# Do we need demographic data to forecast plant population dynamics?

- Andrew T. Tredennick<sup>1\*</sup>, Mevin B. Hooten<sup>2,3,4</sup>, and Peter B. Adler<sup>1</sup>
- <sup>4</sup> Department of Wildland Resources and the Ecology Center, 5230 Old Main Hill, Utah State University,
- 5 Logan, Utah 84322, USA; <sup>2</sup>U.S. Geological Survey, Colorado Cooperative Fish and Wildlife Research Unit,
- <sup>6</sup> Fort Collins, CO 80523, USA; <sup>3</sup>Department of Fish, Wildlife, and Conservation Biology, Colorado State
- 7 University, Fort Collins, CO 80523 USA; <sup>4</sup>Department of Statistics, Colorado State University, Fort Collins,
- 8 CO 80523 USA
- 9 Last compile: July 1, 2016

# Summary

10

12

13

14

15

16

17

18

19

20

21

- 1 Rapid environmental change has generated growing interest in forecasts of future population trajectories. Traditional population models built with detailed demographic observations from one study site can address the impacts of environmental change at particular locations, but are difficult to scale up to the landscape and regional scales relevant to management decisions.
  - 2 An alternative is to build models using population-level data which are much easier to collect than individual-level data over broad spatial scales. However, it is unknown whether models built using population-level data adequately capture the effects of density-dependence and environmental forcing that are necessary to generate skillful forecasts.

<sup>\*</sup>Corresponding author: E-mail: atredenn@gmail.com

- 3 Here, we test the consequences of aggregating individual responses when forecasting the population states and trajectories of four perennial grass species in a semi-arid grassland in Montana, USA. We parameterized two population models for each species, one based on individual-level data (survival, growth and recruitment) and one on population-level data (percent cover), and compared their forecasting skill and forecast horizons with and without the inclusion of climate covariates. For both models we used Bayesian ridge regression to weight the influence of climate covariates for optimal prediction.
- 4 In the absence of climate effects, we found no significant difference between the forecasting skill of models based on individual-level data and models based on population-level data. Climate effects were weak, but increased forecasting skill for two species. Increases in skill with climate covariates were similar between model types.
- 5 For our focal species at this particular location, and using our particular statistical models,
  percent cover models generated forecasts as skillful as those from a demographic model.
  We conclude that models based on aggregated individual-level data offer a practical alternative to data-intensive demographic models when forecasting is the goal.
- Key-words: forecasting, climate change, grassland, integral projection model, population model, statistical regularization, ridge regression

# Introduction

22

23

25

26

27

28

29

30

31

32

- Perhaps the greatest challenge for ecology in the 21st century is to forecast the impacts of envi-
- ronmental change (Clark et al. 2001, Petchey et al. 2015). Forecasts require sophisticated mod-
- eling approaches that fully account for uncertainty and variability in both ecological process and
- 43 model parameters (Luo et al. 2011, but see Perretti et al. 2013). The increasing statistical sophis-
- tication of population models (Rees and Ellner 2009) makes them promising tools for predicting
- the impacts of environmental change on species persistence and abundance. But reconciling the

scales at which population models are parameterized with the scales at which environmental changes play out remains a challenge (Clark et al. 2010, 2012, Freckleton et al. 2011, Queenborough et al. 2011). Most population models are built using demographic data from a single study site because tracking the fates of individuals is so difficult. The resulting models cannot be applied to the landscape and regional scales relevant to decision-making without information about 50 how the estimated parameters respond to spatial variation in biotic and abiotic drivers (Sæther et al. 2007). The limited spatial extent of individual-level demographic datasets constrains our ability to use population models to address applied questions about the consequences of climate 53 change. Aggregate measures of population status, rather than individual performance, offer an intriguing alternative for modeling populations (Clark and Bjørnstad 2004, Freckleton et al. 2011). Population-level data cannot provide inference about demographic mechanisms, but might be sufficient for modeling future population states, especially because population-level data, such as plant percent cover, are feasible to collect across broad spatial extents (e.g., Queenborough et al. 2011). The choice between individual and population-level data involves a difficult trade-off: while individual-level data are necessary for mechanistic models, population-level data enable models that can be applied over greater spatial and temporal extents. An open question is how much forecasting skill is lost when we build models based on population rather than individuallevel data. To date, most empirical population modelers have relied on individual-level data, with few attempts to capitalize on population-level measures. An important exception was an effort by Taylor and Hastings (2004) to model the population growth rate of an invasive species to identify 67 the best strategies for invasion control. They used a "density-structured" model where the state variable is a discrete density state rather than a continuous density measure. Such models do not require individual-level demographic data and can adequately describe population dynamics. Building on Taylor and Hastings (2004), Freckleton et al. (2011) showed that density-structured models compare well to continuous models in theory, and Queenborough et al. (2011) provide

empirical evidence that density-structured models are capable of reproducing population dynamics at landscape spatial scales, even if some precision is lost when compared to fully continuous
models. However, previous tests of density-structured models have yet to assess their ability to
forecast out-of-sample observations, and they have not included environmental covariates, which
are necessary to forecast population responses to climate change.

Addressing climate change questions with models fit to population-level data is potentially problematic. Population growth (or decline) is the outcome of demographic processes such as survival, growth, and recruitment that occur at the level of individual plants. Climate can affect each demographic process in unique, potentially opposing, ways (Dalgleish et al. 2011). These unique 81 climate responses may be difficult to resolve in statistical models based on population-level data where demographic processes are not identifiable. Futhermore, models based on aggregated data may reflect short-term effects of one vital rate more than others whose importance may only emerge over the long-term. For example, a one-year change in a plant species' cover or biomass might reflect individual growth or shrinkage, whereas the long-term trajectory of the population might be more influenced by recruitment. The same is true for density dependence: intraspecific density depedence may act most strongly on vital rates, like recruitment, that are difficult to identify from population-level data. If density dependence and/or important climate effects are missed because of the aggregation inherent in in population-level data, then population models built with such data will make uninformative or unreliable forecasts. 91

We compared the forecasting skill of statistical and population models based on aggregated,
population-level data with the skill of models based on individual-level data. We used a demographic dataset that tracks the fates of individual plants from four species over 14 years to build
two kinds of single-species population models, traditional models using individual growth, survival, and recruitment data and alternative models based on population-level (basal cover) data.
We simulated the models to answer two questions motivated by the fact that the effects of intraspecific competition (density dependence) and interannual weather variability act at the level
of the individual (Clark et al. 2011). First, can population models fit using aggregated individual-

level data (percent cover) adequately capture density dependence to produce forecasts as skillful as those from models fit to demographic data? Second, can population models fit using aggregated data adequately capture the influence of climate on population growth and, in turn, produce forecasts as skillful as those from models fit to demographic data?

