Seed banks in Clarkia xantiana

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Introduction

- ⁴ Seed banks can buffer plant populations against environmental change and stochasticity
- ⁵ (Eager et al. (2014); Paniw et al. (2017)), increase effective population size (Nunney (2002);
- 6 Waples (2006)), and maintain genetic diversity (Mccue and Holtsford (1998)). Dormancy
- ⁷ can affect the outcome of evolution (Heinrich et al. (2018); Ritland (1983)). Theory thus
- 8 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 10 (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 rate is a function of the distribution of fitness (characterized by the variation in fitness), the fitness 14 values, and the rate of seed survivorship. For a given mean fitness, increasing the variance 15 in fitness decreases the optimal germination fraction (Appendix Figure X). Increasing seed 16 survivorship decreases the optimal germination fraction, and the degree to which it does so 17 depends on the probability of a 'good year'. Specifically, as the probability of a high-fitness 18 year decreases, the optimal germination fraction decreases. 19
- 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 (1985b).
- Several studies have examined intraspecific variation in seed dormancy Clauss and Ven-

able (2000); Philippi (1993); ?); ?. The main set of papers I've included are ones that look at variation in germination among Sonoran Desert annuals (Gremer and Venable (2014); 29 Gremer et al. (2016); Venable (2007)). The table lists predictions made by different models. I think examining the following relationships would be good starting points: The correla-31 tion between variance in fitness (seeds/seedling) and germination fraction should be neg-32 ative – this is true under the density-independent and -dependent model. The correlation 33 between seed survivorship and germination fraction should be negative under both a densityindependent and -dependent model but the limit as survivorship approaches 1 differs. Finally, 35 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. 37

38 [intraspecific variation, range dynamics]

Population vital rates are known to vary across *C. xantiana*'s geographic range. Population growth rates determine species abundance and distribution, and are ultimately what limit persistence beyond range edges. Geographic patterns to vital rates have so far been studied to help understand the demography of geography. Seed banks are a strategy that annual plants may use to buffer against environmental variation and may be part of population persistence. I will begin by characterizing geographic variation in belowground vital rates. [What is the geographic pattern to variation in germination or seed survival?] [I think this question could be expanded to make clear predictions and/or address another aspect such as variation in time.]

A previous study suggests that the soil seed bank is important for population dynamics in

Clarkia xantiana (Eckhart et al. (2011)). 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.

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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 (1985a,b).

Seed banks are more likely to be selected in populations which experience higher levels of in
terannual variation in per-capita reproductive success. To investigate this empirical relationship, I will estimate the correlation between interannual variation in per-capita reproductive

success and the proportion of seeds that germinate in the winter immediately following seed

production. I predict that germination is negatively correlated with interannual variation in

per-capita reproductive success.

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

Table 1: Table 1: Models for germination delays: references and predictions

	Density-independent	Density-dependent fit-	Predictive germina-	Structured model
	fitness	ness	tion	
Key theory	Cohen (1966, 1968)	Ellner (1985a,b)	Cohen (1967)	Easterling and Ellner
references				(2000)
Key empirical	Venable (2007)	Gremer and Venable	Gremer et al. (2016)	
tests		(2014)		
Mean of	increase in \bar{Y} increases	increase in \bar{K} can in-		
seed yield	G^*	crease or decrease \hat{G}		
CV of	increasing ρ_Y de-	increasing ρ_K or ρ_C		
seed yield	creases G^*	decreases \hat{G}		
Seed	increasing s decreases	increasing s decreases		
survivorship	G^* ; limit near $s=1$ is	\hat{G} ; limit near $s = 1$ is		
	p	0		

Methods

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Background on study system

Monica Geber and collaborators have collected 12+ years of annual estimates for demographic data on a species of annual plant: survival of seedlings to fruiting adults, fruits per adult plant, and seeds per fruit. As part of the long-term work on *Clarkia xantiana*, there 3 sources of data on the transition between seeds in fruits and seedlings: 1 observational data set and 2 experimental data sets. Here, I present analyses of 1 observational data set and 1 experimental data set.

Starting in 2007, there are (1) estimates of fruits/plant and seeds/fruit that provide an estimate of seed input into a plot and (2) estimates of germinants the following year. For most plots, the number of seeds entering a plot in year t-1 is much greater than the number of seedlings emerging in a plot in year t. However, this is not uniformly true, and there is also experimental data suggesting these seeds may survive in the seed bank for at least 10

years at some locations.

We use two experiments, conducted at non-overlapping points in time, to estimate transitions in the seed bank. From 2006-2010, Geber and collaborators buried seeds in bags and
periodically dug them up to count seedlings and intact, viable seeds. This data estimates
transitions leading to germination or survival of seeds that are 1, 2, and 3 years old. Starting
in 2013, Geber and collaborators placed seeds in pots and counted seedlings. This data estimates transitions of seeds in the soil seed bank but cannot separate germination and survival
in the same way as the first experiment.

Here, we analyze data from an experiment that involved burying seeds in seed bags (2005-2009).

