ORIGINAL PAPER



Reproductive trade-offs in the colorado checkered whiptail lizard (*Aspidoscelis neotesselatus*): an examination of the relationship between clutch and follicle size

Hannah E. Caracalas¹ · S. S. French^{3,4} · S. B. Hudson^{3,4} · B. M. Kluever⁵ · A. C. Webb^{3,4} · D. Eifler⁶ · A. J. Lehmicke⁷ · L. M. Aubry^{1,2}

Received: 15 December 2020 / Accepted: 6 August 2021 © The Author(s), under exclusive licence to Springer Nature Switzerland AG 2021

Abstract

Life history theory predicts that there should be an inverse relationship between offspring size and number, because individuals cannot simultaneously maximize both when resources are limited. Although extensively studied in avian species, the occurrence and determinants of reproductive tradeoffs in oviparous reptiles are far less understood, particularly in parthenogenetic species. We studied this trade-off in the Colorado Checkered Whiptail, Aspidoscelis neotesselatus, a female-only parthenogenetic lizard. Using data previously collected in 2018 and 2019, we tested for clutch and egg size trade-offs and determined whether this relationship could be influenced by female size and aspects of physiological condition. Physiological condition included energy-mobilizing hormone (i.e. corticosterone 'CORT'), oxidative stress (i.e. reactive oxygen metabolites 'ROMs'), and innate immune function (bacterial killing ability 'BKA'). We found the effect of clutch size on follicle size was significant, but not linear. Specifically, follicle size was on average larger in females with clutches of two follicles when compared to clutches of one follicle, but smaller in females with clutches of three when compared to clutches of two. In addition, females that were larger produced larger follicles regardless of clutch size. Neither CORT nor BKA affected the relationship between follicle size and clutch size. However, ROMs did explain variability in this relationship: oxidative stress was more elevated in females that produced larger clutches and larger follicles. We conclude that clutch size and body size are key life history traits that shape follicle size, and that investments into larger clutches and follicle size come at the cost of oxidative damage.

Keywords Clutch size \cdot Glucocorticoids \cdot Immunity \cdot Trade-off \cdot Lizard \cdot Oxidative damage \cdot Reproduction \cdot Whiptail

L. M. Aubry lise.aubry@colostate.edu

Extended author information available on the last page of the article

Introduction

The partitioning of finite resources across a female's reproductive life influences her investment in current and future reproductive success and her survival probability (Williams 1966). A trade-off between the total resources invested per clutch, and the number of clutches produced over reproductive life in particular, has led to the evolution of semelparous versus iteroparous life history strategies (Hughes 2017). Life-history theory further predicts there should be an inverse relationship between offspring size and offspring number because resources are limiting and females cannot simultaneously maximize both (Stearns 1992). These constraints will generally result in an inverse relationship between egg size and number per clutch (Parker and Begon 1986; Dziminski et al 2009), with selection favoring either the production of a few large eggs or many small eggs. Lack's hypothesis (1947) regarding clutch size was the first empirical test of this idea in a bird, the great tit (*Parus major*), given a female's finite egg-carrying capacity and reproductive reserve.

The offspring size vs. number trade-off (Lessels 1991; Stearns 1992) relies on the assumption that larger offspring tend to have higher adult reproductive success and greater chances of survival into adulthood (Ferguson and Fox 1984; Sinervo 1990; Roff 2002). If resources are unlimited, a female should invest more resources per offspring to enhance her own reproductive success (Lack 1947; Smith and Fretwell 1974; Grafen 1988), but if resources are limited, she can only increase the size of individual offspring at a cost to the number of offspring produced. Hence the trade-off between offspring size and number is inevitable in natura (Stearns 1992). Although this trade-off has been extensively studied in avian species (e.g. Rohwer 1988; Blackburn 1991; Williams 2001), we know little about the factors that may govern this relationship in oviparous lizards, apart from an acknowledgement that this trade-off is generally constrained by maternal physiology (Ljungström et al 2016). Clutch sizes vary from one to forty or more among different species of oviparous lizards, capturing a spectrum of r to K-selected species (Pianka 1970). Some species reproduce only once every second or third year, whereas others reproduce yearly and may even lay two or more clutches each year. Substantial spatial and temporal variation in clutch size also exists between species (Stearns 1992).

Aubry et al (2020) studied the trade-off between clutch size and follicle size and its determinants in the Colorado Checkered Whiptail, *Aspidoscelis neotesselatus*, a "female-only" parthenogenetic lizard. Aubry et al (2020) suggest several advantages to studying reproductive trade-offs in this species: egg cell development happens without fertilization by males and offspring are genetically identical to their mother. One can also exclude potential epigenetic effects from males, and assume that the entire investment strategy is driven by the female. In addition, energetically costly reproductive behaviors such as court-ship (Sullivan and Kwiatkowski 2007) and parental care (Case 1978) are minimal in this species. Finally, one can easily and non-invasively use ultrasonography to quantify variability in follicular and egg development (Krawchuk and Brooks 1998; Gilman and Wolf 2007).

