- [Working title]: Intraspecific variation in range-wide seed bank dynamics is not consistent with density-independent model of bet hedging
  - Gregor-Fausto Siegmund and Monica Geber
- Last updated: March 27, 2021

# Writing list

- 1. Update Figure 1 to focus on components of life cycle discussed in this MS.
- 2. Revise Figure 3 to talk about seeds being intact rather than persistent
- 3. Revise description of seed bag experiment to include timing of experiment, temporal focus, and spatial design
- 4. Revise Figure 4 to increase point size
- 5. Revise Figure 5 and 6 to change scale of geometric SD RS axis, change label size in observed vs. predicted plot, change axes in observed vs. predicted plot, move plots of germination probability elsewhere in MS
- 6. Update citation list
- 7. Read Cohen & Ellner to identify the role of complete reproductive failure in the original models for the evolution of bet hedging
- 8. Write about strong and weaker test of the hypothesis (partial pooling, no partial pooling)

- 9. Write paragraph about history of studies of bet hedging via seed bank, with emphasis on how this study addresses this question at an intraspecific level. Lower level of variation in intraspecific germination fraction. (link)
- 21 10. Write and implement model checking process

## Introduction

Seed banks can buffer plant populations against environmental change and stochasticity (Eager et al. (2014); Paniw et al. (2017)), increase effective population size (Nunney (2002); 23 Waples (2006)), and maintain genetic diversity (McCue and Holtsford (1998)). Dormancy can affect the outcome of evolution (Heinrich et al. (2018); Ritland (1983)). Theory thus suggests that seed banks have ecological and evolutionary consequences (Evans and Dennehy (2005)). What drives the evolution of delayed germination? The theory developed by Cohen 28 (1966) frames the problem in the following terms. What is the optimal germination fraction for a given level of interannual variation in fitness and seed survivorship? These models make it clear that the germination fraction that maximizes long-term population growth 31 rate is a function of the distribution of fitness (characterized by the variation in fitness), the fitness values, and the rate of seed survivorship. For a given mean fitness, increasing the variance in fitness decreases the optimal germination fraction. Increasing seed survivorship decreases the optimal germination fraction, and the degree to which it does so depends on the 35 probability of a 'good year'. Specifically, as the probability of a high-fitness year decreases, 36 the optimal germination fraction decreases. Bet hedging should evolve to maximize the longterm geometric population growth rate (as compared to the arithmetic population growth rate) (Cohen (1966, 1968); Ellner (1985a,b)). 39 Cohen (1966) emphasizes the role of particularly bad years. This is highlighted by the 40 inequality in equation (12), which states that for the optimal germination strategy to be 41 bet hedging, it is sufficient that the harmonic mean is less than the survival probability of seeds that do not germinate. The minimum fitness has a strong impact on the harmonic mean of fitness (see definition of harmonic mean on Wikipedia), and a single year of very low fitness would tend to make the harmonic mean small. In particular, the harmonic mean 46 is 0 when any of its values is 0. The probability of complete reproductive failure is thus
47 particularly important. Years with no seedling survival in the plots or no germination are
48 those in which fitness would be zero; there are only 4/20 populations in which this does not
49 happen. We fit two types of models to the aboveground data; one with partial pooling and
50 one with no pooling. The first was an attempt to correct for sampling variation; the latter an
51 attempt to estimate per-capita reproductive success as-is, providing a more extreme estimate
52 of interannual variation but one that reflects the combined effect of sampling variation and
53 true variation.

Revisit theory to see particular role of complete reproductive failure versus low fitness years more generally.

In paragraph on theory, discuss density-independent models Cohen (1966, 1968), densitydependent models Ellner (1985*a*,*b*), predictive germination ?, and unstructured versus structured models ? (Tuljaparkur 1993, Valleriani and Tielborger 2006).

To empirically test this theory, "a density-independent model can be used to check quantitatively the optimality (or evolutionary stability) of a life history trait in a real population, because the density effects are manifested in the measured vital rates" and "to check if a species germination fraction is optimal, one would estimate seed survivorship and the probability distribution of per capita seed yield. From these, the DI model predicts an "optimal" germination fraction, which can be compared with the actual germination fraction." Ellner (1985a).

Several studies have examined intraspecific variation in seed dormancy ????. The main set of papers I've included are ones that look at variation in germination among Sonoran Desert annuals (Gremer and Venable (2014); Venable (2007); ?). The table lists predictions made by different models. I think examining the following relationships would be good starting points: The correlation between variance in fitness (seeds/seedling) and germination fraction should be negative – this is true under the density-independent and -dependent

model. The correlation between seed survivorship and germination fraction should be negative under both a density-independent and -dependent model but the limit as survivorship approaches 1 differs. Finally, the correlation between mean seed yield and germination fraction will be positive if fitness is density-independent but is not necessarily positive if fitness is density-dependent.

#### Paragraph about the history of empirical studies of bet hedging.

77

Population vital rates are known to exhibit intraspecific variation across Clarkia xan-78 tiana's geographic range, including age-specific germination and seed survival (Eckhart et al. 79 (2011)). In a study of C. xantiana population dynamics that identified a decline of long-term 80 stochastic population growth rate from west to east across the range, Eckhart et al. (2011) 81 inferred a decrease in survival through winter  $(s_1)$  and an increase in germination rate  $(q_1)$ 82 of first-year seeds from west to east. Although, Lewis (1962) suggested that Clarkia pop-83 ulations undergo local extinctions and lack a long-lived seed bank, observations of 20 populations across the range are not consistent with that hypothesis (Figure 2). Demographic observations Eckhart et al. (2011) and transplant experiments demonstrate that fitness can exhibit dramatic interannual variation (e.g. 30-fold between a wet and dry year in Geber and Eckhart (2005)).

Seed dormancy and persistence in the soil seed bank may be a bet-hedging strategy
that is favored by environmental uncertainty. If this is the case in *C. xantiana*, we might
expect increased seed survival and decreased germination probabilities in populations with
more variability in per-capita reproductive success. However, broad geographic patterns in
rainfall variability and life history are not consistent with this cursory explanation since
variation in rainfall increases from west to east, and germination probability also increases
Eckhart et al. (2011). We thus sought to understand intraspecific variation in *C. xantiana*seed vital rates in the context of broader life history patterns.

Bet hedging should evolve to maximize the long-term geometric population growth rate

(as compared to the arithmetic population growth rate) (Cohen (1966, 1968); Ellner (1985 a,b)).

Seed banks are more likely to be selected in populations which experience higher levels of
interannual variation in per-capita reproductive success. To investigate this empirical relationship, we estimate the correlation between germination and seed survival, and between
germination variance in per-capita reproductive success. We also examined whether ob-

 $_{103}$  served levels of germination are consistent with germination levels predicted by a density-

independent model of bet hedging.

