INTRODUCTION

Increasing global temperatures in recent years have led to rapidly changing environments around the world, often with profound effect on species and their survival (see Thomas et al., 2004; Walther et al., 2002). As many as 15%–37% of species are projected to be extinct by 2050, with many already experiencing difficulties from climate-induced stress, resulting in range constriction and...
alterations in reproductive phenology (e.g., Barnosky et al., 2011; Chen et al., 2011; Janzen et al., 2018; Miller et al., 2018; Stuart et al., 2004; Thomas et al., 2004; Urban, 2015). These changes call for intensive research on diverse species and ecosystems to deepen our understanding of how organisms respond to rising temperatures, changes in precipitation, resource scarcity, and other rapid shifts in environmental variables affecting different levels of biological organization, from physiology to demography to community and ecosystem function.

Climate change has been shown to increase the intensity, duration, and frequency of droughts worldwide (Allen et al., 2010; Kelley et al., 2015). For instance, the Syrian drought from 2007 to 2010 occurred due to human-induced global climate change and led to mass urbanization and civil war (Kelley et al., 2015). Desertification in the African Sahel has also been connected to climate change and that region is projected to become more arid in the future (Benjaminesen & Hiernaux, 2019; Huang et al., 2016). Similarly, the recent and record-breaking drought of 2012–2015 in California, called the most severe drought in that region in the last century, has been widely attributed to climate change (Griffin & Anchukaitis, 2014, p. 9,017; Mann & Gleick, 2015; Swain et al., 2014; Bales et al., 2018; Sumargo & Cayan, 2018; Asner et al., 2016).

A handful of studies to date have examined the impact of drought on mortality and survival in resident species. Prugh et al. (2018) examined 423 species for several years before and during the drought and classified 25% as ecological “losers” in drought conditions and only 4% as “winners.” Previous research on endangered anurans in California has found that droughts lead to rapid species decline and habitat loss (Drost & Fellers, 1996; Fellers & Drost, 1993). Reptiles and amphibians are especially vulnerable to climate change, particularly in areas where extreme temperature and drought are predicted to become more severe and frequent (Corn, 2005; Thomas et al., 2004; Walls et al., 2013; Walther et al., 2002). These taxa are ectothermic and rely on ambient temperatures to regulate physiological functions. In fact, rising global temperatures already threaten the survival of tropical lizards that often function within a relatively low and narrow range of optimal body temperature (Huey et al., 2009). Furthermore, temperature and precipitation help regulate the breeding cycle of reptiles and amphibians (Araújo et al., 2006; Zani & Stein, 2018). The western side-blotched lizard (Uta stansburiana), for example, experiences delays in breeding due to water deprivation during periods of drought (Zani & Stein, 2018). Similarly, a field study found female rattlesnakes experiencing drought did not lay eggs, whereas experimentally hydrated females did (Capehart et al., 2016). Reptiles and amphibians are also relatively immobile over long distances, unlike birds or some mammals, such that environmental change may pose a particularly significant threat.

Higher temperatures and reduced precipitation not only affect single species, but many species within a community’s food web (Carnicer et al., 2011; Miller et al., 2018; Rosenblatt & Schmitz, 2016). Thus, drought may both directly affect a population’s vital rates and also exert indirect effects through drought effects on prey populations. Both direct and indirect effects can place considerable stress on organisms by increasing mortality and decreasing reproduction and overall body condition (see Allen et al., 2010; Capehart et al., 2016; Urban, 2015; Zani & Stein, 2018). Limited resources may lead to physiological trade-offs between various components of individual fitness (Stearns, 1989). These trade-offs may involve various physiological processes, including growth, reproduction, energy storage, and immune function (Adamo & Lovett, 2011; Bonier et al., 2007; Dupoué et al., 2015; Soler et al., 2003; Werner & Anholt, 1993). Understanding how organisms adapt to ecological pressures such as drought is critical for understanding the mechanisms underlying future impacts of climate change at the level of individual species (Martin et al., 2010; Rohr et al., 2013).

