- Comparison of large-scale citizen science data and
- long-term study data for phenology modeling
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Abstract

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Large-scale observational data from citizen science efforts are becoming increasingly common in ecology, and researchers often choose between these and data from intensive local-scale studies for their analyses. This choice has potential trade-offs related to spatial scale, observer variance, and inter-annual variability. Here we explored this issue with phenology by comparing models built using data from the large-scale, citizen science USA National Phenology Network (NPNUSA-NPN) effort with models built using data from more intensive studies at Long Term Ecological Research (LTER) sites. We built statistical and process based phenology models for species common to each dataset. From these models we compared parameter estimates, estimates of phenological events, and out-of-sample errors between models derived from both NPN-USA-NPN and LTER data. We found that model parameter estimates for the same species were most similar between the two datasets when using simple models, but parameter estimates varied widely as model complexity increased. Despite this, estimates for the date of phenological events and out-of-sample errors were similar, regardless of the model chosen. Predictions for NPN-USA-NPN data had the lowest error when using models built from the NPN-USA-NPN data, while LTER predictions were best made using LTER-derived models, confirming that models perform best when applied at the same scale they were built. This difference is likely due to variation in phenological requirements within species. Models using the USA-NPN dataset can integrate parameters over a large spatial scale while those using an LTER dataset can only estimate parameters for a single location. Accordingly, the choice of dataset depends on the research question. Inferences about species-specific phenological requirements are best made with LTER data, and if NPN-USA-NPN or similar data are all that is available, then analyses should be limited to simple models. Large-scale predictive modeling is best done with the larger-scale NPN-USA-NPN data, which has high spatial representation and a large regional species pool. LTER datasets, on the other hand, have high site
fidelity and thus characterize inter-annual variability extremely well. Future research aimed at
forecasting phenology events for particular species over larger scales should develop models
which integrate the strengths of both datasets.

Keywords: LTER, USA-NPN, forecasting, budburst, flowering, data integration, scale

12 Introduction

Plant phenology, the timing of recurring biological events such as flowering, plays an important role in ecological research extending from local to global scales (Cleland et al., 2007; Richardson et al., 2013; Tang et al., 2016). At large scales , uncertainty in the timing of spring leaf out and fall senescence influence the carbon budget of earth system models, which has implications for correctly accounting for biosphere-atmosphere feedbacks in long-term climate forecasts (Richardson et al., 2012). At smaller scales, species-specific responses to temperature and precipitation can alter flower communities (Diez et al., 2012; CaraDonna et al., 2014; Theobald et al., 2017) and affect the abundance and richness of both pollinators (Ogilvie and Forrest, 2017; Ogilvie et al., 2017) and organisms at higher trophic levels (Tylianakis et al., 2008). Plant phenology models that 51 are robust at multiple ecological scales, or deemed appropriate for a particular scale, are needed to 52 better understand and forecast the timing of key biological events. Many plant phenology studies use intensively collected datasets from a single location over a long time-period by a single research group (Cook et al., 2012; Wolkovich et al., 2012; Iler et al., 2013; Roberts et al., 2015). These datasets have regular sampling and large numbers of samples over long periods of time. As a result, the biological and climatic variability at that site is well rep-57 resented. It is common for phenology models built with observations from a single site to not 58 transfer well to other sites (García-Mozo et al., 2008; Xu and Chen, 2013; Olsson and Jönsson, 2014; Basler, 2016). This lack of transferability can be driven by plasticity in phenology requirements, local adaptation, microclimates, or differences in plant age or population density (Kramer, 1995; Diez et al., 2012). For these reasons, data from a single location is are not adequate for larger scale phenology modeling. Accurately forecasting phenology at larger scales will require models that account for the full range of variation across a species' range (Richardson et al., 2013; Tang et al., 2016; Chuine and Régnière, 2017), which will necessitate the use of data sources beyond traditional single-site studies.

Data from citizen science projects is are becoming increasingly important for ecological research (Kelling et al., 2009; Dickinson et al., 2010; Tulloch et al., 2013). Because this data is these data are often collected by large numbers of volunteers, it is possible to gather data at much larger scales than with individual research teams. A relatively new citizen science project started in 2009, The USA National Phenology Network (NPNUSA-NPN), collects phenology observations from volunteers throughout North America the United States and makes the data openly available (Schwartz et al., 2012). Data from this project has have already been used to study variation in oak phenology at a continental scale (Gerst et al., 2017), develop large-scale community phenology models (Melaas et al., 2016), and forecast long-term phenology trends (Jeong et al., 2013). Large-scale datasets from China and Europe have already contributed considerably to phenological research (Xu and Chen, 2013; Olsson and Jönsson, 2014; Basler, 2016; Zhang et al., 2017), and the NPN-USA-NPN dataset has the potential to meet these needs for North American plant species and communities. However, the features that allow citizen science projects to collect data at large scales can also introduce spatial biases toward cities and easily-accessible ar-80 eas, and variation in sampling effort and observer skill (Dickinson et al., 2010). With hundreds of participants of participants, the potential for variation among observers in their determination of species identification and dating of phenological events is high. While volunteers have been shown to be accurate at distinguishing different leaf and flower stages for plants, (Fuccillo et al., 2015), observations (Fuccillo et al., 2015) and can have high agreement on abundance estimates (Feldman et al., 2018), contributions to USA-NPN are some-