Genetic basis of seed banking; traits associated with seed banks; clines in seed banks/genetic variation. Cline associated with summer temperature Wagmann 2012

Altitudinal clines: more dormancy in seeds produced with lower annual temperature and precipitation Fernandez-Pascual 2012; low elevation pops have high germination fractions but high elevation pops require stratification to attain high germination fractions Gremer 2020, high

the correlation between interannual variation in per-capita reproductive success and the proportion of seeds that germinate in the winter immediately following seed production.

Based on bet hedging models, we predict that germination is negatively correlated with interannual variation in per-capita reproductive success. [What is the relationship between interannual variation in fitness and dormancy? This is a question about whether fitness variation and dormancy are positively correlated as expected? this is what selects for bet hedging.]

A previous study with *Clarkia xantiana* suggests that the soil seed bank is important for population dynamics in *Clarkia xantiana*.

A separate set of seed burial experiments suggests that seeds of *C. xantiana* can remain viable in the soil for at least 10 years (Moeller personal communication). In the study of *C. xantiana* population dynamics that showed a decline of long-term stochastic population growth rate from west to east across the range, Eckhart et al. (2011) inferred a decrease in

survival through winter (s1) and an increase in germination rate (g1) of first-year seeds from west to east.

Population growth rates determine species abundance and distribution, and are ulti-126 mately what limit persistence beyond range edges. Geographic patterns to vital rates have 127 so far been studied to help understand the demography of geography. Seed banks are a strat-128 egy that annual plants may use to buffer against environmental variation and may be part of 129 population persistence. I will begin by characterizing geographic variation in belowground 130 vital rates. [What is the geographic pattern to variation in germination or seed survival?] 131 I think this question could be expanded to make clear predictions and/or address another 132 aspect such as variation in time. 133

Eckhart et al. 2004 conducted a transplant experiment in two years in which seed set varied ten-fold. There are additional studies that did elimination experiments (with other species of Clarkia) and suggested limited role/role for seed bank because plants did or did not come up with following year.

134

135

136

137

145

146

147

148

149

In 13 years of data on 20 populations across the species range, we observed high seedling mortality and/or low germination in at least one year in all but four populations. Although most populations had at least some flowering plants in most years, there were also four populations in which no flowering plants were observed in the entire population. None of these populations, however, went extinct during the period of the study. Seeds can remain viable in the field and laboratory for at least 5 years, contrary to previous suggestions that seeds are unlikely to persist past a few years (Lewis 1962).

See also the comments in Lewis 1953 on lifespan of seeds, absence of dormancy, and cued germination. Lewis 1963 Presidential address at SSE on catastrophic selection is also relevant. Discusses extinction and possible absence of a seed bank in populations of Clarkia species. Specifically mentions a population in the upper Kern Valley that had both whites and pinks; the pinks went extinct during a drought year while the whites maintained their

population size. This work was followed up by others on different species in Clarkia genus. I think its important to consider the possibility that C. xantiana being studied in 1963 was a xantiana/parviflora ssp. which may have different dynamics than the outcrossing xantiana/x-antiana. Furthermore: the upper parts of the range have the highest rates of germination which may also trade off with low survival. It would be interesting to put that observation in the range-wide context!

Knies et al. 2004 studied 2 C. xantiana populations; seeds of outcrosser are heavier than 156 selfers. Identify heavier seeds ( .1 mg heavier at SC than WH population). Seeds from 157 both populations were bullet-shaped but differ in shape of exotesta cells; WH cells are gourd 158 shaped, SC shapes are even width and spherical. Seed size/number trade-off: smaller size for 159 dispersal; greater mass increases germination probability. Smaller seed size also facilitates 160 seed integration into soil seed bank. Small seed size is also correlated with a persistent 161 seed bank (Thompson et al. 1993). Larger seeds may germinate under more conditions and 162 perhaps have smaller seed bank. 163

Moeller et al. 2011 population genetic study.

Eckhart and Geber 1999 identify more populations than those of Lewis, populations that they have since monitored intensively.

Price et al. 1985 followed a population for 2 years, did an elimination experiment. determined a dormant seed bank. Snow 1960 reports eliminating plants and not observing plants the following year.

Geber and Eckhart 2005 observed a 30-fold difference in lifetime fitness [calculated as number of germinants times survival, flower number, fruit set and seed set] between two transplant years (one wet, one dry). Fitness was very low in the dry year of the study (0.02-0.03 seeds per seed planted). They observed a 2-4 fold difference between years in germination, seedling survival, and flower number. They did not observe a difference in fruit or seed set. Western populations showed evidence of regional adaptation: survival and

flower number. Germination declined from the western to eastern transplant sites [in contrast to what we observe here?] Germination is consistently higher in subspecies xantiana than parviflora. From west to east, rainfall declines and becomes more variable. Low survivorship is the product of end-of-season drought, small mammal herbivory (Geber and Eckhart 2005, Benning et al. 2019), competition by higher vegetation cover of grasses (survivorship increases with plant size), fungal rust infection. Plants from western populations are larger. Note that the transplant experiments took place on igneous soils, and later work by Vince has shown that soil type is important.

Eckhart et al. 2010 IJPS: geographic patterns in precipitation, decline from west to east, 184 mirrored geographic pattern in midday water potential (reduced performance). Metasedi-185 mentary soils were more resistant and finer. Soil characteristic curves indicated that metased-186 imentary soils can hold more water but that it becomes unavailable to plants at higher water 187 contents. Higher water potential on steeper, eroding slopes; this soil was softer and coarser. 188 Decline in soil water potential (more negative) decreases plant size and fruit number. How-189 ever, soil water potential is unrelated to seedling survival in the study years. Metasedimen-190 tary rocks may compound effect of decline in precipitation. Metasedimentary rocks may 191 hold more water but it may be more available to plants in igneous soils. 192

Bet hedging should evolve to maximize the long-term geometric population growth rate 193 (as compared to the arithmetic population growth rate) (Cohen (1966, 1968); Ellner (1985 a, b)). Seed banks are more likely to be selected in populations which experience higher levels of in-195 terannual variation in per-capita reproductive success. To investigate this empirical relation-196 ship, I will estimate the correlation between interannual variation in per-capita reproductive 197 success and the proportion of seeds that germinate in the winter immediately following seed 198 production. I predict that germination is negatively correlated with interannual variation in 199 per-capita reproductive success. [What is the relationship between interannual variation in 200 fitness and dormancy? This is a question about whether fitness variation and dormancy are 201

positively correlated as expected? this is what selects for bet hedging.]

