1. INTRODUCTION

Variation in reproductive output is an important determinant of population dynamics, and understanding the drivers of this variation is necessary for predicting future population trends, performing scenario tests (e.g. threat assessment and ranking management actions), and preparing population viability analyses (PVA; e.g. Reed et al. 2009). Reproductive success is influenced by many factors, including climate, individual attributes, and population structure. Understanding the factors that influence reproductive output is essential for effective conservation planning.

The ‘bet hedging’ life history strategy of long-lived iteroparous species reduces short-term reproductive output to minimize the risk of reproductive failure over a lifetime. For desert-dwelling ectotherms living in variable and unpredictable environments, reproductive output is further influenced by precipitation and temperature via effects on food availability and limits on activity. We assembled multiple (n = 12) data sets on egg production for the threatened Agassiz’s desert tortoise Gopherus agassizii across its range and used these data to build a range-wide predictive model of annual reproductive output as a function of annual weather variation and individual-level attributes (body size and prior-year reproductive status). Climate variables were more robust predictors of reproductive output than individual-level attributes, with overall reproductive output positively related to prior-year precipitation and an earlier start to the spring activity season, and negatively related to spring temperature extremes (monthly temperature range in March–April). Reproductive output was highest for individuals with larger body sizes that reproduced in the previous year. Expected annual reproductive output from 1990–2018 varied from 2–5 to 6–12 eggs female⁻¹ yr⁻¹, with a weak decline in expected reproductive output over this time (p = 0.02). Climate-driven environmental variation in expected reproductive output was highly correlated across all 5 Recovery Units for this species (Pearson’s r > 0.9). Overall, our model suggests that climate change could strongly impact the reproductive output of Agassiz’s desert tortoise, and could have a negative population-level effect if precipitation is significantly reduced across the species’ range as predicted under some climate models.

KEY WORDS: Gopherus agassizii · Climate change · Fecundity · Population ecology · Meta-analysis
habitat quality (Powell & Steidl 2000, Blanchfield & Ridgway 2005) and levels of disturbance (Anthony et al. 2014), external pressures such as predation and available forage (Bjurlin & Wiseman 2004, Zanette et al. 2006), individual physiological condition (Hennen 2002, Vleck & Vleck 2002), and genetic differences between populations (Hagerty & Tracy 2010), among others. Many of these factors are driven or influenced by annual weather and changes in climate, which therefore can play a major role in reproductive success. For desert-dwelling animals, water availability can be a critical determinant of reproductive success, both directly via hydration and indirectly via the effects of precipitation on primary production (Beatley 1974, Smith et al. 1995, Wallis et al. 1999). Temperature can also be a critically important driver of reproductive success, especially for ectothermic species, whose bioenergetic processes and activity patterns can be extremely sensitive to alterations in the thermal environment (Clarke & Zani 2012, Zeh et al. 2012).

The impact of short-term climate-driven variation in reproductive output on population growth and extinction risk varies across life history types. With their ‘bet hedging’ life history strategy, long-lived iteroparous species are generally considered to be less vulnerable to short-term disruptions to reproduction than are semelparous species (Stearns 1976, Lovich et al. 2015). Nonetheless, long-lived species must maintain mean reproductive output above a critical minimum threshold for viability, and long-term changes in climate at generational time scales could push reproductive rates below sustainable levels (van de Pol et al. 2010). Furthermore, populations of long-lived species can persist for many years after reproductive rates fall below sustainable levels (Lovich et al. 2018a), often creating a misleading perception of population sustainability. Therefore, it is critical to understand and predict the long-term effects of climate variation on reproductive success for such species.