To determine how seed survival and germination varied among populations of C. xantiana,

we use data collected from a series of seed burial experiments. We started these seed burial

Data

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Seed bag burial experiments

experiments in three subsequent years (2005, 2006, 2007) to obtain multiple estimates for 95 seed survival and germination. 96 In June-July 2005, we collected seeds at each of the 20 populations included in this study. 97 In October 2005, we buried 30.5×5 -cm nylon mesh bags at each population. Each nylon mesh 98 bag contained 100 seeds collected at that population. In January 2006, we removed 10 of 99 these bags and counted the number of germinated seedlings and the number of ungerminated, 100 intact seeds in each bag. We then returned the ungerminated, intact seeds to the resealed 101 bag and returned the bag to the field. In October 2006, we removed these bags and counted 102 the number of ungerminated, intact seeds. We collected the following data: 103

• n_{ijt} = observed count of seeds in the seed bags at the start of the experiment in October

- in the i^{th} bag, from the j^{th} population, in the t^{th} year, assumed to be measured perfectly
- $y_{\text{intact}ijt}$ = observed count of ungerminated, intact seeds in the seed bags in January in the i^{th} bag, from the j^{th} population, in the t^{th} year, assumed to be measured perfectly
- $y_{\text{germ}}ijt$ = observed count of germinated seedlings in the seed bags in January in the i^{th} bag, from the j^{th} population, in the t^{th} year, assumed to be measured perfectly
- $y_{\text{total}ijt}$ = observed count of ungerminated, intact seeds plus germinated seedlings in the seed bags in January in the i^{th} bag, from the j^{th} population, in the t^{th} year, assumed to be measured perfectly
- $y_{\text{surv}ijt}$ = observed count of ungerminated, intact seeds in the seed bags in October in the i^{th} bag, from the j^{th} population, in the t^{th} year, assumed to be measured perfectly

Viability trials

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In the lab, we conducted germination trials and viability assays on subsets of the seeds from each bag to estimate the viability of the ungerminated, intact seeds. First, we placed up to 15 seeds from each bag on to moist filter paper in a disposable cup and observed germination over 10 days; we counted and removed germinants every 2 days.

After 10 days, all remaining ungerminated seeds (up to a total of 10 seeds) were sliced in half and individually placed into the wells of 96-well plates filled with a solution of tetrazolium chloride, which stains viable tissue red. [Eckhart et al. (2011): not all ungerminated seeds were tested; most were] We covered the plates with foil. Each 96-well plate contained seed from at least one bag per population of a given seed-age class. Two or three tests of up to 15 seeds each were conducted for each bag. We checked and counted for viable seeds every 2 days for 10 days.

We collected the following data:

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- $n_{\text{germ}ijt}$ = observed count of seeds at the start of the X^th germination trial for the i^{th} bag, from the j^{th} population, in the t^{th} year, assumed to be measured perfectly
- $y_{\text{germ}ijt}$ = observed count of germinated seedlings in the X^th germination trial for the i^{th} bag, from the j^{th} population, in the t^{th} year, assumed to be measured perfectly
- $n_{\text{viab}}ijt$ = observed count of seeds at the start of the X^th viability trial for the i^{th} bag, from the j^{th} population, in the t^{th} year, assumed to be measured perfectly
- $y_{\text{viab}}ijt$ = observed count of viable seedlings in the X^th viability trial for the i^{th} bag, from the j^{th} population, in the t^{th} year, assumed to be measured perfectly

Seedling survival to fruiting

The data consist of counts of seedlings and fruiting plants in 0.5 m² plots at 20 populations from 2006–present. Each population was visited in February and June to count the number of seedlings and fruiting plants, respectively. Seedlings and plants in each plot are counted by a single person at each visit.

For now, we assume that the data on seedlings is measured perfectly (i.e. no underor over-counts of seedlings). However, there are at least two possible sources of error: (1)
measurement error that arises because we failed to count seedlings that were present and
(2) error that arises because seedlings germinated after we visited the population. Germination phenology varies may vary from year to year but also by geography; higher elevation
populations may have delayed phenology. We may want to develop a model that relates our
estimate of seedlings to the true number of seedlings in a plot because we sometimes observe
more fruiting plants than seedlings. For now, I ignored data that involved undercounting by
filtering out those rows in the dataset.

We assume that the data on fruiting plants is measured perfectly (i.e. we did not underor over-count) because plants stand out from the background vegetation in June. Our model estimates the proportion of seedlings that survive to become fruiting plants. Define:

- n_{ijt} = observed counts of seedlings in the i^{th} plot, from the j^{th} population, from the k^{th} year
- y_{ijt} = observed counts of fruiting plants in the i^{th} plot, from the j^{th} population, from the k^{th} year, assumed to be measured perfectly

Fruits per plant

From 2006–2012, "we recorded...the number of fruits per plant for up to 15-20 plants per 0.5 m²" (Eckhart et al. (2011)). For each plant, we counted the number of undamaged fruits. We then took the damaged fruits and visually stacked them end to end to estimate how many additional undamaged fruits that was equivalent to (e.g. two half fruits corresponded to one undamaged fruit). We used these counts to estimate the number fruits produced per plant.

We seek to estimate (1) the number of fruits produced per plant and (2) the proportion of fruits that are damaged per plant. Define:

- y_{ijk}^{TFE} = observed counts of total fruit equivalents per plant on the i^{th} plant, from the j^{th} population, from the k^{th} year, assumed to be measured perfectly
- n_{ijk} = observed counts of total fruits per plant (sum of y_{ijk}) on the i^{th} plant, from the j^{th} population, from the k^{th} year, assumed to be measured perfectly

Seeds per fruit

From 2006–2012, "we collected one fruit from each of 20-30 haphazardly selected plants distributed across each population (but outside permanent plots, to avoid influencing seed input within them) to estimate the mean number of seeds produced per fruit" (Eckhart et al.