Several factors could affect the potential trade-off between offspring size and offspring number in *A. neotesselatus*: chief among them are a female's size and her physiological state:

(i) Female body size could affect the trade-off between offspring size and offspring number as larger females may have access to additional energetic resources, which they could then invest into follicle development. In both viviparous and oviparous populations of common lizard, larger females were able to produce larger clutches (Qualls and Shine 1998; Horváthová et al. 2013; Recknagel and Elmer 2019), and the selective benefit of larger body size led to an evolutionary increase in female size (Recknagel and Elmer 2019). In Eublepharid geckos (*Eublepharidae*), female body mass and egg mass are related isometrically, with larger females producing larger eggs (Kratochvíl and Frynta 2006). This finding contrasts with most other ectotherms, which show a negative allometry between female size and egg size rather than an isometric relationship (Bauwens and Díaz-Uriarte 1997). In *A. neotesselatus*, which produces small clutches of proportionally large eggs, it is more likely that any additional resources that are put toward egg development would increase the size of the follicle, and not increase the size of the clutch, as suggested by the fractional egg size hypothesis (Ford and Seigel 2010). For *A. neotesselatus*, investing in an additional follicle would require a much larger investment of energy according to this hypothesis, which states that changing clutch size requires a large investment of energy per additional egg (Ford and Seigel 2010).

(ii) Because reproduction is a costly activity, it is expected to decrease other key physiological functions such as immunity; likewise, initiating an immune response is costly and may use up energy that could affect reproductive success (e.g. Adamo et al 2001; French et al 2007b; Martin et al 2008; Knowles et al 2009; Cox et al 2010). Reproduction can involve energy-mobilizing hormones such as corticosterone (CORT), and can lead to the accumulation of reactive oxygen metabolites (ROMs) (Metcalfe and Alonso-Alvarez 2010). Indeed, during reproduction, levels of ROMs may exceed the capacity of antioxidant defense and repair mechanisms (i.e. oxidative stress), leading to oxidative damage of biomolecules (Metcalfe and Alonso-Alvarez 2010). Like any other organism, A. neotesselatus must allocate resources towards competing functions such as reproduction, growth, and maintenance (Stearns 1992). Immunity in particular is expected to limit the resources available for reproduction since immune function is an important aspect of self-maintenance (Sheldon and Verhulst 1996; Lochmiller and Deerenberg 2000; in birds: Hamilton and Zuk 1982; Ilmonen et al 2000; Ardia et al 2003; in mammals: Derting and Compton 2003; Graham et al 2010; and in lizards: Dunlap and Schall 1995; French et al 2007a, b; Cox et al 2010). In addition, because the stress response and corresponding glucocorticoid release can sometimes diminish immune function critical to self-maintenance (Lucas and French 2012), we may observe an increased stress response in females that produce larger clutches, larger follicles, or both. Increased physiological stress is also related to the overproduction of ROMs (Lucas and French 2012), which could be elevated in females that invest in larger clutches and (or) larger follicles. However, these relationships are often context-dependent (Costantini 2016; Smith and French 2017), hence the importance of drawing comparisons between (sub) populations that inhabit different sampling areas.

In this study, we tested whether a clutch-follicle size trade-off exists in a population of *A. neotesselatus* that has been studied at the Fort Carson Army installation. We further investigated whether the potential trade-off changes with (i) body size (snout-vent-length, SVL) or (ii) physiological parameters including energy-mobilizing hormones (CORT), innate immunity (measured with a bacterial killing assay, BKA), and oxidative stress (ROMs). We predicted larger females would not suffer as great a clutch-follicle size trade-off relative to smaller females; we further expected females investing in larger

clutches and (or) larger follicles would show elevated CORT concentrations, lower BKA, and increased oxidative stress (ROMs).

Methods

Field collection

Data were collected from *A. neotesselatus* lizards (Fig. 1) living on the Fort Carson (FC) U.S. Army Installation located in unincorporated El Paso County, Colorado, near the city of Colorado Springs, USA. Fort Carson is 55,000 hectares and also extends into Pueblo and Fremont Counties. *A. neotesselatus* was sampled and surveyed in the northern edge of its range in FC. Lizards were surveyed and sampled across two reproductive seasons in 2018 and 2019. The species is known to occur in several different training areas (TAs) on Fort Carson (Fig. 1), but access to TAs was limited because of military activity. Most surveying focused on TAs 48 and 55, with some data obtained from TA45. The area surveyed across these TAs represented 0.99 ha for TA45 (38.465° N, 104.933° W), 6.12 ha for TA48 (38.467° N, 104.733° W), and 4.85 ha for TA55 (38.398° N, 104.872° W). All TAs were subject to similarly low levels of disturbance from military training activities (i.e. on-foot navigation and orientation).