The field of eco-immunology provides a useful framework with which to study physiological reactions to environmental change in free-living populations (Downs & Stewart, 2014; Martin et al., 2010; Rohr et al., 2013). Immune function is energetically costly to develop and maintain and can shift in response to resource scarcity, water availability, pathogen exposure, and pollution (see Bowden, 2008; Brusch et al., 2019; Georgiev et al., 2016; Lifjeld et al., 2002; Lochmiller & Deerenberg, 2000; McDade et al., 2016; Vermeulen et al., 2015). In particular, innate or nonspecific immunity—the branch of the immune system which is composed of generalist, preexisting mechanisms of defense—may be a particularly useful indicator of population response to chronic environmental stressors since it is (a) not dependent on specific antigens or current infection and (b) can be compared across periods of time or between different environments (Beutler, 2004; Buehler et al., 2008, 2009; Matson et al., 2005; Riera Romo et al., 2016). Furthermore, there is a growing body of literature that demonstrates an important correlation between innate immunity and environmental situations, such as drought, resource availability, and pollution (Bowden, 2008; Brusch et al., 2019; Georgiev et al., 2016; McDade et al., 2016; Vermeulen et al., 2015). For instance, a study on great-tit nestlings found a correlation between higher resource abundance and higher investment in immune function (Buehler et al., 2009). Innate immunity has also been negatively correlated to aridity in wild populations of birds (Horrocks et al., 2015). This relationship is not clearly understood, however, since both resource scarcity and pathogen exposure may mediate the correlation between aridity and immunity (Brusch et al., 2019; Horrocks et al., 2015). Such studies indicate that while numerous environmental factors may affect immune system development, the relationship between environmental change and immune function is unclear and much remains to be learned, especially in wild populations (Brusch et al., 2019; Festa-Bianchet, 1989; Sparkman & Palacios, 2009).

In this seven-year study, we examined the impact of the recent California drought on innate immunity in two sympatric species of semi-aquatic garter snakes, Thamnophis sirtalis and T. elegans from four populations around Eagle Lake in northern California. Short-term variation in precipitation impacts anuran breeding success and thus directly affects the abundance of amphibians in wetland habitats in this system (Miller et al., 2014); indeed, there is a trend across North America
for amphibian communities to be especially sensitive to precipitation immediately prior to breeding, and for warmer winters to have generally negative impacts in western montane habitats (Miller et al. 2018). Previous research has shown that variation in precipitation also has a negative effect on pregnancy and survival rates in garter snakes, most likely via variation in prey availability (Miller et al. 2011; Miller et al., 2014). There is some evidence that differences in prey availability among habitats may explain variation in innate and acquired immunity for snakes at Eagle Lake (Palacios et al., 2011, 2013; Sparkman & Palacios, 2009), but no study has yet tested for temporal patterns in immunity that track differences in climate. Furthermore, it is unclear whether physiological investment in immune function is shaped by climate of the current year or the previous year. It is well-known that reproduction in some species, particularly snakes, may experience time-lagged effects (Bonnet et al., 1998; Gregory & Skebo, 1998; Lourais et al., 2002), and recent research suggests a time-lag effect of pika abundance in response to drought (Johnston et al., 2019). Thus, it is of interest to determine whether immune function can also experience a time-lagged response to drought. To examine the effect of drought conditions on innate immunity, we measured three indices of innate immunity from blood and plasma samples collected in the field: bactericidal competence (BC) of blood, levels of natural antibodies (NABs), and complement-mediated lysis (CL). These indices, developed for domestic birds by Matson et al. (2005) and Matson et al. (2006) and adapted for garter snakes by Sparkman and Palacios (2009), quantify the strength of innate defensive response in several key components of innate immunity. More specifically, the BC assay examines the ability of blood and plasma samples to mobilize against novel pathogen exposure (in this case Escherichia coli), and the NABs and CL assays assess hemagglutination and hemolysis strength of blood plasma after novel exposure to novel foreign red blood cells (Matson et al., 2005, 2006). Combined together, the results of the NABs, CL, and BC assays quantify immune strength for different aspects of the innate immune response and provide an overall indicator of several key aspects of innate immunity in individuals from our study populations across the years of the California drought. We tested for evidence that these three major indices of innate immunity were affected by variation in annual climate (temperature, snow pack precipitation, and drought indices) from 2012–2018. We predicted that immune function would be lower during the years characterized by drought conditions (2012–2015), when prey and moisture availability were low. Furthermore, we tested for evidence of a time-lagged effect of climate on innate immunity, consistent with a capital-breeding reproductive strategy.