Seed dormancy and persistence in the soil seed bank may be a bet-hedging strategy
that is favored by environmental uncertainty. If this is the case in *C. xantiana*, I think we
might expect increased seed survival and decreased germination at the eastern range edge
(precipitation is more variable at eastern populations in winter and spring). This seems to be
opposite of what was observed in the previous study? though I could also be misinterpreting
this. This is could be one reason to revisit this question with a new analysis.

Why using the Clarkia system? We expect to see differences among populations in either seed survivorship or variance in fitness. In the study of *C. xantiana* population dynamics that showed a decline of long-term stochastic population growth rate from west to east across the range, Eckhart et al. 2011 inferred a decrease in survival through winter (s1) and an increase in germination rate (g1) of first-year seeds from west to east. [need to be clear about this] also know that the environment changes across the range; that will affect the mean fitness

I expect bet hedging stages to be sensitive to variation in an environmental cue. For 216 something to be an adaptive strategy, it should respond to variation in the environment to 217 capitalize on good years such as ones with high precipitation. To investigate the sensitivity of vital rates to the environment, I estimate the slope of the relationship between environmental cues and vital rates at individual sites using a random coefficients model that estimates covariation between intercepts and slopes. I predict that I will estimate variation both 221 in the intercept and slope for populations, and that variation in the modeled cue to be 222 determined will be positively related to the estimated slope. What is the relationship 223 between dormancy and environmental cues? This is a question about whether we can make 224 any inferences about the mechanism responsible for bet hedging in this system. 225

# 1 Methods

### 1.1 Clarkia life history

Clarkia xantiana is a winter annual that germinates with late fall and winter rains. In our 226 study region, the Kern Valley in the southern Sierra Nevada Mountains, historically happens 227 between October and late February or early March. Seedlings grow throughout the winter 228 and spring, and surviving plants flower in late spring and early summer, April into early 229 July. Pollinated fruits set seed in the early summer, June to July. Seeds of C. xantiana are produced in early summer, with fruits that dry out and gradually split open. Most seeds 231 appear to be shed from fruits within 3-4 months after production, but can remain on the 232 plant for up to a year. Seeds are small (< 1 mm in width) and have no structures to aid in 233 aerial dispersal. 234

We represent the Clarkia xantiana as a life cycle graph (Figure 1A) that describes tran-235 sitions from October of year t to October of year t+1 in terms of underlying vital rates. 236 The census period occurs when the entire population is seeds, and corresponds to the time 237 at which seed bags are placed into the field (see below). Seeds are grouped into three stages: 238 age 0 seeds, which were produced in the current year; age 1 seeds, which were produced in 239 the previous year; age 2+ seeds, which were produced two or more years ago. Persistence of 240 seeds in the seed bank is represented by transitions from younger to older seeds. Production 241 of new seeds is captured by transition to the age 0 seed state. 242

Transitions in the life cycle graph are the product of age-specific seed survival and germination, and aboveground seedling survival to fruiting, fruit production, and seeds per fruit. Seed-related rates are represented separately for age 0, 1, and 2+ seeds. Germination for each age class is given as  $g_1$ ,  $g_2$ , and  $g_3$ , respectively. Seed survival from seed production to the first October is given as  $s_0$ , and survival from October to February is given as  $s_1$ ,  $s_3$ , and  $s_5$  for age 0, 1, and 2+ seeds, respectively. Survival from February to October is given as  $s_2$ ,  $s_4$ , and  $s_6$  for age 0, 1, and 2+ seeds, respectively. We assume that vital rates remain unchanged after age 2. We also we assume that all plants experience the same vital rates upon germination seed age at germination does not affect seedling survival to fruiting  $(\sigma)$ , fruits per plant (F), or seeds per fruit  $(\phi)$ .

The life cycle graph (Figure 1A) corresponds to the annual projection matrix

$$\mathbf{A} = \begin{bmatrix} s_1 g_1 \sigma F \phi s_0 & s_3 g_2 \sigma F \phi s_0 & s_5 g_3 \sigma F \phi s_0 \\ s_1 (1 - g_1) s_2 & 0 & 0 \\ 0 & s_3 (1 - g_2) s_4 & s_5 (1 - g_3) s_6 \end{bmatrix}$$
(1)

that summarizes transitions between stages.

#### 1.2 Creating the dataset

demography for 20 populations of Clarkia xantiana (Table 1). Specifically, we used ex-255 periments to estimate transitions in the seed bank and surveys to estimate components of 256 per-capita reproductive success. These demographic data have previously been used to test 257 hypotheses about the geography of demography (Eckhart et al. (2011)) and species distri-258 butions (Pironon et al. (2018)). Here, we sought to obtain population-level estimates of 259 germination and seed survival, and yearly estimates of per-capita reproductive success. 260 To estimate transitions in the seed bank, we used observations from a seed bag burial 261 experiment conducted in all populations from 2006-2009 (Figure 3). The experiment has 262 been previously described in Eckhart et al. (2011) and we reanalyze the data here. Geber 263 and collaborators buried seeds in bags and unearthed 1, 2, or 3 years after being buried to 264 count seedlings and intact, viable seeds. The experiment was repeated in 3 consecutive years 265 and ended in 2009. We thus have 3 sets of observations associated with 1 year old seeds, 266

We used field experiments and surveys to assemble observations of below- and above-ground

2 sets of observations associated with 2 year old seeds, and 1 set of observations associated with 3 year old seeds. We use data from the experiment to estimate age-specific germination 268 and seed survival (see Joint model for seed vital rates) but note that we test predictions of bet-hedging theory that are based on an unstructured seed bank and use only the relevant subset of transitions in our analysis (see Computing vital rates).

269

270

271

272

273

274

275

276

277

278

289

281

To estimate per-capita reproductive success, we combine censuses of seedlings and fruiting plants, surveys of fruits per plant, and lab counts of seeds per fruit. To assess the survival of seedlings to fruiting plants, we counted seedlings and fruiting plants in  $30~0.5~\mathrm{m}^2$  permanent plots from 2006–2018 (Eckhart et al. (2011)). To assess seed production by plants that survive to reproduction, we counted fruits per plant on individual plants in permanent plots, and on additional haphazardly chosen plants throughout the population. We then attempted to obtain 20-30 fruits per population, which we used to count seeds per fruit (Eckhart et al. (201.1)).

Table 1: Summary of data sets used to estimate parameters.

