DO LATITUDE, ELEVATION, TEMPERATURE, AND PRECIPITATION INFLUENCE BODY AND CLUTCH SIZES OF FEMALE COMMON FIVE-LINED SKINKS, *PLESTIODON FASCIATUS* (LINNAEUS, 1758)?

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**ABSTRACT:** Common Five-lined Skinks (*Plestiodon fasciatus*) have an extensive distribution that includes much of eastern North America. We examined 490 female specimens (274 with putative clutch sizes) from throughout the range to see if latitude, elevation, mean annual temperature, and/or mean annual precipitation affected body or clutch sizes. We predicted that larger females would produce larger clutches, latitude and elevation would negatively affect both body and clutch sizes, and that temperature and precipitation would exert a positive effect. Our results did not consistently support those predictions. Body size was positively associated with latitude, negatively associated with temperature, and not associated with elevation or precipitation. Clutch size was not related to female body size, but in most instances was positively associated with temperature and precipitation but negatively associated with elevation and latitude. Effectively *K*-selected in the North and *r*-selected in the South, body and clutch sizes in this species appear to be responding to different selective pressures. We evaluated probable causes for the opposite trends in these two life-history traits.

**Key Words:** Bergmann’s rule; *r*- and *K*-selection; Resource rule; Temperature-size rule.

**INTRODUCTION**

Data regarding the effects of latitude, elevation, temperature, or precipitation on body size and reproductive output of squamate reptiles (lizards and snakes) are equivocal. Bergmann’s rule (Bergmann, 1847) states that homeotherms are larger in colder climates. Cooler conditions often reflect increasing latitude or elevation, implying that homeotherms at higher latitudes or elevations are larger. However, responses of ectotherms to these environmental variables vary. For example, Ashton and Feldman (2003) indicated that turtles appear to follow Bergmann’s rule, whereas squamates show an opposite trend (but see the discussions in Angilletta et al., 2004 and Pincheira-Donoso and Meiri, 2013). Contradicting Ashton and Feldman’s (2003) conclusions, Howes and Lougheed (2007) found that male body size in Common Five-lined Skinks (*Plestiodon fasciatus*) increased significantly with increasing latitude, and Aragón and Fitze (2014), in a study of Spanish snakes (*Psammodromus* spp.), found a significant negative association between temperature and snout-vent length (SVL) and a significant positive relationship between elevation and SVL. Further complicating matters is the temperature-size rule (e.g., Angilletta and Dunham, 2003; Stillwell, 2010), which states that cooler conditions can retard growth and delay maturation. Like Bergmann’s rule, it would predict a negative association between temperature and adult size.

An alternative explanation, the resource rule (McNab, 2010), suggests that primary productivity and precipitation and resultant greater resource availability are the primary determinants of animal body sizes. Again, data for ectotherms are equivocal. For example, Pincheira-Donoso and Meiri (2013) indicated that various studies collectively revealed “no support for patterns” and “no signals of processes.” However, Amarello et al. (2010) indicated that body sizes in Western Diamond-backed Rattlesnakes (*Crotalus atrox*) increased in colder and wetter (i.e., more productive) areas, whereas Oufiero et al. (2011) and Volynchik (2012) showed that the size of lizards in the genus *Sceloporus* and that of the Palestine Viper (*Vipera palaestinae*), respectively, increased in...
and arid areas. Tracy (1999) found no association between body size and food abundance in Common Chuckwallas (Sauromalus ater), and Aragón and Fitze (2014) found a relationship between body size and primary productivity in Spanish Sand Racers (Psammodromus spp.) only after correcting for phylogenetic inertia and short-term seasonal variation. Mesquita et al. (2016) examined 674 lizard populations, representing 297 species from 263 sites from throughout the world, and concluded that climatic factors exerted a significant influence on adult female mass, adult female SVL, female SVL at maturity, clutch size, offspring SVL, and relative clutch mass (RCM). However, all regressions showed very low effect sizes, indicative of extensive unexplained variation in the data, some of which was attributed to sampling error common to ecological studies with large data sets.