times made sporadically across seasons, years, and locations. This means that the quanity quantity

and quality of data at a specific site will typically be more variable for citizen science efforts than

for intensive, long-term studies.

In order to accurately model and forecast phenology, it is important to understand how the strengths

and weaknesses of intensive local studies and large-scale citizen science projects influence both our

inferences about biological processes driving phenology (eg. warming requirements for a specific

plant) and our ability to predict future phenology events (eg. forecasting when flowering or leaf

out occurs). Here, we fit a suite of plant phenology models for the budburst and first-flowering

₉₅ phenophases of 24 plant species to data from both the NPN-USA-NPN and a set of intensive long-

term studies from the Long Term Ecological Research (LTER) network. We compare the result-

97 ing models based on both inference about models and parameters and predictions for unobserved

events. We then use this comparison to assess the best methods for both local- and large-scale

99 phenology modeling and to point the way forward for integrating large-scale and local-scale data

to determine the best possible models across scales.

Methods

Datasets

The USA National Phenology Network (USA-NPN) protocol uses status-based monitoring, where

via a phone app or web based interface observers answer 'yes,' 'no,' or 'unsure' when asked if an

individual plant has a specific phenophase present (Denny et al., 2014). Phenophases refer to specific phases in the annual cycle of a plant, such as the presence of emerging leaves, flowers, fruit, or senescing leaves. Sites in the NPN USA-NPN datasets are located across the U.S. and generally 107 clustered around populated areas (Fig. 1). To represent long-term, intensive phenology studies we 108 used four datasets from North America representing three major ecosystem types (Table 1, Fig. 1). 109 All four long-term studies are located in the U.S. and are part of the Long Term Ecological Re-110 search network (LTER). The Harvard Forest and Hubbard Brook Long Term Experimental Forest 111 are located in the northeastern U.S. and are dominated by deciduous broadleaf species. The H.J. 112 Andrews Experimental Forest is a coniferous forest in the coastal range of the western U.S. The 113 Jornada Experimental Range is in the Chihuahua desert of the southwestern U.S.

We downloaded all NPN-USA-NPN observations from 2009, when collections began, to 2016 for 115 the following phenophases: Breaking Leaf Buds, Breaking Needle Buds, Emerging Needles, and 116 Open Flowers (USA National Phenology Network, 2017). The first three phenophases apply to the 117 'leaf out' phase for deciduous broadleafs, evergreen conifers, and pines, respectively. The 'Open 118 Flowers' phenophase refers to fully-open flowers and applies to all angiosperms. Hereafter, we 119 will refer to these as either 'Flowers' for the Open Flower phenophase, or 'Budburst' for all other 120 phenophases. We subset the NPN USA-NPN observations similar to methods outlined in Crimmins 121 et al. (2017). First, 'yes' observations for individual plants were kept only if they were preceded by a 'no' observation within 30 days. Observations for 'Budburst' that were past day of year (DOY) 123 172, and for 'Flowers' that were past DOY 213 were dropped to minimize any influence from outliers. We inferred the observed DOY of each phenophase as the midpoint between each 'yes' observation and the preceding 'no' observation. Finally, only species that had greater than 30 total observations were kept. Crimmins et al. (2017) only kept observations that were preceded by a 'no' within 15 days, and also grouped multiple individuals at single sites to a single observation. We used 30 days to allow for a greater number of species to be compared. We tested the sensitivity of this choice by also performing the analysis using a 15 day cutoff. We chose not to group multiple individuals at a single site to better incorporate intra-site variability.

In the LTER datasets observation metrics varied widely due to different protocols. To match the 132 NPN-USA-NPN data we converted all metrics to binary 'yes' and 'no' observations for each 133 phenophase (see supplementary methods). As with the NPN-USA-NPN data, we inferred the 134 date for each phenophase as the midpoint between the first 'yes' observation and most recent 'no' 135 observation, and only kept species and phenophases phenophase combinations which had at least 136 30 total observations. After data processing there were 38 species and phenophase combinations 137 (with 24 unique species) common to both the NPN-USA-NPN and LTER datasets to use in the 138 analysis (Table 1 & S1). Using a 15 day cutoff in the USA-NPN dataset resulted in 35 unique 139 combinations with 23 species.