# Materials and Methods

# 105 Study site and data

Our demographic data come from a northern mixed grass prairie at the Fort Keogh Livestock and Range Research Laboratory near Miles City, Montana, USA (46° 19' N, 105° 48' W). The 107 dataset is available on Ecological Archives<sup>1</sup> (Anderson et al. 2011), and interested readers should 108 refer to the metadata for a complete description. The site is 800 m above sea level and mean 109 annual precipitation (1878-2009) is 334 mm, with most annual precipitation falling from April 110 through September. The community is grass-dominated, and we focused on the four most abun-111 dant grass species: Bouteloua gracilis (BOGR), Hesperostipa comata (HECO), Pascopyrum 112 smithii (PASM), and Poa secunda (POSE) (Fig. 1). B. gracilis is a warm-season perennial grass, 113 whereas H. comata, P. smithii, and Poa secunda are cool-season perennial grasses. The growing 114 season begins in early spring (typically in April) and lasts through mid-summer (typically in 115 June). 116 From 1932 to 1945, individual plants were identified and mapped annually in 44 1-m<sup>2</sup> quadrats using a pantograph. The quadrats were distributed among six pastures, each assigned a grazing treatment of light (1.24 ha/animal unit month), moderate (0.92 ha/aum), and heavy (0.76 ha/aum) 119 stocking rates (two pastures per treatment). In this analysis, we accounted for potential differences among the grazing treatments, but do not focus on grazing x climate interactions. The an-121 nual maps of the quadrats were digitized and the fates of individual plants tracked and extracted 122 using a computer program (Lauenroth and Adler 2008, Chu et al. 2014). The permanent quadrats 123

<sup>&</sup>lt;sup>1</sup>http://esapubs.org/archive/ecol/E092/143/

have not been relocated, but their distribution in six different pastures means the data represent a broad spatial distribution for the study area. Daily climate data are available for the duration of the data collection period (1932 - 1945) from the Miles City airport, Wiley Field, 9 km from the study site.

We modeled each grass population based on two levels of data: individual and quadrat. The 128 individual data are the "raw" data. For the quadrat-level data we simply sum individual basal 129 cover for each quadrat by species. This is equivalent to a near-perfect census of quadrat percent 130 cover because measurement error at the individual-level is small (Chu and Adler 2015). Based on 131 these two datasets of 13 year-to-year transitions, we can compare population models built using 132 individual-level data and aggregated, quadrat-level data. At the individual level, we explicitly 133 model three vital rates: growth, survival, and recruitment. At the quadrat level, we model popula-134 tion growth as change in percent cover of quadrats with non-zero cover in year t and in year t-1, ignoring within-quadrat extirpation and colonization events because they are very rare in our time series (N=16 and N=13, respectively, across all species). Sample sizes for each species and 137 vital rate model are shown in Table 1.

All R code and data necessary to reproduce our analysis is archived on GitHub as release v1.0<sup>2</sup> (http://github.com/atredennick/MicroMesoForecast/releases). We have also deposited the v1.0 release on Dryad (*link here after acceptance*).

#### 42 Stastical models of vital rates

At both levels of inference (individual and quadrat), the building blocks of our population models are vital rate regressions. For individual-level data, we fit regressions for survival, growth, and recruitment for each species. At the quadrat-level, we fit a single regression model for population growth. We describe the statistical models separately because they required different approaches. For both model types, we fit vital rate models with and without climate covariates. Models with

<sup>&</sup>lt;sup>2</sup>Note to reviewers: so that v1.0 will be associated with the published version of the manuscript, we have re-

climate effects contain five climate covariates that we chose *a priori* based on previous model selection efforts using these data (Chu et al. 2016) and expert advice (Lance Vermeire, *personal communication*): "water year" precipitation at t-2 (lagppt); April through June precipitation at t-1 and t (ppt1 and ppt2, respectively) and April through June temperature at t-1 and t (TmeanSpr1 and TmeanSpr2, respectively), where t-1 to t is the transition of interest. We also include interactions among same-year climate covariates (e.g., ppt1 × TmeansSpr1), resulting in a total of seven climate covariates.

We fit all models using a hierarchical Bayesian approach. We focus on the main process and the model likelihood in what follows (full model descriptions are in the Supporting Information). For the likelihood models,  $\mathbf{y}^X$  is always the relevant vector of observations for vital rate X (X = S, G, R, or P) for survival, growth, recruitment, or population growth). For example,  $\mathbf{y}^S$  is a vector of 0s and 1s indicating whether a genet survives from t to t+1, or not, for all observation years and quadrats. All model parameters are species-specific, but we omit subscripts for species in model descriptions below to reduce visual clutter. For brevity, we only describe models with climate covariates included, but models without climate covariates are simply the models described below with the climate effects removed.

Vital rate models at the individual level We used logistic regression to model the probability that genet i in quadrat q survives from time t to t+1 ( $s_{i,q,t}$ ):

$$y_{i,q,t}^S \sim \text{Bernoulli}(s_{i,q,t}),$$
 (1)

$$logit(s_{i,q,t}) = \beta_{0,t} + \beta_{s,t}x_{i,q,t} + \beta_Q + \mathbf{z}_t'\boldsymbol{\beta}_c + \beta_{d,1}w_{i,t} + \beta_{d,2}(x_{i,q,t}w_{i,q,t}),$$
(2)

where  $x_{i,q,t}$  is the log of genet i basal area at time t,  $\beta_{0,t}$  is a year specific intercept,  $\beta_Q$  is the random effect of quadrat group location,  $\beta_{s,t}$  is the year-specific slope parameter for size,  $\mathbf{z}$  is a vector of p climate covariates specific to year t,  $\beta_c$  is a vector of fixed climate effects of length p,  $\beta_{d,1}$  is the effect of intraspecific crowding experienced by the focal genet at time t ( $w_{i,q,t}$ ), and  $\beta_{d,2}$  is a size by crowding ( $x_{i,q,t}w_{i,q,t}$ ) interaction effect.