Climatic factors and/or resource abundance might not fully explain variations in animal body size, but also could interact with biotic factors, such as competition or predation. For example, in a study of the herpetofauna of southern Florida, Meshaka and Layne (2015) noted that geographic variation in body size in several species was associated with the presence or absence of presumably competing congeners.

Olalla-Tárraga et al. (2006) concluded that “although body size influences the energy balance of thermoregulating ectotherms, inconsistent biogeographical patterns and contrasting associations with energy in lizards and snakes suggest that no single mechanism can explain variation of reptile body size in the northern temperate zone.” Regardless, if environmental factors like latitude, elevation, temperature, or precipitation directly or indirectly affect female body size, we should see a comparable association with clutch size, since larger females produce larger clutches in many reptiles (e.g., Shine, 1988 and references therein). For example, Abell (1999) demonstrated that the Striped Plateau Lizard (Sceloporus virgatus) showed positive associations between body size, clutch size, and precipitation.

We examined 490 female Common Five-lined Skinks, Plestiodon fasciatus (Linnaeus, 1758) from throughout the species’ range to see if latitude, elevation, temperature, or precipitation directly or indirectly affect female body size, we should see a comparable association with clutch size, since larger females produce larger clutches in many reptiles (e.g., Shine, 1988 and references therein). For example, Abell (1999) demonstrated that the Striped Plateau Lizard (Sceloporus virgatus) showed positive associations between body size, clutch size, and precipitation.

These lizards are closely associated with moist wooded areas (Smith, 1946; Hechnar, 1994; Hechnar and M’Clokey, 1998), where they prey on insects and other small invertebrates (e.g., Fitch, 1954; Harding, 1997; Brazeau et al., 2015). Activity varies across the species’ range. In Canadian populations, hibernation may begin as early as late August and extend through late September, with emergence in mid-April to early May (Seburn, 1990; Seburn and Seburn, 1998; Wick, 2004). In Kansas, hibernation typically extends from late September through mid-October to mid- to late-March and early April (Fitch, 1954). Skinks in Georgia generally emerge from hibernation by April (Vitt and Cooper, 1986).

Smith (1946) and Vitt and Cooper (1986) indicated that females reach sexual maturity at two years of age and SVL of 52 mm. Fitch (1954) noted that Kansas females reached maturity at 60 mm SVL, but suggested that sexual maturity varies across the species’ range, with some individuals in southern populations reaching breeding size during their first summer, whereas many lizards in northern regions may not reach that size until their third summer. However, Seburn and Seburn (1998) noted that some individuals in a Canadian population achieved minimum breeding size during their first summer, although those skinks likely did not reproduce until their second summer.

Clutch sizes range from one to 19 eggs (Hechnar and Hechnar, 2019), with numbers reflecting the size, age, and condition of the female (Fitch, 1954; Hechnar and Hechnar, 2005). Smith (1946) and Vitt and Cooper (1986) stated that females have an average gestation period of six to seven weeks. However, Seburn and Seburn (1998) noted that gravid females in a Canadian population retained eggs for 52 days and brooded them for 13 days, whereas Fitch (1954) recorded egg retention of 30–44 days and brooding of 11–32 days in Kansas. Seburn and Seburn (1998) suggested that the differences most likely reflected temperature for a shorter active season in the northern population. Initiation and duration of incubation vary across the species’ range and within populations (Noble and Mason, 1933; Fitch, 1954). Hatching generally occurs by mid-July in Kansas (Fitch, 1954), whereas in Canadian populations, hatching ranges from late July to early August (Seburn and Seburn, 1989; Seburn, 1990). However, Vitt and Cooper (1986) indicated that hatching in Georgia occurs from mid-July through mid-August and Palmer and Braswell (1995) listed hatching dates ranging from early July to early September. Surprisingly, these southeastern records are more similar to that of northern populations than those in Kansas.