The habitat for *A. neotesselatus* included Piñon Pine (*Pinus edulis*), Ponderosa Pine (*Pinus ponderosa*), and mixed oak trees (*Quercus* sp.), as well as the cactus Tree Cholla (*Cylindropuntia imbricata*) and the grass Blue Grama (*Bouteloua gracilis*) which mostly dominated TA45. In TA48, the majority of *A. neotesselatus* were found in a dry creek bed and along its banks. The shrubs found in TA48 included Shadscale (*Atriplex confertifolia*), Four-wing Saltbush (*Atriplex canescens*), James' Seaheath (*Frankenia jamesii*) and Rubber Rabbitbrush (*Chrysothamnus nauseosus*). The secondary vegetation type was One-seeded Juniper (*Juniperus monosperma*) and mixed grassland, located around the edges of the sample area. TA55 was similar to TA48, but with more Juniper trees rather than shrubs dominating the landscape.

Transect surveys were conducted during the morning and early afternoon hours (0900–1300) to seek out and capture individuals. Once caught using a snaring method, lizards were kept in mesh fabric bags in the shade to later be processed. Processing included measuring SVL (mm), unique toe clipping (for new captures), marking a unique color code with non-toxic paint pens, and performing ultrasounds on adults. Ultrasounds were performed in order to assess gravidity, clutch size, and follicular/egg size of adult females. Note that we use the term follicle throughout the manuscript as an inclusive term that encompasses anything from the smallest detectable follicle that could be measured, to the largest clearly defined egg, which ranged from 0.17 to 1.05 cm. Follicle size measurement, i.e. the length along the longest axis, was done using a Sonosite Turbo ultrasound unit with an external linear probe (Sonosite Turbo ultrasound, FUJIFILM SonoSite Inc., Bothell, Washington, USA), and each follicle in a clutch was our sampling unit. Once processed, lizards were released in the same location they were captured.

Blood sample collection

Each captured adult lizard had blood samples taken within the first 5 min. of being pursued for capture, a validated window of time to measure baseline physiological activity in lizards

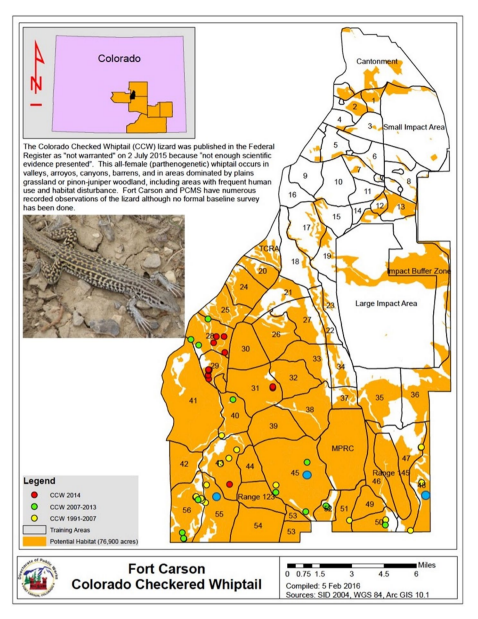


Fig. 1 Surveys of *A. neotesselata* conducted in 1991–2007 (yellow dots), 2007–2013 (green dots), and 2014 (red dots) at Fort Carson, CO. Surveys used in this study (2018–2019) are represented by blue dots, with intensive sampling focused on TA45, TA48, and TA55

(Romero and Reed 2005). If capture attempts exceeded 5 min in duration, blood samples were not taken from that individual lizard to limit the effect of a stress response elicited by the actual pursuit of the animal. Blood samples were taken from the retro-orbital sinus area using heparinized capillary tubes immediately after capture (Maclean et al. 1973). These samples were analyzed to measure baseline levels of physiological activity (Romero and

Reed 2005; Sheriff et al 2011). After collection, blood samples were immediately stored on ice until being centrifuged at 6000 RPM for 10 min to isolate plasma, which was then separated, frozen, and stored at -20 °C for assays at a later time. Samples were only taken during a standardized time frame in order to avoid daily circadian differences in CORT released by stimuli (Maclean et al. 1973; Dallman et al 1987; Jones and Gillham 1988; Romero and Wingfield 2001). This sampling period was 0700–1200 h, and was found to have no relationship between circulating CORT and time of day (Hudson et al 2020).