We follow the approach of Chu and Adler (2015) to estimate crowding, assuming that the crowding experienced by a focal genet depends on distance to each neighbor genet and the neighbor's size, u:

$$w_{i,q,t} = \sum_{k} e^{-\delta d_{ik,q,t}^2} u_{k,q,t}.$$
 (3)

In equation 3,  $w_{i,t}$  is the crowding that genet i in year t experiences from conspecific neighbors in quadrat q. The spatial scale over which conspecific neighbors exert influence on any genet is determined by  $\delta$ . The function is applied for all k conspecific genets that neighbor the focal genet at time t, and  $d_{ik,q,t}$  is the distance between genet i and conspecific genet k in quadrat q. We use regression-specific (survival and growth)  $\delta$  values estimated by Chu and Adler (2015).

We modeled growth as a Gaussian process describing log genet size ( $y_{i,q,t+1}^G$ ) at time t+1 in quadrat q as a function of log size at time t and climate covariates:

$$y_{i,q,t+1}^G \sim \text{Normal}(\mu_{i,q,t+1}, \sigma_{x_{i,q,t+1}}^2 \mathbf{I}),$$
 (4)

$$\mu_{i,q,t+1} = \beta_{0,t} + \beta_{s,t} x_{i,q,t} + \beta_Q + \mathbf{z}_t' \boldsymbol{\beta}_c + \beta_{d,1} w_{i,q,t} + \beta_{d,2} (x_{i,q,t} w_{i,q,t}),$$
(5)

where  $\mu_{i,q,t+1}$  is log of genet *i*s predicted size at time t+1, and all other parameters are as described for the survival regression. We capture non-constant error variance in growth by modeling the variance in the growth regression ( $\sigma_{xi,q,t+1}^2$ ) as a nonlinear function of predicted genet size:

$$\sigma_{x_{i,q,t+1}}^2 = a \exp[b \times \mu_{i,q,t+1}], \tag{6}$$

where  $\mu_{i,q,t+1}$  is log of predicted genet size predicted from the growth regression (Eq. 4), and a and b are constants.

Our data allows us to track new recruits, but we cannot assign a specific parent to new genets.

Therefore, we model recruitment at the quadrat level. We assume the number of individuals,

 $y_{q,t+1}^R$ , recruiting at time t+1 in quadrat q follows a negative binomial distribution:

$$y_{q,t+1}^R \sim \text{NegBin}(\lambda_{q,t+1}, \phi),$$
 (7)

where  $\lambda$  is the mean intensity and  $\phi$  is the size parameter. We define  $\lambda$  as a function of quadrat composition and climate in the previous year:

$$\lambda_{q,t+1} = \tilde{c}_{q,t} \exp\left(\beta_{0,t} + \beta_Q + \mathbf{z}_t' \boldsymbol{\beta}_c + \beta_d \sqrt{\tilde{c}_{q,t}}\right), \tag{8}$$

where  $\tilde{c}_{q,t}$  is effective cover (cm<sup>2</sup>) of the focal species in quadrat q at time t, and all other terms are as in the survival and growth regressions. Effective cover is a mixture of observed cover (c) in the focal quadrat (q) and the mean cover across the entire group  $(\bar{c})$  of Q quadrats in which q is located:

$$\tilde{c}_{q,t} = pc_{q,t} + (1-p)\bar{c}_{Q,t},$$
(9)

where p is a mixing fraction between 0 and 1 that is estimated when fitting the model.

**Population model at the quadrat level** The statistical approach used to model aggregated 203 data depends on the type of data collected. We have percent cover data, which can easily be trans-204 formed to proportion data in our case because plant areas were scaled by plot area. An obvious 205 choice for fitting a linear model to proportion data is beta regression because the support of the 206 beta distribution is (0,1), which does not include true zeros or ones. However, when we used fit-207 ted model parameters from a beta regression in a quadrat-based population model, the simulated 208 population tended toward 100% cover for all species. We therefore chose a modeling approach 209 based on a truncated log-normal likelihood. The model for quadrat cover change from time t to 210 t+1 is

$$y_{q,t+1}^P \sim \text{LogNormal}(\mu_{q,t+1}, \sigma^2)_0^1, \tag{10}$$

$$\mu_{q,t+1} = \beta_{0,t} + \beta_{s,t} x_{q,t} + \beta_Q + \mathbf{z}_t' \boldsymbol{\beta}_c, \tag{11}$$

where  $\mu_{q,t+1}$  is the log of proportional cover in quadrat q at time t+1, and all other parameters are as in the individual-level growth model (Eq. 4) except that x now represents log of proportional cover. The log normal likelihood includes a truncation (subscript 0, superscript 1) to ensure that predicted values do not exceed 100% cover.

Our Bayesian approach to fitting the vital rate models required choosing appro-

## Model fitting and statistical regularization

Model fitting

priate priors for unknown parameters and deciding which, if any, of those priors should be hier-219 archical. For each species, we fit yearly size effects and yearly intercepts hierarchically, where 220 year-specific coefficients were drawn from global distributions representing the mean size effect 221 and intercept. Quadrat random effects were also fit hierarchically, with quadrat offsets being 222 drawn from distributions with mean zero and a shared variance term (independent Gaussian pri-223 ors). Climate effects were modeled as independent covariates whose prior distributions were 224 optimized for prediction using statistical regularization (see Statistical regularization: Bayesian 225 ridge regression below). 226 All of our analyses (model fitting and simulating) were conducted in R (R Core Team 2013). We used the 'No-U-Turn' Hamiltonian Monte Carlo sampler in Stan (Stan Development Team 2014a) to estimate the posterior distributions of model parameters using the package rstan (Stan Development Team 2014b). We obtained posterior distributions for all model parameters 230 from three parallel MCMC chains run for 1,000 iterations after discarding an initial 1,000 iter-231 ations. Such short MCMC chains are possible because the Stan sampler reduces the number of 232 iterations needed to achieve convergence. We assessed convergence visually and checked that 233 scale reduction factors for all parameters were less than 1.1. There were six instances where the 234 scale reduction factor for a particular parameter was greater than 1.1. In those cases, we checked 235 the traceplots of the violating parameter and discovered that one chain, out of three, was poorly 236 behaved. Therefore, we removed that chain from the analysis, leaving two well-mixed and con-237 verged chains for those six parameters. For the purposes of including parameter uncertainty

in our population models, we retained the final 1,000 iterations from each of the three MCMC chains to be used as randomly drawn values during population simulation. We report the posterior mean, standard deviation, and 95% Bayesian Credible Intervals for every parameter of each model for each species in the Supporting Information (Tables S5-S20).

