# Is shrub expansion into grasslands pushed or pulled? A spatial integral projection model for woody plant encroachment

Trevor Drees\*a,b, Brad M. Ochockib, Scott L. Collinsc, and Tom E.X. Millerb

<sup>a</sup>Department of Biology, Penn State University, State College, PA USA
 <sup>b</sup>Program in Ecology and Evolutionary Biology, Department of BioSciences, Rice
 University, Houston, TX USA

<sup>c</sup>Department of Biology, University of New Mexico, Albuquerque, NM USA

July 5, 2022

 $<sup>^*</sup>thd5066@psu.edu\\$ 

# 1 Abstract

Encroachment of shrubs into adjacent grasslands has become an increasingly reported phenomenon across the world, and such encroachment is either pulled forward by high population growth at the low-density encroachment front or pushed forward by higherdensity areas behind the front. However, at sites such as Sevilleta National Wildlife Refuge in central New Mexico, little is known about whether encroachment is pushed or pulled, and the dynamics of encroachment are not well-understood. Here, long-term encroachment of creosotebush (Larrea tridentata), a native perennial shrub, stands in stark contrast with the stagnation in encroachment observed in recent decades. In order to better understand creosotebush encroachment at this site, we quantify it using a spatially structured population model where a wave of individuals travels at a speed governed by both dispersal and density-dependence. Results indicate that population growth rates generally increase with decreasing density, suggesting that encroachment is pulled by 13 individuals at the low-density wave front, and the spatial population model predicts an 14 encroachment rate of less than 2 cm per year. While the predicted rate of encroach-15 ment is consistent with observations over recent decades, it does not explain long-term 16 creosotebush encroachment at the study site, suggesting that this process may occur in pulses when recruitment, seedling survival, or dispersal significantly exceed typical rates. Overall, our work demonstrates that individuals at low densities are likely the biggest contributors to creosotebush encroachment at this site, and that this encroachment is 20 likely a process that occurs in large but infrequent bursts rather than at a steady pace.

# 22 Keywords

- density-dependence, ecotones, woody encroachment, shrubs, integral projection model,
- 24 grassland

<sup>&</sup>lt;sup>1</sup> I am not editing the abstract for now.

# 25 Introduction

The recent and ongoing encroachment of shrubs and other woody plants into adjacent grasslands has caused significant vegetation changes across arid and semi-arid landscapes 27 worldwide (Van Auken, 2000, 2009; Goslee et al., 2003; Gibbens et al., 2005; Parizek et al., 28 2002; Cabral et al., 2003; Trollope et al., 1989; Roques et al., 2001). The process of en-29 croachment generally involves increases in the number or density of woody plants in both time and space (Van Auken, 2000), which can drive shifts in plant community structure 31 and alter ecosystem processes (Schlesinger et al., 1990; Ravi et al., 2009; Schlesinger 32 and Pilmanis, 1998; Knapp et al., 2008). Other effects of encroachment include changes 33 in ecosystem services (Reed et al., 2015; Kelleway et al., 2017), declines in biodiversity 34 (Ratajczak et al., 2012; Sirami and Monadjem, 2012; Brandt et al., 2013), and economic losses in areas where the proliferation of shrubs adversely affects grazing land and pastoral production (Mugasi et al., 2000; Oba et al., 2000). 37 Woody plant encroachment can be studied through the lens of spatial population 38 biology as a wave of individuals that may expand across space and over time (Kot et al., 39 1996; Neubert and Caswell, 2000; Wang et al., 2002; Pan and Lin, 2012). Theory pre-40 dicts that the speed of wave expansion depends on two processes: local demography and dispersal of propagules. First, local demographic processes include recruitment, survival, growth, and reproduction, which collectively determine the rate at which newly colonized locations increase in density and produce new propagules. Second, colonization events are driven by the spatial dispersal of propagules, which is commonly summarized as a 45 probability distribution of dispersal distance, or "dispersal kernel". The speed at which expansion waves move is highly dependent upon the shape of the dispersal kernel, especially long-distance dispersal events in the tail of the distribution (Skarpaas and Shea, 2007). Both demography and dispersal may depend on plant size, since larger plants

often have improved demographic performance and release seeds from greater heights,

leading to longer dispersal distances (Nathan et al., 2011). Accounting for population structure, including size structure, may therefore be important for understanding and predicting wave expansion dynamics (Neubert and Caswell, 2000).

Theory predicts that the nature of conspecific density dependence is another critical 54 feature of expansion dynamics but this is rarely studied in the context of woody plant 55 encroachment. Expansion waves typically correspond to gradients of conspecific density 56 - high in the back and low at the front - and demographic rates may be sensitive to density due to intraspecific interactions like competition or facilitation. If the demographic effects of density are strictly negative due to competitive effects that increase 59 with density then demographic performance is maximized as density goes to zero, at the 60 leading edge of the wave. Under these conditions, the wave is "pulled" forward by indi-61 viduals at the low-density vanguard (Kot et al., 1996), and targeting these individuals and locations would be the most effective way to slow down or prevent encroachment (cite?). However, woody encroachment systems often involve positive feedbacks whereby shrub establishment modifies the environment in ways that facilitate further shrub recruitment. For example, woody plants can modify their micro-climates in ways that 66 elevate nighttime minimum temperatures, promoting conspecific recruitment and sur-67 vival for freeze-sensitive species (D'Odorico et al., 2010; Huang et al., 2020). Positive density dependence (or Allee effects) causes demographic rates to be maximized at higher 69 densities behind the leading edge, which "push" the expansion forward, leading to qualita-70 tively different expansion dynamics (Kot et al., 1996; Taylor and Hastings, 2005; Sullivan 71 et al., 2017; Lewis and Kareiva, 1993; Veit and Lewis, 1996; Keitt et al., 2001). Pushed expansion waves generally have different shapes (steeper density gradients) and slower speeds than pulled waves (Gandhi et al., 2016), and may require different strategies for managing or decelerating expansion (check Taylor and Hastings ref). The potential for positive feedbacks is well documented in woody encroachment systems but it remains unclear whether and how strongly these feedbacks decelerate shrub expansion and influence strategies for management of woody encroachment.

In this study, we linked woody plant encroachment to ecological theory for invasion 79 waves, with the goals of understanding how seed dispersal and density-dependent demog-80 raphy drive encroachment, and determining whether the encroachment wave is pushed or 81 pulled. Throughout the aridlands of the southwestern United States, shrub encroachment 82 into grasslands is well documented (D'Odorico et al., 2012) but little is known about the 83 dispersal and demographic processes that govern it. Our work focused on encroachment of creosotebush (Larrea tridentata) in the northern Chihuahuan Desert. Expansion of this species into grasslands over the past 150 years has been well documented, leading to 86 decreased cover of Bouteloua eriopoda, the dominant foundation species of Chihuahuan 87 desert grassland (Gardner, 1951; Buffington and Herbel, 1965; Gibbens et al., 2005). 88 As in many woody encroachment systems, creosotebush expansion generates ecotones marking a transition from dense shrubland to open grassland, with a transition zone in between where shrubs can often be found interspersed among grasses (Fig. 1).

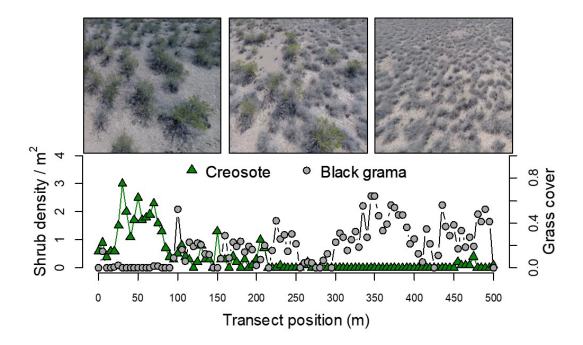


Figure 1: Example of an ecotone transect at Sevilleta LTER, spanning gradients of creosotebush and black grama grass. Photo credits: TEX Miller

Historically, creosotebush encroachment into grasslands is believed to have been 92 driven by a combination of factors including overgrazing, drought, variability in rainfall, 93 and suppression of fire regimes Moreno-de las Heras et al. (2016). These shrubs are also thought to further facilitate their own encroachment through positive feedbacks (Grover and Musick, 1990; D'Odorico et al., 2012) by modifying their environment in ways that favor continued growth and recruitment, including changes to the local micro-climate (D'Odorico et al., 2010) and rates of soil erosion (Turnbull et al., 2010). Such positive 98 feedback also involve suppression of herbaceous competitors, reducing competition as well as the amount of flammable biomass used to fuel the fires that keep creosotebush 100 growth in check (Van Auken, 2000). We hypothesized that, given potential for positive feedback mechanisms, the rarity of conspecifics at the low-density encroachment front may depress demographic performance and generate pushed-wave dynamics. 103