Parameter data	Description	Data set	Time span
SEED VITAL RATES	_		_
Seed survival and germination	Seed bag burial	$\mathbf{Y}_1$	2006-2009
Seed viability	Viability trials	$\mathbf{Y}_2$	2006-2009
SEEDLING SURVIVAL		_	
Seedling survival to fruiting	Field surveys	$\mathbf{Y}_4$	2006-2019
FRUITS PER PLANT		_	
Total fruit equivalents per plant	Field surveys	$\mathbf{Y}_5$	2006-2012
Undamaged and damaged fruits per plant	Field surveys	$\mathbf{Y}_{6}$	2013-2019
Total fruit equivalents per plant	Extra plots	$\mathbf{Y}_7$	2006-2012
Undamaged and damaged fruits per plant	Extra plots	$\mathbf{Y}_8$	2013-2019
SEEDS PER FRUIT		_	
Seeds per undamaged fruit	Lab counts	$\mathbf{Y}_9$	2006-2019
Seeds per damaged fruit	Lab counts	$\mathbf{Y}_{10}$	2013-2019

#### 1.3 Model

We use observational and experimental data from 20 populations of *Clarkia xantiana* to estimate transition probabilities across the life cycle. We fit multilevel models to obtain population-specific estimates for belowground vital rates, and year- and population-specific estimates for aboveground vital rates. Because we were interested in describing the life histories of individual populations, we built separate models for each population. Our general approach applies a common model structure to partially pool observations from each population.

We first explicitly describe our formulation in terms linear mixed models before defining the joint posterior (Evans et al. (2010); Ogle and Barber (2020)). We assume that the latent mean of observations in year j at a population k,  $\theta_{jk}$ , is drawn from a normal distribution with mean  $\theta_{0,k}$  and variance  $\sigma_j^2$ .

$$\theta_{jk} = \theta_{0,k} + \epsilon_{(jk)}. \tag{2}$$

Our model includes a population-level intercept  $\theta_{0,k}$  and random effects  $\epsilon_{(jk)}$ . The random effects can be written as  $\epsilon_{(jk)} \sim N(0,\varsigma^2)$ . For the moment, we focus on describing the hierarchical structure of the model but note that we use link functions for transformation to parameters that are appropriate for the likelihoods we use to model different sets of observations (e.g. binomial for seed bag experiments; Poisson for counts of seed per fruit). We note that such a linear mixed effects model with random intercepts for years is one method commonly used to model interannual variation in demographic rates (e.g. Metcalf et al. (2015)). Using hierarchical centering, the same model is rewritten as

$$\theta_{jk} = \alpha_{(jk)}. (3)$$

The mean  $\theta_{jk}$ , is now drawn from a normal distribution with mean  $\alpha_{(jk)}$  and variance  $\sigma_j^2$ . We place a prior on  $\alpha_{(jk)}$  such that  $\alpha_{(jk)} \sim N(\theta_{0,k}, \varsigma^2)$ . The expressions are related by  $\alpha_{(jk)} = \theta_{0,k} + \epsilon_{(jk)}$ . We thus draw year-level means from the population-level means.

For a single population (ie. suppressing subscript k), we write the posterior proportional to the joint distribution as

$$[\theta_j, \theta_0, \sigma_j^2, \varsigma^2 | y_{ij}] \propto [y_{ij} | \theta_j, \sigma_j^2] [\theta_j | \theta_0, \varsigma^2] [\theta_0] [\sigma_j^2] [\varsigma^2]. \tag{4}$$

The distribution of the observations  $y_{ij}$  is conditional on the year-specific parameters  $\theta_j$  and  $\sigma_j^2$ . In turn, the year-specific parameter  $\theta_j$  is conditional on the population-specific parameters  $\theta_0$  and  $\varsigma^2$ . We placed priors on all parameters found only on the right hand side of conditional statements  $(\theta_0, \sigma_j^2, \varsigma^2)$ . In practice, we implemented this model by specifying the population- and year-levels of the model with normal distributions; for example,  $[\theta_j|\theta_0, \varsigma^2]$  is  $\theta_j \sim N(\theta_0, \varsigma^2)$ . The model thus describes a structure in which years are nested within populations.

# 1.4 Model statements, implementation, and fitting

We include the expression for the posterior proportional to the joint distribution, and corresponding directed acyclic graphs, in Appendix: Joint Posterior. Priors for all parameters are defined in Table Priors. We applied the following principles for specifying priors: (1) we used weakly informative priors that avoided placing probability mass on biologically implausible values (Gelman Lemoine (2019); Wesner and Pomeranz (2020)), (2) we placed positive, unbounded priors on variance components (REF), (3) we conducted prior predictive checks to assess the scale of priors after parameter transformation (Gabry et al. (2019); Hobbs and Hooten (2015); Wesner and Pomeranz (2020)), and (4) we simulated prior predictive distributions to confirm that the joint likelihood generated data within the observed range (Conn et al. (2018); Gabry et al. (2019); Hobbs and Hooten (2015)). We provide additional detail regarding our choice of priors in Appendix: Priors.

We prepared data for analysis using the tidyverse and tidybayes packages (CITE) in R

VERSION; CITE. We wrote, fit all models, and estimated posterior distributions using

JAGS VERSION with rjags (Plummer 2016). We randomly generated initial conditions for

all parameters with a prior by drawing from the corresponding probability distribution in R

before passing the initial values to rjags. We ran three chains for XX,000 iterations. The first

XX,000 samples were discarded as burn-in and we sampled the following XX,000 iterations.

We did not thin the chains (Elderd and Miller (2015)).

We assessed convergence of the MCMC samples with visual inspection of trace plots, by calculating the Brooks-Gelman-Rubin diagnostic (R-hat), and by calculating the Heidelberg-Welch diagnostic (Elderd and Miller (2015)). The Gelman-Rubin diagnostic is used to assess convergence between chains and the Heidelberg-Welch for stationarity within chains. Trace plots for all chains, histograms of R-hat, and the percentage of chains that passed the HW test are shown in the appendix.

To evaluate our model's fit to the data, we performed model checks that are described in
full in Appendix: Model Checking. We used our posterior distribution to simulate replicate
datasets based on the parameters of our model. We compared samples from the simulated
datasets to the real, observed datasets using both graphical, visual checks and by calculating
Bayesian p-values for test statistics calculated for the observed and simulated data. In the
following section, we describe how we used the models we fit to obtain the parameters that

describe the *Clarkia* life history. While we do not perform model checks for these derived quantities (e.g. winter seed survival accounting for the combined effect of seed decay and loss of viability) because we combine the output of multiple models, the model checks are still essential to determine whether our inferences are reasonable.