Statistical regularization: Bayesian ridge regression For models with climate covariates, our objective is to model the response of our focal grass species to interannual variation in 244 climate, even if those responses are weak. Therefore, we avoid selecting among models with all 245 possible combinations of climate covariates, and instead use Bayesian ridge regression to regu-246 late, or constrain, the posterior distributions of each climate covariate (Gerber et al. 2015, Hooten 247 and Hobbs 2015). Ridge regression is a specific application of statistical regularization that seeks 248 to optimize model generality by trading off bias and variance. As the name implies, statistical reg-249 ularization involves the use of a regulator that constrains an optimization. The natural regulator in 250 a Bayesian application is the prior on the coefficient of interest. Each of our statistical models in-251 cludes the effects of climate covariates via the term  $\mathbf{z}_t'\boldsymbol{\beta}_c$  with prior  $\boldsymbol{\beta}_c \sim \text{Normal}(\boldsymbol{\mu}_{\beta_c}, \sigma_{\beta_c}^2 \mathbf{I})$ . Be-252 cause we standardized all climate covariates to have mean zero and variance one, we set  ${m \mu}_{eta_c}=0$ 253 and let  $\sigma_{\beta_c}^2$  serve as the regulator that shrinks covariates toward zero – the smaller the prior vari-254 ance, the more the posteriors of  $\beta_c$  are shrunk toward zero, and the stronger the penalty (Hooten 255 and Hobbs 2015). 256 To find the optimal penalty (i.e., optimal value of the hyperparameter  $\sigma_{\beta_c}^2$ ), we fit each statistical 257 model with a range of values for  $\sigma_{\beta_c}^2$  and compared predictive scores from leave-one-year-out 258 cross-validation. We performed the grid search over 24 values of  $\sigma_{\beta_c}^2$ , ranging from  $\sigma_{\beta_c}^2=0.01$ 259 to  $\sigma_{\beta_c}^2=2.25$ . For each statistical model and each species, we fit  $13\times 24=312$  iterations of 260 the model fitting algorithm to search  $\sigma^2_{\beta_c}$  for the optimal value (13 years to leave out for cross-26 validation and 24 values of  $\sigma_{\beta_c}^2$ ) – a total of 4,992 model fits. We calculated the log pointwise 262 predictive density (lppd) to score each model's ability to predict the left-out data (Gelman et 263 al. 2014). Thus, for training data  $y_{train}$  and held-out data  $y_{hold}$  at a given value of  $\sigma_{\theta}^2$  across all

MCMC samples s=1,2,...,S and all hold outs of data from year t to year T, and letting  $\theta$ represent all unknowns, lppd is

$$lppd_{CV} = \sum_{t=1}^{T} log_e \int [y_{t,hold}|\theta] [\theta|y_{train}] d\theta,$$
(12)

268 and computed as

$$\sum_{t=1}^{T} \log_e \left( \frac{1}{S} \sum_{s=1}^{S} (y_{t,hold} | \theta_{ts}) \right). \tag{13}$$

We chose the optimal prior variance for each species-statistical model combination as the one that produced the highest *lppd* and then fit each species-statistical model combination using the full data set for each species and the optimal prior variance. We calculated the *lppd* from posterior samples using the algorithm from Vehtari et al. (2016).

## 74 Population models

Using samples from the posterior distribution of the vital rate statistical models, it is straightforward to simulate the population models. We used an Integral Projection Model (IPM) to simulate populations based on individual-level data (Ellner and Rees 2006) and a quadrat-based version of an individually-based model (Quadrat-Based Model, QBM) to simulate populations based on quadrat-level data. We describe each in turn.

Integral projection model We use a stochastic IPM (Rees and Ellner 2009) to simulate our focal populations based on the vital rate regressions described above. In all simulations, we ignore the random year effects so that interannual variation is driven solely by climate. We fit the random year effects in the vital rate regressions to avoid over-attributing variation to climate covariates. Our IPM follows the specification of Chu and Adler (2015) where the population of species j is  $n(u_j, t)$ , giving the density of sized-u genets at time t. Genet size is on the natural log scale, so that  $n(u_j, t)du$  is the number of genets whose area (on the arithmetic scale) is between  $e^{u_j}$  and  $e^{u_j+du}$ . The function for any size v at time t+1 is

$$n(v_j, t+1) = \int_{L_j}^{U_j} k_j(v_j, u_j, \bar{w}_j(u_j)) n(u_j, t) du_j,$$
(14)

where  $k_j(v_j, u_j, \bar{w_j})$  is the population kernel that describes all possible transitions from size u to v and  $\bar{w}_i$  is a scalar representing the average intraspecific crowding experienced by a genet of size  $u_i$  and species j. The integral is evaluated over all possible sizes between predefined lower (L) and upper (U) size limits that extend beyond the range of observed genet sizes. 292 The IPM is spatially-implicit, thus, we cannot calculate neighborhood crowding for specific 293 genets  $(w_{ij})$ . Instead, we use an approximation  $(\bar{w}_i)$  that captures the essential features of neigh-294 borhood interactions (Adler et al. 2010). This approximation relies on a 'no-overlap' rule for 295 conspecific genets to approximate the overdispersion of large genets in space (Adler et al. 2010). 296 The population kernel is defined as the joint contributions of survival (S), growth (G), and recruit-297 ment (R):

$$k_j(v_j, u_j, \bar{w}_j) = S_j(u_j, \bar{w}_j(u_j))G_j(v_j, u_j, \bar{w}_j(u_j)) + R_j(v_j, u_j, \bar{w}_j),$$
(15)

and adding in newly recruited (R) individuals of an average sized one-year-old genet for the focal 301 species. Note the S, G, and R are incorporated in the IPM using the fitted vital rate regressions. 302 Our statistical model for recruitment (R, described above) returns the number of new recruits 303 produced per quadrat. Following previous work (Adler et al. 2012, Chu and Adler 2015), we 304 assume that fecundity increases linearly with size  $(R_j(v_j, u_j, \bar{w}_j) = e^{u_j} R_j(v_j, \bar{w}_j))$  to incorporate 305 the recruitment function in the spatially-implicit IPM. 306 We used random draws from the final 1,000 iterations from each of three MCMC chains for each vital rate regression to carry-through parameter uncertainty into our population models. At 308 each time step, we drew the full parameter set (climate effects and density-dependence fixed 309 effects) from a randomly selected MCMC iteration. Relatively unimportant climate covariates

which means we are calculating growth (G) for individuals that survive (S) from time t to t+1

300 299 (those that broadly overlap 0) will have little effect on the mean of the simulation results, but can contribute to their variation. To retain temporal variation associated with random year effects, we used posterior estimates of the mean temporal effect and the standard deviation of that effect to generate a random year effect for unobserved years. That is, for some future year T, the intercept is  $\beta_0, T \sim \text{Normal}(\beta_0, \sigma_{\beta_0})$  and the effect of size is  $\beta_s, T \sim \text{Normal}(\beta_s, \sigma_{\beta_s})$ .