141 Models

It is common to fit multiple plant phenology models to find the one that best represents a specific species and phenophase (Chuine et al., 2013). For each of the 38 species and phenophase combinations in the five datasets (NPN-USA-NPN and four LTER datasets), we fit eight phenology models (Table 2). The *Naive model* uses the mean DOY from prior observations as the estimated DOY.

The *Linear model* uses a regression with the mean spring (Jan. 1 - March 31) temperature as the independent variable and DOY as the response variable. For the six remaining models, the general

form is based on the idea that a phenological event will occur once sufficient thermal forcing units, F^* , accumulate from a particular start day of the year (t_1) . Forcing units are a transformation of the daily mean temperature and are calculated differently for each model (Table 2). The start day can either be estimated or fixed. For the Growing Degree Day (GDD) model, forcing units are the total 151 degrees above the threshold T_{base} (Réaumur, 1735; Wang, 1960; Hunter and Lechowicz, 1992). 152 The Fixed GDD model uses the same form but has fixed values for start day $(t_1 = \text{Jan } 1)$ and 153 temperature threshold (T- $T_{base} = 0$ °C). The Alternating model has a variable number of required 154 forcing units defined as a function of the total number of days below 0°C since Jan. 1 (number 155 of chill days - NCD). The Uniforc model is like the GDD model but with the forcing units trans-156 formed via a sigmoid function (Chuine, 2000). These models are some of the most commonly used 157 in phenology research and serve as a suitable baseline for comparing the NPN and LTER datasets. 158

We also fit two models that attempt to capture spatial variation in phenological requirements. The

first spatial model, MI, is an extension of the GDD model which adds a correction in the required

forcing using the photoperiod (L) (Blümel and Chmielewski, 2012). The second, the *Macroscale Species-specific Budburst model* (MSB), uses the mean spring temperature as a linear correction

on the total forcing required in the *Alternating model* (Jeong et al., 2013). Since there is little

to no spatial variation in the LTER datasets, we fit the two spatial models to data from the NPN

USA-NPN only. We compared the resulting parameters, estimates, and errors for the NPN-derived

USA-NPN derived MI and MSB models to their non-spatial analogs (the GDD and Alternatingmodels, respectively) for each species and phenophase in the LTER data.

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We extracted corresponding daily mean temperature for all NPN USA-NPN and LTER observa-

tions from the gridded PRISM dataset using the latitude and longitude of the site associated with
each observation (PRISM Climate Group, 2004). We parameterized all models using differential
evolution to minimize the root mean square error (RMSE) of the estimated DOY of the phenological event. Differential evolution is a global optimization algorithm which uses a population of
randomly initialized models to find the set of parameters that minimize the RMSE (Storn and Price,
175 1997). Confidence intervals for parameters were obtained by bootstrapping, in which individual
models were re-fit 250 times using a random sample, with replacement, of the data. We made
predictions by taking the mean DOY estimated from the 250 bootstrapped iterations. A random
subset consisting of 20% of observations from each species and phenophase combination was held
out from model fitting for later evaluation.

180 Analysis

As described above, we fit two sets of models for each species and phenophase: one set of models 181 parameterized using only NPN-USA-NPN data, and one set parameterized using only LTER data 182 (with the exception of the M1 and MSB models, see above). To compare the inferences about 183 process made by the two datasets, we compared the distribution of each parameter between LTER 184 and NPN-derived USA-NPN derived models for each species and phenophase combination. Using 185 the mean value of each bootstrapped parameter, we also calculated the coefficient of determi-186 nation (R^2) between LTER and NPN-derived USA-NPN derived models among the 38 species-187 phenophases. In three cases where a species phenophase combination occurred in two LTER sites (Budburst for Acer saccharum, Betula alleghaniensis, and Fagus grandifolia in the Harvard and Hubbard Brook datasets) they were compared separately to the NPN-USA-NPN data.

Models with different parameter values, and even entirely different structures, can produce similar estimates for the date of phenological events (Basler, 2016). Therefore, to compare the predictions and potential forecasts for models fit to the different datasets, we compared the estimated DOY predicted by the LTER and NPN-USA-NPN derived models for all held out observations. For each of the eight models, we calculated the coefficient of determination (*R*²) between LTER and NPN-derived USA-NPN derived estimates for estimates made at the four LTER sites and across all NPN-USA-NPN sites.