2 | MATERIALS AND METHODS

2.1 | Study system

Two species of garter snakes (Thamnophis elegans and T. sirtalis) were studied from four populations (M1, M3, M4, and L1)—see Matson et al., 2006 and Gangloff et al., 2020 for population details—at a well-known study system around Eagle Lake in Northern California. T. elegans and T. sirtalis are both found in the seasonally flooded, high-elevation meadows (M1, M3, and M4) around Eagle Lake and rely primarily on a diet of anurans, leeches, and occasionally small mammals—with a particular reliance on the Sierran tree frog (Pseudacris sierra) (Kephart, 1982). However, only T. elegans are commonly found along the rocky lakeshore of Eagle Lake (L1 in this study), and these populations are primarily dependent on shoals of small minnows such as tui chub (Gila bicolor) and speckled dace (Rhinichthys osculus) as prey (Kephart, 1982).

During the severest years of the California drought (2012–2015), several populations of garter snakes in our study experienced severe decline. Data from our long-term studies indicate that 7 of the 15 total study populations in the area were extirpated, predominantly those around the lakeshore (A. M. Bronikowski, unpublished data). During the years of the drought, decreased snowfall and spring precipitation caused lake levels at Eagle Lake to decrease rapidly (D. Willis, personal communication, 6 May 2019), exposing large muddy banks which increased the distance between sheltering/basking and foraging areas and thus increasing snake exposure to predators. In addition, prey populations have also fluctuated substantially for meadow populations; for instance, in 2014 amphibian breeding was impeded in several meadows due to severely dry conditions in the normally flooded meadows.

2.2 | Data collection

Snakes were surveyed during the month of June during the summers of 2012 to 2018. All snakes were caught by hand and bled intravenously through the tail. Less than 1% bodyweight of blood (50 to 300 μl) was collected with heparinized syringes and kept on ice until centrifuged to separate the cellular components of blood from plasma, which was stored in liquid nitrogen for later use in immunocompetence assays. Sample sizes for each assay are as follows: T.elegans: BC n = 405, NABs n = 359, CL n = 294; T. sirtalis: BC n = 170, NABs n = 188, CL n = 163. For overall sample sizes by year and population see Table 1.

Body weight, sex, species, and date of capture were recorded for each snake. Snout-to-vent length (SVL) was also collected and can be considered a proxy for snake age since body length is correlated with age in this system (Bronikowski & Arnold, 1999). We also recorded presence/absence of tail trematode infection by looking for swelling, pinkness, or other abnormalities on the tail (Uhrig et al., 2015).

2.3 | Climate data

Climate data for California Climate Division 3 were obtained from the National Oceanic and Atmospheric Administration (NOAA) National Climate Data Center (NCDC) (Vose et al., 2014). For this study, we examined levels of snow telemetry (SNOTEL), a measure of snow pack,
snow water content, and snow precipitation, and Palmer Hydrological Drought Index (PHDI), a long-term measure of drought and aridity that also accounts for soil moisture, ground water, and temperature (Jacobi et al., 2013; Palmer, 1965). SNOTEL data were obtained from the nearest station, located on Mt. Adin, and the average of March and April values for each year were used in the analysis. April estimates for PHDI were used, and spring precipitation was defined as the cumulative precipitation between January and May which, along with snow smelt, is the primary source of spring water availability in the montane meadow habitats. We also obtained minimum, maximum, and mean monthly June temperatures, which correspond to the time of peak snake activity and field data collection. We also tested for the possibility of a one-year time-lag for the effect of temperature and precipitation on immunity. This was done by creating a field that assigned the climate values of the previous year (year – 1) for each individual entry. Annual climate variables are shown in Figure 1.