Quad-based model To simulate our quad-based model (QBM), we iterate the quadrat-level statistical model (Eqs. 9-10). We use the same approach for drawing parameter values as described for the IPM. After drawing the appropriate parameter set, we calculate the mean response (log cover at t+1 is  $\mu_{t+1}$ ) according to Eq. 10. We make a random draw from a [0,1] truncated lognormal distribution with mean equal to  $\mu_{t+1}$  from Eq. 10 and the variance estimate from the fitted model. We project the model forward by drawing a new parameter set (unique to climate year and MCMC iteration) at each timestep. Random year effects are included as described above for the IPM.

#### 324 Model validation

To test each model's ability to forecast population states, we made out-of-sample predictions 325 using leave-one-year-out cross validation. For both levels of modeling and for models with and 326 without climate covariates, we fit the vital rate models using observations from all years except 327 one, and then used those fitted parameters in the population models to perform a one-step-ahead forecast for the year whose observations were withheld from model fitting. Within each observation year, several quadrats were sampled. We made predictions for each observed quadrat in 330 each focal year, initializing each simulation with cover in the quadrat the previous year. Because 331 we were making quadrat-specific predictions, we incorporated the group random effect on the 332 intercept for both models. We repeated this procedure for all 13 observation years, making 100 333 one-step-ahead forecasts for each quadrat-year combination with parameter uncertainty included 334 via random draw from the MCMC chain as described above. As described above, year-specific 335

parameters for left-out data were drawn from the posterior distribution of the mean intercept. This cross-validation procedure allowed us to compare accuracy and precision of the two mod-337 eling approaches (IPM versus QBM) with and without climate covariates. We first calculated 338 the median predicted cover across the 100 simulations for each quadrat-year and then calculated 339 forecast skill as the correlation  $(\rho)$  between forecasts and observations. We calculated forecast 340 error as mean absolute error (MAE) between forecasts and observations. We compared  $\rho$  and 341 MAE between model types and within model types between models with and without climate 342 covariates using one-sided t tests with adjusted degrees of freedom following Wilcox (2009) and 343 standard errors calculated using the HC4 estimator of Cribari-Neto (2004). Statistical tests were 344 conducted using algorithms from Ye et al. (2015). 345

#### 46 Forecast horizons

An important feature of any forecasting model is the rate at which forecast skill declines as the time between an observation and a forecast increases; the so-called ecological forecast horizon 348 (Petchey et al. 2015). To assess the forecast horizons of our models, we initiate the forecast 349 model with the population state at some time t and make sequential forecasts of the population 350 at times  $t+1, t+2, \ldots, t+T$  where T is the maximum number of years between the initial 351 year and the final year of our observations. For example, if we initialize the forecast model with 352 percent cover in 1940, we are able to make five forecasts up to the year 1945. Forecast models 353 are not re-initialized with observations between years. Thus, in our current example, the model 354 forecast for percent cover in 1941 has a forecast horizon of one year, the forecast in 1942 has a 355 forecast horizon of two years, and so on. We performed these simulations for all model types 356 (IPM with/without climate; QBM with/without climate) using mean parameter values for all 357 possible initial years. For a given forecast horizon, we averaged the correlation between forecasts 358 and observations. Note that these forecasts are all made using in-sample data because we used 359 model fits from the full data set. Nonetheless, these simulations offer insight into the differences among model forecast horizons.

## 2 Results

The IPM and QBM generated one-step-ahead forecasts of similar skill for out-of-sample obser-363 vations, with an average correlation between predictions and observations ( $\rho$ ) of 0.72 across all 364 models and species (Fig. 2). Without climate covariates, the accuracy of forecasts from the IPM 365 were not statistically greater than the accuracy of forecasts from the QBM (Fig. 2) and overall er-366 ror was similar (mean absolute error; Fig. S1, Supporting Information). With climate covariates, 367 the best out-of-sample predictive model (highest lppd) for each species and vital rate typically 368 resulted from highly constrained priors on the climate effects (Fig. S2, Supporting Information). 369 Thus, the posterior distributions of climate effects included in our models overlapped zero and 370 generally shrunk toward zero, though for some species-vital rate combinations, important effects (80% credible interval does not include zero) did emerge (Fig. 3). Despite the weak climate effects, including climate covariates did increase the accuracy of fore-373 casts for two species: B. gracilis and Poa secunda (Fig. 2). However, only for B. gracilis were 374 the skill increases statistically significant at  $\alpha = 0.05$  for the IPM  $(t_{(279)} = 1.70, P = 0.045)$  and 375 the QBM ( $t_{(279)} = 1.80$ , P = 0.037). Similarly, forecast error decreased significantly with the in-376 clusion of climate covariates for the B. gracilis IPM  $(t_{(280)} = -3.72, P = 0.029)$  and QBM  $(t_{(280)} = -3.72, P = 0.029)$ 377 = -3.34, P < 0.0001), and for the *Poa secunda* IPM ( $t_{(196)} = -1.90$ , P < 0.0001) and QBM ( $t_{(196)} = -1.90$ ). 378 -2.47, P = 0.007) (Fig. S2, Supporting Information). In no case did including climate covariates 379 significantly decrease forecast skill (Table S21), despite small changes in the mean skill (Fig. 2). 380 IPM forecasts were significantly more accurate than the QBM in only one case (Fig. 2): forecast 38 accuracy of P. smithii percent cover from an IPM with climate covariates was greater than the 382 accuracy from the QBM with climate covariates ( $t_{(215)} = 1.92$ , P = 0.028). However, adding 383 climate covariates decreased the skill of both models, and the difference between the IPM and 384 QBM emerges only because skill decreased less for the IPM than the QBM. Results from all 385

pairwise statistical tests are shown in Table S22 of the Supporting Information.

With climate covariates included, the accuracy of both models' forecasts declined as the forecast horizon increased, but they did so at similar rates (Fig. 4). The only exception is for *Poa secunda*, where QBM forecast accuracy remained steady as the forecast horizon increases, whereas IPM forecast accuracy declined (Fig. 4).