95

96

97

99

101

102

We used a combination of observational and experimental data from shrub ecotones 104 in central New Mexico to parameterize a spatial integral projection model (SIPM) that 105 predicts that speed of encroachment (m/yr) resulting from lower-level demographic and 106 dispersal processes. Our data came from demographic surveys and experimental trans-107 plants along replicate ecotone transects spanning a gradient of shrub density, and seed 108 drop experiments to estimate the properties of the dispersal kernel. We focused on wind 109 dispersal of seeds, since little is known about the natural history of dispersal in this 110 system and the seeds lack rewards to attract animal dispersers. We also used re-surveys 111 of permanents transects as an independent measure of encroachment that provided a 112 benchmark against which to evaluate model predictions. The SIPM accounts for size-113 structured demography of creosotebush, allows us to test whether shrub expansion is 114 pulled by the low-density front or pushed from the high-density core, and identifies the 115 local (demographic) and spatial (seed dispersal) life cycle transitions that most strongly contribute to expansion speed<sup>2</sup>. We address the following specific questions: 117

- 1. What is the nature of conspecific density dependence in demographic vital rates along shrub encroachment ecotones? Is encroachment pulled by the individuals at the front or pushed by individuals behind it?
- 2. What is the seed dispersal kernel for this species and how does this vary with maternal plant size?
- 3. What is the predicted rate of expansion from the SIPM and which lower-level processes most strongly affect the expansion speed?
  - 4. How does the observed rate of encroachment in the recent past compare to model predictions?

125

126

 $<sup>^2</sup>$  we will need to stay consistent with the language of encroachment/expansion/invasion. For now I am swicting a lot.

# 27 Materials and methods

# 28 Study species

Creosotebush Larrea tridentata is a perennial, drought-resistant shrub that is native to 129 the arid and semiarid regions of the southwestern United States and northern Mexico. 130 High-density areas of creosotebush consist largely of barren soil between plants due to 131 the "islands of fertility" these shrubs create around themselves (Schlesinger et al., 1996; 132 Reynolds et al., 1999), though lower-density areas will often contain grasses in the inter-133 shrub spaces (Fig. 1). In our northern Chihuahuan desert study region creosotebush 134 reproduces sexually, with numerous small yellow flowers giving rise to highly pubescent 135 spherical fruits several millimetres in diameter; these fruits consist of five carpels, each 136 of which contains a single seed. Seeds are dispersed from the parent plant by gravity and 137 wind, with the possibility for seeds to subsequently be transported by animals or water 138 (Maddox and Carlquist, 1985). In other regions, this species also reproduces asexually 139 and can give rise to long-lived clonal stands (Vasek, 1980), but this does not occur in our 140 study region. The foliage is dark green, resinous, and unpalatable to most grazing and 141 browsing animals (Mabry et al., 1978). 142

#### 143 Study site

We conducted our work at the Sevilleta National Wildlife Refuge (SNWR), a Long-Term Ecological Research (SEV-LTER) site in central New Mexico. The refuge exists at the intersection of several eco-regions, including the northern Chihuahuan Desert, Great Plains grassland, and steppes of the Colorado Plateau. Annual precipitation is approximately 250 mm, with the majority falling during the summer monsoon season from June to September. The recruitment events that facilitate creosotebush expansion are thought to be highly episodic (Peters and Yao, 2012), and this may be linked to fluctuations in monsoon precipitation (Boyd and Brum, 1983; Bowers et al., 2004). Monsoon precipitation

tion during the study years (2013-2017) was [summarise climate data].

## 153 Demographic data

#### 154 Ecotone transects

155

173

174

175

work was conducted at four sites in the eastern part of SNWR<sup>3</sup> (one site was initiated 156 in 2013 and the other three in 2014), with three transects at each site. All transects 157 were situated along a shrubland-grassland ecotone so that a full range of shrub densities 158 was captured: each transect spanned core shrub areas, grassland with no or few shrubs, 159 and the transition between them. Lengths of these transects varied from 200 to 600 m. 160 determined by the strength of vegetation transition since "steep" transitions required less 161 length to capture the full range of shrub density. We quantified shrub density in 5-meter "windows" along each transect, including all 163 shrubs within one meter of the transect on either side (shrubs that partially overlapped 164 with the census area were included). Densities were quantified once for each transect 165 (in 2013 or 2014) and were assumed to remain constant for the duration of the study, 166 a reasonable assumption for a species with very low recruitment and very high survival of established plants. Given the population's size structure, we weighted the density of 168 each window by the sizes of the plants, which we quantified as volume (cm<sup>3</sup>). Volume 169 was calculated as that of an elliptic cone:  $V_i = \frac{\pi h}{3} \frac{lw}{4}$  where l, w, and h are the maximum 170 length, maximum width, and height, respectively. Maximum length and width were 171 measured so that they were always perpendicular to each other, and height was measured

We collected demographic data during early June of every year from 2013-2017. This

in the window.

from the base of the woody stem at the soil surface to the tallest part of the shrub. The

weighted density for a window was then expressed as log(volume) summed over all plants

<sup>&</sup>lt;sup>3</sup>would a map be helpful?

#### 176 Observational census

At approximately 50-m intervals along each transect we tagged up to 10 plants for an-177 nual demographic census and recorded their local (5-m resolution) window so that we 178 could connect individual demographic performance to local density. These tagged shrubs 179 were revisited every June and censused for survival (alive/dead), size (width, length, and 180 height, as above), flowering status, and fertility of flowering plants (numbers of flower-181 buds, flowers, and fruits). In instances where shrubs had large numbers of reproductive 182 structures that would be difficult to reliably count (a large shrub may have thousands 183 of flowers or fruits), we made counts on a fraction of the shrub and extrapolated to es-184 timate whole-plant reproduction. Creosotebush does not have one discrete reproductive event per year; instead, flowering may occur throughout much of the warm season. By 186 combining counts of buds, flowers, and fruits we intended to capture a majority of the 187 season's reproductive output, assuming that all buds and flowers will eventually become 188 fruits. Our measurements of reproductive output are therefore conservative and may un-189 derestimate total seed production for an entire transition year. Each year, we searched 190 for new recruits within one m on either side of the transect. New recruits were tagged 191 and added to the demographic census. The observational census included a total of 522 192 unique individuals. 193

#### 194 Transplant experiment

We conducted a transplant experiment in 2015 to test how shrub density affects seedling survival. This approach complemented observational estimates of density dependence and filled in gaps for a part of the shrub life cycle that was rarely observed due to low recruitment. Seeds for the experiment were collected from plants in our study population in 2014. Seeds were germinated on Pro-Mix potting soil (Quakertown, PA) in Fall 2014 and seedlings were transferred to 3.8 cm-by-12.7 cm cylindrical containers and maintained in a greenhouse at Rice University. Seedlings were transported to SNWR

202 and transplanted into the experiment during July 27-31, 2015. Transplant timing was intended to coincide with the monsoon season, when most natural recruitment occurs.

The transplant experiment was conducted at the same four sites and three transects 204 per site as the observational demographic census, where we knew weight shrub densities 205 at 5-m window resolution. We established 12 1-m by 1-m plots along each transect. 206 Plots were intentionally placed to capture density variation: four plots were in windows 207 with zero shrubs, four plots were placed in the top four highest-density windows on the 208 transect, and the remaining four plots were randomly distributed among the remaining 209 windows with weighted density greater than zero. Plots were placed in the middle of 210 each 5-m window (at meter 2.5) and were divided into four 0.5-m by 0.5-m subplots. We 211 divided each subplot into nine squares (0.125-m by 0.125-m) and recorded ground cover 212 of each square as one of the following categories: bare ground, creosotebush, black grama 213 (B. eriopoda), blue grama (B. qracilis), other grass, or "other". Each subplot received one transplanted shrub seedling, for a total of 48 transplants per transect, 144 transplants 215 per site, and 576 transplants in the entire experiment. Each site was set up on a different 216 day and there was a significant monsoon event after the third and before the fourth 217 site. This resulted in differential mortality that appears to be related to site (captured 218 as a statistical random effect) but more likely reflects the timing of the monsoon event relative to planting (moist soil likely promoted transplant survival). We revisited the 220 transplant experiment on October 24, 2015 to survey mortality. After that first visit, 221 transplants were censused along with the naturally occurring plants each June, following 222 the methods described above. 223

#### 224 Demographic analysis

We fit statistical models to the demographic data and used AIC-based model selection to evaluate empirical support for alternative candidate models. The top statistical models were then used as the vital rate sub-models of the SIPM, so there is a strong connection between the statistical and population modeling, as is typical of integral projection modeling. Our analyses focused on the following demographic vital rates: survival, growth, probability of flowering, fertility (flower and fruit production), seedling recruitment, and seedling size. Most of these vital rates were modeled as a function of plant size, and all of them included the possibility of density dependence.

