decisions will affect their experimental inference. There are three basic experimental designs (see figure 1 which are detailed in brief below:

Manipulation of Temperature intensity and photoperiod:

The most simple design to test for a temperature x light interaction is to manipulate only one axis of each cue, for example, temperature intensity and light period. This would be implemented by applying high and low forcing treatments at constant temperatures, and crossing these with a long and short photo-period treatment at a constant light intensity (figure 1a).

The main advantage of this design is that is it simple to implement, maintains the orthogonality among the four treatment combinations (warm/short, warm/long, cool/short, cool/long) and allows for a very straight forward interpretation of the predictors. However, in nature, plants experience substantial diurnal temperature variation, and growth chambers experiments without a diurnal thermoperiod would be a poor approximation of natural conditions. Several studies indicated that species may infact be responding to the differences between day and night temperatures (Erwin & Heins, 1995), which raises futher questions about the utility experiments lacking thermoperiodicty for understanding the ecology and evolution of play in nature. On account of these shortcomings, it has become quite standard for researchers to include a diurnally varying thermo-period for forcing treatments. For example, rather than static high/low forcing treatments of 20°C and 16°C, researchers would use, a high forcing treatment of 24°/16°C and low 20°/12°C (day/night). (Ettinger et al.).

Coupled photo- and thermo-periodicity:

Incorporating diurnal thermo-periodicity into experimental design require that researcher decide how to vary this periodicity relative to the experiments photo-period cycle. The vast majority of published experiments employ a design in which photo-and thermo-periodicity are synchronized as in (figure 1a) (Ettinger et al.). This design is perhaps the most intuitive, one, following the fact the photo- and thermo-period tend to co-vary in nature (FindCitation). This approximation of natural conditions may make this design the most useful for extrapolating inferences between experimental and observational studies. However, while it appears, to maintain orthogonality between treatments when considering temperature intensity alone, but, in this case, the forcing response is a product of both the intensity of the temperature cue and the duration of its exposure (growing degree hours). Thus, the coupling of photo- and thermo-period results in non-orthogonality between the four treatment

combinations (see figure 2a). In this scenario, the effect of photo-period cannot be neatly distinguished from the effects of increased thermo-period under the longer day treatments, making it impossible to accurately evaluate the relative importance of these cues, which is often a a major goal of growth chamber experiments. This results in spurious interpretation of experimental result.

Decoupled photo- and thermo-periodicity:

An alternative experimental design that incorporates thermo-periodicity into a temperature x photoperiod interaction experiment is decouples the experimental thermo- and photoperiods. The would be achevied by varying thermo-period on a constant schedule across all four treatment combinations (figure 1c). Though this experiment set up has not be widely, if ever, employed in growth chamber experiments (Ettinger et al.), this design restores full orthogonality to growing degree hour sums that make up the forcing treatment (figure 2a).

But this obvious advantage for reasonible statistical comparisons introduces a new biological artifact that may bias experimental results. Evidence from horticulture studies have demonstrated that cell growth is most sensitive to temperature fluctuation at the beginning of the photoperiod(Erwin, 1998). This suggests that dawn temperatures may be disproportionally strong driver biological activity in plants. For example, it has been shown that that increasing temperatures in the first two hours of the photoperiod was almost as effective for stimulating shoot elongation as similar temperature increases for the whole photoperiod (Erwin, 1998). By de-coupling thermo-period from the photo-period manipulations, experimenters by neccesity introduce an assymetry between the photoperiod treatments relative to the thermoperiod. In our example in 1c), the long and short day photoperiod treatments have the same 12 thermoperiodicity cycle, but in the long day treatments, the plant does not experience "day time" temperatures until after they encounter "day light", while the plants in the short day treatment experience day time warming before light.

Given the demonstrated importance of the interaction between temperature and photoperiod at dawn, it could be suggested that a design that the one depicted in figure 1d in and improved version of a uncouple thermo and photo period design, because the temporal relationship between dawn light and temperature is standardized across all treatments, but this again produces an unreasilstic comparision with nature.

As we have shown above, each design introduces its own biological or statistical artifacts that will bias he results of the experiments. We cannot conclusively solve the problems of true inference through modifying experimental designs, but rather, we must thoughtful incorporate the implications of these design choices into our statistical metrics and interpretation of results. While we cannot conclusively suggest a optimal experimental design, we expect that because of its history and relationship to field conditions, a coupled photo-and thermo- period design will remain the most popular choice for phenological experiments in artificial environments. In the following section, we will use a public dataset to demonstrate how mathematical principles can be applied to tease apart the artifact introduced by this inherently imperfect experimental design and make more meaningful inferences about the cue interactions.