The threatened Agassiz’s desert tortoise Gopherus agassizii (USFWS 1990) is a long-lived desert-dwelling ectotherm whose physiology and behavior are strongly influenced by environmental conditions such as precipitation and temperature (Nagy & Medica 1986, Zimmerman et al. 1994, Lovich et al. 2012). As is many other reptiles, reproduction in the desert tortoise is seasonal, and larger females tend to produce more eggs than females with smaller body size (Turner et al. 1986, Mueller et al. 1998). However, body size typically only explains a small proportion of variation in reproductive output, and thus other exogenous factors also drive reproduction for this species (Wallis et al. 1999). Temperature appears to be a proximate cue for physiological changes and behavior related to the reproductive cycle as well as a determining factor for the length of the nesting season (Rostal et al. 1994, Lovich et al. 2012). Additionally, as an obligate herbivore, many aspects of desert tortoise ecology tied to fecundity, growth, and survivorship are positively correlated with the diversity and annual biomass of vegetation communities (Longshore et al. 2003, Medica et al. 2012). These processes are highly dependent on precipitation, which is extremely variable in the desert ecosystems occupied by the desert tortoise (Noy-Meir 1973, Beatley 1974). In fact, several analyses of desert tortoise egg production at long-term study sites have detected effects of temperature and precipitation on total annual reproductive output (Wallis et al. 1999, Hennen 2002, Nussear et al. 2012, Lovich et al. 2012). While these studies provide important insight into the effects of climate on annual variation in reproduction, detecting long-term trends in reproductive output is also a crucial component of conservation efforts. Recovery efforts for this species are focused on increasing tortoise abundance (USFWS 2011); therefore, an understanding of the factors that influence desert tortoise fecundity, on an annual scale as well as long-term trends, is fundamental to informing these efforts. Furthermore, with climate regimes across the species’ range expected to exhibit rapid shifts in coming decades (Seager et al. 2007, Barrows 2011), understanding the range-wide climatic drivers of reproductive success in this species is critical for assessing its future conservation outlook.

The objective of this study was to understand the endogenous (e.g. body size) and exogenous (e.g. climate) drivers of reproductive output (total number of eggs female$^{-1}$ yr$^{-1}$) for Agassiz’s desert tortoise, and to build a predictive model of egg production as a function of environmental conditions across the range of this species. Our study included multiple years of sampling at study sites across a large extent of the species’ range. We summarized our data at the level of the US Fish and Wildlife designated Recovery Unit (RU) for the tortoise, as this not only reflects a tangible metric for management of the species, but the RUs are delineated based on estimations of genetic structure of desert tortoise populations (Hagerty & Tracy 2010, USFWS 2011). Based on a priori knowledge of this species (see Section 2.1) and other turtle and tortoise species (Ernst & Lovich 2009), we hypothesized that increased female body size (midline carapace length, MCL) would have a
positive effect on reproductive output. We also hypothesized that higher total precipitation in the year leading up to the start of the nesting season would improve forage quality and availability during both vitellogenesis and egg production processes and thereby have a positive effect on reproductive output. Finally, we predicted that years with an earlier start to the activity season (driven by high spring temperatures early in the activity season) would be conducive to higher reproductive output, primarily due to enabling more females to produce multiple clutches (Wallis et al. 1999).

2. MATERIALS AND METHODS

2.1. Desert tortoise reproductive cycle

The reproductive cycle of the female Agassiz’s desert tortoise starts in late summer/early fall with follicular enlargement during vitellogenesis (coinciding with peak air temperatures; Rostal et al. 1994) (Fig. 1). This is followed by a fall mating period before activity decreases and tortoises seek winter hibernacula (Nussear et al. 2007). Upon emergence from hibernacula in the spring, when air temperatures increase to within the typical range of activity for this species (February–April; Nussear et al. 2007), ovulation and spring mating occur followed by oviductal shelling and tortoise nesting (Rostal et al. 1994). Females typically produce 1–3 clutches over the course of the nesting season, which generally occurs from May–July (although late April lay dates have been observed; Turner et al. 1986, 1984); females occasionally forgo reproduction. Following the end of the nesting period in mid- to late summer, there is a brief latent period before follicular enlargement begins and the cycle repeats.

2.2. Egg production data

Desert tortoise annual egg production data were acquired from 12 study sites distributed across the range of the species (Fig. 2). Data were collected from 1997–2018 as part of 7 separate studies (Table 1; some studies included >1 site), although data were not collected at all sites in each year (see Text S1 in the Supplement at www.int-res.com/articles/supp/n044p217_supp.pdf for site-specific information). Study sites were located within 4 of the 5 revised RUs for the desert tortoise: Upper Virgin River, Northeastern Mojave, Eastern Mojave, and Colorado Desert (USFWS 2011). Reproductive data in the Western Mojave RU were not available for this study, and applicability of resulting predictions may be limited in that region due to climatic differences between the Western Mojave and the rest of the range. Because the range of Agassiz’s desert tortoise extends over most of the Mojave Desert and the western portion of the Sonoran Desert, it encompasses broad climatic and elevational gradients (Nussear & Esque 2019).

In all studies used in our analyses, radio-transmitters were attached to the carapace of females and VHF radio-telemetry was used to relocate animals to assess reproductive status. Egg production was...
determined from X-radiographs taken at mostly weekly or biweekly intervals (depending on the study) using a portable X-ray machine between April and July or August of each year (Fig. S1). In addition, tortoises were weighed with spring scales (g) and their MCLs were measured using calipers (mm). Because multiple data sets based on differing protocols were collated for this analysis, we lacked necessary measurements on each individual for estimating a body score index (i.e. mass/volume; Wallis et al. 1999) and data collection precluded the adoption of visual body condition scoring into the standard sampling protocol (Lamberski 2013). A nesting event was recorded if a female previously observed with eggs was observed without eggs during a subsequent X-ray session. For a more detailed description of field methods see Lovich et al. (2015) and Nussear et al. (2012).