The alternative hypotheses of pushed versus pulled wave expansion rest on how the 233 rate of population increase  $(\lambda)$ , derived from the combination of all vital rates, respond 234 to density. We were particularly interested in whether demographic performance was 235 maximized as local density goes to zero (pulled) or at non-zero densities behind the 236 wave front (pushed). To flexibly model density dependence and detect non-monotonic 237 responses, we used generalized additive models in the R package 'mgcv' (Wood, 2017). 238 For each vital rate, we fit candidate models with or without a smooth term for local 239 weighted density (among other possible covariates). To avoid over-fitting, we set the 240 'gamma' argument of gam() to 1.8, which increases the complexity penalty, results in 241 smoother fits (Wood, 2017), and makes our approach more conservative (other gamma 242 values yielded qualitatively similar results). We pooled data across transition years for 243 analysis. All models included the random effect of transect (12 transects across 4 sites); 244 we did not attempt to model both site and transect-within-site random effects due to the low numbers of each. All vital rate functions used the natural logarithm of volume (cm<sup>3</sup>) 246 as the size variable and the sum of log(volume) as the weighted density of a transect 247 window. 248

Survival We modeled survival or mortality in year t+1 as a Bernoulli random variable with three candidate models for survival probability. These included smooth terms for initial size in year t only (1), initial size and weighted density (2), and both smooth terms plus an interaction between initial size and weighted density (3). We analyzed survival of experimental transplants and observational census plants together in the same analyses,

with a fixed effect of transplant status (yes/no) included in all candidate models. Since 254 recruits and thus mortality events were both very rare in the observational survey, this 255 approach allowed us to "borrow strength" over both data sets to generate a predictive 256 function for size- and possibly density-dependent survival while statistically accounting 257 for differences between experimental and naturally occurring plants. Because we had 258 additional, finer-grained cover data for the transplant experiment that we did not have for 259 the observational census, we conducted an additional stand-alone analysis of transplant 260 survival that explored the influence of covariates at multiple spatial scales (Appendix). 261

**Growth** We modeled size in year t+1 as a Gaussian random variable. There were nine 262 candidate models for growth. The simplest model (1) defined the mean of size in year 263 t+1 as a smooth function of size in year t and constant variance. Models (2) and (3) 264 had constant variance but the mean included smooth terms for initial size and weighted 265 density (2) or both smooth terms plus an interaction between initial size and weighted 266 density (3). Models 4-6 had the same mean structure as 1-3 but defined the standard 267 deviation of size in year t+1 as a smooth function of initial size. Models 7-9 mirrored 4-6 268 and additionally included a smooth term for weighted density in the standard deviation. 269 Modeling growth correctly is important because it defines the probability of any future 270 size conditional on current size, a critical element of the IPM transition kernel. We 271 verified that the AIC-selected model described the data well by simulating data from it 272 and comparing the moments (mean, variance, skewness, and kurtosis) of simulated and 273 real data. 274

Flowering and fruit production We modeled shrub reproductive status (vegetative or flowering) in year t as a Bernoulli random variable with three candidate models for flowering probability. These included smooth terms for current size (in year t) only (1), size and weighted density (3), and both smooth terms plus an interaction between size and weighted density. We modeled the reproductive output of flowering plants (the sum

of flowerbuds, open flowers, and fruits) in year t as a negative binomial random variable.

There were three candidate models for mean reproductive output that corresponded to

the same three candidates for flowering probability.

Recruitment and recruit size We modeled seedling recruitment in each transect 283 window as a binomial random variable given the number of total seeds produced in that 284 window in the preceding year. There were two candidate models, with and without 285 an influence of weighted density on the per-seed recruitment probability. To estimate 286 window-level seed production, we used the best-fit models for flowering and fruit produc-287 tion and applied this to all plants in each window that we observed in our initial density 288 surveys. We assume that recruits come from the previous year's seeds and not from a 289 long-lived soil seed bank. 290

We modeled recruit size as a Gaussian-distributed random variable and fit four candidate models including an influence of weighted density on mean, variance, both, and neither.

# 294 Density-dependent IPM

The size- and density-dependent statistical models comprised the sub-models of a density dependent Integral Projection Model (IPM) that we used to evaluate how the shrub population growth rate responded to con-specific density; we present this non-spatial model before layering on the spatial dynamics generated by seed dispersal. A basic density-independent IPM predicts the number of individuals of size x' at time t+1 (n(x',t+1)) based on a projection kernel (K) that gives the rates of transition from sizes x to x' from times t to t+1 and is integrated over the size distribution from the minimum (L) to maximum (U) sizes. In a density-dependent IPM, components of the

projection kernel may respond to population abundance and structure:

305

$$n(x',t+1) = \int_{L}^{U} K(x',x,\tilde{n}(t))n(x,t) dx \tag{1}$$

Here,  $\tilde{n}(t)$  is some function of population structure n(x,t) such as the total density of conspecifics  $(\tilde{n}(t) = \int n(x,t) dx)$  or, as in our case, total density weighted by size 306  $(\tilde{n}(t) = \int x n(x,t) dx)$ . For simplicity, in the analyses that follow we do not model density as a dynamic state variable; instead, we treat density as a static covariate  $(\tilde{n}(t) = \tilde{n})$  and 308 evaluate the IPM at a range of density values. As in our statistical modeling, the size 309 variable of the IPM (x, x') was  $log(cm^3)$ . 310 For our model, the size- and density-dependent demographic transitions captured by 311 the projection kernel include growth or shrinkage (q) from size x to x' conditioned on 312 survival (s) at size x (combined growth-survival function  $G(x', x, \tilde{n}) = g(x', x, \tilde{n})s(x, \tilde{n})$ ), and the production of new size-x' individuals from size-x parents  $(Q(x', x, \tilde{n}))$ . Repro-314 duction reflects the probability of flowering at size x(p), the number of seeds produced 315 by flowering plants (d), the per-seed probability of recruitment (r), and the size distri-316 bution of recruits (c). Collectively, the rate at which x-sized individuals produce x'-sized 317 individuals at density  $\tilde{n}$  is given by the combined reproduction-recruitment function  $Q(x', x, \tilde{n}) = p(x, \tilde{n})d(x, \tilde{n})r(\tilde{n})c(x', \tilde{n})$ . Thus, we can express the projection kernel as:

$$K(x\prime, x, \tilde{n}) = G(x\prime, x, \tilde{n}) + Q(x\prime, x, \tilde{n})$$
(2)

For analysis, we evaluated the IPM kernel over a range of local densities from the min-321 imum to the maximum of weighted density values from the 5-meter windows ( $0 \leq \tilde{n} \leq$ 322  $\tilde{n}_{max}$ ). At each density level, we discretized the IPM kernel into a  $200 \times 200$  approximat-323 ing matrix and calculated the asymptotic growth rate  $\lambda(\tilde{n})$  as its leading eignevalue. We 324 extended the lower (L) and upper (U) integration limits to avoid unintentional "eviction" 325 using the floor-and-ceiling method (Williams et al., 2012).

We sought to characterize the shape of density dependence: whether fitness declined monotonically or not with increasing density. We quantified uncertainty in the density-dependent growth rate  $\lambda(\tilde{n})$  by bootstrapping our data. For each bootstrap, we randomly sampled 75% of our demographic data, re-ran the statistical modeling and model selection, and used the top vital rate models to generate  $\lambda(\tilde{n})$  for that data subset. We repeated this procedure for 500 bootstrap replicates.

#### 333 Dispersal modelling

327

328

329

330

331

332

344

WALD dispersal model Dispersal kernels were calculated using the WALD, or Wald 334 analytical long-distance dispersal, model that uses a mechanistic approach to predict 335 dispersal patterns of plant propagules by wind. The WALD model, which is based in fluid 336 dynamics, can serve as a good approximation of empirically-determined dispersal kernels 337 (Katul et al., 2005; Skarpaas and Shea, 2007) and may be used when direct observations 338 of dispersal are not available. Under the assumptions that wind turbulence is low, wind 339 flow is vertically homogenous, and terminal velocity is achieved immediately upon seed 340 release, the WALD model simplifies a Lagrangian stochastic model to create a dispersal 341 kernel that estimates the likelihood a propagule will travel a given distance (Katul et al., 342 2005). Our dispersal kernel takes the form of the inverse Gaussian distribution

$$p(r) = \left(\frac{\lambda'}{2\pi r^3}\right)^{\frac{1}{2}} \exp\left[-\frac{\lambda'(r-\mu')^2}{2\mu'^2 r}\right]$$
(3)

that is a slight adaptation<sup>4</sup> from equation 5b in Katul et al. (2005), using r to denote dispersal distance. Here,  $\lambda'$  is the location parameter and  $\mu'$  is the scale parameter, which depend on environmental and plant-specific properties of the study system. (We use  $\lambda'$  for consistency with notation in related papers, but  $\lambda'$  the dispersal location parameter should not be confused with  $\lambda$  the geometric growth rate.) The location and scale parameters are defined as  $\lambda' = (H/\sigma)^2$  and  $\mu' = HU/F$ ; these are functions of the height

<sup>&</sup>lt;sup>4</sup>unclear what this refers to

H of seed release, wind speed U at seed release height, seed terminal velocity F, and the turbulent flow parameter  $\sigma$  that depends on both wind speed and local vegetation roughness. We parameterized the WALD dispersal kernel using windspeed data from the SEV-LTER weather station nearest our study site (Moore and Hall, 2022) and seed terminal velocity data from laboratory-based seed-drop experiments. Methods for our seed data collection and technical details of dispersal kernel modeling are provided in Appendix A.