#### 1.5 Computing vital rates

#### 1.5.1 Belowground vital rates

We used the age-specific germination probabilities, survival function, and viability estimates to account for viability in estimates for the probability of germination and survival. We first discretized the survival function to times at which we observed germination and counted seeds (January and October). Estimates of survival over these intervals are the probability that a seed remains intact, but does not account for loss of viability. Next, we used viability 351 estimates from October to calculate viability for January by interpolation (Figure 3D). We tested the viability of seeds in October, and were thus able to estimate the proportion of 353 viable seeds (Figure 3B; filled points). We inferred the viability of intact seeds in January 354 by assuming that seeds lost viability at a constant rate (exponential decay). Further, we 355 interpolated between estimates by assuming that viability changed at a constant rate between 356 years, and that all seeds were viable at the start of the experiment (Figure 3B; open points). 357 We combined the discretized survival function and viability estimates to construct a 358 survival function for the probability that a seed remains intact and viable (Table 2, col-359 umn X). Specifically, we multiplied the posteriors of the discretized survival and viability 360 estimates. Because we combined estimates, some portions of the posterior for seed survival 361 probability was than 1, especially for later seed ages. We restricted the posterior to be less 362 than 1 by truncating the distribution and resampling to redistribute the probability mass. 363 We take this step to retain parameter uncertainty about survival probability in cases where combining the estimates implies a high probability of survival. The survival function for viable seeds  $(\phi)$  is composed of estimates of persistence over time  $(\theta.)$ , estimates of viability  $(\nu.)$ , and estimates of germination conditional on persistence  $(\gamma.)$ .

We used the discretized survival function and age-specific germination probability to 368 obtain the estimates of germination and seed survival required to test predictions from bet-369 hedging theory. Table ?? defines the age-specific germination probabilities and survival 370 probabilities for the structured model in Eckhart et al. (2011) in terms of the survival func-371 tion and age-specific germination probabilities. Figure 3E-F illustrate the relationship among 372 the various probabilities of germination and seed survival. Estimates from the seed bag ex-373 periment correspond to the probability of germination or survival conditional on persistence 374 (e.g.  $\gamma_1$ ). Multiplying these estimates by the probability of persistence up to a certain time 375 gives the unconditional probability (e.g.  $\theta_1 \times \gamma_1$ ). Finally, the probability conditional on per-376 sistence and viability is estimated by incorporating loss of viability into the survival function 377 (e.g.  $\gamma_1/\phi_1$ ), and defines the parameters in the structured population model.

Table 2: Seed persistence and viability in the soil seed bank

	Persistence	Persistence & viability
Time $(x_i)$	$S(x_i)$	$S(x_i)$
$\operatorname{Oct}_0$	$\theta_0$	$\phi_0 = \theta_0$
$ m Jan_{1,total}$	$ heta_1$	$\phi_1 = \theta_1(\gamma_1 + (1 - \gamma_1)\nu_1^{1/3})$
$\operatorname{Jan}_{1,\operatorname{intact}}$	$ heta_2$	$\phi_2 = \theta_2 \nu_1^{1/3}$
$\operatorname{Oct}_1$	$ heta_3$	$\phi_3 = \theta_3 \nu_1$
$Jan_{2,total}$	$ heta_4$	$\phi_4 = \theta_4(\gamma_2 + (1 - \gamma_2)\nu_1(\nu_2/\nu_1)^{1/3})$
Description	Parameter	Probability
July-October	$s_0$	
October-January	$s_1$	$\mid \phi_1 \mid$
1-year old germination	$g_1$	$\gamma_1/\phi_1$
January-October	$s_2$	$\phi_3/\phi_2$
October-January	$s_3$	$\phi_4/\phi_3$

379

#### 1.5.2 Per-capita reproductive success

In order to make our analysis comparable to previous empirical studies of bet hedging,
we calculated per-capita reproductive success as the product of the probability of seedling
survival to fruiting, fruits per plant, and seeds per fruit. We thus calculate per-capita
reproductive success as the number of seeds produced per seedling, on average (e.g. Gremer
and Venable (2014); Venable (2007)).

We used a consistent method to estimate seedling survival to fruiting throughout the experiment, and use the population- and year-level means  $(\mu_{S,jk})$  in our calculation. Because we estimated fruit production in 2 different ways during the study, we chose to use total fruit equivalents (TFE) per plant as our common estimate of fruit production. From 2006–2012, we used  $\mu_{TFE,jk}$ ) as estimated in the statistical model. From 2013–2018, we used the ratio of seeds per damaged to undamaged fruit to calculate a proportion of damaged fruits to add to undamaged fruit counts, as in

$$TFE = undamaged fruits + \frac{seeds per damaged fruit}{seeds per undamaged fruit} \times damaged fruits.$$
 (5)

We used posterior distributions for population- and year-level parameters (e.g.  $\mu_{\text{US},jk}$ ) for these calculations and obtained estimates of  $\mu_{\text{TFE},jk}$ ) for 2013–2018. Finally, we used estimates of seeds per undamaged fruit  $(\mu_{\text{US},jk})$  as our estimate of seeds per fruit.

In terms of parameters from our statistical models, per-capita reproductive success  $F_{jk}$  at population j in year k is calculated as

$$F_{jk} = \phi_{jk} \times \lambda_{\text{TFE},jk} \times \lambda_{\text{US},jk}, \tag{6}$$

where

$$\phi_{jk} = \text{logit}^{-1}(\mu_{S,jk})$$

$$\lambda_{\text{TFE},jk} = \exp(\mu_{\text{TFE},jk})$$

$$\lambda_{\text{US},jk} = \exp(\mu_{\text{US},jk}).$$
(7)

Our multilevel models for aboveground vital rates pooled data more strongly in years with 398 relatively little data. A benefit of this approach is that it implicitly corrects for variation 399 in sample size. While this is beneficial for distinguishing between spurious estimates and 400 true temporal variation, it may also underestimate variation in reproductive success. At 401 the extreme, estimates in years without any data are pooled to the population-level means. 402 Years with zero seedling survivorship would have estimates for fruits per plant that are pooled 403 towards the population-mean (because there were no fruiting plants on which to count fruits). 404 Because estimates of fruits per plant are based on surveys of the whole population, we are 405 confident that per-capita reproductive success was 0 in years in which we observed no fruits 406 per plant and obtained no counts of seeds per fruit (years with no observations for fruits 407 per plant and seeds per fruit in Figure 2). We also considered a second, less conservative 408 estimate of reproductive success in which we assumed that per-capita reproductive success 409 was 0 in years with no seedlings germinated or survived in our permanent plots. This is 410 equivalent to basing our calculations of per-capita reproductive success on a model with no 411 pooling, as the posterior mode of seedling survivorship to fruiting should be very small (0) 412 in years without any seedlings surviving.