### 2.3. Climate data

Monthly climate data, that is mean daily maximum temperature ($T_{\text{max}}$), mean daily minimum temperature ($T_{\text{min}}$), and total precipitation (mm), were obtained from the PRISM database (PRISM Climate Group 2019; 800 m resolution). The monthly temperature range (MTR) was derived by subtracting $T_{\text{min}}$ from $T_{\text{max}}$. Daily temperatures were obtained from Daymet (Thornton et al. 2018; 1 km$^2$ resolution). We used Daymet data to determine the initiation of the spring activity period (activity–temperature threshold), which we defined as the first calendar day on which the maximum daily

| Site                   | Survey period | No. of years | Individual tortoises | Total observations | Source                        |
|------------------------|---------------|--------------|----------------------|--------------------|-------------------------------|
| Area 31                | 1998–2000     | 3            | 14                   | 34                 | Nussear et al. (2012)        |
| Barrow                 | 1997–1999     | 3            | 7                    | 12                 | Lovich et al. (1999)        |
| Bird Spring Valley     | 1997–2002     | 6            | 71                   | 216                | Nussear et al. (2012),      |
|                        |               |              |                      |                    | P. A. Medica unpubl. data   |
| Cima                   | 1997–2000     | 4            | 28                   | 69                 | Lovich et al. (1999),       |
|                        |               |              |                      |                    | P. A. Medica unpubl. data   |
| Cottonwood             | 2015–2016     | 2            | 7                    | 10                 | Lovich et al. (2018b)       |
| Lake Mead              | 1998–1999     | 2            | 20                   | 35                 | Nussear et al. (2012)       |
| Mesa                   | 1997–2000     | 4            | 18                   | 49                 | Lovich et al. (1999, 2012, 2015) |
| Orocopia               | 2017–2018     | 2            | 4                    | 7                  | Lovich et al. (2019)        |
| Pahcoo Flat            | 1998–2000     | 3            | 6                    | 13                 | Nussear et al. (2012)       |
| Piute Valley           | 1998–2002     | 5            | 37                   | 75                 | P. A. Medica unpubl. data   |
| Redcliffs              | 1998          | 1            | 10                   | 10                 | T. C. Esque unpubl. data    |
| Sandstone Mountain     | 1999–2000     | 2            | 4                    | 6                  | Nussear et al. (2012)       |
| Summary                | 1997–2018     | 37           | 226                  | 536                |                               |
temperature reached or exceeded 25°C for 5 consecutive days (based on Zimmerman et al. 1994, which indicated minimal tortoise activity below 25°C). Inclusion of the activity–temperature threshold variable was based on findings by Wallis et al. (1999) that the earlier in the season a female developed a clutch of eggs, the greater the total annual reproductive output due to the potential for development of multiple clutches. The climate variables used for further analysis were not highly correlated during the study period (Pearson’s |r| < 0.7).

2.4. Identifying appropriate climate ‘windows’

We used moving window analyses to determine optimal time windows for modeling egg production (‘climate windows’; implemented in the ‘climwin’ package for R; Bailey & van de Pol 2016, van de Pol et al. 2016). For each climate variable (T\textsubscript{max}, MTR, and precipitation) and for each reproductive process (reproductive probability [RP], egg production [EP]; see below), we assessed all possible climate windows in the 18 mo period leading up to the 1 May initiation of the nesting period in 1 mo increments (570 models: 3 climate variables × 190 climate windows). Variables were aggregated as the mean (for temperature variables) or sum (for precipitation) of all values within the climate window. We chose an 18 mo maximum time lag because current-year fecundity may plausibly be affected by current as well as previous-year vegetation availability (via capital energy reserves; Henen 2004), which could be driven by precipitation from the previous winter. We did not conduct a climate window analysis for the activity–temperature threshold variable, which was based on a well-specified a priori hypothesis (see Section 2.3).