#### 358 Spatial integral projection model

365

We used a spatial integral projection model to piece together seed dispersal and densitydependent demography, and generate predictions for the rate of shrub expansion that
results from this combination of local and spatial processes. The spatially explicit model
builds upon the non-spatial model (Eq. 1) and adds a spatial variable (z, z') such
that demographic transitions occur across both time and space according to a combined
demography-dispersal kernel  $\tilde{K}$ :

$$n(x\prime,z\prime,t+1) = \int_{-\infty}^{+\infty} \int_{L}^{U} \tilde{K}(x\prime,x,z\prime,z,\tilde{n}(z,t)) n(x,z,t) \, dx \, dz \tag{4}$$

Here,  $\tilde{K}(x', x, z', z, \tilde{n}(z, t))$  is the rate of transition from size x and location z to size x'366 and location z' given density  $\tilde{n}(z,t)$  at location z. As before,  $\tilde{n}$  is a function of pop-367 ulation structure - in our model, weighted local density - but here integrated over an 368 explicit competitive "neighborhood":  $\tilde{n}(z,t) = \int_{z-h}^{z+h} \int_{L}^{U} x n(x,z,t) \, dx \, dz$  where h repre-369 sents neighborhood size in the units of z. 370 Given that the shrub population at this site is approximately homogeneous perpen-371 dicular to the direction of encroachment, expansion is modelled as a wave moving in one 372 dimension. A spatial integral projection model (SIPM) is used to estimate the speed at 373 which encroachment occurs; such a model incorporates the effects of variation in traits like plant size that stage-structured models, such as those described in Neubert and Caswell (2000), do not capture. According to Jongejans et al. (2011), a general SIPM can be formulated as

$$\mathbf{n}(x_2, z_2, t+1) = \iint \tilde{K}(x_2, x_1, z_2, z_1) \mathbf{n}(x_1, z_1, t) dx_1 dz_1$$
 (5)

where  $x_1$  and  $x_2$  are locations of individuals of a particular size before and after one unit of 379 time, and  $z_1$  and  $z_2$  are the respective sizes. The vector **n** indicates the population density 380 of each size, and  $\tilde{K}$  is a kernel that combines dispersal with demography. Though this 381 SIPM represents a continuous spectrum of shrub sizes and densities, it was implemented 382 by discretising the above integral with a 200 x 200 matrix, as this makes calculations 383 significantly more tractable. 384 Movement of the wave is determined by the components of the combined disper-385 sal/demography kernel  $\tilde{K}$ , which is of the same form as that used in Jonjegans et al. 386 (2011). Here, 387

$$\tilde{K}(x_2, x_1, z_2, z_1) = K(x_2 - x_1)Q(z_2 - z_1) + \delta(x_2 - x_1)G(z_2 - z_1)$$
(6)

and K is the dispersal kernel, Q a reproduction function, G a growth function, and  $\delta$ 389 the Dirac delta function. G is derived from the model for annual growth ratio, and Q is 390 derived from the reproductive structures model as well as other factors including number 391 of seeds per reproductive structure, probability of recruitment from seed, and recruit 392 size. Both G and Q give the probability of transition between sizes; in the case of G, this 393 is the probability of growing from one specific size to another, and in the case of Q the 394 probability that an individual of a specific size produces a recruit of a specific size. The 395 product of K and Q represents the production and dispersal of motile propagales, while 396 the product of G and  $\delta$  represents the growth of sessile individuals. 397

Assuming strictly negative density dependence (i.e., pulled waves), the speed of the

398

moving wave can be calculated as

418

419

420

$$c^* = \min_{s>0} \left[ \frac{1}{s} \ln(\rho_s) \right] \tag{7}$$

where s is the wave shape parameter and  $\rho_s$  is the dominant eigenvalue of the kernel  $\mathbf{H_S}$ 402 (Jongejans et al., 2011). The kernel  $\mathbf{H_S}$  is defined as

$$\mathbf{H_S} = M(s)Q(z_2 - z_1) + G(z_2 - z_1) \tag{8}$$

where M(s) is the moment-generating function of the dispersal kernel (Jongejans et al., 2011). For one-dimensional dispersal, this moment-generating function can be estimated as

$$M(s) = \frac{1}{N} \sum_{i=1}^{n} I_0(sr_i)$$
(9)

where r is the dispersal distance for each observation, and  $I_0$  is the modified Bessel function of the first kind and zeroth order (Skarpaas and Shea, 2007). In order to obtain M, numerous dispersal distances were simulated from the dispersal kernel K(r) described in the previous section, with over 2000 replications for each shrub height increment of 1 cm. This was performed over the range from the lowest possible dispersal height to the maximum shrub height. Once M(s) was obtained for dispersal at each shrub height,  $H_S$  and  $c^*$  were calculated for each value of s; this was done for values of s ranging from 0 to 2, as it is this range in which  $c^*$  occurs.

Estimates of the wavespeed were bootstrapped for a total of 1000 replicates. Each bootstrap replicate recreated size- and density-dependent demographic models using 80% resampling on the original demographic data, and recreated dispersal kernels also using 80% resampling on the wind speeds and seed terminal velocities. Between replicates, the structure of the demographic models was kept constant, though coefficient estimates were not; this approach, while effectively ignoring model uncertainty, has the benefit of

increasing computational efficiency, which is especially useful given the time-consuming
nature of numerically estimating the many dispersal kernels used in the model.

#### 424 Encroachment re-surveys

We recorded shrub percent cover along two permanent 1000-m transects that spanned the shrub-grass ecotone, from high to low to near-zero shrub density. These surveys were conduced in summer 2001 and again in summer 2013 to document change in creosotebush abundance and spatial extent. At every 10 meters, shrub cover was recorded in nine cover classes (<1%, 1–4%, 5–10%, 10–25%, 25–33%, 33–50%, 50–75%, 75–95%, >95%). For visualization, we show midpoint values of these cover classes at each meter location for both transects and years.

# $_{\scriptscriptstyle 432}$ Results

# 433 Size and density dependent demography

Demographic data from naturally occurring and transplanted individuals revealed strong size- and density-dependence in demographic vital rates. For most sizes and vital rates, local density had negative demographic effects. Statistical support for size- and density-dependence is provided in Table XX, which provides AIC rankings for candidate models based on the completed (not bootstrapped) data set.

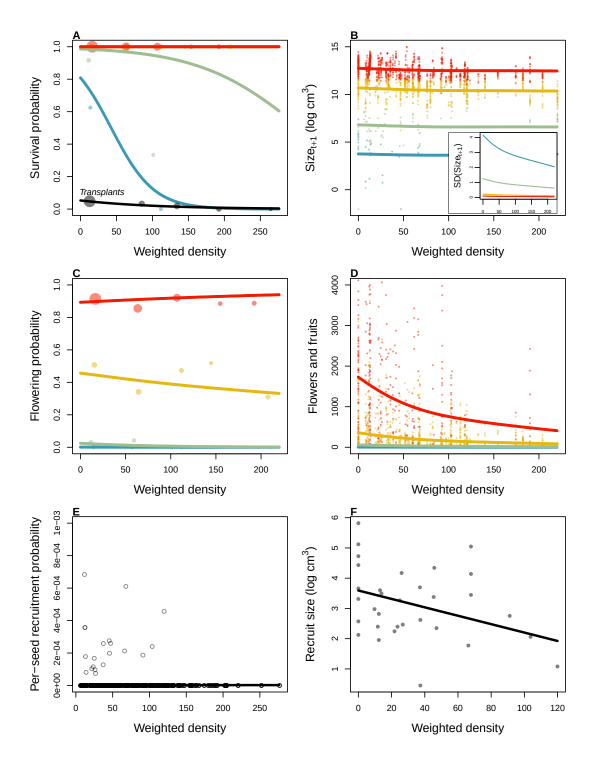


Figure 2: Size- and density-dependence in demographic vital rates. **A** Probability of survival from natural population census and transplant experiment (black line), **B** Mean and variance (inset) of size conditional on previous size, **C** Probability of flowering, **D** Flower and fruit production, **E** Probability of recruitment per seed, **F** Recruit size. In  $\mathbf{A}-\mathbf{E}$ , colored lines indicate four size groups (red is largest, blue is smallest), discretized for data visualization only. In all panel, weighted density is the sum of all plant sizes  $log(cm^3)$  within the same 5-m window as the census individual.