Fitch (1954) recorded hatching SVL of 23–27 mm, with rapid growth to ~50 mm SVL prior to hibernation in Kansas. Canadian hatchlings grew less rapidly (0.18–0.26 mm/day vs. ~0.5 mm/day in Kansas), but maximum SVL entering hibernation was 48 mm (Seburn, 1990). Adult growth rates in Canada were 0.02 mm/day in males and 0.08 mm/day in females. Howes and Lougheed (2007) found a significant sex-specific latitudinal cline in sexual size dimorphism (SSD), with male body size increasing at higher latitudes while female body size remained relatively constant.

Based on data presented by Fitch (1954) and Hechnar (2005, 2019), we predicted that larger females would bear larger clutches. Based on the inverse of Bergmann’s rule (e.g., Ashton and Feldman, 2003 — but see Howes and Lougheed, 2007) and the resource rule (but see discussions in Pincheira-Donoso and Mcleroy, 2013 and Mesquita et al., 2016), we predicted that latitude and elevation would negatively affect both body and clutch sizes and that temperature and precipitation would exert a positive effect.

MATERIALS AND METHODS

We examined preserved museum specimens (Appendix 1) from Alabama, Arkansas, Delaware, Georgia, Illinois, Florida, Kansas, Kentucky, Louisiana, Massachusetts, Michigan, Minnesota, Mississippi, Missouri, North Carolina, Ohio, Oklahoma, Ontario (Canada), South Carolina, Ohio, Tennessee, Texas, Wisconsin, and Virginia (Fig. 1). We made incisions through the abdominal wall from the xiphoid process to the pelvic girdle. After determining the sex of each specimen, we eliminated males and considered only females in subsequent analyses. We
counted yolked ovarian (= vitellogenic) follicles (based on yellowish coloration) and oviductal eggs, and measured snout-vent length (SVL) to the nearest 0.1 mm using a digital caliper (Mitutoyo Corp., Aurora, IL). We acquired elevation, temperature, and precipitation data from the local weather stations nearest to the sites where specimens had been collected (https://www.idcide.com/weather/index.htm).

We defined four categories of females for our analyses: Category 1 = all females; Category 2 = all females with SVL ≥ 33.2 mm (the smallest female in our sample with evidence of vitellogenesis) with either yolked ovarian follicles or oviductal eggs and with clutch sizes determined solely by the presence of oviductal eggs (to eliminate the possibility that not all vitellogenic follicles develop into oviductal eggs; e.g., Ramirez-Pinilla et al., 2014). We used StatView 5.0 (SAS Institute Inc., Cary, North Carolina) for all statistical tests. Means are reported ± SE. For all tests, α = 0.05. To assure normality, we log_{10}-transformed all data. Because environmental variables were not independent and we wanted to assess the relative effects of each factor, we ran simple regression analyses for SVL versus clutch size and for both SVL and clutch size versus latitude (°), elevation (m), mean annual temperature (°C), and mean annual precipitation (cm).

**RESULTS**

Means and ranges of body sizes for all four categories of specimens and of clutch sizes for categories 2–4 are in Table 1, which also lists means and ranges of environmental variables (latitude, elevation, mean annual temperature, mean annual precipitation) for each category. All environmental variables were significantly correlated. Mean annual temperature and precipitation were negatively associated with both latitude and elevation, mean annual temperature and precipitation were positively associated, and latitude and elevation also were positively associated (all \( P < 0.0001 \)).

Associations of body size (SVL) with clutch size and of SVL and clutch size with environmental variables (latitude, elevation, mean annual temperature, mean annual precipitation) for each category.

**Table 1.** Means (± 1 SE) and ranges (in parentheses) for SVL (mm) and clutch size in four categories of female Common Five-lined Skinks (*Plestiodon fasciatus*) and the environmental variables (latitude, elevation, mean annual temperature, mean annual precipitation) applicable to each category. Category 1 = all females with SVL ≥ 26.8 mm; Category 2 = all females with SVL ≥ 26.8 mm and with yolked ovarian follicles or oviductal eggs; Category 3 = females with SVL > 52 mm and with either yolked ovarian follicles or oviductal eggs; Category 4 = females with SVL > 52 mm and only oviductal eggs. Clutch-size data were not analyzed and are omitted for Category 1. Clutch sizes based on either yolked ovarian follicles or oviductal eggs are marked with an asterisk (*; see text); those based solely on the presence of oviductal eggs are marked with a double asterisk (**).