We used logistic regression to model RP and Poisson regression to model EP (using total annual egg counts for all females that produced ≥1 clutch). The ‘baseline’ models for both reproductive processes (upon which climate variables were subsequently added) included female body size (MCL). We compared all models to the baseline (MCL-only) model using Akaike’s information criterion corrected for small sample size (AICc). To guard against inflated Type I error due to multiple comparisons, we used a permutation test (100 randomizations) implemented in ‘climwin’ that iteratively re-ran the climate window analysis with randomized dates (Bailey & van de Pol 2016, van de Pol et al. 2016). The top model (‘optimal’ climate window) was then compared to the null distribution of ΔAICc scores to calculate the likelihood that its ΔAICc score occurred by chance (PΔAICc). We only interpreted climate covariates with PΔAICc < 0.05.

2.5. Predictive model of desert tortoise reproductive success

We fitted our final egg production model using a Bayesian hierarchical modeling framework, which enables researchers to fit complex models that integrate multiple ecological processes and sources of error (Clark 2005). The hierarchical model we constructed consisted of an RP process model and an EP process model. In this framework, individuals either reproduce or not each year (with probability determined on the basis of the RP model), and the total number of eggs produced each year by each reproductive tortoise is determined on the basis of the EP model. We modeled RP as a Bernoulli process and EP as a Poisson process, analogous to a ‘zero inflated Poisson’ model (Zuur et al. 2009). We rescaled all continuous covariates (MCL, climate variables; see Table 2) to have a mean of 0 and standard deviation of 1.

The expected probability of reproducing for each individual each year was modeled as a logit-linear function of the observed reproductive status of individuals in the year prior, MCL, cumulative prior-year precipitation, and mean MTR from March–April of the current year. Mean egg production for each tortoise in each year (conditional upon being reproductive) was modeled as a log-linear function of MCL, cumulative precipitation, and the activity–temperature threshold. Specifically, we modeled the RP and EP processes using the following equations:

**RP process:**

\[
\text{logit}(pR_{ij}) = \text{logit}(bpR) + \beta_1 r_{s,i-1} + \beta_2 cp_{ij} + \beta_3 mcl_{ij} + \beta_4 mtr_{ij}
\]

**EP process:**

\[
\text{log}(EE_{ij}) = \text{log}(BE) + \beta_5 cp_{ij} + \beta_6 mcl_{ij} + \beta_7 att_{ij}
\]

where \(pR_{ij}\) is the probability of individual \(i\) reproducing in year \(j\), \(bpR\) is the baseline probability of reproducing (logit scale), \(r_{s,i-1}\) is the observed or estimated (latent) reproductive status of individual \(i\) in year \(j - 1\), \(cp_{ij}\) is cumulative prior-year precipitation for individual \(i\) in year \(j\), \(mcl_{ij}\) is the MCL for individual \(i\) in year \(j\), \(mtr_{ij}\) is the mean MTR for individual \(i\) in year \(j\), \(EE_{ij}\) is the expected number of eggs produced by individual \(i\) in year \(j\), \(BE\) is the baseline number of eggs pro-
duced annually by reproductive females (log scale), \( \text{att}_{ij} \) is the activity–temperature threshold date (see Table 2), and \( \beta_{1-7} \) represent regression coefficients. For Bayesian parameter estimation, \( b_{pR} \) was assigned a vague Beta(1,1) prior and \( BE \) was assigned a Uniform(1,15) prior. All other regression coefficients were assigned Gaussian priors with a mean of 0 and variance of 10.

Models were fitted in R v.3.6.0 using a Markov Chain Monte Carlo algorithm implemented in the JAGS software (Plummer 2017, R Core Team 2019), which was called from R using the ‘runjags’ package (Denwood 2016). We ran 3 independent chains for 25,000 iterations each after a burn-in sample of 2000, and stored every 10th run for a total of 7500 joint posterior samples. Model convergence for each parameter was inspected by visually examining traceplots and by examining the Gelman-Rubin and the Heidelberger and Welch convergence diagnostic tests (Heidelberger & Welch 1983, Brooks & Gelman 1998). All code and data for fitting this model are available at https://github.com/kevintshoemaker/TortoiseEggModel.

2.6. Model validation

To assess goodness-of-fit, we simulated multiple additional data sets based on parameter values drawn from the fitted joint posterior distribution. In order to quantify whether the observed data could plausibly have been generated from our best fit model, we computed the root mean squared error (RMSE) for the simulated data sets and contrasted these values with RMSEs computed for the observed data. We calculated a Bayesian posterior predictive p-value (metric of goodness-of-fit) as the fraction of instances in which the RMSE values from simulated data sets exceeded the RMSE values computed using the observed data set (Kéry 2010). In general, a p-value close to 0.5 generated under these methods demonstrates that the model’s predictions are close to the observed data (Kéry 2010, Gelman et al. 2013).