Survival Among naturally occurring plants, survival of large, established individuals was very high (Fig. 2A). We observed relatively few mortality events and nearly all of these were among new recruits. The probability of survival at these small sizes declined with increasing density. Survival of transplants was very low, lower even than survival of similarly-sized, naturally occurring recruits (Fig. 2B). However, the transplant results support the general pattern of negative density dependence in survival. Among the 20 survivors, 15 of them occurred in transect windows below the median of weighted shrub density.

SHORT PARAGRAPH SUMMARIZING SMALLER-SCALE ANALYSIS IN AP-PENDIX.

Growth Current size was strongly predictive of future size, as expected, and there was weak negative density dependence in mean future size conditioned on current size (Fig. 2C). However, there was a stronger signal of density dependence in the variance of future size (Fig. 2C, inset). Plants at low density exhibited greater variance in growth trajectories and this was especially true at the smallest sizes. Thus, large increases in the size of new recruits were most likely to occur under low-density conditions.

Flowering and fruit production Flowering probability was strongly size-dependent and and very weakly sensitive to local density (Fig. 2D). However, fertility of flowering plants was strongly negative density dependent, with greatest flower and fruit production by the largest plants at the lowest densities, and vice versa (Fig. 2E).

Recruitment and recruit size We observed 32 natural recruitment events along our transects during the study years and our estimate recruitment rate, given total expected seed production in each window preceding the recruitment year, was very low (2.47×10<sup>-6</sup>, 2F). While most recruitment events occurred at low density, this is also where most seed production was concentrated (Fig. 2E) and low-density windows were over-represented

relative to high density. For these reasons we were more likely to observe recruiment
events at low density. Controlling for sampling effort and seed production, the statistical
models indicated that our data were most consistent with a constant, density-independent
recruitment rate (Table XX). However, the mean size of new recruits declined significantly
with local density (Fig. 2F).

Population growth rate As expected based on the vital rate results, the asymptotic population growth rate  $\lambda$  declined monotonically with density (Fig. 3). This was true across all bootstrap replicates, indicating high certainty that shrub fitness is maximized at zero density and thus that the expansion wave is "pulled". Mean growth rate at low density was 3% per year, with bootstrap uncertainty spanning 1–6%. At high density in the core of the expansion wave, population growth rates approached  $\lambda = 1$ , indicating population stasis driven by near-perfect survival and extremely rare recruitment.

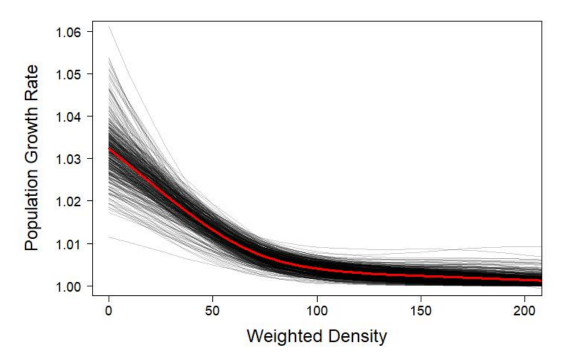


Figure 3: Density dependence in the asymptotic population growth rate ( $\lambda$ ). Lines show bootstrap replicates sub-sampled from the full demographic data set. Weighted density is the sum of all plant sizes  $log(cm^3)$  within 5-m windows.

## 476 Seed dispersal

As the speed of encroachment is quite limited, so is the extent of wind dispersal. Long 477 distance dispersal events, while more common for taller shrubs than their shorter coun-478 terparts, are still uncommon overall. For the tallest shrub height of 1.98 m, only 0.32% 479 of propagules exceed a dispersal distance of 5 m, and 0.02% exceed 10 m. At 1 m, or ap-480 proximately half the tallest shrub height, long distance dispersal is even less likely, with 481 0.0046% of propagules exceeding a dispersal distance of 5 m and 0.0009% exceeding 10 m. 482 Given that the median shrub height is only 0.64 m, the occurrence of long-distance wind 483 dispersal in most of the shrub population is highly improbable, and the few instances in which it occurs will only be limited to the tallest shrubs. Thus, as Figure 4 demonstrates,

shorter dispersal distances dominate; even for the tallest shrub, 81% of seeds fall within only a metre of the plant, and this percentage increases as shrub height decreases. Dispersal kernels have their highest probability density at dispersal distances between 2 and 8 cm from the shrub; here, as shrub height increases, the most probable dispersal distance slightly increases while maximum probability density decreases. Regardless of the shrub height, most dispersal will occur very close to the plant, though increases in shrub height dramatically increase the likelihood of dispersal at longer distances. It is clear that the shape of the height-dependent dispersal kernel K(r) varies greatly among the shrub population given the large range of shrub heights observed; shrubs at lower heights have more slender kernels with most of the seeds dispersing closer to the plant, while taller shrubs have kernels with much fatter tails and are more capable of longer-distance dispersal.

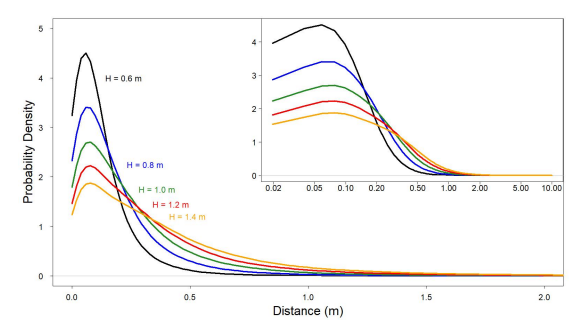


Figure 4: Dispersal kernels, with each colour representing a selected shrub height. The inset plot is the same as the large plot, though with a logarithmic x-axis to more easily show differences in dispersal probability at smaller distances.

# $_{ ext{\tiny 498}}$ Expansion speed

The speed of encroachment at the study site as estimated by the SIPM is rather slow; as 499 can be seen in Figure 5, the low-density wavefront moves at approximately 0.5 cm/yr un-500 der normal conditions and at 1 cm/yr under the best seedling survival conditions observed 501 in the dataset. These improved conditions were observed due to above-average rainfall 502 that occurred after greenhouse-grown seedlings were transplanted to the site. Population 503 growth in this low-density region of the moving wave is also low, with a geometric growth 504 rate of  $\lambda \approx 1.006$  and even lower rates of growth the higher-density regions behind; in 505 the higher-survival scenario the maximum rate increases to  $\lambda \approx 1.013$ , with growth still 506 decreasing as density increases. For both scenarios, the decrease in population growth rate with increasing density was monotonic across the range of observed standardised 508 densities, as is shown in Figure 5. This suggests that an Allee effect is likely not present 509 in this population, as the highest rate of population growth is found at the lowest density 510 vanguard of the encroaching population. Thus, the conditions necessary for equation 9 511 to be valid are satisfied, and these wavespeeds are applicable for a pulled-wave scenario in which no Allee effects are present. 513

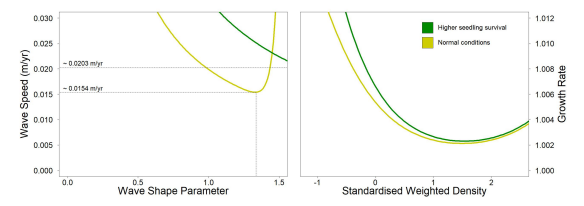


Figure 5: Estimated encroachment wave speeds (left) and geometric rates of population growth (right) for higher post-rainfall seedling survival and normal conditions.

#### 14 Transect re-surveys

Re-surveys along two permanent transects revealed virtually no change the in the creosote expansion wave over 12 years (Fig. 6). There were local changes in percent cover: on average cover increased by XX% between surveys. However, there was no clear indication that the leading edge of the creosote shrubland has advanced (the modest right-ward shift on both transects is within the range of measurement error).

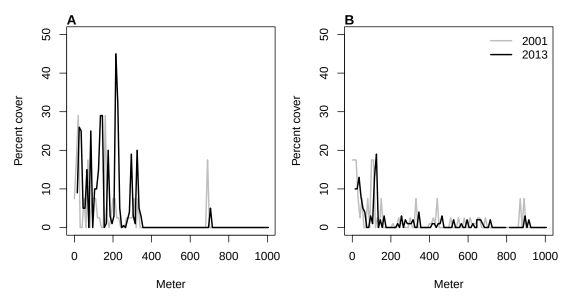


Figure 6: Re-surveys of shrub cover along two permanent trasects (A,B) surveyed in 2001 and 2013.