(2011)). In the field, we collected fruits that were undamaged. In the lab, we broke open the fruits to count the number of seeds per fruit. For each population in each year, we attempted to obtain 20-30 counts of seeds produced per undamaged fruit.

From 2013—present, we collected one undamaged and one damaged fruit from each of 2030 haphazardly selected plants distributed across each population. The plants were outside
permanent plots to avoid affecting seed input. We used these fruits to estimate the mean
number of seeds produced by undamaged and damaged fruits. In the lab, we broke open
the fruits to count the number of seeds per fruit. For each population in each year, we
attempted to obtain 20-30 counts of seeds produced per undamaged fruit and 20-30 counts
of seeds produced per damaged fruit.

We seek to estimate the number of seeds per undamaged fruit. Define:

- y_{ijk}^{und} = observed counts of seeds in the i^{th} undamaged fruit, from the j^{th} population, from the k^{th} year, assumed to be measured perfectly
 - λ_{jk} = true, unobserved mean number of seeds per undamaged fruit from the j^{th} population, from the k^{th} year

Model framework

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We use observational and experimental data from 20 populations of *Clarkia xantiana* to estimate transition probabilities across the life cycle. We obtain population-specific estimates for belowground vital rates, and to obtain year- and population-specific estimates for aboveground vital rates. We use these transition probabilities to analyze correlations between germination probability and variance in per-capita reproductive success, correlation between germination probability and seed survival, and to compare the optimal germination fraction from a density-independent model to the observed germination fraction.

Parameter estimates for belowground transitions

To estimate population-specific estimates for belowground vital rates, we use data from seed burial experiments in the field and seed viability trials in the lab. We combine these data to infer seed survival across different parts of the year and germination. I estimate probabilities of success using the data, and compose these estimates to obtain transition probabilities for the life history of *Clarkia xantiana*. I provide details of this approach in the **Appendix on Conditional Probability**. Briefly, I estimate probabilities of success $(\theta_1, \theta_2, \theta_3, \theta_4, \theta_5, \nu_1, \nu_2)$ using data from seed burial experiments and viability trials. I use these probabilities of success to compose transition probabilities (s_1, s_2, s_3, g_1) .

Figure 1 illustrates the relationship between the data and the estimated probability of success. There are two boxes: one for the seed bag experiment and one for the viability trials.

In the seed bag experiment, I split January into two steps, one for just before germination and one for just after. Solid arrows represent estimated probabilities and are labeled with corresponding parameters.

The probability that seeds from the start of the experiment remain intact in January 205 is represented as θ_1 . In January, all seeds are intact (this includes viable and non-viable seeds). We estimate the probability of a seedling emerging, conditional on being intact as θ_2 . I assume that there is no decay during germination (i.e., seed loss is instantaneous in January). The number of intact seeds before germination is equal to the number of seeds and seedlings after germination. At this point, the seeds transition into one of four possible 210 states. Intact and viable seeds may have (1) germinated or (2) not germinated and remain 211 dormant. All (3) other intact seeds are non-viable because (4) seeds that were not viable 212 could not have germinated. Finally, we represent the probability of a seed being intact in 213 October, conditional on being intact in January as θ_3 . 214

We use the viability trials to estimate the probability of viability (ν_1) for a seed that is

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intact in October of year t + 1. I need to make some assumptions in order to incorporate the loss of viability into the model. I assume that viability is lost at a constant rate, and that germination removes some number of seeds from the pool of viable seeds but does not change the rate of decay. Some fraction of the total seeds in January pre-germination is viable $(\nu_1^{1/3})$ and some of those viable seeds germinate.

The seed burial experiments and viability trials thus provide information about the fate of seeds in the seed bank. We define s_1 as the probability of a seed being intact and viable from October in year t to January in year t+1. We define g_1 as the probability of germination for a seed that is intact and viable. We define s_2 as the probability of survival from January to October in year t for a seed that was intact and viable in January. We define s_3 as the probability of survival from October in year t to January in year t+1 for a seed that was intact and viable in October. Mathematically, we write each of the transition probabilities as follows:

$$s_{1} = \theta_{1} \times (\theta_{2} + (1 - \theta_{2}) \times \nu_{1}^{1/3})$$

$$g_{1} = \frac{\theta_{2}}{1 - (1 - \nu_{1}^{1/3}) \times (1 - \theta_{2})}$$

$$s_{2} = \theta_{3} \times \nu_{1}^{2/3}$$

$$s_{3} = \frac{\theta_{4} \times (\theta_{5} + (1 - \theta_{5}) \times \nu_{2}^{1/3})}{s_{1} \times (1 - g_{1}) \times s_{2}}$$

$$(1)$$

Parameter estimates for aboveground transitions

To estimate obtain year- and population-specific estimates for aboveground vital rates, we use data from annual observational surveys of 20 *Clarkia xantiana* populations. We use data from surveys of plots to estimate seedling survival to fruiting (ϕ) . We also estimate fruits per plant (parameter) and seeds per fruit (parameter). We combine these data to obtain annual estimates of per-capita reproductive reproductive success.

Models

Model for seed burial experiment data

The models below represent the joint likelihood for data from the seed bag experiments. In each model, we obtain the population-level posterior distribution probability of success by marginalizing across years and taking the inverse logit.