### 2.4 Immunological assays

A natural antibody (NABs) and complement-mediated lysis (CL) assay, developed by Matson et al., (2005), and modified for garter snakes by Sparkman and Palacios (2009) was performed to assess hemagglutination and hemolysis strength of blood plasma. Serial twofold dilutions of 10 µl plasma were made with Dulbecco’s phosphate-buffered saline 14,190,144 (Thermo Fisher Scientific, Santa Barbara, USA). Sheep red blood cells (SRBC, cat # SBA-050) in Alsevers was centrifuged four times for 10 min at 500 g and rinsed with PBS, and 10 µl 2% SRBC dilution was pipetted into each well (Hemostat Laboratories, Dixon, USA). Plates were incubated at 28°C for 60 min and then scored immediately for lysis and agglutination. On each plate, 1,000 µl PBS and 10 µl SRBC served as a negative control. Scores were given for the highest well that displayed hemagglutination or lysis respectively, and intermediate wells were given half scores.

Bactericidal competence was assessed using a method developed by Matson et al. (2006) and adapted for garter snakes by Sparkman and Palacios (2009). One pellet of dehydrated *Escherichia coli* ATCC 8,739 was dissolved in 40 ml PBS and incubated at 34°C for 30 min (American Type Culture Collection, Manassas, USA). For each *T. elegans* plasma sample, a 1:80 plasma to PBS dilution was made, whereas for *T. sirtalis*, a 1:83 dilution was prepared. A 1:25 dilution the *E. coli* dilution was made with PBS and 10 µl of this solution was added to each plasma dilution. Samples were assayed in

#### TABLE 1

| Year | L1 | M1 | M3 | M4 | TOTAL |
|------|----|----|----|----|-------|
| T. elegans | 2012 | 0  | 23 | 8  | 2     | 10    |
|       | 2013 | 20 | 22 | 14 | 4     | 60    |
|       | 2014 | 10 | 16 | 3  | 6     | 35    |
|       | 2015 | 27 | 23 | 19 | 6     | 75    |
|       | 2016 | 20 | 24 | 20 | 15    | 79    |
|       | 2017 | 20 | 21 | 19 | 8     | 68    |
|       | 2018 | .  | 3  | 12 | 6     | 18    |
| T. sirtalis | 2012 | .  | 3  | 12 | 6     | 18    |
|       | 2013 | .  | 4  | 11 | 13    | 28    |
|       | 2014 | .  | 0  | 20 | 0     | 20    |
|       | 2015 | .  | 6  | 6  | 3     | 15    |
|       | 2016 | .  | 10 | 14 | 6     | 30    |
|       | 2017 | .  | 12 | 18 | 1     | 31    |
|       | 2018 | .  | 19 | 20 | 16    | 55    |

Cells with ‘.’ indicate that no snakes were captured from that population in that year.
duplicate using 50 µl aliquots on each plate and sterilizing a spreading slide with 70% EtOH in between application. Plates were left to dry for 20 min and then incubated overnight at 37°C, after which colonies were counted manually. The intra-assay coefficient of variation for each assay/species is as follows: BC—

\[ T. \text{elegans} \] 13.9, \[ T. \text{sirtalis} \] 14.6; NABs—\[ T. \text{elegans} \] 10.8, \[ T. \text{sirtalis} \] 7.5; CL—\[ T. \text{elegans} \] 11.8, \[ T. \text{sirtalis} \] 11.5.