# Discussion

521

522

523

524

525

The slow movement of the encroaching creosotebush wave at the Sevilleta LTER site can likely be contributed to a combination of three factors: short dispersal distances with extremely limited long-distance dispersal events, very low probability of recruitment from seed, and high seedling mortality. These three barriers, when combined, form a formidable challenge to the establishment of new shrubs at the low-density front of the wave. First, a seed must travel far enough to avoid competition with the parent

shrub, which is unlikely given the dispersal kernels shown in Figure 2. Even if the seed 527 manages to be dispersed this far, its chances of becoming a seedling are low. Caching and consumption by seed-eaters such as a variety of seed-harvesting ants (Whitford, 1978; 529 Whitford et al., 1980; Lei, 1999) and the kangaroo rat Dipodomys merriami (Chew and 530 Chew, 1970) decreases the amount of seeds available for germination. However, reduc-531 tion in germination caused by destruction of seeds may be partly mitigated by the more 532 favourable germination conditions that these seeds can experience when cached under-533 ground (Chew and Chew, 1970). Many of the remaining seeds will still fail to germinate, and in the unlikely event that germination does occur, seedlings will likely die given 535 the high rates of mortality observed in smaller shrubs. Such high rates of creosotebush 536 seedling mortality have been observed in other studies as well (Boyd and Brum, 1983; 537 Bowers et al., 2004), probably due to a combination of herbivory, competition, and abiotic 538 stresses.

However, as low as they are, the wavespeed estimates given in this paper are still 540 conservative estimates for reasons mostly related to dispersal. First, it is important to 541 note that the dispersal kernels used here, while they account for variation in factors 542 such as wind speed and terminal velocity, may underestimate the distances that shrub 543 propagules travel. Because the WALD model assumes that terminal velocity is reached immediately upon seed release, seeds in the estimate thus take a shorter time to fall 545 and have less time to be transported by wind, and the true frequency of long-distance 546 dispersal events may thus be greater than what is estimated here. Second, dispersal at the 547 study site could occur through additional mechanisms other than wind. For example, 548 secondary dispersal through runoff from significant rainfall events can transport seeds (Thompson et al., 2014), and given that long-distance dispersal by bird and subsequent 550 species divergence is thought to be responsible for creosotebush being in North America 551 in the first place (Wells and Hunziker, 1976), short-distance dispersal by other animals 552 at the study site likely occurs. As mentioned above, seeds are transported by seedharvesting ants and granivorous mammals, where they are often stored in caches that can be appreciable distances from the parent shrubs. Whether transportation occurs via ant or rodent, creosotebush seeds can be moved significantly further than wind alone can, though many of these seeds are eventually consumed.

554

556

557

578

579

Despite the more conservative estimates our model yields, the estimated rate of dis-558 persal in creosotebush populations at the Sevilleta National Wildlife Refuge is consistent 559 with observations from the past 50-60 years, as crossotebush expansion during this time 560 has been minimal (Moreno-de las Heras et al., 2016). However, it cannot explain the 561 long-term increases in creosotebush cover at the study site, as total encroachment over 562 the past 150 years is much greater than what would be expected given the encroachment 563 rates derived by our models. Such a discrepancy is likely due to much of the expansion 564 occurring in an episodic fashion, with short times during which rapid encroachment oc-565 curs due to favourable environmental conditions. This could be due in part to seedling recruitment, which is a factor that strongly limits creosotebush expansion, being rare 567 and episodic. For example, Allen et al. (2008) estimate that a major recruitment event 568 occurred at this site in the 1950s, which is supported by photographic evidence from 569 Milne et al. (2003) of a drought-driven expansion during this time. Moreno-de las Heras 570 et al. (2016) estimate that after this expansion, several smaller creosotebush recruitment 571 events occurred in decadal episodes. However, such events can be highly localised and may not necessarily occur at the low-density front of encroachment, which could explain 573 how these recruitment events can still coexist with lack of encroachment in the recent 574 past. 575

Overall, our observations and model highlight three aspects of creosotebush encroachment that should be the focus of future studies seeking to obtain better estimates of encroachment rates. First, negative density dependence in survival, growth, and reproduction is demonstrated, along with size dependence. The clear dependence on size and conspecific density suggests that they both should be considered when estimating cre-

osotebush expansion and quantifying the demographic variation that contributes to it. 581 Second, wind dispersal in these shrubs is quite limited; though the dispersal kernels seen here are typical in the sense that they are characterised by high near-plant dispersal and 583 exceptionally low long-distance dispersal, the scale across which such dispersal occurs 584 is small, with most seeds landing within only 1 m of the shrub. Wind dispersal alone 585 may be an underestimate of the true amount of dispersal occurring, and future work 586 should seek to incorporate the effects of dispersal by runoff and animals so that a more 587 representative model of total dispersal can be obtained. Finally, encroachment is slow or 588 even stagnates, but only most of the time. Though our encroachment speed estimates 589 are representative of creosotebush populations for most years, the significant expansion 590 seen over larger time scales suggests that there is episodic expansion in other years; while 591 our model is consistent with the recent stagnation in creosotebush encroachment at the 592 Sevilleta LTER site, a model that also includes interannual variability in factors such as survival and recruitment would be able to better account for instances of episodic population expansion that are characteristic of this location.

# 596 Acknowledgements

# 597 Author contributions

# 598 Data accessibility

# References

Allen, A., W. Pockman, C. Restrepo, and B. Milne. 2008. Allometry, growth and population regulation of the desert shrub Larrea tridentata. Functional Ecology pages 197–204.

Bowers, J. E., R. M. Turner, and T. L. Burgess. 2004. Temporal and spatial patterns in

- emergence and early survival of perennial plants in the Sonoran Desert. Plant Ecology
- 605 **172**:107–119.
- Boyd, R. S., and G. D. Brum. 1983. Postdispersal reproductive biology of a Mojave Desert
- population of Larrea tridentata (Zygophyllaceae). American Midland Naturalist pages
- 608 25-36.
- Brandt, J. S., M. A. Haynes, T. Kuemmerle, D. M. Waller, and V. C. Radeloff. 2013.
- Regime shift on the roof of the world: Alpine meadows converting to shrublands in
- the southern Himalayas. Biological Conservation 158:116–127.
- Buffington, L. C., and C. H. Herbel. 1965. Vegetational changes on a semidesert grassland
- range from 1858 to 1963. Ecological monographs 35:139–164.
- Bullock, J. M., S. M. White, C. Prudhomme, C. Tansey, R. Perea, and D. A. Hooftman.
- 2012. Modelling spread of British wind-dispersed plants under future wind speeds in
- a changing climate. Journal of Ecology **100**:104–115.
- 617 Cabral, A., J. De Miguel, A. Rescia, M. Schmitz, and F. Pineda. 2003. Shrub encroach-
- ment in Argentinean savannas. Journal of Vegetation Science 14:145–152.
- 619 Chew, R. M., and A. E. Chew. 1970. Energy relationships of the mammals of a desert
- shrub (Larrea tridentata) community. Ecological Monographs pages 2–21.
- 621 D'Odorico, P., J. D. Fuentes, W. T. Pockman, S. L. Collins, Y. He, J. S. Medeiros,
- S. DeWekker, and M. E. Litvak. 2010. Positive feedback between microclimate and
- shrub encroachment in the northern Chihuahuan desert. Ecosphere 1:1–11.
- D'Odorico, P., G. S. Okin, and B. T. Bestelmeyer. 2012. A synthetic review of feedbacks
- and drivers of shrub encroachment in arid grasslands. Ecohydrology 5:520-530.
- 626 Gandhi, S. R., E. A. Yurtsev, K. S. Korolev, and J. Gore. 2016. Range expansions
- transition from pulled to pushed waves as growth becomes more cooperative in an

- experimental microbial population. Proceedings of the National Academy of Sciences
- **113**:6922-6927.
- 630 Gardner, J. L. 1951. Vegetation of the creosotebush area of the Rio Grande Valley in
- New Mexico. Ecological Monographs 21:379–403.
- 632 Gibbens, R., R. McNeely, K. Havstad, R. Beck, and B. Nolen. 2005. Vegetation changes
- in the Jornada Basin from 1858 to 1998. Journal of Arid Environments 61:651–668.
- 634 Goslee, S., K. Havstad, D. Peters, A. Rango, and W. Schlesinger. 2003. High-resolution
- images reveal rate and pattern of shrub encroachment over six decades in New Mexico,
- USA. Journal of Arid Environments **54**:755–767.
- 637 Grover, H. D., and H. B. Musick. 1990. Shrubland encroachment in southern New Mexico,
- USA: an analysis of desertification processes in the American Southwest. Climatic
- change 17:305-330.
- 640 Hsieh, C.-I., and G. G. Katul. 1997. Dissipation methods, Taylor's hypothesis, and
- stability correction functions in the atmospheric surface layer. Journal of Geophysical
- Research: Atmospheres **102**:16391–16405.
- 643 Huang, H., L. D. Anderegg, T. E. Dawson, S. Mote, and P. D'Odorico. 2020. Crit-
- ical transition to woody plant dominance through microclimate feedbacks in North
- American coastal ecosystems. Ecology **101**:e03107.
- Jongejans, E., K. Shea, O. Skarpaas, D. Kelly, and S. P. Ellner. 2011. Importance of
- individual and environmental variation for invasive species spread: a spatial integral
- projection model. Ecology **92**:86–97.
- 649 Katul, G., A. Porporato, R. Nathan, M. Siqueira, M. Soons, D. Poggi, H. Horn, and
- 650 S. A. Levin. 2005. Mechanistic analytical models for long-distance seed dispersal by
- wind. The American Naturalist 166:368–381.