To estimate θ_1 , I fit the following model for bag i in year j in population k.

$$[\boldsymbol{\mu}, \boldsymbol{\sigma}, \boldsymbol{\alpha}, \boldsymbol{\theta} | \boldsymbol{n}, \boldsymbol{y_{total}},] \propto \prod_{k=1}^{K} \prod_{j=1}^{J} \prod_{i=1}^{I} \text{binomial}(\boldsymbol{y_{ijk}^{tot}} | n_{ijk}, \text{logit}^{-1}(\alpha_{ijk}))$$

$$\times \text{normal}(\alpha_{ijk} | \mu_{jk}, \sigma_{jk})$$

$$\times \text{normal}(\mu_{jk} | \mu_{0,k}, \sigma_{0,k})$$

$$\times \text{uniform}(\sigma_{jk} | 0, 100)$$

$$\times \text{normal}(\mu_{0,k} | 0, 100) \text{uniform}(\sigma_{0,k} | 0, 100).$$

$$(2)$$

To estimate θ_2 , I fit the following model for bag i in year j in population k.

$$[\boldsymbol{\mu}, \boldsymbol{\sigma}, \boldsymbol{\alpha}, \boldsymbol{\theta} | \boldsymbol{n}, \boldsymbol{y}_{\mathbf{germ}},] \propto \prod_{k=1}^{K} \prod_{j=1}^{J} \prod_{i=1}^{I} \operatorname{binomial}(y_{ijk}^{\text{germ}} | y_{ijk}^{\text{total}}, \operatorname{logit}^{-1}(\alpha_{ijk}))$$

$$\times \operatorname{normal}(\alpha_{ijk} | \mu_{jk}, \sigma_{jk})$$

$$\times \operatorname{normal}(\mu_{jk} | \mu_{0,k}, \sigma_{0,k})$$

$$\times \operatorname{uniform}(\sigma_{jk} | 0, 100)$$

$$\times \operatorname{normal}(\mu_{0,k} | 0, 100) \operatorname{uniform}(\sigma_{0,k} | 0, 100).$$

$$(3)$$

To estimate θ_3 , I fit the following model for bag i in year j in population k.

$$[\boldsymbol{\mu}, \boldsymbol{\sigma}, \boldsymbol{\alpha}, \boldsymbol{\theta} | \boldsymbol{n}, \boldsymbol{y}_{\text{germ}},] \propto \prod_{k=1}^{K} \prod_{j=1}^{J} \prod_{i=1}^{I} \text{binomial}(y_{ijk}^{\text{oct}} | y_{ijk}^{\text{total}} - y_{ijk}^{\text{germ}}, \text{logit}^{-1}(\alpha_{ijk}))$$

$$\times \text{normal}(\alpha_{ijk} | \mu_{jk}, \sigma_{jk})$$

$$\times \text{normal}(\mu_{jk} | \mu_{0,k}, \sigma_{0,k})$$

$$\times \text{uniform}(\sigma_{jk} | 0, 100)$$

$$\times \text{normal}(\mu_{0,k} | 0, 100) \text{uniform}(\sigma_{0,k} | 0, 100).$$

$$(4)$$

Model for viability trial data

Each bag i from population j in year k had n trials. The problem is most bags only had 2 trials so it's difficult to estimate a variance. I want to estimate a bag-specific viability because that is what I would use in the seed bag survival and germination model to put a 232 bag-specific viability rather than a population-specific viability. I fit the following model for trial h for bag i at population j in year k:

For the germination trials, I estimated a probability of germination for each bag, θ^{g} . Because the goal of the lab experiments was to provide an estimate of viability in each bag, I was not interested in modeling the viability in each bag as coming from a population of viabilities at the population level. I provided each parameter with an uninformative beta (1,1) prior. I did not apply hyperpriors because (see above).

$$[\boldsymbol{\theta}^{\mathbf{g}}|\boldsymbol{n}^{\mathbf{g}},\boldsymbol{y}^{\mathbf{g}}] \propto$$

$$\prod_{j=1}^{J} \prod_{i=1}^{I} \prod_{h=1}^{H} \prod_{k=1}^{K} \operatorname{binomial}(y_{hijk}^{\mathbf{g}}|n_{hijk}^{\mathbf{g}},\theta_{ijk}^{\mathbf{g}}) \operatorname{beta}(\theta_{ijk}^{\mathbf{g}}|1,1)$$
(5)

I started considering applying the probabilities from the conditional probability tree.

This might make sense but then I realize that I've gone down this rabbit hole before. Not

all of the seeds that didn't germinate were tested. But I'm not sure how else to control for that besides....

Where P(V) = P(V - Gc) * P(Gc) + P(V - G) * P(G)

If there are missing data, it might make sense to estimate a per-population/year viability from hyperpriors. In other words, we would a distribution for theta defined by hyperpriors set at the population level. Each population would get its own set of hyperpriors. Then when we are looking to get the viability of a given bag in a given year, if the data are missing we would draw from the hyperpriors for a population mean rather than propagate NAs.

The dataset has NAs in the viability stain column. These are there because the viability trial started with 0 seeds. Viability trials started with 0 seeds when all/most of the seeds in the germination trials germinated. The estimated probability for the (0,NA) set is 0.5; this is because the result provides no new information. One approach might be to throw these rows out entirely or to just take the germination probablity as the true probability. However, because the probabliity of being viable if not germinating is multiplied by the probablility of not germinating, the effect is quite small when most of the seeds germinate.