| Environmental Variables | Category 1 \( (n = 489) \) | Category 2 \( (n = 274) \) | Category 3 \( (n = 212) \) | Category 4 \( (n = 156) \) |
|-------------------------|----------------------------|--------------------------|--------------------------|--------------------------|
|                         | 56.9 ± 0.4 (26.8–85.2)     | 58.9 ± 0.5 (32.2–75.0)   | 62.8 ± 0.4 (52.1–75.0)   | 62.5 ± 0.4 (53.1–75.0)   |
|                         |                            | 12.4 ± 0.3 (5–26)*       | 12.2 ± 0.3 (5–26)*       | 10.0 ± 0.2 (5–15)**      |
| Latitude (°N)           | 34.6 ± 0.2 (26.7–45.6)     | 33.9 ± 0.3 (29.0–45.3)   | 34.4 ± 0.3 (29.6–45.3)   | 35.1 ± 0.4 (29.8–45.3)   |
| Elevation (m)           | 130.0 ± 5.1 (0.9–715.0)    | 106.5 ± 6.7 (0.9–684.0)  | 109.2 ± 7.1 (0.9–591.6)  | 120.7 ± 8.3 (0.9–591.6)  |
| Mean Annual Temperature (°C) | 16.1 ± 0.2 (4.2–24.1) | 16.7 ± 0.3 (5.5–21.1) | 16.2 ± 0.3 (5.5–21.1) | 15.5 ± 0.4 (5.5–21.1) |
| Mean Annual Precipitation (cm) | 125.5 ± 1.1 (70.8–175.1) | 127.8 ± 1.6 (74.2–167.6) | 126.3 ± 1.8 (74.2–167.6) | 123.3 ± 2.0 (74.2–167.6) |

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al precipitation) are in Table 2. Body and clutch sizes were not significantly related in any category (Fig. 2). Body sizes (Fig. 3) were positively associated with latitude, negatively associated with mean annual temperature, and not significantly associated with elevation in all categories; body sizes were negatively associated with mean annual precipitation only for Category 4, although relationships approached significance for Categories 1 and 3. Clutch sizes (Fig. 4) were negatively associated with latitude and elevation but positively associated with mean annual temperature for Categories 2 and 3 but not 4. The association with mean annual precipitation was significant for Category 2, approached significance for Category 3, but was not significant for Category 4.

### DISCUSSION

**Environmental variables.**—Relationships between the environmental variables examined in this study were largely as expected. The negative relationship between temperature and both latitude and elevation has been well documented (e.g., Birdsall et al., 2017). Similarly, Zhao and Khalil (1993) demonstrated that precipitation and temperature are frequently and positively associated (although warm summers in temperate zones are typically dryer), and higher latitudes or elevations often are inversely associated with precipitation. The positive relationship between latitude and elevation could merely reflect sampling bias, as the lowest elevations in our sample were largely in the southern Coastal Plain and northern sampling sites were generally far from coastal areas.

### TABLE 2.

Results of simple regression analyses of log$_{10}$-transformed SVL (mm) versus clutch size and both SVL (mm) and clutch size versus latitude ($\degree$), elevation (m), mean annual temperature ($\degree$C), and mean annual precipitation (cm) in female Common Five-lined Skinks (*Plestiodon fasciatus*). Significant relationships are indicated with an asterisk (*). Category 1 = all females with SVL $\geq$ 26.8 mm; Category 2 = all females with SVL $\geq$ 26.8 mm and with yolked ovarian follicles or oviductal eggs; Category 3 = females with SVL $> 52$ mm and with either yolked ovarian follicles or oviductal eggs; Category 4 = females with SVL $> 52$ mm and only oviductal eggs.