We also directly evaluated model performance using the four combinations of models and observed data: A) LTER-derived models predicting LTER observations, B) NPN-derived USA-NPN derived 199 models predicting LTER observations, C) LTER-derived models predicting NPN-USA-NPN obser-200 vations, and D) NPN-derived models predicting NPN observations. Within each of these scenarios, 201 we calculated the RMSE of the held-out observations for each species, phenophase, and model 202 type.-USA-NPN derived models predicting USA-NPN observations. Using the RMSE values, we 203 calculated two different metrics to compare from held out observations, we compared the perfor-204 mance of LTER and NPN-derived USA-NPN derived models on different data types. The first 205 metric focuses in two different ways. First, we focused on local-scale prediction by comparing 206 the fits calculating the difference in the RMSE of LTER and NPN-derived models on LTER 207 observations: RMSE_A - RMSE_B. The second metric focuses USA-NPN derived models solely 208 with LTER observations. Secondly, to focus on large-scale prediction by comparing the fits of 209 LTER and NPN-derived models on the NPN data: RMSE_C – RMSE_D. These metrics we calculated the difference in RMSE using solely USA-NPN data. These differences were calculated for each of the model types and 38 species-phenophase combinations. Negative values indicate that LTER-

derived models perform better, while positive values indicate that the NPN-derived USA-NPN derived model performed better. We used a t-test to test the difference from zero in these values. In the three cases where the same species and phenophase combination occurred in two LTER sites, we made the LTER-LTER comparison (scenario A) within each site, not across sites, to focus on local scale prediction when LTER data is are available. Absolute RMSE values are provided in the 217 supplement (Fig. \$1-\$3\$5-\$7). 218 We performed all analysis using both the R and Python programming languages (R Core Team, 219 2017; Python Software Foundation, 2018). Primary R packages used in the analysis included 220 dplyr (Wickham et al., 2017), tidyr (Wickham and Henry, 2018), ggplot2 (Wickham, 2016), lubri-221 date (Grolemund and Wickham, 2011), prism (Hart and Bell, 2015), raster (Hijmans, 2017), and 222 sp (Pebesma and Bivand, 2005). Primary Python packages included SciPy (Jones et al., 2001), 223 NumPy (Oliphant, 2006), Pandas (McKinney, 2010), and MPI for Python (Dalcin et al., 2011). 224 Code to fully reproduce this analysis is available on GitHub (https://github.com/sdtaylor/phenology_dataset_study 225 and archived on Zenodo (https://doi.org/10.5281/zenodo.1256705)

227 Results

Throughout the analysis no significant difference was found between a 30 day or 15 day threshold between the first 'yes' and most recent 'no' observation in the USA-NPN dataset. Results presented here reflect the 30 day cutoff, see the supplemtary figures S2-S4 for matching figures using a 15 day cutoff. The best matches between parameter estimates based on NPN-USA-NPN and LTER data were the *Fixed GDD model* ($R^2 = 0.49$) and the *Linear model* ($R^2 = 0.39$ for β_1 and -0.05

for β_2). The parameters for all other models had R^2 values <0 indicating that the relationship was worse than no relationship between the parameters (but with matching mean parameter values across the two sets of models) (Fig. 2). The *Naive model* showed a distinct late bias in mean DOY estimates for phenological events, likely resulting from the LTER datasets being mostly in the northern United States compared to the site locations of the NPN-USA-NPN dataset (Fig. 2). 237 The large outlier for the *Fixed GDD model* is *Larrea tridentata*; this species' flower phenology is 238 largely driven by precipitation, which is not considered in the Fixed GDD model (Beatley, 1974). 239 While the *Fixed GDD* and *Linear* models showed reasonable correspondence between parameter 240 estimates, all parameters for individual species and phenophase combinations had different distri-241 butions between NPN-USA-NPN and LTER-derived models (Fig. 86-87S10-S11). 242

When comparing estimates of phenological events between the two sets of models, many NPN USA-NPN and LTER models produced similar estimates (Fig. 3). The Fixed GDD model had the highest correlation between the two models sets at NPN-USA-NPN sites ($R^2 = 0.82$), while the GDD, M1, and Uniforc models had the highest correlation at LTER sites ($R^2 = 0.51, 0.52$, and 0.51, respectively). Comparing models with spatial corrections to the non-spatial alternatives, the MSB (an extension of the Alternating model with a spatial correction based on mean spring temperature, 248 see Table 2 and Methods) improved the correlation between the two datasets over the Alternating 249 model. The MSB model improved the R^2 from 0.36 to 0.45 at LTER sites, and from -0.23 to -0.15 250 at NPN-USA-NPN sites. The M1 model (an extension of the GDD model with a spatial correction 251 based on day length) improved the correlation over the GDD model only slightly at LTER sites 252 (from 0.51 to 0.52) and did not improve the correlation at NPN-USA-NPN sites. 253