#### **Analysis**

#### Correlation between germination probability and seed survival

Increased seed survivorship is predicted to decrease the optimal germination probability Cohen (1966); Ellner (1985a). I assessed whether the observed germination probability was negatively correlated with seed survival (Gremer and Venable (2014)). I calculated the 416 probability that seeds which do not germinate in January remain in the seed bank until the 417 following January  $(s_2s_3)$ . I obtained the posterior distribution for the correlation between 418 germination and seed survival by calculating the correlation of  $g_1$  and  $s_2s_3$  at each iteration 419 of the MCMC output (Hobbs and Hooten (2015), p 194-5). Results of this analysis are shown 420 in Figure 4. Bet hedging models predict that germination probability should be negatively 421 correlated with seed survival; 95% credible intervals that do not overlap zero provide support 422 for this prediction. The bottom panel shows the posterior distribution of correlation between 423 the probability of germination and seed survival.

# Correlation between germination probability and variance in per-capita reproductive success

Increased variance in per-capita reproductive success is predicted to decrease the optimal germination probability (Cohen (1966); Ellner (1985a)). I assessed whether the observed germination probability was negatively correlated with variance in per-capita reproductive success (Venable (2007)).

To calculate the temporal variation in per-capita reproductive success for each population,
I sampled the posterior distribution of reproductive success for each year and calculated
the geometric SD of per capita reproductive success. I obtained the sample correlation
of germination and geometric SD of per capita reproductive success at each iteration of
the MCMC output (Hobbs and Hooten (2015), p 194-5). Bet hedging models predict that

germination probability should be negatively correlated with temporal variance in fitness;

95% credible intervals that do not overlap zero provide support for this prediction. The

geometric SD of per capita reproductive success was calculated as exp(SD (log (per capita
reproductive success+0.5))) (Venable (2007)). Results of this analysis are shown in Figures 5

and 6.

#### Density-independent model for germination probability

We use estimates of seed survival and reproductive success to investigate the adaptive value of delayed germination (Gremer and Venable (2014)). We parameterize a model of population growth rate and calculate the optimal germination strategy for different combinations of seed survival and reproductive success. We describe *C. xantiana*'s life cycle and calculate population growth rate with the equation:

Seed survival rates  $(s_0, s_1, s_2, s_3)$  are population-level estimates. Per capita reproductive

$$\lambda = g_1 Y(t) s_0 s_1 + (1 - g_1) s_2 s_3 \tag{8}$$

success (Y(t)) is calculated as the product of seedling survival to fruiting, fruits per plant, and seeds per fruit (equation (6)). Temporal variation is incorporated into the model by varying the per-capita reproductive success, Y(t), between years. 442 For each population, I numerically calculate the optimal germination probability for 443 the observed variation in reproductive success and seed survival. In each case, I use the posterior mode of the parameter estimates in the equation for density-independent growth 445 (equation (8)). I resampled the posterior modes of per-capita reproductive success (Y(t))446 to obtain a sequence of 1000 years. I used this same sequence of Y(t) and the seed sur-447 vival probabilities to calculate long-term stochastic population growth rates  $(\lambda_s)$  at each 448 germination probability along an evenly spaced grid of possible germination probabilities 440

(G) between 0 and 1. The optimal germination probability is estimated as the value of G that maximizes the geometric mean of the population growth rate. I repeat the simulations 50 times for each population, resampling the sequence of per-capita reproductive success, Y(t), each time. I then calculated the mean of the optimal germination fractions.

Models in which per-capita reproductive success is density-independent predict that germination probability should respond to variance in fitness (Cohen (1966)). To evaluate the
density-independent model, I compared modeled germination probabilities to predicted germination optima. I plot this comparison in Figure 5 and 6. The dotted line indicates a 1:1
relationship between observations and predictions. Values below the line indicate that the
model predicts higher germination probabilities than observed; values above the line would
indicate that the model predicts lower germination probabilities than observed.

# Results

#### Correlation between germination probability and seed survival

I examined the correlation between germination probability and seed survival in the seed bank. Results of this analysis are shown in Figure 4. The bottom panel shows the posterior distribution of correlation between modeled germination probability and the probability of seed survival; the 95% credible interval for the correlation overlaps 0, suggesting that there is no correlation between germination and seed survival.

# Correlation between germination probability and variance in per-capita reproductive success

I examined the correlation between germination probability and variance in per-capita reproductive success. Results of this analysis are shown in Figure 5 and 6. The bottom left panel shows the posterior distribution of correlation between modeled germination probability and geometric SD in per-capita reproductive success. Setting years without any observed plants to have a fitness of zero increases the range of the geometric standard deviation in reproductive success (compare panels A in Figure 5 and 6). However, for both calculations of per capita reproductive success, the median correlation is slightly positive and the 95% credible interval overlaps 0.

#### Optimal germination probability predicted by a density-independent model

Optimal germination probabilities were less than 1 in all populations when we assumed that
years without plants had zero fitness, but not when we used the partially pooled estimates
of per-capita reproductive success (Figure 5 and 6). In both cases, predictions from the
density-independent model overestimated the probability of germination (points fall below
the 1:1 line).

# Discussion

179 . . .

# **Figures**

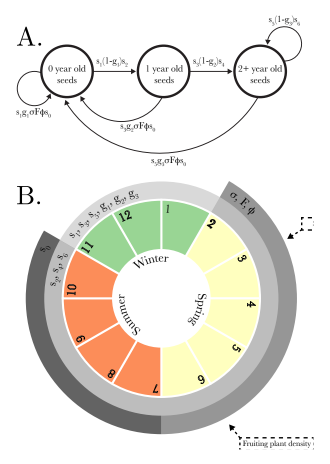


Figure 1: Diagram representations of Clarkia life cycle. Note: Bottom panel needs to be edited so that the dotted lines on the outside of the figure are removed.

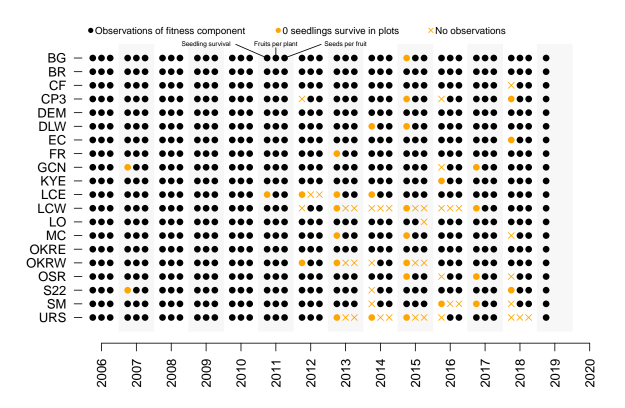


Figure 2: Summary of the aboveground observations, low fitness, and no observations.