| Variables                          | Categories 1 & 2 ($n = 489$, $n = 274$ for clutch size) | Category 3 ($n = 212$) | Category 4 ($n = 156$) |
|------------------------------------|--------------------------------------------------------|-------------------------|-------------------------|
|                                    | $r^2$         | P          | $r^2$         | P          | $r^2$         | P          |
| Log(SVL)/log(Clutch size)          | 0.005         | 0.2387     | 0.007         | 0.2299     | 0.001         | 0.7185     |
| Log(SVL)/log(Latitude)             | 0.041         | <0.0001*   | 0.044         | 0.0020*    | 0.101         | <0.0001*   |
| Log(SVL)/log(Elevation)            | 0.0001        | 0.8119     | 0.001         | 0.6526     | 0.018         | 0.0967     |
| Log(SVL)/log(Mean annual temperature) | 0.043          | <0.0001*   | 0.033         | 0.0076*    | 0.074         | 0.0006*    |
| Log(SVL)/log(Mean annual precipitation) | 0.007          | 0.0593     | 0.015         | 0.0726     | 0.066         | 0.0013*    |
| Log(Clutch size)/log(Latitude)     | 0.073         | <0.0001*   | 0.065         | 0.0002*    | 0.009         | 0.2268     |
| Log(Clutch size)/log(Elevation)    | 0.053         | <0.0001*   | 0.065         | 0.0002*    | 0.009         | 0.2328     |
| Log(Clutch size)/log(Mean annual temperature) | 0.055       | <0.0001*   | 0.048         | <0.0013*   | 0.007         | 0.3130     |
| Log(Clutch size)/log(Mean annual precipitation) | 0.016         | 0.0383*    | 0.018         | 0.0532     | 0.00006       | 0.9656     |

**Fig. 2.** Simple regression graphs of log(clutch size) versus log(SVL) in female Common Five-lined Skinks (*Plestiodon fasciatus*). Numbers represent categories: Category 2 = all females with with either yolked ovarian follicles or oviductal eggs; Category 3 = females with either yolked ovarian follicles or oviductal eggs and SVL $> 52$ mm; Category 4 = females with SVL $> 52$ mm and with clutch sizes determined solely by the presence of oviductal eggs. No relationships were significant (Table 2).
Trends in body size.—Unlike Lowes and Lougheed (2007), who found that male body size increased with latitude (following Bergmann’s rule, but contrary to the conclusions of Ashton and Feldman, 2003) but that female body size was relatively constant throughout the species’ range, we found considerable variation in female body size. The negative association between SVL and temperature and the positive association of SVL and latitude lends support for the contention that female skinks either follow Bergmann’s rule, contrary to Ashton and Feldman’s (2003) analysis and our prediction, or that the temperature-size rule might apply. The equivocal associations between SVL and mean annual precipitation could suggest that the expected effects of precipitation were at least to some degree offset by factors that could include seasonal variation in rainfall (e.g., Aragón and Fitze, 2014) and its relationship to growth or perhaps some level of productivity despite cooler temperatures and less rainfall in northern parts of the species’ range. Alternatively, effective microhabitat selection by northern females (e.g., Hecnar, 1994; Hecnar and M’Closkey, 1998; see also below) could compensate for presumably inhospitable climatic conditions. We suggest that the relatively limited elevational range represented in our samples limited our ability to detect any associations with elevation.

Tentative support for the temperature-size rule is evident in minimum body size or age at sexual maturity. The smallest female with yolked ovarian follicles (from Texas) had a SVL of 33.2 mm, which was considerably
gin vitellogenesis during their first full year of life. That said, the smallest female in our sample (from Arkansas) with oviductal eggs had a SVL of 53.1 mm, and all of the smaller females with yolked follicles were from the southern half of the species’ range, where maturation apparently is achieved at smaller body sizes.