When comparing the prediction accuracy on held-out data, NPN-derived USA-NPN derived mod-

els made more accurate predictions for held-out NPN-USA-NPN observations, and LTER-derived models performed better on held-out LTER observations (all *p* < 0.001, Fig. 4). The *Naive* and *Linear* models had the largest differences between the two model sets, while the *Fixed GDD model* had relatively similar errors when evaluated on both NPN-USA-NPN and LTER held-out observations. Although the *Fixed GDD model* had the highest agreement in accuracy between NPN USA-NPN and LTER-derived models, it was not the best performing model overall. The *GDD* and *Uniforc* models commonly made the best predictions, having the lowest RMSE in 23% and 40% of cases among NPN-derived USA-NPN derived models, and 42% and 32% of cases among LTER-derived models, respectively (Fig. 81 & \$2\$5 & \$6\$).

4 Discussion

Data used to build phenology models typically falls into two categories: intensive long-term data 265 with long time-series at a small number of locations (e.g., LTER data in this study), and large-266 scale data with less intensive sampling at hundreds of locations (e.g., NPN-USA-NPN data) (Table 267 3). This data scenario-a small amount of intensive data and a large amount of less intensive 268 data-is common in many areas of science and makes it necessary to understand how to choose 269 between, or combine, data sources (Hanks et al., 2011). We explored this issue for phenology 270 modeling in relation to making predictions and inferring process from models. For inference we 271 found that models based on different data sources resulted in different parameter estimates for all but the simplest models. For prediction we found that models fit to different data sources tended to make similar predictions, but that models better predicted out-of-sample data from the data type to which they were fit. These results are consistent with other research showing that
phenology model performance decreases when transferring single-site models to other locations
(García-Mozo et al., 2008; Xu and Chen, 2013; Basler, 2016), and with the call for models that
better incorporate spatial variation in phenology requirements (Richardson et al., 2013; Chuine and
Régnière, 2017). Understanding and making predictions for the phenology of a single location is
best served by intensive local-scale data, when available, but large-scale datasets work better for
extrapolating phenology predictions across a species range. Thus, the best choice of both data and
models depends on the desired research goals.

In this study, parameter estimates differed widely within the same phenology model when fit to the two different types of data, except for the simplest process-oriented model: the Fixed GDD 284 (Fig. 2). These differences may be caused by a variety of factors that have different implica-285 tions for interpreting process-oriented models and their parameters. First, the differences could 286 result from limitations in the sampling of the NPN dataset, USA-NPN dataset, such as irregular 287 sampling of the same location within or between seasons, leading to less accurate parameter es-288 timates. If this is the case, it would suggest that using LTER data is ideal for making inferences 289 about plant physiology, and that focusing on the Fixed GDD model is best for making inferences 290 when NPN data is USA-NPN data are all that is available. Second, spatial variation (eg. from 291 local adaptation, acclimation, microclimates, or plant age) in phenology requirements could drive 292 these differences, because NPN data integrates and drivers could contribute to these differences 293 (Diez et al., 2012; Zhang et al., 2017). Model built using USA-NPN data integrate over that spatial 294 variation, while models built using LTER data only estimates estimate the phenological requirements for a specific site. In this case, NPN USA-NPN data would provide a better estimate of

the general phenological requirements of a species, but LTER data would provide a more accurate understanding for a single site. The best solution to this issue would be the development of models that accurately incorporate spatial variation, such as including genetic variation between different populations (Chuine and Régnière, 2017), although localized models could also be generated when large-scale predictions are unnecessary. Third, these differences could result from issues 301 with model identifiability. Since: since different parameter values can yield nearly identical es-302 timates of phenological events, parameter estimates can differ between datasets even when the 303 underlying processes generating the data are the same. Information about which of these issues 304 may be causing the differences between datasets can be explored using these analyses the analyses 305 in the current study, as will be explained below. 306