### 2.5 Statistical analyses

We tested for correlations between our indices of innate immunity and climate using simple correlation analysis. We conducted a mixed model analysis of covariance (ANCOVA) for each immune variable for both \[ T. \text{sirtalis} \] and \[ T. \text{elegans} \]. All four assumptions for ANCOVA were met (but see discussion of multicollinearity below). The dependent variables—bactericidal competence (BC), natural antibodies (NABs), and complement-mediated lysis (CL)—were tested for effects of SVL, body condition (defined as the residuals of the regression of mass on SVL), day of year, PHDI, SNOTEL, mean/max/min temperature and spring precipitation as covariates and reproductive status (male/nongravid female/gravid female), source population (L1/M1/M3/M4), and parasites (presence/absence of tail trematodes) as fixed effects. We also tested for interactions between climate variables and source population. Year was included in each model as a random effect, as climate variables were not independent among years, and we made no a priori predictions for our models regarding which years were drought and which were nondrought. All variables were added one by one in a stepwise fashion and those with \( p \geq .2 \) were dropped from the model, as this indicated that they explained little or no variation in immune function. Note that some of our temperature variables were strongly correlated (see Results). To confirm that the order in which variables were introduced did not affect our outcome and to address any potential concerns with multicollinearity, we performed this process by adding and removing variables in different orders such that every possible combination of variables was ultimately considered. Regardless of the model selection method, we arrived at the same top models. Note that none of the top models contained combinations of explanatory variables that exhibited collinearity. Significance was assessed at an \( \alpha \)-level of \( p = .05 \). Significant differences among groups were assessed post hoc using Tukey’s HSD test. All analyses were conducted with JMP 14.1.0 (SAS Institute, Inc.) and figures were constructed using Prism 8.0.2 (GraphPad Software, Inc.).

### 3 RESULTS

There were significant positive correlations between each of our three indices of innate immunity. As previously documented (Sparkman & Palacios, 2009), NABs were significantly positively correlated to CL titers in both species (\[ T. \text{elegans} \] : \( r = 0.73, p < .0001 \); \[ T. \text{sirtalis} \] : \( r = 0.74, p < .0001 \)). Similarly, though the relationship was less pronounced, both NABs and CL were both also significantly positively correlated to BC in both species (\[ T. \text{elegans} \] NABs: \( r = 0.41, p < .0001 \); \[ T. \text{sirtalis} \] NABs: \( r = 0.45, p < .0001 \); \[ T. \text{elegans} \] CL: \( r = 0.38, p < .0001 \); \[ T. \text{sirtalis} \] CL: \( r = 0.34, p < .0001 \)). Pairwise correlations among all environmental variables indicated that only the three temperatures variables were significantly correlated to each other (\( r = 0.95–0.99, all \ p < .0001 \)). All other pairwise correlations were nonsignificant (\( p > .1 \)).

The final model for each \[ T. \text{elegans} \] index of immunity included source population and year as a random effect (Table 2). Both BC and NABs were significantly correlated to spring precipitation of the former year (Figure 2a, b; Table 2). No climate variables were significantly correlated to CL; however, PHDI was retained in the model as it showed a \( p < .1 \) (Table 2). Differences among populations were largely consistent across immune indices, with L1 and M1 consistently at the high end, M4 intermediate, and M3 consistently at the low end (Figure 3a–c). Additionally, reproductive status (i.e., gravid female, nongravid female, or male) was a significant effect in the final model for NABs and CL, with gravid females having significantly lower NABs and CL scores than both nongravid females and males (Table 2; Figure 4).

As with \[ T. \text{elegans} \], the final model for each \[ T. \text{sirtalis} \] index of immunity included source population and year as a random effect (Table 2). In our first round of analyses, it appeared that the only relationship between annual climate and innate immunity in \[ T. \text{sirtalis} \] occurred with respect to precipitation in the current year and NABs. However, we noted that 2014 (the most severe drought year) was an unusual year in that we only had \[ T. \text{sirtalis} \] represented from a single population, which appeared to be exhibiting unusually high BC values (see Figure 2d). Thus, we also ran our analyses excluding 2014 and found that in this case BC, like NABs, exhibited a significant, positive correlation with precipitation the current year (Table 2, Figure 2d–e). Consistent with \[ T. \text{elegans} \], M1 consistently had the highest levels for each measure of innate immunity, whereas both M4 and M3 levels were low (Figure 3). There was no trend for reduced immunity in gravid \[ T. \text{sirtalis} \] females for any of the three indices.

Other variables considered—SVL, body condition, day of year, SNOTEL, mean/min/max temperature, incidence of parasites, and interactions between climate variables and source population—did not have \( p < .2 \) in any model for any immune index for either species.