Our seemingly counterintuitive findings of larger mean adult female body size in northern areas, where productivity appears to be lower than in the south, and smaller minimum body sizes at sexual maturity of females in the south could be explained by the effects of interspecific competition or greater predation pressures. In their study of the southern Florida herpetofauna, Meshaka

Fig. 4. Simple regression graphs of log(clutch size) versus the log10 of four environmental variables in female Common Five-lined Skinks (*Plestiodon fasciatus*). Numbers represent categories: Category 2 = all females with with either yolked ovarian follicles or oviductal eggs; Category 3 = females with either yolked ovarian follicles or oviductal eggs and SVL > 52 mm; Category 4 = females with SVL > 52 mm and with clutch sizes determined solely by the presence of oviductal eggs. Significant relationships are marked with an asterisk (*). See also Table 2.

smaller than expected. Although we cannot rule out the possibility that follicles do not fully develop in younger females or that the yellowish coloration of follicles in some females was an artifact of preservation, the color of follicles in the smaller lizards was consistent with that of obvious vitellogenic follicles in larger specimens. Fitch (1954) indicated that females reach maturity in their second spring, and Vitt and Cooper (1986) stated that sexual maturity was achieved at SVL = 52 mm in the southeastern United States. However, based on the sizes of the smallest individuals with yolked ovarian follicles and growth data presented by Fitch (1954) and Seburn (1990), our data suggest that at least some females begin vitellogenesis during their first full year of life. That said, the smallest female in our sample (from Arkansas) with oviductal eggs had a SVL of 53.1 mm, and all of the smaller females with yolked follicles were from the southern half of the species’ range, where maturation apparently is achieved at smaller body sizes.

Our seemingly counterintuitive findings of larger mean adult female body size in northern areas, where productivity appears to be lower than in the south, and smaller minimum body sizes at sexual maturity of females in the south could be explained by the effects of interspecific competition or greater predation pressures. In their study of the southern Florida herpetofauna, Meshaka
and Layne (2015), contrary to expectations, encountered larger individuals of several species farther north on the Florida Peninsula. The common element in these instances was a reduction in the number of putative interspecific competitors. If such interspecific competition has a negative effect on body size in Common Five-lined Skinks, that effect would be least evident in the most northerly populations, which have no sympatric congeners (e.g., Powell et al., 2016), and most intense in the southern portions of the species’ range, where a number of sympatric congeners (P. anthracinus, P. inexpectatus, P. laticeps) could be competing for many of the same essential resources. That also could explain a rapid attainment of sexual maturity followed by stunted adult body sizes in the south. Conversely, individuals in the north can grow larger than would otherwise be expected because of a release from competition, even if the rate of growth is slower than in southern populations because of colder temperatures and shorter growing seasons. Similarly, northern populations are likely subjected to lower predation pressure from a less diverse suite of predators (e.g., Sidorovich, 2011). Although we cannot rule out differences in genetic factors (Howes et al., 2006), a reduction in competition, predation, or both could result in enhanced survival, longer periods for growth (even at slower rates), and larger average adult body size.

Also, although our data suggest that environmental factors can influence body size, albeit in a direction opposite to our predictions and those of Ashton and Feldman (2003), we suggest that any conclusions could be compromised by the effective exploitation of microhabitats by northern females (e.g., Hecnar, 1994; Hecnar and M’Closkey, 1998), especially since our environmental data did not reflect the exact localities, much less the microhabitats, where specimens had been collected. Common Five-lined Skinks are closely associated with moist wooded areas throughout their range (e.g., Fitch, 1954; Hecnar, 1994; Environment Canada, 2014). By restricting activity to microhabitats (e.g., rotting logs) that provide ample food and minimize the potential negative effects of cooler, drier macroclimatic conditions, skinks might avoid the effects on growth normally associated with such environmental factors.

Finally, hatching size, which we did not assess, can dramatically affect adult sizes. Vitt and Cooper (1986) found significant differences in hatching sizes among clutches in the southeastern United States. Such variability within a geographic subset of the species’ total distribution suggests that considerable variation in body sizes could exist throughout the entire range of the species. If true, that variability could obscure or confound trends attributable to climatic differences.