To assess model predictive performance, we ran a 10 fold cross-validation procedure in which one-tenth of individual tortoises were held out from the model fitting process in turn and the observed number of eggs was compared with the predicted number of eggs for all held-out observations. We used area under the curve (AUC; for the RP process) and \( R^2 \) (for the EP process) metrics to summarize model predictive performance across all holdout samples. Predictive performance for the EP process was only evaluated for observations with \( \geq 1 \) egg produced. To evaluate the relative contributions of climate variables and individual-level variables to predictive performance, we repeated the cross-validation procedure for (1) a reduced model with only individual-level covariates and (2) a reduced model with only climate covariates.

2.7. Model interpretation and predictions

We interpreted covariate effects using 3 sets of partial dependence plots: one depicting the probability of reproducing across the full range of all model covariates, one depicting the total egg production, conditional on reproducing, and the final set depicting the total expected number of eggs (accounting for both reproductive processes).

2.8. Predictive maps

Predictive maps of annual egg production potential were calculated using an 800 × 800 m grid matching the PRISM data grid over the range of desert tortoise habitat (Carter et al. 2020) using parameters from the best model and raster products of all input data spanning the study area. The mean tortoise MCL used for predictions was calculated from 4579 females using the average from this study, as well as several study sites across the range (Table S1). Annual maps were created for the time period from 1990–2018. To account for the prior year reproduction effect on the reproductive process, we conducted a 10 yr ‘burn in’, where we set all reproduction values for the initial year to a probability of 1, and cycled the reproductive process (according to the model parameters) over 10 yr prior to calculating the first reproductive process used in map production. The annual egg production for each successive year was then taken as the reproductive process of the prior and current year, along with the egg laying process calculated for the current year. Averaged egg potential values within the 5 RUs (Fig. 2; USFWS 2011) were aggregated to create RU-based estimates of average annual egg production potential. Finally, we computed the Pearson correlation in climate-driven annual variation between each pair of RUs.

3. RESULTS

Across all studies, we obtained data on reproductive output for a total of 226 unique females and 536 unique individual–year combinations (Fig. S2).
3.1. Climate variable selection and ‘climate window’ analyses

Both cumulative precipitation and MTR had meaningful effects on the RP and EP processes, while \( T_{\max} \) had no meaningful effect on either. However, the coefficient on MTR in the EP model was not significant and was therefore excluded from our final Bayesian predictive model in favor of the Daymet activity–temperature threshold variable. The optimal climate windows which were incorporated into our final model were cumulative precipitation from 14–2 mo before 1 May (13 mo; March–March) of the nesting year and monthly temperature range from 2–1 mo before 1 May (March–April) of the nesting year (Table 2, Fig. 1).

3.2. Predictive model of desert tortoise reproductive success

The fitted Bayesian hierarchical model confirmed the importance of all factors included in the linear model, as none of the 95% credible intervals for regression coefficients overlapped zero (Table 3). The coefficient estimates indicated positive effects for cumulative precipitation and MCL on both the RP and EP processes, a negative effect of MTR on the RP process, and a negative effect of the calendar day of first activity (activity–temperature threshold) on the RP process. Variables were standardized prior to the analysis (see Section 2.5); therefore, relative contributions of each variable are interpreted based on the magnitude of the resulting coefficient value. The intercept of the RP model was estimated at 66% and interpreted as the baseline probability of reproducing for an average female in an average year. Cumulative precipitation had the highest magnitude coefficient value in the RP process (\( \beta = 0.62 \)), resulting in the expected probability of reproducing increasing from 0.76–0.99 as cumulative precipitation increased from 30.2–637.1 mm (Fig. 3). March–April MTR (\( \beta = -0.41 \)) also had a significant effect on the RP process, resulting in expected probability of reproduction decreasing from 0.89–0.66 as temperature range increased from 9.6–19.2°C. Finally, 2 individual-level attributes — MCL (\( \beta = 0.34 \)) and prior-year reproductive status (\( \beta = 1.46 \)) — affected the RP process, with probability of reproduction increasing from 0.76–0.96 over the observed range of MCL (180–289 mm) and increasing from 0.66–0.89 for females that reproduced in the previous year (Fig. 3). For the EP process, the baseline number of eggs was estimated as 6.5 eggs per average female in an average year. Cumulative precipitation (\( \beta = 0.15 \)) had the strongest influence on the EP process, with mean number of eggs expected to increase from 5–13 eggs with an increase from 30.2–637.1 mm precipitation. The second most influential covariate on the EP process was MCL (\( \beta = 0.12 \)), with mean number of eggs expected