### 4 DISCUSSION

Previous work has shown that environmental characteristics, including climate, pathogen intensity, resource abundance, and pollution can impact indices of innate immunity (Bowden, 2008; Brusch et al., 2019; Georgiev et al., 2016; McDade et al., 2016; Vermeulen et al., 2015). Consequently, we predicted that annual climatic variation associated with drought, which has already been shown to decrease body condition and reproductive success in reptiles, would result in a decrease in innate immunity in garter snakes (Allen et al., 2010; Capehart et al., 2016; Urban, 2015; Zani & Stein, 2018).
TABLE 2  Results of mixed model analysis of immune variables with year included as a random effect

| Species           | Dependent variable          | Explanatory variable | df  | F    | p     |
|-------------------|-----------------------------|----------------------|-----|------|-------|
| T. elegans        | Bactericidal competence     | Source Population    | 3,401 | 10.73  | <.0001 |
|                   | Precipitation - 1           |                      | 1,4  | 12.24 | .020  |
| Natural antibodies| Source Population           |                      | 3,342 | 8.09   | <.0001 |
|                   | Precipitation - 1           |                      | 1,2  | 72.48 | .027  |
|                   | Reproductive Status         |                      | 2,222 | 7.46   | .001  |
| Complement-mediated lysis | Source Population |                      | 3,283 | 3.55   | .15   |
|                   | PHDI                        |                      | 1,4  | 4.58  | .093  |
|                   | Reproductive Status         |                      | 2,275 | 9.13   | .0001 |
| T. sirtalis       | Bactericidal competence     | Source Population    | 1,110 | 10.67  | <.0001 |
|                   | Precipitation               |                      | 1,2  | 25.99 | .033  |
| Natural antibodies| Source Population           |                      | 3,161 | 7.72   | .0002 |
|                   | Precipitation               |                      | 1,10 | 6.83  | .027  |
| Complement-mediated lysis | Source Population |                      | 3,157 | 5.38   | .002  |

Consistent with our expectations, we found that, in addition to showing consistent patterns of variation both within and across populations, innate immunity was associated with annual spring precipitation in both species of sympatric garter snake in our study—albeit in a species-specific manner.

We tested for the influence of seven regional climate variables on innate immunity, including annual indices of snowpack; spring precipitation; mean, maximum, and minimum June temperature; and a drought index, PHDI. Only spring precipitation, however, explained significant variation in immune function in our study. Two of three measures of immunity—BC and NABs—were positively correlated with spring precipitation in T. elegans and T. sirtalis (though note that BC was only significant for the latter when unusually high values during 2014 were excluded from the analysis). Interestingly, while BC and NABs were significantly associated with spring precipitation in the current year for T. sirtalis, T. elegans exhibited a time-lagged pattern, such that immunity was correlated to spring precipitation of the previous year (Table 2; Figure 2). Reproductive investment in garter snakes and other species may depend primarily on the resource capital acquired a year prior (Bonnet et al., 1998; Gregory & Skebo, 1998; Lourdais et al., 2002; Sabrina et al., 2009). The finding that both BC and NAB levels in T. elegans were significantly explained by precipitation levels of the previous year indicates that stored capital may also influence immune function in some species (Table 2). To our knowledge, this is the first time a time-lagged effect of climate has been demonstrated in immunological—or indeed, any physiological—study in the wild. This finding complements other recent work in pika (Ochotona princeps) showing the delayed effect of drought on population mortality (Johnston et al., 2019) and calls for investigation into investment in immune function as a possible mechanism for this effect.

We are uncertain whether populations of the two species in our study differ in the extent to which they are capital or income breeders. It is clear that T. elegans demography is very sensitive to the conditions of the previous year such as variation in annual spring precipitation, which may limit access to water or decrease prey availability. This in turn may lead to reductions in probability of individual reproduction, rates of growth, individual condition, and survival the following year (Dinsmore, 2019; Miller et al., 2011; Miller et al., 2018). In contrast, there is some evidence that although T. sirtalis are also sensitive to conditions in the previous year, they are less so. For example, over-winter survival is not correlated with conditions in the previous summer and nearly all adult females reproduce every year regardless of conditions (Dinsmore, 2019). We are currently pursuing additional analyses of climate effects on survival, reproduction, and stress physiology that will be of great interest to combine with our findings regarding the presence of a time-lagged effect of climate on T. elegans but not T. sirtalis.