Blood plasma analysis

Corticosterone In order to determine CORT concentrations, enzyme-linked immunosorbent assay (ELISA) kits were validated and used with blood plasma (10 uL), following Hudson et al (2020). This colorimetric ELISA is based on competitive binding between sheep polyclonal antibodies and plasma hormone that takes place on a donkey anti-sheep immunoglobulin (IgG) microtiter plate. We assayed each sample in duplicate across seven 96-well plates according to manufacturer guidelines (Product # ADI-901–097; Enzo Life Sciences, Farmingdale, NY). Assay sensitivity was 27 pg/ml with a mean intra-assay coefficient of variation (CV) of 1.46% and an inter-assay CV of 12.19%.

Bacterial killing assay To assess the inherent immune function for this species, BKA was measured using a validated volume of blood plasma (11 uL) following Hudson et al (2020). Using the protocol outlined in French and Neuman-Lee (2012), a 1:2 plasma dilution was combined with CO_2 -independent media (plus 4 nM l-glutamine), 10⁴ colony producing units of *Escherichia coli* (EPowerTM Microorganisms #483–581-1, ATCC 8739, MicroBioLogics, St. Cloud, MN, USA), and agar broth on a 96-well microplate. Included were both negative (media and no plasma or bacteria) and positive (media and bacteria with no plasma) controls to account for potential growth and to ensure that there was no contamination. The plate was incubated for 12 h at 37 °C and absorbance per well was measured with a microplate reader at 300 nm (xMark; BioRad Benchmark, Hercules, CA, USA). Bactericidal ability was then calculated as 1 – (absorbance of sample/absorbance of positive controls)×100. We ran 7 microplates in total with a mean intra-assay CV of 2.82% and an inter-assay CV of 7.88%.

Reactive oxygen metabolites Oxidative status was measured using reactive oxygen metabolites (ROMs). Circulating ROMs were quantified using a d-ROMs test kit (MC435, Diacron International, Italy) which discerns levels of hydroperoxides that oxidize an alkyl-substituted aromatic amine (A-NH₂). Plasma was diluted in an acidic buffered solution (5 μ 1: 100 μ 1) following "endpoint" mode manufacturer instructions that were adjusted for a 96-well microplate (Hudson et al 2020), and incubated for 90 min at 37°C. Values were calculated as absorbance change relative to the standard. The mean intra-assay CV over 8 plates was 3.26% and the inter-assay CV was 3.33%.

Statistical analyses

All analyses and visual representation of data were performed using default packages in RStudio (version 1.2.5001, R Core Team 2019) and additional packages: 'dplyr' (Wickham 2020) and 'ggplot2' (Wickham 2016). Because we had to account for "nested" measurements of follicle size within clutches, we included a random effect of clutch identity in our models (Bolker et al 2009) and used linear mixed models with a gaussian distribution and an identity link to model the relationship between covariates of interest (e.g. SVL, CORT)

and follicle size, using the 'nlme' package ("lme" function, Pinheiro et al 2021). We built models that systematically included clutch size as a categorical independent variable, and where follicle size was the dependent variable.

To address our first question, we investigated the relationship between clutch size (independent variable) and follicle size (dependent variable) while accounting for sampling month (i.e. May, June, or July). Past work (Hudson et al 2020; Aubry et al 2020) indicates that variability in clutch size is best captured by the month within which animals are sampled, with reproductive activity peaking in June and slowing down in July. Accounting for the sampling month in our analysis will help control for variability in reproductive effort, as well as various stages of investment into both clutch size and follicle size. We tested for normality of model residuals using Shapiro's test (Royston 1982) and for homogeneity of variance using Levene's test (i.e. homoscedasticity, Hines and Hines 2000).

We tested for differences in follicle size on clutches of various sizes (one to three) while accounting for female size (SVL) using linear mixed models. We then investigated the effect of CORT, BKA, and ROMs on the relationship between clutch size and follicle size with the same approach. SVL, CORT and ROMs were log-transformed to un-skew their distribution and normalize residuals.

Model assumptions of normality of residuals and homoscedasticity were tested using appropriate statistics and are reported in the result section below. We adopted a significance level alpha of 0.05 for statistical significance in all analyses.

Results

We collected data from 454 follicles and 291 clutches. We obtained measures of CORT for 123 samples (9 from TA45, 82 from TA48, and 32 from TA55), BKA for 141 samples (11 from TA45, 96 from TA48, and 34 from TA55), and ROMs for 170 samples (16 from TA45, 102 from TA48, and 52 from TA55).

Clutch size and follicle size trade-off

We detected an effect of clutch size on follicle size (Table 1, n=454) where follicle size was on average larger in females with clutches of size two when compared to clutches of size one (*p-value* < 0.001, Fig. 2) or three (*p-value* < 0.001, Fig. 2). Although the effect of capture month was significant as well, its interaction with clutch size did not impact follicle size (Table 1). Model residuals were normally distributed (Shapiro–Wilk normality test, W=0.983, *p-value* = 0.263) and variances were homoscedastic across clutch sizes (Bartlett test of homogeneity of variances, Bartlett's $K^2=0.750$, df=2, *p-value* = 0.687).