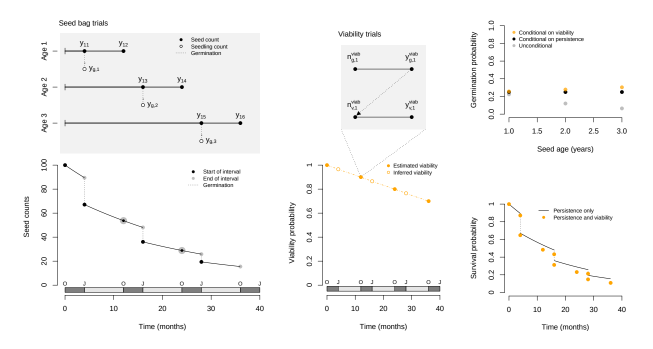


Figure 3: Summary of the seed bag burial experiments and viability trials. Figure will be labeled as (A-B: seed bag trials, C-D: viability trials, E-F: germination probability and survival probability. Add month markers to the y-axis in panels B, D, F. (A) The gray panel contains a graphical representation of the seed bag trials. Seeds were buried at the start of each experiment (100 seeds in month 0). Seed bags were unearthed and intact seeds  $(y_{\cdot\cdot\cdot})$  and germinants  $(y_{g,\cdot\cdot})$  counted. The graph below the panel shows a hypothetical survival function associated with persistence of seeds in the soil seed bank. (B) The gray panel contains a graphical representation of the viability trials. Seeds were tested in two rounds; germination trials were performed and then some or all of the ungerminated seeds were tested for viability. The graph below the panel shows hypothetical data from a series of viability trials and the interpolated, inferred viabilities at times when viability was unobserved. (C) Age-specific germination probably is summarized in three ways. (D) The graph shows the survival function for persistence of seeds in the soil seed bank (black line) and the estimated discrete survival probabilities for persistence and viability of seeds (orange points).

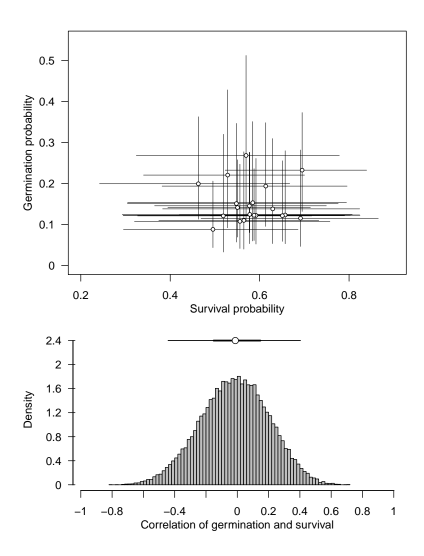


Figure 4: The top panel shows the observed germination probability plotted against probability of seed survival. The bottom panel shows the posterior distribution of correlation between observed germination probability and the probability of seed survival.

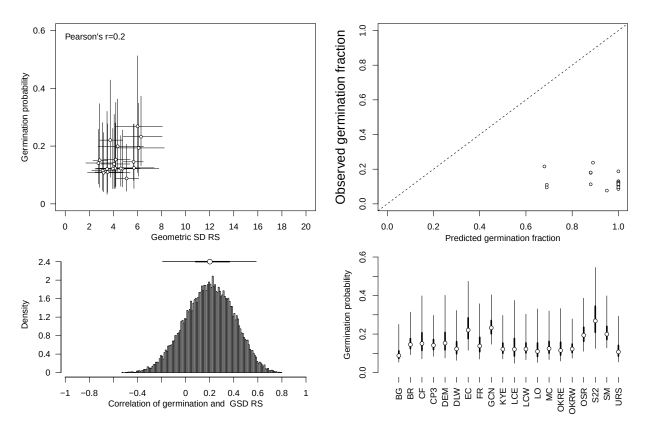


Figure 5: (A) The top left panel shows the observed germination probability plotted against the temporal variation in per capita reproductive success. (B) The bottom left panel shows the posterior distribution of correlation between observed germination probability and geometric SD of per capita reproductive success. (C) The top right panel shows observed germination probability plotted against the optimal germination probability predicted by a density-independent model. For each population, the observed germination probability is the obtained from the model for seed bank vital rates. Each point is the population-specific median of the posterior of  $g_1$  for a model fit to data from seed bag experiments from 2006–2009. Data was pooled across years. The dotted line indicates a 1:1 relationship between observations and predictions. Values below the line indicate that the model predicts higher germination probabilities than observed; values above the line would indicate that the model predicts lower germination probabilities than observed.

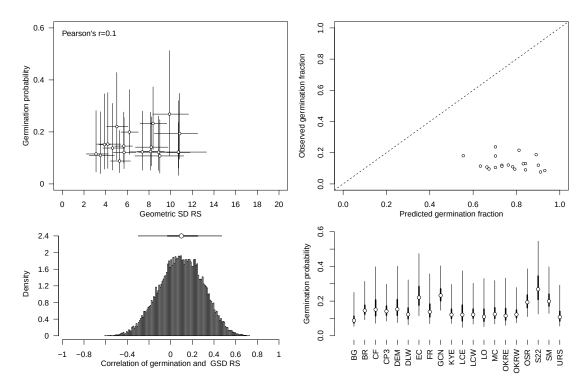


Figure 6: Results with low fitness years set to 0. (A) The top left panel shows the observed germination probability plotted against the temporal variation in per capita reproductive success. (B) The bottom left panel shows the posterior distribution of correlation between observed germination probability and geometric SD of per capita reproductive success. (C) The top right panel shows observed germination probability plotted against the optimal germination probability predicted by a density-independent model. For each population, the observed germination probability is the obtained from the model for seed bank vital rates. Each point is the population-specific median of the posterior of  $g_1$  for a model fit to data from seed bag experiments from 2006–2009. Data was pooled across years. The dotted line indicates a 1:1 relationship between observations and predictions. Values below the line indicate that the model predicts higher germination probabilities than observed; values above the line would indicate that the model predicts lower germination probabilities than observed.