Despite substantial differences in parameter estimates, LTER and NPN-derived-USA-NPN derived 307 models produced similar estimates for phenological events in most cases (Fig. 3). This greater 308 correspondence between predictions than parameters suggests that more complex models may have 309 identifiability issues. For example, two GDD models with parameters of $t_1=1$, F=10, $T^*_{base}=0$ 310 and $t_1=5$, F=5, $T^*T_{base}=0$ produce nearly identical estimates in many scenarios. This possibility 311 is supported by the fact that the highest correlation between parameter estimates is seen in models 312 with only 1 or 2 parameters. In addition, bootstrap results for more complex models suggest 313 a high degree of variability in parameter estimates and potentially multiple local optima in fits 314 to both NPN-USA-NPN and LTER data (Fig. \$6-\$7\$10-\$11). Finally, parameter estimates of 315 more complex models are also not consistent among models for the same species when comparing multiple LTER datasets (Fig. \$4-\$5\$8-\$9). These results are consistent with research showing that models estimating failed to estimate the starting day of warming accumulation solely from

budbreak time-seriesfailed to accurately infer the internal phenology described in the models, thus
producing parameter estimates which were not biologically realistic (Chuine et al., 2016). Basler

(2016) suggests that the key component in phenology models is the thermal forcing, with additional
parameters being sensitive to over-fitting. Here, our simplest model, the *Fixed GDD model* which
uses only a warming component, had the highest correlation among parameters between LTER and
NPN-USA-NPN datasets. In combination with this previous research the aforementioned studies,
our results warrant caution-indicate that caution is warranted in interpreting parameter estimates
from complex phenology models regardless of the data source used for fitting the models.

While more complex phenology models appear to have identifiability issues, there is also evidence that they capture useful information, beyond the Fixed GDD model, based on their ability to make 328 out-of-sample predictions. Based on the RMSE, the GDD and Uniforc models produce the best 329 out-of-sample predictions for the majority of species and phenophases at both NPN-USA-NPN 330 and LTER datasets (Fig. \$1 & \$2\$5 & \$6). This demonstrates that the more complex models 331 are capturing additional information about phenology, and that some of the differences between 332 datasets result from differences in either the scales or the sampling of the data. Spatial variation 333 in phenological requirements is known to exist in plants (Zhang et al., 2017). In combination 334 with our results showing observed differences in parameter estimates between LTER sites (Fig. 335 \$4-\$5\$8-\$9), this suggests that variation in phenological requirements across the the range is likely 336 important. However, the models that attempted to address this by incorporating spatial variation 337 did not yield improvements over their base models in our analyses. Specifically, correspondence between parameter estimates (Fig. 2), estimates of phenological events (Fig. 3), and out-of-sample error rates (Fig. 4) for the MSB and M1 models were essentially the same as the Alternating and GDD models, respectively. This lack of improvement from incorporating spatial variation could be caused either by models not adequately capturing the process driving the spatial variation, the NPN USA-NPN dataset having biases from variation in sampling effort and/or spatial auto-correlation, or some combination of these factors. Basler (2016) used the M1 model to predict budburst for six species across Europe and found it was generally among the best models in terms of RMSE, albeit never by more than a single day. Their result was strengthened by having a 40-year time-series across a large region. Chuine and Régnière (2017) listed the incorporation of spatial variation in warming requirements in models as a primary issue in future phenology research. Large-scale phenology datasets, like NPNUSA-NPN, will be key in addressing this and other phenological research needs.

In addition to exploring differences between phenology datasets our analysis provides guidance on which models to use when making predictions at a local scale using models built from large scale 352 data, or vice versa. Among the eight models tested the Uniforc model performed the best overall in the cross dataset comparison in terms of pearson correlation and RMSE (Table S2). One exception was the Fixed GDD model had a slightly higher pearson value when using LTER derived models to 355 make predictions for USA-NPN data. When the scale is the same between predictions and training 356 data (ie. using LTER derived models to make predictions for LTER data) the Uniforc, GDD, and 357 M1 models performed equally well, though the Uniforc had slightly better RMSE (Table S2). The 358 best model for specific species and phenophases varied, but was commonly the Uniforc and GDD 350 models (Figs. S5-S6). 360

In conclusion, our results suggest that both LTER and NPN-USA-NPN data provide valuable information on plant phenology. Models built using both data sources yield effective predictions