| Covariate                                                                 | Description                                                                 | Source                                                                 |
|--------------------------------------------------------------------------|-----------------------------------------------------------------------------|-----------------------------------------------------------------------|
| **RP process**                                                           |                                                                             |                                                                      |
| Reproductive status \((rs_{ij−1}; \text{latent variable})\)            | Yes/No nesting observed in year prior; recorded annually for each tortoise\(^a\) | See Table 1                                                           |
| Cumulative precipitation \((cp_{ij})\)                                  | Cumulative precipitation over the 13 mo (March–March) prior to the nesting season | PRISM Climate Group (2019); 800 m                                      |
| Midline carapace length \((mcl_{ij})\)                                 | The length of the carapace measured annually by caliper for each tortoise (mm) | See Table 1                                                           |
| Monthly temperature range \((mtr_{ij})\)                               | The average monthly temperature range (mean daily maximum – mean daily minimum) of the 2 mo leading to the nesting season (March–April) | PRISM Climate Group (2019); 800 m                                      |
| **EP process**                                                           |                                                                             |                                                                      |
| Cumulative precipitation \((cp_{ij})\)                                  | Cumulative precipitation over the 13 mo (March–March) prior to the nesting season | PRISM Climate Group (2019); 800 m                                      |
| Midline carapace length \((mcl_{ij})\)                                 | The length of the carapace measured annually by caliper for each tortoise (mm) | See Table 1                                                           |
| Activity-temperature threshold \((att_{ij})\)                          | Calendar day of first 5 d period where the maximum temperature is above 25°C | Daymet: Thornton et al. (2018); 1 km                                    |

\(^a\)Instances of unknown prior-year reproductive status were estimated as latent parameters in our model
to increase from 4.6–9.2 eggs with an MCL increase from 180–289 mm. The activity–temperature threshold ($\beta = -0.07$) had a weak negative effect on the EP process, with egg production expected to decrease from 8–5 eggs as the date of reaching the activity–temperature threshold shifts from late January to mid-June (Fig. 3).

### 3.3. Model convergence and performance evaluation

The potential scale reduction factor and Heidelberger and Welch diagnostics confirmed that all chains converged on the posterior distribution (Gelman et al. 2013). The posterior predictive check resulted in a relatively even division of RMSE values above and below the 1:1 line (Bayesian posterior predictive $p = 0.7$), suggesting that the model could plausibly have generated the observed data (Gelman 2013). From 10-fold cross-validation (leaving out entire individuals and thereby eliminating any information about prior reproductive status), we showed that model performance was generally weak for predicting reproductive status.
(AUC = 0.67; 95% CI: 0.65–0.7). When trained to the full data set, AUC increased to 0.75 (95% CI: 0.72–0.79). For reproductive individuals, our model explained 30% of the observed variance in egg production for withheld individuals ($R^2$; 95% CI: 0.28–0.31). When climate variables (precipitation, MTR, activity–temperature threshold) were excluded from the predictive model, AUC for the RP process decreased from 0.67–0.54 (little better than random), and median $R^2$ for the EP process decreased from 0.30–0.23 in cross-validation. When individual-level covariates (MCL, prior-year reproduction) were excluded from the model, AUC for the RP process increased from 0.67–0.69 and median $R^2$ for the EP process decreased from 0.30–0.24 in cross-validation.

### 3.4. Egg potential per RU

Egg potential maps indicated that egg production ranges from low values (e.g. 2–5 eggs female$^{-1}$ yr$^{-1}$) in years with unfavorable climate conditions for reproductive processes to average egg production values of 6–12 eggs female$^{-1}$ yr$^{-1}$ in favorable conditions. These values differed among the 5 RUs, and among years, and some RUs benefited more from improved conditions. For example, increases in egg production relative to the lowest year were highest for the Northeastern Mojave and Upper Virgin River RUs in both moderate and high years, while increases in the more southern RUs were more modest (Fig. 4). Further, there were spatial differences in egg production potential that remained lower even in the highest years (e.g. the southernmost portion of the distribution, as well as areas near Death Valley and the Owens Valley of California) (Fig. 4—High). Annual variation in egg production potential indicates a generally declining trend over time ($p = 0.02$), with the notable exception of a single year (2004) in which reproductive potential was higher in all RUs (Fig. 4). Annual climate-driven variation in expected egg production was highly correlated among RUs, with pairwise correlation ranging from $|r| = 0.91$ (Western Mojave vs. Upper Virgin River) to $|r| = 0.97$ (Eastern Mojave vs. Western Mojave).