Population models built using individual-level data provide inference on demographic processes,

# Discussion

but they can only forecast future population states across the (typically limited) spatial extent of the observations. Population-level data are much easier to collect across broad spatial extents, 394 so models built using such data offer an appealing alternative to traditional population models 395 (Queenborough et al. 2011). However, density-structured models rely on the aggregation of 396 individual-level data. Given that individuals, not populations, respond to intraspecific competition 397 and weather (Clark et al. 2011), can models based on population-level metrics generate forecasts 398 that are as skillful as those generated from models based on individual-level data? Are models 399 based on population-level metrics as sensitive to climate forcing as models based on individual-400 level data? 401 Our comparison of a traditional, demographic population model without environmental forc-402 ing (the IPM) to an equivalent model inspired by density-structured models (the QBM) showed 403 that, generally, IPM forecasts of out-of-sample plant population states were no more accurate than forecasts from the QBM (Fig. 2; 'no-climate' bars). This result overturned our expectation that the IPM would out-perform the QBM, thanks to its mechanistic representation of the perennial life cycle, and confirms theoretical (Freckleton et al. 2011) and empirical work (Taylor and Hastings 2004, Queenborough et al. 2011) showing that density-structured models can be useful 408 surrogates for demographic models when the goal is to estimate or forecast population states over 409 large spatial extents.

We also expected the inclusion of environmental forcing to reveal more differences between the models. Interannual variation in weather can affect vital rates in different ways (Dalgleish et al. 2011), so, estimates of climate effects on plant population growth may be biased or nonidentifiable when the underlying statistical model is fit using population-level data that integrates 414 over the potentially unique climate responses of individual vital rates. We found some evidence 415 that the QBM failed to detect climate effects for three species (B. gracilis, H. comata, and Poa 416 secunda), where important climate effects were identified in the individual vital rate models but 417 not in the percent cover model (Fig. 3). However, for the two species where including climate 418 covariates increased forecast accuracy (B. gracilis and Poa secunda), forecast skill (Fig. 2) and 419 error (Fig. S2) were equivalent between the IPM and QBM. 420 The higher accuracy of the IPM and QBM with climate covariates for B. gracilis and Poa se-421 cunda highlights the advantage of contemporary modeling and variable selection approaches 422 such as ridge regression and LASSO over techniques that would exclude "non-significant" effects from final models. Ridge regression allows researchers to retain covariates whose effects may be difficult to identify in noisy data or short time series. This is especially important when forecasting the impacts of climate variability, where it is important to include to effects of forcing variables (e.g., temperature and precipitation) even of such effects are difficult to identify. Indeed, 427 we failed to detect strong climate effects in the QBM for B. gracilis and Poa secunda, but includ-428 ing climate covariates still improved forecasting skill (Fig. 2). If a species is truly unresponsive 429 to a given climate variable, statistical regularization techniques will shrink the mean and variance 430 of a covariate estimate toward zero (Hooten and Hobbs 2015). Of course, no matter what model 431 selection approach is adopted, a critical step is identifying the appropriate candidate covariates, 432 which we attempted to do based on our knowledge of this semi-arid plant community. However, 433 the climate covariates we chose required aggregating daily weather data over discrete time peri-434 ods. It is possible that we did not choose the optimal time periods over which to aggregate. New 435 methods using functional linear models (or splines) may offer a data-driven approach for identi-436 fying the appropriate time periods over which to aggregate to produce a tractable set of candidate

climate variables (Teller et al. 2016).

We also expected IPM forecast accuracy to decline at a lower rate than the QBM as the forecast 439 horizon increased. In principle, more mechanistic models should produce better predictions, 440 especially under novel conditions (Evans 2012, Schindler and Hilborn 2015). In our case, the 441 IPM explicitly models the influence of weather on recruitment and survival, effects that may be 442 poorly represented in the QBM because recruitment and survival mainly affect small plants that 443 contribute little to year-to-year changes in percent cover. Over time, of course, the addition and subtraction of small plants can have large effects on population growth, so explicitly modeling 445 these effects could contribute to a longer forecast horizon. However, we found no evidence for a 446 difference between the IPM and QBM forecast horizons (Fig. 4). In conclusion, we found that models based on individual-level demographic data generally failed to generate more skillful population forecasts than models based on population-level data. This 449 finding runs counter to our expectations, but is consistent with recent theoretical (Freckleton et 450 al. 2011) and empirical work (Queenborough et al. 2011). Moreover, for the two species where 451 including climate covariates improved forecast skill, both models generated equally skillful one-452 step-ahead forecasts. Thus, we conclude that models based on population-level data, rather than 453 individual-level data, may be adequate for forecasting the states and dynamics of plant popu-454 lations. Unfortunately, our analysis, where climate effects were relatively unimportant in vital 455 rate regressions, did not allow us to sufficiently test our prediction that individual-level data is 456 neccessary to generate skillful forecasts if different vital rates respond to climate in unique, po-457 tentially opposing, ways. Nonetheless, our results are encouraging for the use of easy-to-collect 458 population-level data for forecasting the state of plant populations. 459

# Acknowledgments

This work was funded by the National Science Foundation through a Postdoctoral Research
Fellowship in Biology to ATT (DBI-1400370), award MSB-1241856 to MBH, and a CAREER

award to PBA (DEB-1054040). We thank the original mappers of the permanent quadrats in

Montana and the digitizers in the Adler lab, without whom this work would not have been possible. Informal conversations with Stephen Ellner, Giles Hooker, Robin Snyder, and a series of
meetings between the Adler and Weecology labs at USU sharpened our thinking. Brittany Teller
provided comments that improved our manuscript. Compute, storage and other resources from
the Division of Research Computing in the Office of Research and Graduate Studies at Utah
State University are gratefully acknowledged. Any use of trade, firm, or product names is for
descriptive purposes only and does not imply endorsement by the U.S. government.

# 471 Tables

Table  $\underline{\text{1: Description of data.}}$  The observations span 13 year-to-year transitions.