Math

Our analysis is based on results from a large scale growth chamber experiment by Flynn & Wolkovich (2018), in which in addition to chilling and provenance, fully factorial light and temperature manipulations were applied to twig cuttings from 28 woody species. The authors determined a mean budburst sensitive to temperature of -9.5, and a mean sensitivity to photo-period of -4.5 with a weak, negative interaction between the two cues. We set out to calculate how much of the reported photo-period response could in actuality be driven by the latent differences in thermo-period between the long and short photo-period treatments.

- 1. State some assumptions. IE forcing is dominant cue
- 2. Do some math
- 3. Make a prediction.

1 Validation

- 1. How do we know if this is a good prediction? We are essentially predicting the expected difference between experimental design two and three.
- 2. We have 2 dataset. A subset of a Flynn & Wolkovich (2018) study were sampled at Harvard Forest, were forced at 18 day 12 night, with photoperiod treatments of 8 and 12 hours.
- 3. Another experiment by Buonaiuto and Wolkovich (unpublished data) also included these treatment levels, But, decoupled thermo-and photo-periodicity. What a happy accident.
- 4. We reanalyzed these datasets to compare the effects of coupling vs. uncoupling, see 2.

2 Wrap up

- 1. We can't say which design is absolutely best.
- 2. We expect design 2 will remain popular. This is okay because we have shown the design flaws are easiest to overcome mathematically.
- 3. We externally validated the math
- 4. Future studies should do this.

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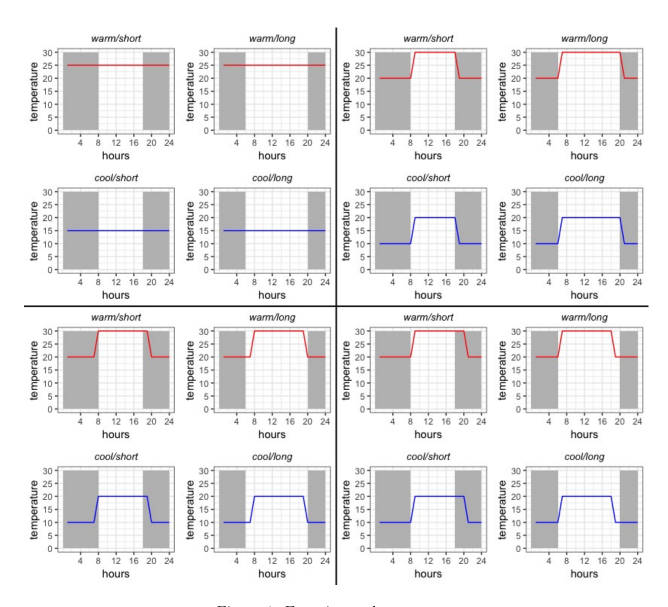


Figure 1: Experimental treatments

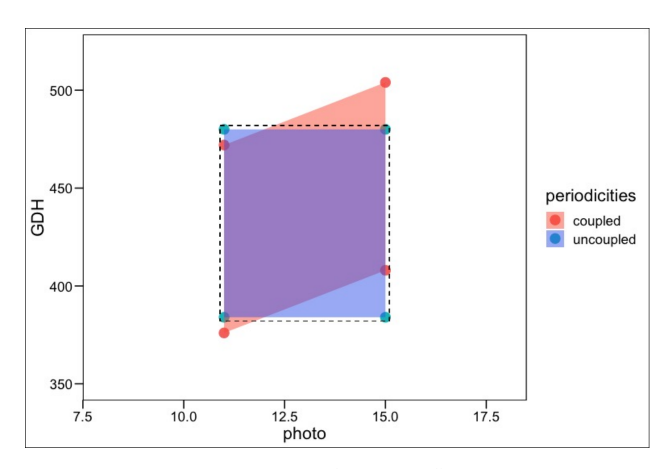


Figure 2: Experimental comparison all species

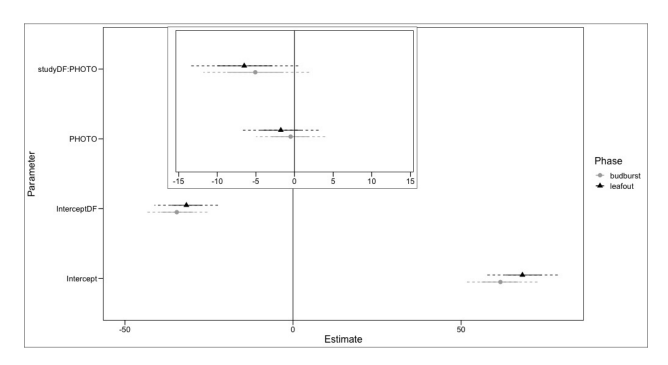


Figure 3: Matching species only