### 4. DISCUSSION

We found that increased reproductive output was associated with increased prior-year cumulative precipitation and an earlier (warmer) start to the nesting season. Interestingly, with climate change likely to result in warmer climates and potentially increased water deficits, our model suggests that earlier activity periods (resulting in increased reproductive output) could offset the negative impact of reduced precipitation. However, if climate change results in reduced forage availability (e.g. via reduced precipitation), tortoises may forgo additional reproduction attempts even if temperatures permit a longer nesting season (Henen 2002). Furthermore, earlier activity seasons could disrupt the temporal synchrony between important events that must occur (e.g. forage availability, vitellogenesis) in order for tortoises to successfully reproduce or for hatchlings to survive (Post & Forchhammer 2008). Accurate projection of tortoise reproductive output under future non-analogue climate conditions will require further investigation into the potential for phenological mismatch or other complex interactive effects of precipitation and temperature on egg production.

The relationship we identified between cumulative prior-year precipitation and reproductive output almost certainly relates to the role of precipitation in regulating forage availability (Beatley 1974, Medica et al. 2012). The positive relationship between precipitation, and thus resource availability, and egg production observed in our study has also been observed in previous studies (Turner et al. 1986, Wallis et al. 1999, Nussear et al. 2012, Lovich et al. 2015).
Although many previous studies have focused on the importance of winter precipitation in the Mojave and western Sonoran desert ecosystems (e.g., Wallis et al. 1999, Nussear et al. 2012, Lovich et al. 2015), our model results suggest that cumulative precipitation across all seasons (13 mo prior to the nesting season) may be important for reproductive success in Agassiz’s desert tortoise. While not as productive as winter annuals, there is a late summer flora of Mojave annuals in years of higher summer precipitation (Beatley 1974), and this additional forage, as well as additional hydration, may provide important resources that influence reproduction, and may become more important if climate change involves changes in the timing of precipitation. Our results suggest that forage availability may not only influence egg development and follicle maintenance during the spring nesting season, but it may also influence the number of follicles that reach ovulatory size in late summer/early fall (Rostal et al. 1994). Incorporating a seasonal measure of female body condition into future analyses may provide further insight into the role of sum-

![Fig. 4. Range-wide spatiotemporal projections of annual egg production for Agassiz’s desert tortoise Gopherus agassizii. (A) Maps of expected egg potential for the years with the lowest, average (‘medium’), and highest mean annual predicted egg potential (range-wide) for 1990−2018. (B) Mean annual predicted egg potential by Recovery Unit (CO: Colorado Desert; WM: Western Mojave; EM: Eastern Mojave; NEM: Northeastern Mojave; UVR: Upper Virgin River) for the same years depicted in (A), where the 25th, 50th, and 75th quartiles are given by the white bar within the violin plot representing the range of the data. (C) Averaged annual egg potential plotted by year for the 5 Recovery Units](image)
mer vs. winter precipitation on reproductive output. For example, Averill-Murray et al. (2018) found that body condition of female Sonoran desert tortoises *Gopherus morafkai*, a closely related species, was positively related to multiple aspects of reproductive output, including follicle growth, clutch size, and egg width.

Our findings that the majority of tortoises (66%) are predicted to reproduce in an average year and that tortoises exhibit substantial reproductive output even under unfavorable conditions (62% of study animals reproduced in 1997, one of the least productive years of our study period; Fig. 4), are consistent with a bet-hedging strategy in which reproductive output is sparingly spread over many years to maximize recruitment probability in an unpredictable environment (Henen 2004, Lovich et al. 2015). Furthermore, our finding of a positive correlation between reproductive status in current and prior years may suggest that high forage availability in one year affords Agassiz’s desert tortoises the ability to spread reproductive effort over more than one year. Confirming this hypothesis will require additional studies.

Our study, consistent with Wallis et al. (1999) and Lovich et al. (2012), suggests that desert tortoises can (when they are provisioned with sufficient resources for greater output) produce more eggs in years with warmer springs, possibly due to the potential for accelerated egg development and because more females are able to produce multiple clutches due to a longer nesting season. Lovich et al. (2012) observed that the development of an initial clutch was affected by spring temperatures, with clutch development delayed in cooler springs relative to warmer springs. Findings by Wallis et al. (1999) demonstrated that the earlier in the season a female desert tortoise developed an initial clutch, the greater the annual reproductive output, likely due to the potential to produce multiple clutches. In fact, previous studies have suggested that differences in total egg production among individuals are driven largely by the number of clutches laid (Turner et al. 1986, Wallis et al. 1999, but see Mueller et al. 1998). The tendency to produce multiple clutches over the course of a season is another aspect of the bet-hedging strategy used by this species (e.g. spreading the risk temporally and spatially; Ennen et al. 2017). Given that sex of the offspring in tortoises is controlled by nest temperatures (Rostal et al. 2002), the partitioning of eggs into multiple clutches may also be a factor in maintaining balanced sex ratios in populations. Further research is needed to better understand how climate change is likely to affect this complex strategy and the potential for producing multiple clutches. While not included here, we hope to investigate the factors influencing the partitioning of and timing of reproductive output into multiple clutches in future analyses of these data.