Species	Vital Rate Model	Num. Obs.	Num. Quadrats
B. gracilis	Growth	5670	29
	Survival	10102	33
	Recruitment	304	33
	Percent cover	281	29
H. comata	Growth	1990	16
	Survival	3257	18
	Recruitment	304	18
	Percent cover	171	17
P. smithii	Growth	8052	19
	Survival	11344	19
	Recruitment	304	19
	Percent cover	217	19
Poa secunda	Growth	3018	18
	Survival	4650	18
	Recruitment	304	18
	Percent cover	197	18

# Figures

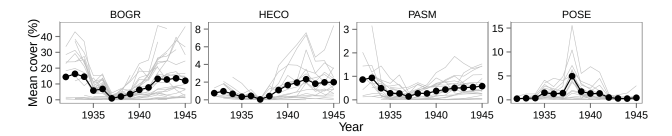


Figure 1: Time series of average percent cover over all quadrats for our four focal species: *Bouteloua gracilis* (BOGR), *Hesperostipa comata* (HECO), *Pascopyrum smithii* (PASM), and *Poa secunda* (POSE). Light grey lines show trajectories of individual quadrats. Note the different y-axis scales across panels.

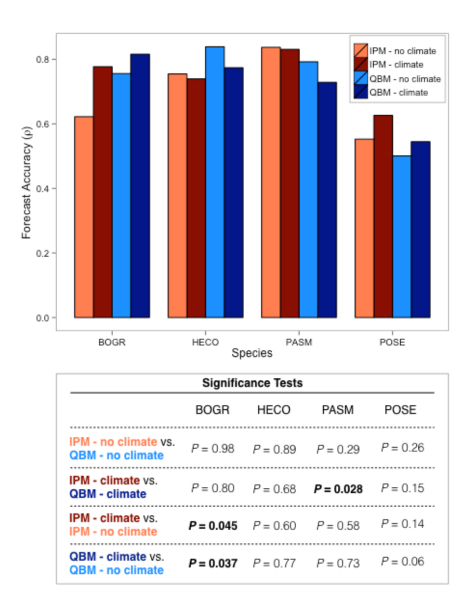


Figure 2: Comparisons of one-step-ahead, out-of-sample forecast accuracy between the IPM and QBM models with and without the inclusion of climate covariates. For each comparison, *P*-values are from one-sided *t* tests designed to assess whether the first model in the comparison statement had higher accuracy than the second model in the comparison statement (see details in Table S22). In no case did adding climate covariates decrease forecast accuracy (Table S21). Species codes are as in Fig. 1.

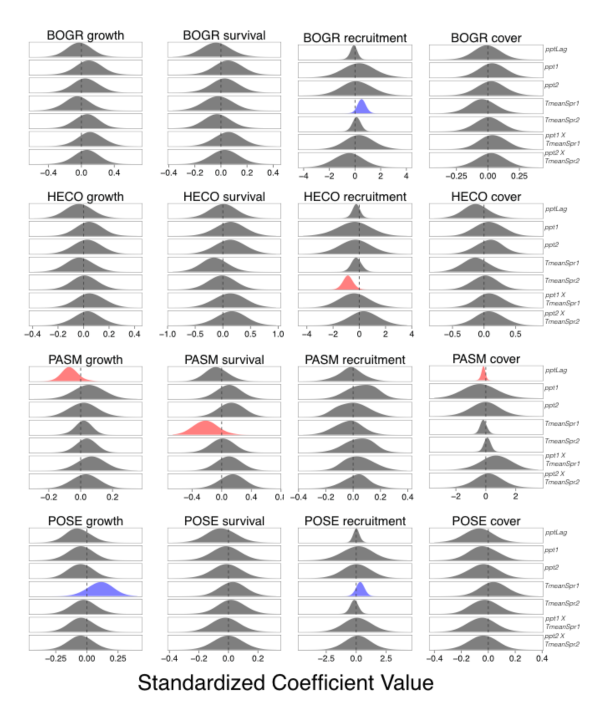


Figure 3: Posterior distributions of climate effects ( $\beta_C$ ) for each species and vital rate statistical model. Because our priors were constrained via ridge-regression, we highlight climate effects whose 80% credible intervals do not overlap zero (red for negative coefficients, blue for positive coefficients). Kernel bandwidths of posterior densities were adjusted by a factor of 4 for visual clarity. Species codes are as in Fig. 1.

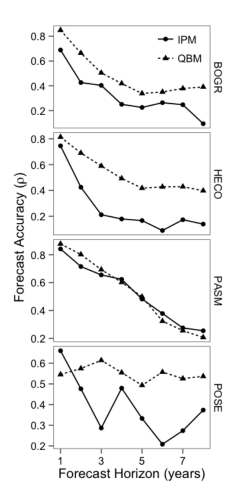


Figure 4: The forecast horizons for both models with climate covariates included. Points show the average accuracy ( $\rho$ ) across all forecasts at a given time horizon. We only examine the forecast accuracy of models with climate covariates included because in no case did including climate covariates significantly decrease accuracy (see Fig. 2). Species codes are as in Fig. 1.

# References

- Adler, P. B., H. J. Dalgleish, and S. P. Ellner. 2012. Forecasting plant community impacts of
- climate variability and change: when do competitive interactions matter? Journal of Ecology
- 476 100:478-487.
- Adler, P. B., S. P. Ellner, and J. M. Levine. 2010. Coexistence of perennial plants: An embarrass-
- ment of niches. Ecology Letters 13:1019–1029.
- Anderson, J., L. Vermeire, and P. B. Adler. 2011. Fourteen years of mapped, permanent quadrats
- in a northern mixed prairie, USA. Ecology 92:1703.
- <sup>481</sup> Chu, C., and P. B. Adler. 2015. Large niche differences emerge at the recruitment stage to stabi-
- lize grassland coexistence. Ecological Monographs 85:373–392.
- Chu, C., K. M. Havstad, N. Kaplan, W. K. Lauenroth, M. P. McClaran, D. P. Peters, L. T. Ver-
- meire, and P. B. Adler. 2014. Life form influences survivorship patterns for 109 herbaceous
- perennials from six semi-arid ecosystems. Journal of Vegetation Science 25:947–954.
- <sup>486</sup> Chu, C., A. R. Kleinhesselink, K. M. Havstad, M. P. McClaran, D. P. Peters, L. T. Vermeire,
- 487 H. Wei, and P. B. Adler. 2016. Direct effects dominate responses to climate perturbations in
- grassland plant communities. Nature Communications 7.
- <sup>489</sup> Clark, J. S., and O. N. Bjørnstad. 2004. Population time series: Process variability, observation
- errors, missing values, lags, and hidden states. Ecology 85:3140–3150.
- <sup>491</sup> Clark, J. S., D. M. Bell, M. H. Hersh, M. C. Kwit, E. Moran, C. Salk, A. Stine, D. Valle, and K.
- <sup>492</sup> Zhu. 2011. Individual-scale variation, species-scale differences: Inference needed to understand
- diversity. Ecology Letters 14:1273–1287.
- <sup>494</sup> Clark, J. S., D. M. Bell, M. Kwit, A. Stine, B. Vierra, and K. Zhu. 2012. Individual-scale infer-
- ence to anticipate climate-change vulnerability of biodiversity. Philosophical Transactions of the
- 496 Royal Society B: Biological Sciences 367:236–246.
- <sup>497</sup> Clark, J. S., D. Bell, C. Chu, B. Courbaud, M. Dietze, M. Hersh, J. HilleRisLambers, I. Ibáñez, S.
- LaDeau, S. McMahon, J. Metcalf, J. Mohan, E. Moran, L. Pangle, S. Pearson, C. Salk, Z. Shen,
- 499 D. Valle, and P. Wyckoff. 2010. High-dimensional coexistence based on individual variation: a
- synthesis of evidence. Ecological Monographs 80:569–608.
- Clark, J. S., S. R. Carpenter, M. Barber, S. Collins, A. Dobson, J. A. Foley, D. M. Lodge, M.
- Pascual, R. Pielke, W. Pizer, C. Pringle, W. V. Reid, K. A. Rose, O. Sala, W. H. Schlesinger, D.
- H. Wall, and D. Wear. 2001. Ecological forecasts: an emerging imperative. Science (New York,
- 504 N.Y.) 293:657-660.
- <sup>505</sup> Cribari-Neto, F. 2004. Asymptotic inference under heteroskedasticity of unknown form. Compu-
- tational Statistics and Data Analysis 45:215–233.
- Dalgleish, H. J., D. N. Koons, M. B. Hooten, C. A. Moffet, and P. B. Adler. 2011. Climate influ-
- ences the demography of three dominant sagebrush steppe plants. Ecology 92:75–85.
- Ellner, S. P., and M. Rees. 2006. Integral projection models for species with complex demogra-
- phy. The American naturalist 167:410–428.