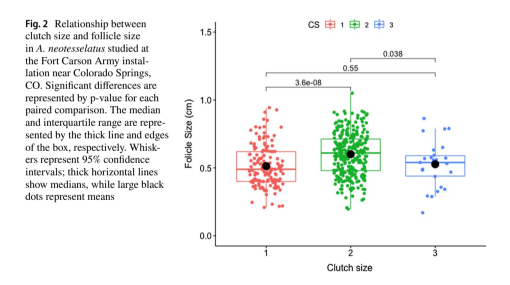
Effect of body size on clutch-follicle size trade-off

Females that were larger produced larger follicles regardless of clutch size (*p-value* < 0.001, n=291, Fig. 3). Follicle size was larger in clutches of size two when compared to other clutch sizes, independently of SVL (Fig. 3). Although SVL did affect follicle size, its interaction with clutch size did not (Table 2).

			•	
Fixed effects	Value	Std. Error	t-value	p-value
Month (-Intercept)	-0.0575	0.0225	-2.5518	0.0113
Month * (Clutch size of 2)	0.0183	0.0373	0.4909	0.6239
Month * (Clutch size of 3)	0.1175	0.1184	0.9926	0.3219
Clutch size of 1	0.6214	0.0477	13.0314	< 0.0001
Clutch size of 2	0.6919	0.0618	11.1871	< 0.0001
Clutch size of 3	0.4481	0.2099	2.1345	0.0338
Random effect	Intercept	Residual		
Std. deviation (Clutch Identity)	0.1321	0.0840		

 Table 1
 Table of results testing for the effect of clutch size (independent variable ranging from one to three) and month of capture 'Month' (independent variable) on follicle size (dependent variable) while accounting for nested measurements of follicle size within clutch (random effect of clutch identity)

We report regression estimates (Value), standard error (Std. Error), t-statistic and p-value for each effect, as well as the standard deviation of the intercept and residuals of the random effect of clutch identity; degrees of freedom = 238.



Effect of physiological state on clutch-follicle size trade-off

The effect of CORT and its interaction with clutch size did not affect follicle size (Table 3). Similarly, the effect of BKA and its interaction with clutch size did not affect follicle size (Table 4). ROMs and its interaction with clutch size did have significant effects on follicle size (Table 5, Fig. 4). Specifically, oxidative stress was higher in females with larger clutches (two or three follicles) and follicles of larger size, when compared to females that produced a single clutch with one small follicle (Fig. 4).

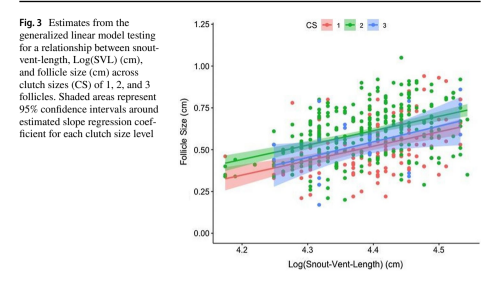


 Table 2
 Linear mixed model testing for the effects of clutch size ' (independent variable ranging from one to three), snout-vent-length 'SVL' (independent variable), and their interaction on follicle size (dependent variable) while accounting for nested measurements of follicle size within clutch (random effect of clutch identity)

Fixed effects	Value	Std. Error	t-statistic	p-value
log(SVL)(-Intercept)	0.8667	0.1664	5.2086	< 0.001
log(SVL) * (Clutch size of 2)	0.1879	0.2231	0.8419	0.4007
log(SVL) * (Clutch size of 3)	-0.1657	0.5593	-0.2962	0.7673
Clutch size of 1	- 3.2944	0.7297	-4.5149	< 0.001
Clutch size of 2	-0.7106	0.9779	-0.7267	0.4681
Clutch size of 3	0.7760	2.4513	0.3166	0.7519
Random effect	Intercept	Residual		
Std. deviation (Clutch Identity)	0.1099	0.0852		

We report regression estimates (Value), standard error (*Std. Error*), *t-statistic* and *p-value* for each effect, as well as the standard deviation of the intercept and residuals of the random effect of clutch identity; degrees of freedom = 288

Discussion

Given limited resources, theory predicts that a mother cannot simultaneously increase offspring size and offspring number, meaning that a trade-off between these two traits is inevitable because of physiological constraints (Lessels 1991; Stearns 1992). The existence of a size–number trade-off is strongly supported by both theoretical models and empirical data (Stearns 1992). Our results did not point to a trade-off between clutch size and follicle size, but rather to an optimum level of investment for clutches of size two, which also happens to be the most frequent clutch size amongst our observations. In addition, both clutch size and body size were particularly important factors in

Table 3 Linear mixed model testing for the effects of clutch size (independent variable ranging from one to three), log(CORT) (pg/mL) (independent variable), and their interaction on follicle size (dependent variable) while accounting for nested measurements of follicle size within clutch (random effect of clutch identity)