for phenological events, but parameter estimates from the two data sources differ and models from each source best predict that data source's phenology events. The primary difference in the datasets is spatial scale, but due to trade-offs in data collection efforts, the larger scale NPN data has USA-NPN data have shorter time-series, less site fidelity and other differences from the intensively collected LTER data (Table 3). These differences can be strengths or potential limitations. 367 Observers sampling opportunistically allows the NPN USA-NPN dataset to have a large spatial 368 scale, but also leads to low site fidelity which limits the ability to measure long-term trends at 369 local scales (Gerst et al., 2016). Tracking long-term trends is the major strength of LTER data, 370 but having a relatively small species pool limits its their use in species-level predictive modeling. 371 Due to these differences, the best data source for making predictions depends on the scale at which 372 the predictions are being made. Identifying the most effective data sources for different types and 373 scales of analysis is a useful first step, but the ultimate solution to working with diverse data types 374 is to focus on integrating all types of data into analyses and forecasts (Hanks et al., 2011; Melaas 375 et al., 2016). Our results suggest that methods that can learn from the intensive information avail-376 able in LTER data in regions where it is they are available, and simultaneously use large-scale 377 data to capture spatial variation in phenological requirements will help improve our ability to un-378 derstand and predict phenology. Data integration efforts should also leverage data from remote 379 sensing sources such as the PHENOCAM network or satellite imagery, which have both a large 380 spatial extent and high temporal resolution (?)(Peng et al., 2017; Richardson et al., 2018a,b). Data 381 integration provides the potential to use data from many sources to produce the best opportunity 382 for accurate inference about, and forecasting of, the timing of biological events.

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Dataset Name	Habitat	Phenological Event (Num. Species) Reference	Reference
Harvard Forest	N.E. Deciduous Forest	Budburst (17) Flowering (7)	(O'Keefe, 2015)
Jornada Experimental Range	Chihuahuan Desert	Flowering (2)	
H.J. Andrews Experimental Forest N.W. Wet Coniferous Forest Budburst (5) Flowering (4)	N.W. Wet Coniferous Forest	Budburst (5) Flowering (4)	(Schulze, 2017)
Hubbard Brook	N.E. Deciduous Forest	Budburst (3)	(Bailey, 2018)

 Table 1: LTER datasets used in the analysis

DOY Estimator	Forcing Equations	Total	Reference
		Parameters	8
\overline{DOX}	ı	1	-
$\sum_{t=0}^{DOY} R_f(T_i) \geq F^*$	$R_f(T_i) = min(T_i, 0)$, $R_f(T_i) = max(T_i, 0)$.	1	-(Réaumur, 1735; Wang, 1960; Hunter and Lechowicz, 1992)
$DOY = eta_1 + eta_2 T_{mean}$	•	2	
$\sum_{I=t_1}^{DOY} R_f(T_i) \geq F^*$	$R_f(T_i) = \max(T_i - T^*, 0) R_f(T_i) = \max(T_i - T_{base}, 0)$	3	-(Réaumur, 1735; Wang, 1960; Hunter and Lechowicz, 1992)
$\sum_{I=t_1}^{DOY} R_f(T_i) \geq (rac{L_i}{24})^k F^*$	$R_f(T_i) = \max(T_i - T^*, 5) R_f(T_i) = \max(T_i - T_{base}, 5)$	4	(Blümel and Chmielewski, 2012)
$ \begin{array}{l} DOY \\ t=0 \\ t=0 \end{array} R_f(T_i) \ge a + be^{cNCD(t)} $	$R_f(T_i) = max(T_i - 5, 0)$	8	(Cannell and Smith, 1983)
$R_f(T_i) \ge \mathcal{L} + be^{cNCD_i} + dT_{mean}$	$R_f(T_i) = max(T_i - 5, 0)$	4	(Jeong et al., 2013)
$\sum_{l=t_1}^{DOY} R_f(T_l) \geq F^*$	$R_f(T_i) = rac{1}{1+e^{b(T_i-c)}}$	4	(Chuine, 2000)

Table 2: Phenology models used in the analysis

		<u>LTER</u>	$\underline{\frac{NPN\underbrace{USA-NPN}}{NPN\underbrace{NSA-NPN}}}$
	Time-series length	High	Low
	Spatial extent	Low	High
570	Local species representation	High	Low
	Regional/Continental species representation	Low	High
	Number of observers	Low	High
	Site fidelity	High	Low

Table 3: Attributes of the two datasets used in this study. Bold text indicates an attribute is expected

to increase over time.

Figure 1: Locations of National Phenology Network sites used (black points) and Long Term

Ecological Research sites (labeled circles), with greyscale showing elevation.

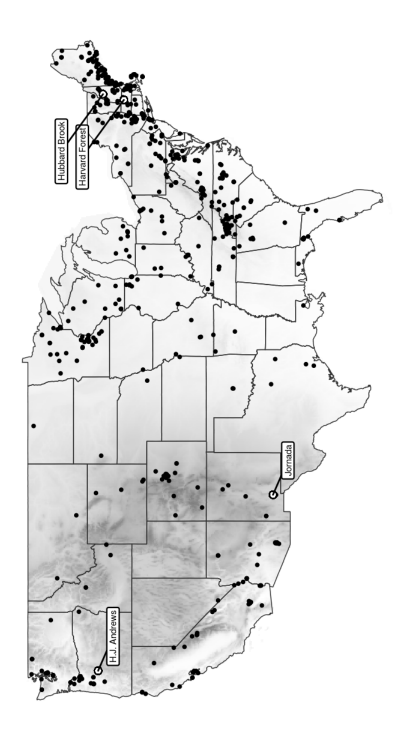


Figure 1: .