- Evans, M. R. 2012. Modelling ecological systems in a changing world. Philosophical transac-
- tions of the Royal Society of London. Series B, Biological sciences 367:181–190.
- Freckleton, R. P., W. J. Sutherland, A. R. Watkinson, and S. A. Queenborough. 2011. Density-
- structured models for plant population dynamics. American Naturalist 177:1–17.
- <sup>515</sup> Gelman, A., J. Hwang, and A. Vehtari. 2014. Understanding predictive information criteria for
- 516 Bayesian models. Statistics and Computing 24:997–1016.
- Gerber, B. D., W. L. Kendall, M. B. Hooten, J. A. Dubovsky, and R. C. Drewien. 2015. Optimal
- population prediction of sandhill crane recruitment based on climate-mediated habitat limitations.
- Journal of Animal Ecology 84:1299–1310.
- Hooten, M. B., and N. T. Hobbs. 2015. A guide to Bayesian model selection for ecologists. Eco-
- logical Monographs 85:3–28.
- Lauenroth, W. K., and P. B. Adler. 2008. Demography of perennial grassland plants: Survival,
- life expectancy and life span. Journal of Ecology 96:1023–1032.
- Luo, Y., K. Ogle, C. Tucker, S. Fei, C. Gao, S. LaDeau, J. S. Clark, and D. S. Schimel. 2011.
- Ecological forecasting and data assimilation in a data-rich era. Ecological Applications 21:1429–
- 526 1442.
- Perretti, C. T., S. B. Munch, and G. Sugihara. 2013. Model-free forecasting outperforms the
- 528 correct mechanistic model for simulated and experimental data. Proceedings of the National
- Academy of Sciences of the United States of America 110:5253–5257.
- Petchey, O. L., M. Pontarp, T. M. Massie, S. Kéfi, A. Ozgul, M. Weilenmann, G. M. Palamara, F.
- Altermatt, B. Matthews, J. M. Levine, D. Z. Childs, B. J. McGill, M. E. Schaepman, B. Schmid,
- P. Spaak, A. P. Beckerman, F. Pennekamp, and I. S. Pearse. 2015. The ecological forecast hori-
- zon, and examples of its uses and determinants. Ecology Letters 18:597–611.
- Queenborough, S. A., K. M. Burnet, W. J. Sutherland, A. R. Watkinson, and R. P. Freckleton.
- 2011. From meso- to macroscale population dynamics: A new density-structured approach.
- 536 Methods in Ecology and Evolution 2:289–302.
- R Core Team. 2013. R: A language and environment for statistical computing.
- Rees, M., and S. P. Ellner. 2009. Integral projection models for populations in temporally varying
- environments. Ecological Monographs 79:575–594.
- 540 Schindler, D. E., and R. Hilborn. 2015. Prediction, precaution, and policy under global change.
- 541 Science 347:953–954.
- Stan Development Team. 2014a. Stan: A C++ Library for Probability and Sampling, Version
- 543 2.5.0.
- Stan Development Team. 2014b. Rstan: the R interface to Stan, Version 2.5.0.
- 545 Sæther, B. E., S. Engen, V. Grøtan, W. Fiedler, E. Matthysen, M. E. Visser, J. Wright, A. P.
- Møller, F. Adriaensen, H. Van Balen, D. Balmer, M. C. Mainwaring, R. H. McCleery, M. Pampus,
- and W. Winkel. 2007. The extended Moran effect and large-scale synchronous fluctuations in the
- size of great tit and blue tit populations. Journal of Animal Ecology 76:315–325.

- Taylor, C. M., and A. Hastings. 2004. Finding optimal control strategies for invasive species: a density-structured model for Spartina alterniflora. Journal of Applied Ecology 41:1049–1057.
- Teller, B. J., P. B. Adler, C. B. Edwards, G. Hooker, R. E. Snyder, and S. P. Ellner. 2016. Linking demography with drivers: climate and competition. Methods in Ecology and Evolution 7:171–553 183.
- Vehtari, A., A. Gelman, and J. Gabry. 2016. Efficient implementation of leave-one-out cross-validation and WAIC for evaluating fitted Bayesian models. ArXiv preprint.
- Wilcox, R. R. 2009. Comparing Pearson Correlations: Dealing with Heteroscedasticity and Nonnormality. Communications in Statistics - Simulation and Computation 38:2220–2234.
- Ye, H., R. J. Beamish, S. M. Glaser, S. C. H. Grant, C.-h. Hsieh, L. J. Richards, J. T. Schnute, and G. Sugihara. 2015. Equation-free mechanistic ecosystem forecasting using empirical dynamic modeling. Proceedings of the National Academy of Sciences 112:E1569–E1576.