Fixed effects	Value	Std. Error	t-value	p-value
Log(CORT) (-Intercept)	0.0087	0.0067	1.2969	0.2034
Log(CORT) * (Clutch size of 2)	-0.0190	0.0134	-1.4141	0.1612
Log(CORT) * (Clutch size of 3)	-0.0391	0.2204	-0.1776	0.8595
Clutch size of 1	0.4928	0.0372	13.2438	< 0.001
Clutch size of 2	0.6153	0.0422	14.5924	< 0.001
Clutch size of 3	0.5318	0.1815	2.9293	0.0044
Random effect	Intercept	Residual		
Std. deviation (Clutch Identity)	0.1410	0.1056		

We report on regression estimates (Value), standard error (*Std. Error*), *t-statistic* and *p-value* for each effect, as well as the standard deviation of the intercept and residuals of the random effect of clutch identity; degrees of freedom = 82.

 Table 4
 Linear mixed model testing for the effects of clutch size (independent variable ranging from one to three), BKA (%) (independent variable), and their interaction on follicle size (dependent variable) while accounting for nested measurements of follicle size within clutch (random effect of clutch identity)

Fixed effects	Value	Std. Error	t-value	p-value
BKA (-Intercept)	-0.0006	0.0007	-0.8720	0.3883
BKA * (Clutch size of 2)	-0.0009	0.0011	-0.8671	0.3881
BKA * (Clutch size of 3)	0.0011	0.0026	0.4214	0.6745
Clutch size of 1	0.5452	0.0438	12.4496	< 0.001
Clutch size of 2	0.6805	0.0463	14.6812	< 0.001
Clutch size of 3	0.4868	0.1902	2.5596	0.0121
Random effect	Intercept	Residual		
Std. deviation (Clutch Identity)	0.1268	0.0982		

We report on regression estimates (Value), standard error (*Std. Error*), *t-statistic* and *p-value* for each effect, as well as the standard deviation of the intercept and residuals of the random effect of clutch identity; degrees of freedom=93.

determining follicle size (Figs. 2 and 3, respectively), and a reproductive investment into larger clutches and follicle size came at the cost of oxidative damage (Fig. 4).

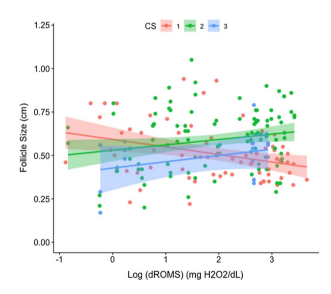
Specifically, our findings indicate that clutch size and follicle size were not inversely, nor linearly related in *A. neotesselatus*. Instead, follicles were larger in intermediate clutch sizes (i.e. two follicles) than those that were smaller and larger (i.e. one or three follicles). Among larger clutches of size three, we detect a trade-off between clutch size and follicle size, which suggests that this level of reproductive investment into the largest possible clutch comes at the cost of follicle quality/size, which is consistent with other reptile studies (Rowe 1994; Olsson and Shine 1997; Uller and Olsson 2009). Still,

Fixed effects	Value	Std. Error	t-value	p-value
Log(ROMs) (-Intercept)	-0.0401	0.0173	-2.3147	0.0244
Log(ROMs) * (Clutch size of 2)	0.0734	0.0267	2.7513	0.0070
Log(ROMs) * (Clutch size of 3)	0.0376	0.0686	0.5482	0.5847
(Clutch size of 1)	0.5859	0.0395	14.8505	0.0000
(Clutch size of 2)	0.5267	0.0459	11.4674	0.0000
(Clutch size of 3)	0.5286	0.1581	3.3423	0.0011
Random effect	Intercept	Residual		
Std. deviation (Clutch Identity)	0.1356	0.0912		

Table 5 Linear mixed model testing for the effects of clutch size (independent variable ranging from one to three), ROMs (mg H2O2/dL) (independent variable), and their interaction on follicle size (dependent variable) while accounting for nested measurements of follicle size within clutch (random effect of clutch identity)

We report on regression estimates (Value), standard error (*Std. Error*), *t-statistic* and *p-value* for each effect, as well as the standard deviation of the intercept and residuals of the random effect of clutch identity; degrees of freedom = 108.

Fig. 4 Estimates from the linear model testing for a relationship between oxidative damage (Log(ROMs) in mg H2O2/dL) and follicle size (cm) across clutch sizes (CS) of 1, 2, and 3 follicles. Shaded areas represent 95% confidence intervals around estimated slope regression coefficient for each clutch size level



the observed non-linear relationship between clutch size and follicle size is inconsistent with theoretical predictions (Lessels 1991; Stearns 1992).