A figure that would help is one that plots the predicted vs. observed proportions for the germination and viability experiments for one population. Maybe start with partial pool with direct parameterization of theta via a beta prior on each bag. The goal of the experiment is bag-specific viability. Reasons to include population would be to

s1 We want this estimate to include seeds that germinated and seeds that were intact and viable. First, we estimate the probability of a seed being intact to January as $p^{intact} = (y^{germ} + y^{jan1})/n^{total}$. Second, we estimate the probability of a plant emerging, conditional on being intact as $p^{germ} = y^{germ}/(y^{germ} + y^{jan1})$. We then calculate

g1 First, we estimate the probability of a plant emerging, conditional on being intact as $p^{germ} = y^{germ}/(y^{germ} + y^{jan1}).$

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s2 First, we estimate the probability of a seed being intact in October, conditional on

being intact in January $p = y^{oct1}/y^{jan1}$.

s3 First, we make use the probabilities for s_1 , g_1 and s_2 (described above) to normalize the event space. Second, we estimate the probability of a seed being intact to January as $p^{intact} = (y^{germ2} + y^{jan2})/n^{total}$. Finally, we estimate the probability of a plant emerging, conditional on being intact as $p^{germ2} = y^{germ2}/(y^{germ2} + y^{jan2})$.

These models relate the average seed survival in each population to the seed survival 267 observed in a given year through a linear model. The function has a parameter for the 268 average seed survival in each population, and a parameter for the seed survival of each 269 population in each year. I treated seed survival as a binomially-distributed random variable 270 because the data come from an experiment in which we buried a known number of seeds 271 and counted seeds. Thus for the ith observation, I parameterized a binomial distribution in 272 terms of a probability and a known number of trials. I sampled a 95% credible interval from 273 the posterior predictive distribution for seed survival for each population and compared it 274 to the seed burial experiments to assess whether the model from the seed burial experiments 275 could predict the seed survival in seed burial experiments. 276

I fit this model with JAGS in R.

Seedling survival to fruiting

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We estimated survival across all populations taking into account both temporal and between-278 population variability with the following model. In this model, $\alpha_{0,j}^S$ is the logit mean survival 279 probability at population $j,\ \beta^S_{jk}$ are independent identically distributed random variables 280 drawn from normal distributions with mean 0 and population-specific temporal variance 281 parameters σ_j^S . Writing the population-specific logit survival as a fixed effect means that 282 each population parameter estimate is estimated separately with no shared variance term. 283 The population-specific temporal variance parameter is written as a random effect, which 284 means that each population has has year components that are drawn from a distribution 285

with a shared variance term. I estimated the probability of surviving to fruiting using data from plots at populations in different years:

$$[\boldsymbol{\alpha_0^S}, \boldsymbol{\beta^S}, \sigma^S | \boldsymbol{n}, \boldsymbol{y}] \propto \prod_{i=1}^{I} \prod_{j=1}^{J} \prod_{k=1}^{K} \text{binomial}(y_{ijk} | n_{ijt}, f(\alpha_{0,j}^S, \beta_{jk}^S))$$

$$\times \text{normal}(\beta_{jk}^S | 0, \sigma_j^S)$$

$$\times \text{normal}(\alpha_{0,j}^S | 0, 10) \text{uniform}(\sigma_j^S | 0, 100)$$

$$(6)$$

where

$$f(\alpha_{0,j}^S, \beta_k^S)) = logit^{-1}(\alpha_{0,j}^S + \beta_{jk}^S)$$
(7)

Fruits per plant

To assess what probability distribution to use when fitting this model, I fit a power model with an intercept to the mean and variance using the 'nls' function in R, which returned an exponent of 1.99. The fit is close to quadratic which means a negative binomial is likely to be an appropriate distribution (Lindn and Mntyniemi (2011)). We estimated fruits per plant across all populations taking into account both temporal and between-population variability with the following model. I first worked only with data on total fruit equivalents on a plant (2006-2012). I estimated total fruit equivalents per plant as:

$$[\boldsymbol{\alpha_0^F}, \boldsymbol{\beta^F}, \sigma^F | \boldsymbol{n}, \boldsymbol{y}] \propto \prod_{i=1}^{I} \prod_{j=1}^{J} \prod_{k=1}^{K} \text{negative binomial}(y_{ijk}^{\text{TFE}} | f_1(\boldsymbol{\alpha}_{0,j}^F, \boldsymbol{\beta}_{jk}^F), \kappa^F))$$

$$\times \text{normal}(\boldsymbol{\alpha}_{0,j}^F | 0, 10)$$

$$\times \text{normal}(\boldsymbol{\beta}_{jk}^F | 0, \sigma_j^F)$$

$$\times \text{normal}(\boldsymbol{\sigma}_j^F | 0, 100)$$
(8)

where

$$f_1(\alpha_{0,j}^F, \beta_{jk}^F)) = \lambda_{jk}^F = \exp(\alpha_{0,j}^F + \beta_{jk}^F)$$
 (9)

negative binomial
$$(y_{ijk}^{\text{TFE}}|\frac{\kappa^F}{\kappa^F + \lambda_{jk}^F}, \kappa^F)$$
 (10)

where the negative binomial is parameterized with probability parameter p and dispersion parameter r [negative binomial(p,r)]. In this case $p = \frac{\kappa}{\kappa + \mu}$.