- Keitt, T. H., M. A. Lewis, and R. D. Holt. 2001. Allee effects, invasion pinning, and species' borders. The American Naturalist 157:203–216.
- Kelleway, J. J., K. Cavanaugh, K. Rogers, I. C. Feller, E. Ens, C. Doughty, and N. Sain-
- tilan. 2017. Review of the ecosystem service implications of mangrove encroachment
- into salt marshes. Global Change Biology 23:3967–3983.
- 657 Knapp, A. K., J. M. Briggs, S. L. Collins, S. R. Archer, M. S. BRET-HARTE, B. E.
- Ewers, D. P. Peters, D. R. Young, G. R. Shaver, E. Pendall, et al. 2008. Shrub
- encroachment in North American grasslands: shifts in growth form dominance rapidly
- alters control of ecosystem carbon inputs. Global Change Biology 14:615–623.
- 661 Kot, M., M. A. Lewis, and P. van den Driessche. 1996. Dispersal data and the spread of
- invading organisms. Ecology 77:2027–2042.
- Lei, S. A. 1999. Ecological impacts of Pogonomyrmex on woody vegetation of a Larrea-
- Ambrosia shrubland. The Great Basin Naturalist pages 281–284.
- 665 Lewis, M., and P. Kareiva. 1993. Allee dynamics and the spread of invading organisms.
- Theoretical Population Biology 43:141–158.
- 667 Mabry, T. J., J. H. Hunziker, D. Difeo Jr, et al. 1978. Creosote bush: biology and
- chemistry of Larrea in New World deserts. Dowden, Hutchinson & Ross, Inc.
- 669 Maddox, J. C., and S. Carlquist. 1985. Wind dispersal in Californian desert plants:
- experimental studies and conceptual considerations. Aliso: A Journal of Systematic
- and Evolutionary Botany 11:77–96.
- Milne, B. T., D. I. Moore, J. L. Betancourt, J. A. Parks, T. W. Swetnam, R. R. Par-
- menter, and W. T. Pockman. 2003. Multidecadal drought cycles in south-central New
- Mexico: Patterns and consequences. Oxford University Press: New York, NY.

- 675 Moore, D., and K. Hall, 2022. Meteorology Data from the Sevilleta
- National Wildlife Refuge, New Mexico. Environmental Data Initiative.
- https://doi.org/10.6073/pasta/d56307b398e28137dabaa6994f0f5f92.
- 678 Moreno-de las Heras, M., L. Turnbull, and J. Wainwright. 2016. Seed-bank structure
- and plant-recruitment conditions regulate the dynamics of a grassland-shrubland Chi-
- huahuan ecotone. Ecology 97:2303–2318.
- 681 Mugasi, S., E. Sabiiti, and B. Tayebwa. 2000. The economic implications of bush
- encroachment on livestock farming in rangelands of Uganda. African Journal of Range
- and Forage Science **17**:64–69.
- Nathan, R., G. G. Katul, G. Bohrer, A. Kuparinen, M. B. Soons, S. E. Thompson,
- A. Trakhtenbrot, and H. S. Horn. 2011. Mechanistic models of seed dispersal by wind.
- Theoretical Ecology 4:113–132.
- Neubert, M. G., and H. Caswell. 2000. Demography and dispersal: calculation and
- sensitivity analysis of invasion speed for structured populations. Ecology 81:1613-
- 689 1628.
- Oba, G., E. Post, P. Syvertsen, and N. Stenseth. 2000. Bush cover and range condition
- assessments in relation to landscape and grazing in southern Ethiopia. Landscape
- ecology 15:535-546.
- Pan, S., and G. Lin. 2012. Invasion traveling wave solutions of a competitive system
- with dispersal. Boundary Value Problems **2012**:120.
- Parizek, B., C. M. Rostagno, and R. Sottini. 2002. Soil erosion as affected by shrub
- encroachment in northeastern Patagonia. Rangeland Ecology & Management/Journal
- of Range Management Archives **55**:43–48.

- Peters, D. P., and J. Yao. 2012. Long-term experimental loss of foundation species:
- consequences for dynamics at ecotones across heterogeneous landscapes. Ecosphere
- **3**:1–23.
- Ratajczak, Z., J. B. Nippert, and S. L. Collins. 2012. Woody encroachment decreases
- diversity across North American grasslands and savannas. Ecology 93:697–703.
- Raupach, M. 1994. Simplified expressions for vegetation roughness length and zero-
- plane displacement as functions of canopy height and area index. Boundary-Layer
- 705 Meteorology **71**:211–216.
- 706 Ravi, S., P. D'Odorico, S. L. Collins, and T. E. Huxman. 2009. Can biological invasions
- induce desertification? The New Phytologist 181:512–515.
- Reed, M., L. Stringer, A. Dougill, J. Perkins, J. Atlhopheng, K. Mulale, and N. Favretto.
- <sup>709</sup> 2015. Reorienting land degradation towards sustainable land management: Linking
- sustainable livelihoods with ecosystem services in rangeland systems. Journal of envi-
- ronmental management 151:472–485.
- Reynolds, J. F., R. A. Virginia, P. R. Kemp, A. G. De Soyza, and D. C. Tremmel. 1999.
- Impact of drought on desert shrubs: effects of seasonality and degree of resource island
- development. Ecological Monographs **69**:69–106.
- Roques, K., T. O'connor, and A. R. Watkinson. 2001. Dynamics of shrub encroach-
- ment in an African savanna: relative influences of fire, herbivory, rainfall and density
- dependence. Journal of Applied Ecology **38**:268–280.
- Schlesinger, W. H., and A. M. Pilmanis. 1998. Plant-soil interactions in deserts. Biogeo-
- chemistry **42**:169–187.
- Schlesinger, W. H., J. A. Raikes, A. E. Hartley, and A. F. Cross. 1996. On the spatial

- pattern of soil nutrients in desert ecosystems: ecological archives E077-002. Ecology 721 **77**:364–374.
- Schlesinger, W. H., J. F. Reynolds, G. L. Cunningham, L. F. Huenneke, W. M. Jarrell,
- R. A. Virginia, and W. G. Whitford. 1990. Biological feedbacks in global desertification. 724
- Science 247:1043-1048. 725
- Sirami, C., and A. Monadjem. 2012. Changes in bird communities in Swaziland savannas 726
- between 1998 and 2008 owing to shrub encroachment. Diversity and Distributions 727
- **18**:390-400. 728

722

- Skarpaas, O., and K. Shea. 2007. Dispersal patterns, dispersal mechanisms, and invasion 729
- wave speeds for invasive thistles. The American Naturalist 170:421-430. 730
- Sullivan, L. L., B. Li, T. E. Miller, M. G. Neubert, and A. K. Shaw. 2017. Density depen-
- dence in demography and dispersal generates fluctuating invasion speeds. Proceedings 732
- of the National Academy of Sciences 114:5053-5058. 733
- Taylor, C. M., and A. Hastings. 2005. Allee effects in biological invasions. Ecology 734
- Letters 8:895-908. 735
- Thompson, S. E., S. Assouline, L. Chen, A. Trahktenbrot, T. Svoray, and G. G. Katul. 736
- 2014. Secondary dispersal driven by overland flow in drylands: Review and mechanistic 737
- model development. Movement ecology 2:7. 738
- Trollope, W., F. Hobson, J. Danckwerts, and J. Van Niekerk. 1989. Encroachment and 739
- control of undesirable plants. Veld management in the Eastern Cape pages 73-89.
- Turnbull, L., J. Wainwright, and R. E. Brazier. 2010. Changes in hydrology and erosion
- over a transition from grassland to shrubland. Hydrological Processes: An Interna-742
- tional Journal 24:393-414. 743