Our findings prompt the question of how spring precipitation might be affecting innate immunity, if this correlation reflects causation. Climate change may affect organisms either directly, through effects on survival, growth, reproduction and/or dispersal, or indirectly, via impacts on community dynamics, such as competitive and predator-prey interactions, pathogen-host relationships, and habitat characteristics including vegetation and hydrology (reviewed in Blaustein et al., 2010). In our system, we have ample evidence that water availability significantly affects the abundance of anuran prey, such that the driest years may be characterized by little to no prey availability (Miller et al., 2011, Miller et al., 2014). However, beyond evaluations of tail trematode infections, we know little regarding the complex nature of predatory and pathogenic interactions that could be exerting stress in concert with reduced prey in our system (Bowden, 2008; Brusch et al., 2019; Georgiev et al., 2016; McDade et al., 2016; Vermeulen et al., 2015; Zani & Stein, 2018). Regardless of the mechanism, reduced innate immunity could have detrimental consequences for survival and population persistence (Charbonnel et al., 2008; Møller & Saino, 2004). Lower immunity during dry years or years following dry years could be the result of a life-history
trade-off, where investment in other factors such as reproduction or growth is being prioritized over physiological correlates of survival, such as immune function (Sheldon & Verhulst, 1996). Recent work in frogs (*Rana pipiens*), for instance, has shown evidence of trade-offs between growth, survival, and immune function in the context of rapid pond drying (Brannelly et al., 2019). Alternatively, if resources are sufficiently limited during dry years, we might expect reduced investment in all measurable life-history outcomes, such that individuals are not only suffering reduced immune function, but also experiencing reductions in other critical life-history traits such as growth and reproduction.

While previous work has shown variation in immune function associated with variation in resource availability, pathogen exposure, or other mediating ecological variables (Buehler et al., 2009; Horrocks et al., 2015; Pigeon et al., 2013; Sandland & Minchella, 2003; Vermeulen et al., 2015), obtaining high-quality data on spatial and temporal heterogeneity in environmental variables is in fact one of the major challenges in the field of eco-immunology—or

**FIGURE 2** Relationship between indices of innate immunity and spring precipitation from 2012 to 2018. *T. elegans* (a–c) and *T. sirtalis* (d–f). Though the analysis presented in Table 2 is conducted with precipitation as a continuous covariate and year as a random effect, least square means (cm) and standard errors for each year from the final model are depicted here with respect to spring precipitation of the previous year for *T. elegans* and the current year for *T. sirtalis*. Note that *T. sirtalis* data from 2014 were excluded from the final analysis (see Results), but mean 2014 values are indicated with an open marker. Asterisks indicate significant correlations.
“macroimmunology” as it has been recently termed (Becker et al. 2020). We are currently pursuing a fully linked analysis exploring repeated measures of immune function and other physiological and genetic endpoints in relationship to growth, reproduction and survival, which will help illuminate the complex effects of climate and the interplay of variables at each level of biological organization. The question of whether climate influences immune function directly, or via community dynamics, however, will require intensive and strategic field work in the future that quantifies temporal and spatial changes in local climate, hydrology, vegetation, prey availability, additional pathogens, and predator-mediated stressors. We would anticipate that immune function as a whole is in fact a complex function of many or all of these factors, including climate; but which, if any, prove to more strongly and/or directly impact some indices of immune function than others must be ascertained through additional study.