Seeds per fruit

To assess what probability distribution to use when fitting this model, I fit a power model with an intercept to the mean and variance using the nls function in R, which returned an exponent of 1.38. The fit is greater than linear but less than quadratic which means that neither a Poisson nor negative binomial are likely to be entirely appropriate distributions for the data (Lindn and Mntyniemi (2011)). I might try the parameterization in that reference but for now I am using the negative binomial because the data are overdispersed. We estimated seeds per fruit across all populations taking into account both temporal and between-population variability with the following model. Here, I used data from undamaged fruits from the years 2006-2012. I estimated seeds per fruit as:

$$[\boldsymbol{\alpha_0^P}, \boldsymbol{\beta^P}, \sigma^P | \boldsymbol{n}, \boldsymbol{y}] \propto \prod_{i=1}^{I} \prod_{j=1}^{J} \prod_{k=1}^{K} \text{negative binomial}(y_{ijk}^{\text{und}} | f_1(\alpha_{0,j}^P, \beta_{jk}^P), \kappa^P))$$

$$\times \text{normal}(\alpha_{0,j}^P | 0, 10)$$

$$\times \text{normal}(\beta_{jk}^P | 0, \sigma_j^P)$$

$$\times \text{normal}(\sigma_j^P | 0, 100)$$

$$(11)$$

where

$$f_1(\alpha_{0,j}^P, \beta_{jk}^P)) = \lambda_{jk}^P = \exp(\alpha_{0,j}^P + \beta_{jk}^P)$$
 (12)

negative binomial
$$(y_{ijk}^{\text{und}}|\frac{\kappa^P}{\kappa^P + \lambda_{jk}^P}, \kappa^F P)$$
 (13)

where the negative binomial is parameterized with probability parameter p and dispersion parameter r [negative binomial(p,r)]. In this case $p = \frac{\kappa}{\kappa + \mu}$.

Analysis

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 (1985b)). I assessed whether the observed germination probability was negatively correlated with variance in per-capita reproductive success (Venable (2007)). Per-capita reproductive success F_{jk} at population j in year k was calculated at the per year and per population level as follows:

$$F_{jk} = \phi_{jk} \times \lambda_{jk}^F \times \lambda_{jk}^P \tag{14}$$

where

$$\phi_{jk} = \operatorname{logit}^{-1}(\alpha_{0,j}^{S} + \beta_{jk}^{S})$$

$$\lambda_{jk}^{F} = \exp(\alpha_{0,j}^{F} + \beta_{jk}^{F})$$

$$\lambda_{jk}^{P} = \exp(\alpha_{0,j}^{P} + \beta_{jk}^{P})$$
(15)

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 293 the geometric SD of per capita reproductive success. For each population, I calculated the 294 correlation between germination and variance in per-capita reproductive success with the 295 posterior distribution for the geometric SD of per capita reproductive success and the poste-296 rior distribution of germination probability from model XX. Using this approach, I obtained 297 a distribution of correlation estimates. Results of this analysis are shown in Figure 2. Bet 298 hedging models predict that germination probability should be negatively correlated with 299 temporal variance in fitness; 95% credible intervals that do not overlap zero provide support 300 for this prediction.

Correlation between germination probability and seed survival

Increased seed survivorship is predicted to decrease the optimal germination probability Cohen (1966); Ellner (1985b). I assessed whether the observed germination probability was negatively correlated with seed survival (Gremer and Venable (2014)). I calculated seed 304 survival as s_2s_3 as the product of these vital rates is the probability that seeds which do 305 not germinate in January remain in the seed bank until the following January. I used the 306 posteriors of g_1 and s_2s_3 to calculate the correlation between germination and seed survival. 307 Using this approach, I obtained a distribution of correlation estimates. Results of this 308 analysis are shown in Figure 3. Bet hedging models predict that germination probability 309 should be negatively correlated with seed survival; 95% credible intervals that do not overlap 310 zero provide support for this prediction. 311

Density-independent model for germination probability

We used 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 use the following equation to describe *Clarkia xantiana*'s life cycle and calculate population growth rate:

$$\lambda_j = s_1 g_1 Y(t) s_0 + s_1 (1 - g_1) s_2 \tag{16}$$

The parameters in this equation were fit in models corresponding to equations (??), (??), and (??). Seed survival rates (s_0, s_1, s_2, s_3) are population-level estimates. Per capita reproductive success (Y(t)) is calculated as the product of seedling survival to fruiting, fruits per plant, and seeds per fruit (equation (14)). Variation is incorporated into the model by varying per-capita reproductive success, Y(t), between years.

I numerically calculated the optimal germination probability for the observed level of 317 variation in reproductive success and seed survival in each population. For each population, 318 I randomly selected values 1000 from the posterior distribution for reproductive success. I 319 used this same sequence of Y(t) and the observed seed survival probabilities to calculate 320 long-term stochastic population growth rates (λ_s) at each germination probability along an 321 evenly spaced grid of possible germination probabilities (G) between 0 and 1. The optimal 322 germination probability is estimated as the value of G that maximized geometric mean of the 323 population growth rate. I repeated the simulations 50 times for each population, resampling 324 from the posterior distribution for reproductive success each time. I calculated the mean of 325 the optimal germination fractions. 326

Models in which per-capita reproductive success is density-independent predict that germination probability should respond to variance in fitness (Cohen (1966)). To evaluate a density-independent model for germination probability, I compared observed germination probability to predicted germination optima. I plot this comparison in Figure 4. The dotted line indicates a 1:1 relationship between observations and predictions. Values below the