Although progression in reproductive effort over the sampling period could confound this finding, the above relationship between clutch size and follicle size held true independently of reproductive stage. Indeed, the interaction between sampling month and clutch size did not have a significant effect on follicle size, meaning that seasonal progression did not affect the relationship that exists between clutch and follicle size. However, sampling month did have an effect on follicle size, suggesting that we did capture a change in follicle size with month of sampling. The observed non-linear relationship between follicle size and clutch size was mostly driven by samples collected in June, but there was no difference in follicle size across clutches of different sizes in July, and only a marginal difference in follicle size between clutches of size one and clutches of size two (where follicles were significantly larger in clutch of size two); results not presented herein for conciseness. Total investment per clutch, and the trade-off between follicle size and number, can shift among years within a single reptile population, depending upon a mother's provisioning rate (e.g. Olsson and Shine 1997). Additional years of data collection will help determine whether the observed bell-shaped pattern (Fig. 2) holds true, and whether intermediate clutches of size two maximize follicle size *in natura*.

We detected an effect of body size on follicle size within larger females (i.e. longer SVL) that was independent of clutch size which is consistent with other reptile studies (Ford and Siegel 1989; Qualls and Shine 1998; Horváthová et al. 2013; Recknagel and Elmer 2019). Larger females tend to have access to greater resources, and thus are more likely to allocate energy towards reproduction, resulting in larger clutches and/or larger follicles. Our findings suggest that larger *A. neotesselatus* females able to invest in larger clutches were also able to produce larger follicles, without having to compromise, suggesting that access to resources and female size are key in determining female reproductive success.

Findings regarding the effects of physiological state on the observed clutch-follicle size trade-off were mixed. We detected no relationship between CORT and follicle size, no matter the clutch size considered, indicating no difference in stress hormones between females that invested in large clutch sizes and females that invested in large follicle sizes. Aubry et al (2020) found that females who invested more energy into reproduction and produced larger clutch sizes often suffered higher CORT levels. However, focusing on a subset of the data for which we had follicle size measurements, we found that the relationship between clutch size and follicle size was not affected by energy-mobilization via CORT release.

Similarly, the effect of BKA on follicle size was not significant across clutches of various sizes, suggesting no difference in innate immunity between females that invested in large clutch sizes or large follicle size compared to other females. Aubry et al (2020) observed that an increase in reproductive investment correlated with a decrease in innate immunity, though only in lizards sampled from TA45, a difference in reproduction-immune trade-offs that is rarely documented in lizards (but see Lucas and French 2012). A decrease in immune function is seen in other animals that heavily invest in reproduction (Bonneaud et al 2003; French et al 2007a), yet the lack of a trade-off in our study suggests that more fine scale investment in clutch size and follicle size may not affect innate immunity, and vice versa.

As expected, we detect a relationship between oxidative stress and follicle size across clutches of various sizes. Aubry et al (2020) observed on a larger dataset that females with larger clutch sizes suffered greater oxidative stress. Of note, this comparison included females that were not gravid. Oxidative stress is the rate at which biomolecular oxidative damage is generated, which results from a complex interaction between compounds that oxidize (e.g. free radicals) and compounds that protect against oxidation (antioxidants) (Costantini and Verhulst 2009). Evolutionary ecologists have mainly studied the relationship between oxidative damage and life history trade-offs within the context of ageing theories, which involve compromises between current reproduction, future reproduction, and survival. A large body of work suggests that oxidative stress may be connected to reproduction, but findings have been quite mixed within and across taxa and reproductive strategies (Blount et al 2016). Our results suggest that such compromises are expressed when investigating the relationship between clutch size, follicle size, and oxidative stress (Fig. 4).

Because shifts in physiological state can affect individual reproduction and survival, which collectively drive population growth, it is important to understand the relationships that exist between physiological state and fitness components such as clutch size and egg size, which both influence reproductive success. If for example, elevations on oxidative stress markers such as reactive oxygen metabolites lead to tissue damage, then elevated physiological stress could result in decreased survival within populations (Lucas and French 2012).

With additional years of data collection on this species, we hope to bring survival probability into this equation in an effort to holistically quantify the trade-offs that exist between key fitness components such as reproduction, growth, maintenance and survival (Aubry et al 2019). A mechanistic understanding of how physiology and demographic rates are connected, along with a population-level assessment for the various sub-populations located at Fort Carson (which encompasses a large portion of the species narrow range), will help define a sound conservation plan for *A. neotesselatus*.

Acknowledgements We would like to thank Roger Payton and FC's Directorate of Public Works, Conservation Branch, for logistical and administrative assistance in facilitating our research. We would like to thank Dr. Sara Bombaci for serving on HC's Honors Thesis and providing valuable feedback on this manuscript. We would like to thank Dr. Maria Eifler and all of the volunteers and technicians who helped collect this data, including students from the CSU-ESA SEEDS Chapter.