## References

- Cohen, D. 1966. Optimizing reproduction in a randomly varying environment. Journal of
  Theoretical Biology, **12**:119–129.
- Cohen, D. 1968. A General Model of Optimal Reproduction in a Randomly Varying Environment. The Journal of Ecology, **56**:219.
- Conn, P. B., D. S. Johnson, P. J. Williams, S. R. Melin, and M. B. Hooten. 2018. A guide to Bayesian model checking for ecologists. Ecological Monographs, 88:526–542.
- Eager, E. A., R. Rebarber, and B. Tenhumberg. 2014. Modeling and Analysis of a Density-
- Dependent Stochastic Integral Projection Model for a Disturbance Specialist Plant and
- Its Seed Bank. Bulletin of Mathematical Biology, **76**:1809–1834.
- Eckhart, V. M., M. A. Geber, W. F. Morris, E. S. Fabio, P. Tiffin, and D. A. Moeller.
- 2011. The Geography of Demography: Long-Term Demographic Studies and Species
- Distribution Models Reveal a Species Border Limited by Adaptation. The American
- 492 Naturalist, **178**:S26–S43.
- Elderd, B. D. and T. E. X. Miller. 2015. Quantifying demographic uncertainty: Bayesian
- methods for Integral Projection Models (IPMs). Ecological Monographs, pages 15–1526.1.
- Ellner, S. 1985a. ESS germination strategies in randomly varying environments. I. Logistic-
- type models. Theoretical Population Biology, **28**:50–79.
- Ellner, S. 1985b. ESS germination strategies in randomly varying environments. II. Recip-
- rocal Yield-Law models. Theoretical Population Biology, **28**:80–116.
- Evans, M. E. K. and J. J. Dennehy. 2005. Germ Banking: Bet-Hedging and Variable Release
- From Egg and Seed Dormancy. The Quarterly Review of Biology, 80:431–451.

- Evans, M. E. K., K. E. Holsinger, and E. S. Menges. 2010. Fire, vital rates, and population
- viability: a hierarchical Bayesian analysis of the endangered Florida scrub mint. Ecological
- 503 Monographs, **80**:627–649.
- Gabry, J., D. Simpson, A. Vehtari, M. Betancourt, and A. Gelman. 2019. Visualization
- in Bayesian workflow. Journal of the Royal Statistical Society: Series A (Statistics in
- society), **182**:389–402.
- 507 Geber, M. A. and V. M. Eckhart. 2005. EXPERIMENTAL STUDIES OF ADAPTATION
- 508 IN clarkia xantiana:II. FITNESS VARIATION ACROSS A SUBSPECIES BORDER.
- Evolution, **59**:521–531. \_eprint: https://onlinelibrary.wiley.com/doi/pdf/10.1111/j.0014-
- 3820.2005.tb01012.x.
- Gremer, J. R. and D. L. Venable. 2014. Bet hedging in desert winter annual plants: optimal
- germination strategies in a variable environment. Ecology Letters, 17:380–387.
- Heinrich, L., J. Müller, A. Tellier, and D. Živković. 2018. Effects of population- and seed
- bank size fluctuations on neutral evolution and efficacy of natural selection. Theoretical
- Population Biology, **123**:45–69.
- 516 Hobbs, N. T. and M. B. Hooten. 2015. Bayesian models: a statistical
- primer for ecologists. Princeton University Press, Princeton, New Jersey.
- Http://newcatalog.library.cornell.edu/catalog/9201001.
- Lemoine, N. P. 2019. Moving beyond noninformative priors: why and how to
- choose weakly informative priors in Bayesian analyses. Oikos, 128:912–928. \_eprint:
- https://onlinelibrary.wiley.com/doi/pdf/10.1111/oik.05985.
- Lewis, H. 1962. Catastrophic Selection as a Factor in Speciation. Evolution, 16:257–271.
- Publisher: [Society for the Study of Evolution, Wiley].

- McCue, K. A. and T. P. Holtsford. 1998. Seed bank influences on genetic diversity in the
- rare annual Clarkia springvillensis (Onagraceae). American Journal of Botany, 85:30–36.
- eprint: https://bsapubs.onlinelibrary.wiley.com/doi/pdf/10.2307/2446551.
- Metcalf, C. J. E., S. P. Ellner, D. Z. Childs, R. Salguero-Gómez, C. Merow, S. M. McMahon,
- E. Jongejans, and M. Rees. 2015. Statistical modelling of annual variation for inference
- on stochastic population dynamics using Integral Projection Models. Methods in Ecology
- and Evolution, **6**:1007–1017.
- Nunney, L. 2002. The Effective Size of Annual Plant Populations: The Interaction of a Seed
- Bank with Fluctuating Population Size in Maintaining Genetic Variation. The American
- Naturalist, **160**:195.
- ogle, K. and J. J. Barber. 2020. Ensuring identifiability in hierarchical
- mixed effects Bayesian models. Ecological Applications, **30**:e02159. \_eprint:
- https://esajournals.onlinelibrary.wiley.com/doi/pdf/10.1002/eap.2159.
- Paniw, M., P. F. Quintana-Ascencio, F. Ojeda, and R. Salguero-Gómez. 2017. Accounting for
- uncertainty in dormant life stages in stochastic demographic models. Oikos, **126**:900–909.
- Pironon, S., J. Villellas, W. Thuiller, V. M. Eckhart, M. A. Geber, D. A. Moeller, and
- M. B. García. 2018. The 'Hutchinsonian niche' as an assemblage of demographic niches:
- implications for species geographic ranges. Ecography, 41:1103–1113.
- Ritland, K. 1983. The joint evolution of seed dormancy and flowering time in annual plants
- biving in variable environments. Theoretical Population Biology, 24:213–243.
- Venable, D. L. 2007. Bet hedging in a guild of desert annuals. Ecology, 88:1086–1090.
- Waples, R. S. 2006. Seed Banks, Salmon, and Sleeping Genes: Effective Population Size

- in Semelparous, Age-Structured Species with Fluctuating Abundance. The American
- <sup>547</sup> Naturalist, **167**:118.
- <sup>548</sup> Wesner, J. S. and J. P. Pomeranz. 2020. Choosing priors in Bayesian ecological models by
- simulating from the prior predictive distribution. preprint, Ecology.

# Supplementary material

### Data summary.

```
Summary tables for all datasets used in the manuscript. The document summarizes the types
of data collected. The document provides a table summarizing each dataset (e.g. sample
size per each site and year). Link to document: https://github.com/gregor-fausto/
clarkiaSeedBanks/blob/master/products/tables/data-summary.pdf
```

# Joint posterior.

```
Expression for the posterior proportional to the joint distribution, and corresponding directed
acyclic graphs. Link to document: https://github.com/gregor-fausto/clarkiaSeedBanks/
blob/master/products/appendices/appendix-joint-posteriors/appendix-joint-posteriors.
pdf
```

#### Priors.

```
Explanation of priors. Link to document: https://github.com/gregor-fausto/clarkiaSeedBanks/
blob/master/products/appendices/appendix-priors/appendix-priors.pdf
```

#### Model checks.

```
Model checks, including visual posterior predictive checks and assessments with Bayesian p-
values for test statistics. Link to document: https://github.com/gregor-fausto/clarkiaSeedBanks/
blob/master/products/appendices/appendix-model-checks/appendix-x-model-checks.

pdf
```