- Van Auken, O. 2009. Causes and consequences of woody plant encroachment into western
- North American grasslands. Journal of environmental management **90**:2931–2942.
- Van Auken, O. W. 2000. Shrub invasions of North American semiarid grasslands. Annual
- review of ecology and systematics **31**:197–215.
- Vasek, F. C. 1980. Creosote bush: Long-lived clones in the Mojave Desert. American
- Journal of Botany **67**:246–255.
- Veit, R. R., and M. A. Lewis. 1996. Dispersal, population growth, and the Allee ef-
- fect: dynamics of the house finch invasion of eastern North America. The American
- 752 Naturalist 148:255–274.
- Wang, M.-H., M. Kot, and M. G. Neubert. 2002. Integrodifference equations, Allee
- effects, and invasions. Journal of mathematical biology 44:150–168.
- Wells, P. V., and J. H. Hunziker. 1976. Origin of the creosote bush (Larrea) deserts of
- southwestern North America. Annals of the Missouri Botanical Garden pages 843–861.
- Whitford, W., E. Depree, and P. Johnson. 1980. Foraging ecology of two chihuahuan
- desert ant species: Novomessor cockerelli and Novomessor albisetosus. Insectes Sociaux
- **27**:148–156.
- Whitford, W. G. 1978. Structure and seasonal activity of Chihuahua desert ant commu-
- nities. Insectes Sociaux **25**:79–88.
- Wiernga, J. 1993. Representative roughness parameters for homogeneous terrain.
- Boundary-Layer Meteorology **63**:323–363.
- Williams, J. L., T. E. Miller, and S. P. Ellner. 2012. Avoiding unintentional eviction
- from integral projection models. Ecology **93**:2008–2014.
- 766 Wood, S. 2017. Generalized Additive Models: An Introduction with R. 2 edition.
- 767 Chapman and Hall/CRC.

# Appendix A: Dispersal kernel modeling

WALD dispersal kernel In order to create the dispersal kernel, we first take the wind speeds at measurement height  $z_m$  and correct them to find wind speed U for any height H by using the logarithmic wind profile  $^5$ 

$$U = \frac{1}{H} \int_{d+z_0}^{H} \frac{u^*}{K} \log\left(\frac{z-d}{z_0}\right) dz \tag{A1}$$

given in Bullock et al. (2012) equation 6, with the notation slightly modified. Here, 773 z is the height above the ground, K is the von Karman constant, and  $u^*$  is the friction 774 velocity. The zero-plane displacement d and roughness length  $z_0$  are surface roughness 775 parameters that, for a grass canopy height h above the ground, are approximated by  $d \approx 0.7h$  and  $z_0 \approx 0.1h$ . These estimates are from Raupach (1994) for a canopy area index  $\Lambda = 1$  in which the sum of grass canopy elements is equal to the unit area being 778 measured. A 0.15 m grass height at our study site gives d = 0.105 and  $z_0$ , which are 779 suitable approximations for grassland (Wiernga, 1993). Calculations of  $u^*$  were done 780 using equation A2 from Skarpaas and Shea (2007), in which 781

$$u^* = KU_m \left[ \log \left( \frac{z_m - d}{z_0} \right) \right]^{-1} \tag{A2}$$

and  $U_m$  is the mean wind velocity at the measurement height  $z_m$ . Values for the turbulent flow parameter  $\sigma$  were then calculated using the estimate made by Skarpaas and Shea (2007) in their equation A4, where

$$\sigma = 2A_w^2 \sqrt{\frac{K(z-d)u^*}{C_0 U}} \tag{A3}$$

and  $C_0$  is the Kolmogorov constant.  $A_w$  is a constant that relates vertical turbulence to friction velocity and is approximately equal to 1.3 under the assumptions of above-

<sup>&</sup>lt;sup>5</sup> We need to describe and cite the wind data used here.

canopy flow made by Skarpaas and Shea (2007), based off calculations from Hsieh and Katul (1997). In addition, the assumption that z = H was made in order to make the calculation of  $\sigma$  more feasible.<sup>6</sup>

The values from the previous three equations give us the necessary information to 792 calculate  $\mu'$  and  $\lambda'$ , thus allowing us to create the WALD distribution p(r). However, the 793 base WALD model does not take into account variation in wind speeds or seed terminal 794 velocities, which limits its applicability in systems where such variation is present. In 795 order to account for this variation, we integrate the WALD model over distributions of these two variables using the same method as Skarpaas and Shea (2007). Additionally, 797 the WALD model assumes seed release from a single point source, which is not realistic 798 for creosote bush; because seeds are released across the entire height of the shrub rather 799 than from a point source, we integrated p(r) across the uniform distribution from the 800 grass canopy height to the shrub height. Thus, under the assumptions that the height at which a seed is located does not affect its probability of being released and that seeds 802 are evenly distributed throughout the shrub, this gives the dispersal kernel K(r), where 803

$$K(r) = \iiint p(F)p(U)p(z)p(r) dF dU dz$$
 (A4)

and p(F) and p(U) are the PDFs of the terminal velocity F and wind speed U, respectively, and p(z) is the uniform distribution from h to H.

Dispersal data collection The distribution p(F) in the integral above was constructed using experimentally determined seed terminal velocities. This was done by using laboratory-based seed release experiments with a high-speed camera and motion tracking software to determine position as a function of time. We then used the Levenberg-Marquardt algorithm to solve a quadratic-drag equation of motion for F. Before seeds were released, they were dried, dyed with yellow fluorescent powder, and then

804

805

806

<sup>&</sup>lt;sup>6</sup> Can you describe this assumption in biological terms?

put against a black background to improve visibility and make tracking easier. While the powder added mass to the seeds, this added mass only yielded an approximately 2.5% increase, likely having little effect on terminal velocities. Measurements were conducted for 48 seeds that were randomly chosen from a seed pool derived from different plants, and then an empirical PDF of terminal velocities was constructed using the data. Constructing p(U) involved creating an empirical PDF of hourly wind speeds using data from a Sevilleta LTER meterological station (Five Points), the station closest to our transects. We used wind speed data collected from 1988 to 2010.7

<sup>&</sup>lt;sup>7</sup> Most SEV data sets have a doi, so ideally we should cite the wind speed data.

# 821 Appendix B: Model selection results

surv	df	dAIC
$\tilde{\text{size}} + \text{transplant} + \text{size:transplant} + (1 \text{transect})$	11.50	1.72
~size + transplant + density + size:transplant + density:transplant +	13.19	0.19
(1 transect)		
~size + transplant + density + size:transplant + density:transplant +	14.22	0.00
size:density + size:transplant:density + $(1 $ transect)		

Table B1: AIC model selection for survival probability.

mean(size)	sd(size)	df	dAIC
$\tilde{size} + (1 transect)$	~1	3.00	1024.88
$\tilde{size} +  ext{density} + (1  ext{transect})$	~1	8.50	977.23
$\tilde{size} + density + size: density + (1 transect)$	~1	10.47	975.17
$\tilde{size} + (1 transect)$	~size	9.65	146.23
$\tilde{size} + density + (1 transect)$	~size	16.24	19.45
$\tilde{size} + density + size: density + (1 transect)$	~size	18.55	19.62
$\tilde{size} + (1 transect)$	~size + density	10.40	115.52
$\tilde{size} + density + (1 transect)$	~size + density	18.97	0.08
$\tilde{size} + density + size: density + (1 transect)$	~size + density	21.33	0.00

Table B2: AIC model selection for mean and variance of future size

Pr(Flowering)	df	dAIC
$\tilde{size} + (1  ext{transect})$	5.78	0.63
$\tilde{size} +  ext{density} + (1  ext{transect})$	6.80	2.32
$\tilde{size} + density + size: density + (1 transect)$	7.24	0.00

Table B3: AIC model selection for flowering probability.

No. fruits	df	dAIC
$\tilde{size} + (1  ext{transect})$	14.25	71.99
$\tilde{size} + density + (1 transect)$	5.52	0.00
$\tilde{size} + density + size:density + (1 transect)$	6.23	0.37

Table B4: AIC model selection for fruit number.

Pr(Recruitment)	df	dAIC
~(1 transect)	6.57	0.00
$\tilde{density} + (1 transect)$	7.39	0.93

Table B5: AIC model selection for recruitment probability.

mean(size)	sd(size)	df	dAIC
$\sim$ (1 transect)	~1	2.00	2.90
$\tilde{density} + (1 transect)$	~1	4.42	0.00
$\tilde{}(1 { m transect})$	$^{\sim}  m density$	3.00	4.74
$\tilde{density} + (1  ext{transect})$	$\tilde{density}$	5.56	1.21

Table B6: AIC model selection for mean and variance of recruit size.