The positive relationship we detected between female body size and reproductive output has been well established for many turtles (Landers et al. 1980, Gibbons 1982, Congdon & Gibbons 1985), including the desert tortoise (Mueller et al. 1998, Wallis et al. 1999, Nussear et al. 2012). Medica et al. (2012) demonstrated that tortoise growth is dependent upon the amount of vegetation available, which in turn is highly correlated with the amount of seasonal rainfall. Thus, changes in climate that negatively impact vegetation growth could inhibit tortoise body size, which would likely cause a delay and ultimately a decrease in egg production. This suggests that changes in precipitation may cause long-term effects on reproductive output beyond the immediate impacts documented in this study. For example, climate forecasts also predict more extreme events (with respect to both temperature and precipitation; Archer & Predick 2008), which can promote invasive plant species (e.g. *Bromus* spp.) with known harmful consequences for tortoise growth and nutrition (Tracy et al. 2006, Drake et al. 2016), as well as altered fire cycles that can lead to large-scale changes in habitat condition (D’Antonio & Vitousek 1992, Van Linn et al. 2013, Horn & St. Clair 2017).

Our study detected important range-wide spatial differences in predicted reproductive rates for Agassiz’s desert tortoise. For example, our model indicates reduced reproductive capacity in southern portions of the range (Fig. 4), where habitat is also predicted to be decreasing (Barrows 2011), suggesting that those populations may experience heightened conservation challenges. In the northern portions of the range there is differential predicted reproductive potential in the western and eastern extents; the northeastern regions are predicted to have higher reproduction, while the northwestern have lower predicted rates, and less pronounced response to years with higher precipitation (Fig. 4). Our finding that predicted egg potential is higher in the northeastern portion of the range is consistent with long-term population monitoring trends, which indicate that desert tortoise density is decreasing across the range, with the exception of populations in the Northeastern RU (Allison & McLuckie 2018). While our study found substantial spatial variation in expected reproductive output among the 5 RUs for this species, expected climate-driven temporal variation in egg production vary...
was highly correlated across the 5 RUs. This was not unexpected, as while the 5 RUs generally represent the genetic differentiation of tortoise populations across the range is relatively low (Hagerty & Tracy 2010).

Ultimately, addressing range-wide management questions for Agassiz’s desert tortoise and other species will require models that integrate spatiotemporal drivers of variation for all demographic processes to produce realistic expectations of species responses to environmental change (Averill-Murray et al. 2013, Zylstra et al. 2013, Campbell et al. 2015, 2018, Barrows et al. 2016). These efforts would be better if informed by additional data expressing mechanistic processes across landscapes (Kearney & Porter 2004); however, sufficient knowledge of these processes (even in this well studied species) is generally lacking (Guisan & Thuiller 2005) and warrants further consideration (Buckley et al. 2010). For example, a spatial PVA for the Sonoran desert tortoise included well-supported estimates of spatial variation in several key demographic parameters (Campbell et al. 2018). However, despite finding that fecundity was a key determinant of population dynamics in their system, the authors lacked sufficient data to estimate spatial variation in reproductive output (Campbell et al. 2018).

Conservation of threatened and endangered species is best done when a landscape perspective of the threats, habitat condition, and demographics of the local populations are considered in one overarching management framework. Conservation efforts for the desert tortoise are becoming increasingly coordinated (Averill-Murray et al. 2012), and the need for range-wide demographic information is of critical importance for management decisions and conservation planning efforts for this and other species facing unprecedented anthropogenic disturbance and climate change (Turner et al. 1995). The present research provides a model that can estimate a key aspect of the demographic process at a range-wide scale, and can be modified to provide insights into future climates. This information, along with similar efforts targeting other demographic processes, has the potential to aid in management of the desert tortoise and other species.

Data accessibility. All code and data for fitting this model are available at https://github.com/kevinshoemaker/TortoiseEggModel. Desert tortoise egg production data sets collected as part of several studies associated with the manuscript can also be accessed at Puffer et al. (2020). These data are published following US Geological Survey data policies.

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