Figure 2: Comparisons of parameter estimates between NPN USA-NPN and LTER derived models. Each point represents a parameter value for a specific species and phenophase, and is the mean value from 250 bootstrap iterations. The black line is the 1:1 line. The R^2 is the coefficient of determination, which can be negative if the relationship between the two parameter sets is worse than no relationship but with the same mean values.

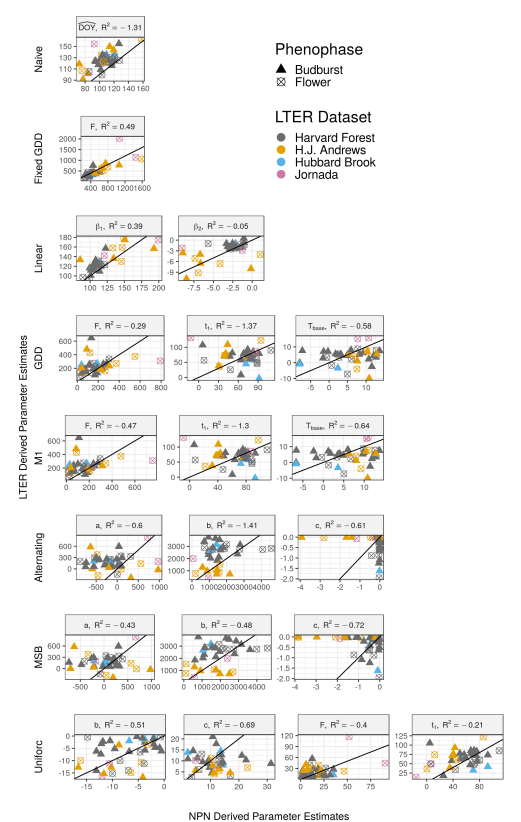


Figure 2: .

Figure 3: Comparison of predicted day of year (DOY) of all phenological events between NPN

USA-NPN and LTER-derived models. Top panels show comparisons at LTER sites and bottom

panels show comparisons at NPN-USA-NPN sites. Each point is an estimate for a single held-out

observation. Colors indicate observations for a single species and phenophase combination.

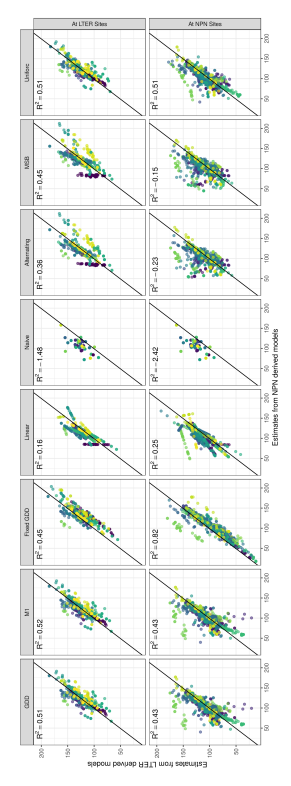


Figure 3: .

Figure 4: Differences in prediction error between NPN-USA-NPN and LTER-derived models.

Density plots for comparisons of predictions on LTER data (top row) and NPN-USA-NPN data

(bottom row). Each plot represents the difference between the RMSE for LTER-derived model

and the NPN-derived-USA-NPN derived model, meaning that values less than zero indicate more

accurate prediction by LTER-derived models and values greater than zero indicate more accurate

prediction by NPN-derived models. *p* <0.001 for all t-tests. Differences are calculated pairwise for

the 38 species/phenophase comparisons.

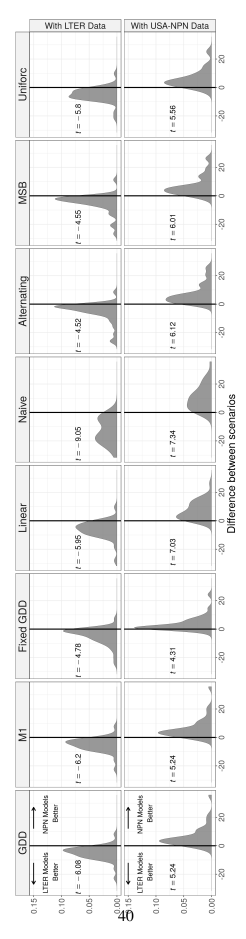


Figure 4: .