Our study system is a powerful one in which to continue to explore these questions, due to the long-term nature of the data we do have on well-differentiated populations of garter snakes distributed across the landscape—and indeed, source population was a significant effect in the final model for each immune variable for both *T. elegans* and *T. sirtalis* in our study (Table 2). Population differences in immunity may have been due to variations in habitat quality and prey availability between our population sites. Of the three populations from which *T. sirtalis* were sampled, all three immune indices were highest at M1 meadow (Figure 4). Compared with the other two population sites, this meadow is the largest and is populated by both minnows and anurans. It has the lowest elevation, is closest

**FIGURE 3** Differences among populations with respect to NABs, CL, and BC for *T. elegans* (a–c) and *T. sirtalis* (d–f). Least square means and standard errors of the means are shown. Different letters indicate significant differences between groups.
to the lakeshore, and is fed by deep streams, arguably making it the most ideal habitat for the snakes. *T. elegans* were sampled from a lakeshore study site (L1) in addition to the three meadow sites. In general, we found that immune function was highest at the lakeshore site, with M1 meadow a close second (Figure 3). The lakeshore habitat may offer the most consistent food source, since drought levels may not have had as severe of an impact on the abundance of minnows in the lake. In general, these findings are compelling in that they suggest that regardless of the plasticity that each population may show in innate immunity across years, there may also be fixed differences between habitats that have independently resulted in consistent patterns of innate immunity in the two snake species that reside there. Whether this reflects plastic responses to habitat characteristics for both species, or selection resulting in convergent genetic differentiation in immune function remains to be determined.

Neither body size/age nor condition appeared to affect these indices of immune function in this study, which is largely consistent with prior work in this system (Palacios et al. 2011; Palacios et al., 2013; Palacios & Bronikowski, 2017—but see Sparkman & Palacios, 2009 for a relationship between NABS/CL and SVL). However, our results do suggest that gravid *T. elegans* females have lower innate immunity than nongravid females and males, although this pattern was significant only for NAB and CL scores (Table 2, Figure 4), and was not present in *T. sirtalis*. Previous work in this system has shown reduced adaptive immunity for gravid females in this system (Palacios & Bronikowski, 2017). Similar trends have been observed in mammals and birds, since females experience physiological trade-offs during reproduction, and may invest less energy into immunity due to the high energetic costs of reproduction (Festa-Bianchet, 1989; Neggazi et al., 2016; but see Neggazi et al., 2016). Our results are also consistent with other work on reptiles that has found that reproductive female lizards experience severe decline in body condition during drought (Zani & Stein, 2018). However, it is worth noting that even in this study of two closely related species living in sympatry, only one showed lower immune function during reproduction, indicating that the ability to predict reduced immune function in gravid individuals in any system without prior empirical evidence to that effect may be limited.

In this study, we provide evidence that drought conditions may lead to a reduction in immune function in a similar but distinctive manner in multiple populations of two closely related species living in the same habitats and experiencing the same fluctuations in annual climate. In addition, we provide evidence of reliance on capital for immune investment in *T. elegans*, reminiscent of a capital-breeding strategy (Gregory & Skebo, 1998; Lourdais et al., 2002). Together, these findings have important implications for the impact of climate change-induced drought on the physiological adaptation and survival of reptiles and other species. As droughts become more frequent and severe, threatened species may undergo physiological challenges and incur physiological costs that leave them vulnerable to diseases and parasites, leading to mortality and population decline. Our work brings us closer to understanding the complex dynamics of this system in free-ranging species, with compelling ramifications for future research on the mechanisms underlying species-specific responses to climate change worldwide.

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CONFLICT OF INTEREST
The authors declare no conflict of interest.

AUTHOR CONTRIBUTIONS
Lucia L. Combrink: Conceptualization (supporting); Data curation (lead); Formal analysis (equal); Visualization (equal); Writing-original
data (lead); Writing-review & editing (lead). Anne M. Bronikowski: Funding acquisition (equal); Methodology (supporting); Writing-review & editing (supporting). David A. W. Miller: Funding acquisition (equal); Writing-review & editing (supporting). Amanda M. Sparkman: Conceptualization (lead); Data curation (supporting); Formal analysis (lead); Funding acquisition (equal); Investigation (lead); Methodology (lead); Project administration (lead); Software (lead); Supervision (lead); Visualization (equal); Writing-original draft (supporting); Writing-review & editing (supporting).

DATA AVAILABILITY STATEMENT
Data can be accessed from the Dryad Digital Repository https://doi.org/10.5061/dryad.xksn02vcj (Combrink et al., 2020).

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