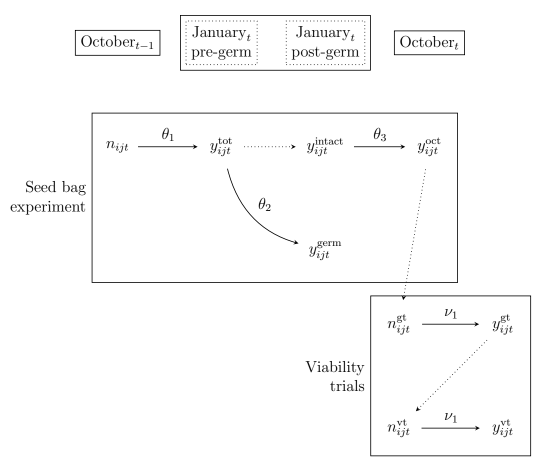
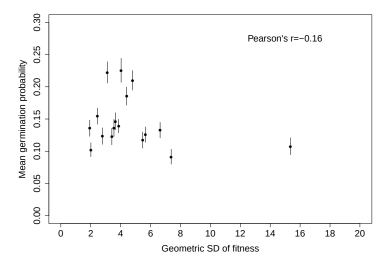


Figure 1: Diagram of data from the seed bag experiments and viability trials.

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

Figures



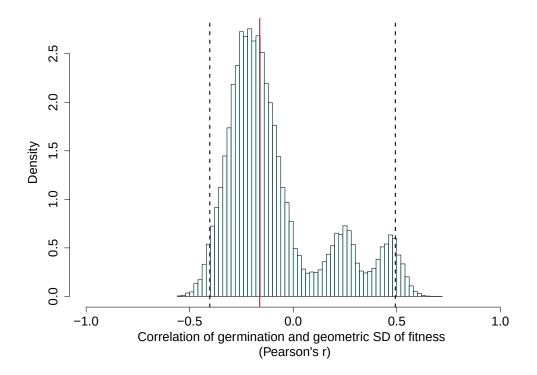
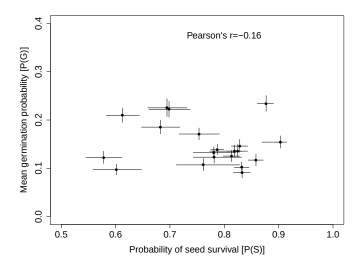


Figure 2: The top panel shows the observed germination probability plotted against the temporal variation in per capita reproductive success. The bottom panel shows the posterior distribution of correlation between observed germination probability and geometric SD of per capita reproductive success; the median correlation is negative (-0.16) but the 95% credible interval overlaps 0.



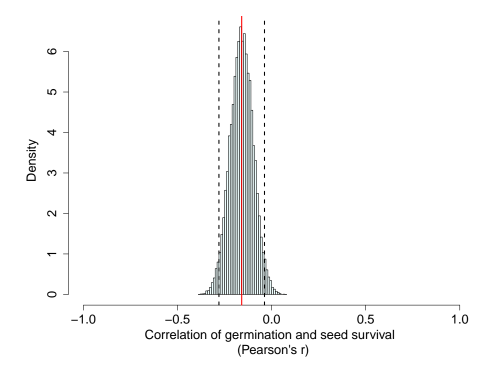


Figure 3: The top panel shows the observed germination probability plotted against probability of seed survival. The bottom panel shows the posterior distribution of correlation between observed germination probability and the probability of seed survival; the median correlation is negative (-0.16) and the 95% credible interval does not overlap 0.

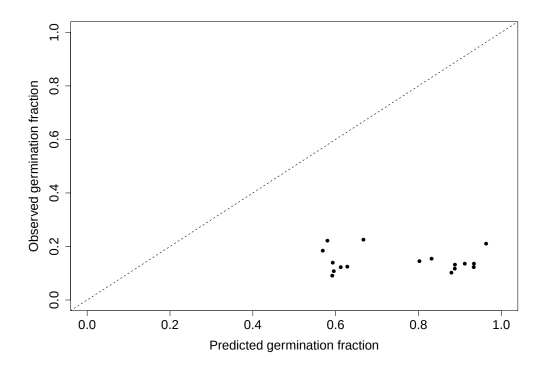


Figure 4: Observed germination probability plotted against the optimal germination probability predicted by a density-independent model. For each population, the observed germination probability is the obtained from the model for seed bank vital rates. Each point is the population-specific median of 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

- 335 M. J. Clauss and D. L. Venable. Seed Germination in Desert Annuals: An
- Empirical Test of Adaptive Bet Hedging. The American Naturalist, 155(2):168–
- 186, Feb. 2000. ISSN 0003-0147, 1537-5323. doi: 10.1086/303314. URL
- https://www.journals.uchicago.edu/doi/10.1086/303314.
- D. Cohen. Optimizing reproduction in a randomly varying environment. Journal of Theoret-
- $ical\ Biology,\ 12(1):119-129,\ {
 m Sept.}\ 1966.\ ISSN\ 00225193.\ doi:\ 10.1016/0022-5193(66)90188-$
- 3. URL https://linkinghub.elsevier.com/retrieve/pii/0022519366901883.
- D. Cohen. Optimizing Reproduction in a Randoruly Varying Environment when a Corre-
- lation May Exist between the Conditions at the Time a Choice has to be Made and the
- Subsequent Outcome. Journal of Theoretical Biology, 16:1–14, 1967.
- D. Cohen. A General Model of Optimal Reproduction in a Randomly Varying Environment.
- The Journal of Ecology, 56(1):219, Mar. 1968. ISSN 00220477. doi: 10.2307/2258075.
- URL https://www.jstor.org/stable/2258075?origin=crossref.
- E. A. Eager, R. Rebarber, and B. Tenhumberg. Modeling and Analysis of a Density-
- Dependent Stochastic Integral Projection Model for a Disturbance Specialist Plant and
- 350 Its Seed Bank. Bulletin of Mathematical Biology, 76(7):1809–1834, 2014. ISSN 15229602.
- doi: 10.1007/s11538-014-9978-y.
- M. R. Easterling and S. P. Ellner. Dormancy strategies in a random environment: Comparing
- structured and unstructured models. Evolutionary Ecology Research, 2:387–407, 2000.
- V. M. Eckhart, M. A. Geber, W. F. Morris, E. S. Fabio, P. Tiffin, and D. A. Moeller.
- The Geography of Demography: Long-Term Demographic Studies and Species Distribu-
- tion Models Reveal a Species Border Limited by Adaptation. The American Naturalist,