Trends in clutch size.—That larger females did not produce larger clutches contradicts our prediction and the fact that most reptiles (e.g., Shine, 1988) follow that pattern. Hecnar and Hecnar (2019) documented a significant increase in clutch size with increasing female body size. Brown (1992) examined 16 P. fasciatus clutches from North Carolina and determined that the largest females tended to produce the most eggs. Vitt and Cooper (1986) documented a similar trend in the southeastern U.S., although the pattern was less evident than in many other species of lizards (i.e., Fitch, 1970). We initially thought that the most logical explanation for why our data did not follow that trend was that many of our putative clutch sizes were based on yolked ovarian follicles, some in very small females, and that vitellogenic follicles do not necessarily translate into ovq (e.g., Ramírez-Pinilla et al., 2014). Fitch (1970) listed a maximum clutch size of 18 and Hecnar and Hecnar (2019) recorded a clutch of 19, but we found 13 individuals with more than 18 yolked ovarian follicles, although our largest observed clutch size based solely on oviducal eggs was 15. Nevertheless, the relationship between clutch size and female SVL remained insignificant when we excluded females with SVL < 52 mm or when based solely on oviducal egg counts (Table 2). Trends observed by Brown (1992) and Vitt and Cooper (1986) in the southeastern U.S. might have been obscured by other climatic or life-history variables in other parts of the species’ range (see also the discussion in Mesquita et al., 2016) or unknown genetic variables (e.g., Howes et al., 2006) might apply.

In a review of the literature, Hecnar and Hecnar (2019) found no significant relationship between either latitude or longitude and mean clutch size and they suggested that clutch size appeared to be a “generally stable trait across the species geographic range.” In our study, the relationship between clutch size and the abiotic environmental factors generated two strikingly different results. Relationships in line with our predictions were evident for categories 2 and 3, although contrary to findings of Pincheira-Donoso and Meiri (2013), whereas regressions for category 4, derived solely from oviducal egg counts in females with SVLs > 52 mm and most like those reported in Hecnar and Hecnar (2019), with data derived from actual clutches, were not significant. These data suggest that females with clutches that are not fully developed are strongly affected by the productivity associated with warmer, wetter southern areas (i.e., the resource rule). However, final investment in a clutch, as measured by the number of oviducal eggs, appears to be controlled by some other factor or suite of factors, which could include any number of life-history variables (e.g., Mesquita et al., 2016) or something as mundane as sampling bias (i.e., fewer records from northern and eastern portions of the range where skinks are less abundant). We also cannot rule out egg size (and consequently hatching size), which could be affected by both female body size and productivity. If true, abiotic factors would affect potential clutch size, whereas actual clutch size could be determined by an interplay of female body size and energy.

r- and K-selection.—Body sizes of female Common Five-lined Skinks tend to increase with latitude and, although Hecnar and Hecnar (2019) showed a positive albeit insignificant relationship between latitude and clutch sizes, our data, which included samples from many more southerly locations, showed a trend of decreasing clutch sizes with increasing latitude. This could reflect selection for two different regionally dependent life-history strategies. A continuum in life-history strategies variously embodies r- and K-selected strategies (e.g., Pianka, 1970). In P. fasciatus, two different suites of traits apparently depend on latitude, the reasons for which may be found in an interaction of regionally specific biotic and abiotic factors. We suggest that a combination of competitive release, reduced predation pressure, and temperature-size associations result in northern populations adhering to a predominantly K-selected strategy of delayed sexual maturity and larger adult body size. Smaller clutch sizes and presumably larger egg sizes would be associated with high juvenile survivorship if these large females live longer than their southern counterparts. Conversely, the temperature-size rule would select for early maturity at smaller body sizes in the south. We suspect that, despite
greater resource abundance, the presence of congeneric competitors and greater predation pressure limits growth and lowers adult survivorship. Such selective pressures would favor r-selection in southern populations and would also explain the apparently intense pressure on large clutches.

Although our effect sizes are low, we suggest that this is the result of the "noise" generated by a relatively large dataset with considerable and inevitable variation among individuals and populations. Nevertheless, the evident geographic patterns associated with female body and clutch sizes in Common Five-lined Skinks are consistent with predictions of r- and K-selection. The selective pressures behind the geographic trends in body size might tentatively be ascribed to a combination of the presence or absence of competitors, greater or lesser predation pressures, and effects of the temperature-size rule. This tentative explanation does not contradict the resource rule with respect to greater productivity in warmer areas but does address the effects resulting from the variable availability of resources on growth and the concomitant diversion of energy to clutch size in skinks that are sympatric with many competitors and predators versus those that are not.

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