- 178(S1):S26-S43, Oct. 2011. ISSN 0003-0147, 1537-5323. doi: 10.1086/661782. URL

 https://www.journals.uchicago.edu/doi/10.1086/661782.
- S. Ellner. ESS germination strategies in randomly varying environments. II.
- Reciprocal Yield-Law models. Theoretical Population Biology, 28(1):80–116,
- Aug. 1985a. ISSN 00405809. doi: 10.1016/0040-5809(85)90023-1. URL
- https://linkinghub.elsevier.com/retrieve/pii/0040580985900231.
- 363 S. Ellner. ESS germination strategies in randomly varying environments. I.
- Logistic-type models. Theoretical Population Biology, 28(1):50-79, Aug.
- 1985b. ISSN 00405809. doi: 10.1016/0040-5809(85)90022-X. URL
- https://linkinghub.elsevier.com/retrieve/pii/004058098590022X.
- M. E. K. Evans and J. J. Dennehy. Germ Banking: BetHedging and Variable Release From
- Egg and Seed Dormancy. The Quarterly Review of Biology, 80(4):431-451, 2005. ISSN
- 369 0033-5770. doi: 10.1086/498282.
- 370 J. R. Gremer and D. L. Venable. Bet hedging in desert winter annual plants:
- optimal germination strategies in a variable environment. Ecology Letters, 17
- 372 (3):380–387, Mar. 2014. ISSN 1461023X. doi: 10.1111/ele.12241. URL
- 373 http://doi.wiley.com/10.1111/ele.12241.
- J. R. Gremer, S. Kimball, and D. L. Venable. Within-and among-year germination in Sonoran
- Desert winter annuals: bet hedging and predictive germination in a variable environment.
- Ecology Letters, 19(10):1209–1218, Oct. 2016. ISSN 1461023X. doi: 10.1111/ele.12655.
- URL http://doi.wiley.com/10.1111/ele.12655.
- L. Heinrich, J. Mller, A. Tellier, and D. ivkovi. Effects of population- and seed bank size
- fluctuations on neutral evolution and efficacy of natural selection. Theoretical Popula-

- tion Biology, 123:45-69, 2018. ISSN 10960325. doi: 10.1016/j.tpb.2018.05.003. URL
 https://doi.org/10.1016/j.tpb.2018.05.003.
- A. Lindn and S. Mntyniemi. Using the negative binomial distribution to model overdispersion in ecological count data. *Ecology*, 92(7):1414–1421, July 2011. ISSN 0012-9658. doi: 10.1890/10-1831.1. URL http://doi.wiley.com/10.1890/10-1831.1.
- 385 K. A. Mccue and T. P. Holtsford. Seed bank influences on genetic diversity in the rare
 386 annual Clarkia springvillensis (Onagraceae). American Journal of Botany, 85(1):30–36,
 387 1998. ISSN 00029122. doi: 10.2307/2446551.
- L. Nunney. 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(2):195, 2002. ISSN 00030147. doi: 10.2307/3079137. URL

 https://www.jstor.org/stable/10.2307/3079137.
- M. Paniw, P. F. Quintana-Ascencio, F. Ojeda, and R. Salguero-Gmez. Accounting for uncertainty in dormant life stages in stochastic demographic models. *Oikos*, 126(6):900–909, 2017. ISSN 16000706. doi: 10.1111/oik.03696.
- T. Philippi. Bet-Hedging Germination of Desert Annuals: Variation Among Populations and Maternal Effects in Lepidium lasiocarpum. *The American Naturalist*, 142 (3):488–507, Sept. 1993. ISSN 0003-0147, 1537-5323. doi: 10.1086/285551. URL https://www.journals.uchicago.edu/doi/10.1086/285551.
- K. Ritland. The joint evolution of seed dormancy and flowering time in annual plants living
 in variable environments. Theoretical Population Biology, 24(3):213–243, 1983. ISSN
 10960325. doi: 10.1016/0040-5809(83)90026-6.

- D. L. Venable. Bet hedging in a guild of desert annuals. *Ecology*, 88(5):1086–1090, May 2007.
- ISSN 0012-9658. doi: 10.1890/06-1495. URL http://doi.wiley.com/10.1890/06-1495.
- 404 R. S. Waples. Seed Banks, Salmon, and Sleeping Genes: Effective Population Size
- in Semelparous, Age-Structured Species with Fluctuating Abundance. The Ameri-
- can Naturalist, 167(1):118, 2006. ISSN 00030147. doi: 10.2307/3491252. URL
- https://www.jstor.org/stable/10.2307/3491252.