Authors' contributions HEC, SSF and LMA conceived the study; HEC and LMA analyzed the data and wrote the manuscript; DE, HEC, field technicians and volunteers collected the data; SSF, SBH, ACW conducted the lab work; BMK and AJL facilitated field work and research activities; all authors provided feedback on earlier versions of the manuscript. This research predominantly funded by Fort Carson by way of a US Fish and Wildlife Cooperative Agreement F17AC00326. This research was also supported in part by the US Department of Agriculture, Animal and Plant Health Inspection Service, Wildlife Services, National Wildlife Research Center. The findings and conclusions in this publication have not been formally disseminated by the US Department of Agriculture and should not be construed to represent any agency determination or policy. Field methods were approved through Colorado State University IACUC 18–7772A. R code is available upon request; please contact senior author LMA.

Declarations

Conflict of interest The authors declare no conflicts of interest.

References

- Adamo SA, Jensen M, Younger M (2001) Changes in lifetime immunocompetence in male and female Gryllus texensis (formerly G. integer): trade-offs between immunity and reproduction. Anim Behav 62(3):417–425
- Ardia DR, Schat KA, Winkler DW (2003) Reproductive effort reduces long-term immune function in breeding tree swallows (Tachycineta bicolor). Proceed Royal Soc Lond Series b: Biol Sci 270(1525):1679–1683
- Aubry LM, Eifler D, Utsumi K et al (2019) Demographic assessment of the triploid parthenogenetic lizard Aspidoscelis neotesselatus at the northern edge of its range. Herpetol Conserv Biol 14(2):411
- Aubry LM, Hudson SB, Kluever BM et al (2020) Competing reproductive and physiological investments in an all-female lizard, the Colorado checkered whiptail. Evol Ecol 34:999–1016
- Bauwens D, Díaz-Uriarte R (1997) Covariation of life-history traits in lacertid lizards: a comparative study. Am Nat 149:91–111
- Blackburn TM (1991) An interspecific relationship between egg size and clutch size in birds. Auk 108(4):973–977
- Blount JD, Vitikainen EI, Stott I et al (2016) Oxidative shielding and the cost of reproduction. Biol Rev 91(2):483–497

- Bolker BM, Brooks ME, Clark CJ et al (2009) Generalized linear mixed models: a practical guide for ecology and evolution. Trends Ecol Evol 24(3):127–135
- Bonneaud C, Mazuc J, Gonzalez G et al (2003) Assessing the cost of mounting an immune response. Am Nat 161(3):367–379
- Case TJ (1978) Endothermy and parental care in the terrestrial vertebrates. Am Nat 112(987):861-874
- Costantini D (2016) Oxidative stress ecology and the d-ROMs test: facts, misfacts and an appraisal of a decade's work. Behav Ecol Sociobiol 70(5):809–820
- Costantini D, Verhulst S (2009) Does high antioxidant capacity indicate low oxidative stress? Funct Ecol 23:506–509
- Cox RM, Parker EU, Cheney DM et al (2010) Experimental evidence for physiological costs underlying the trade-off between reproduction and survival. Funct Ecol 24:1262–1269
- Dallman MF, Akana SF, Cascio CS et al (1987) Regulation of ACTH secretion: variations on a theme of B. In: Proceedings of the 1986 Laurentian hormone Conference. Academic Press 43:113–173
- Derting TL, Compton S (2003) Immune response, not immune maintenance, is energetically costly in wild white-footed mice (Peromyscus leucopus). Physiol Biochem Zool 76(5):744–752
- Dunlap KD, Schall JJ (1995) Hormonal alterations and reproductive inhibition in male fence lizards (Sceloporus occidentalis) infected with the malarial parasite Plasmodium mexicanum. Physiol Zool 68(4):608–621
- Dziminski MA, Vercoe PE, Roberts JD (2009) Variable offspring provisioning and fitness: a direct test in the field. Funct Ecol 23(1):164–171
- Ferguson GW, Fox SF (1984) Annual variation of survival advantage of large juvenile side-blotched lizards, Uta stansburiana: its causes and evolutionary significance. Evolution 38:342–349
- Ford NB, Seigel RA (1989) Relationships among body size, clutch size, and egg size in three species of oviparous snakes. Herpetologica 45(1):75–83
- Ford NB, Seigel RA (2010) An experimental test of the fractional egg size hypothesis. Herpetologica 66(4):451-455
- French SS, Neuman-Lee LA (2012) Improved ex vivo method for microbiocidal activity across vertebrate species. Biology Open 1(5):482–487
- French SS, DeNardo DF, Moore MC (2007a) Trade-offs between the reproductive and immune systems: facultative responses to resources or obligate responses to reproduction? Am Nat 170(1):79–89
- French SS, Johnston GIH, Moore MC (2007b) Immune activity suppresses reproduction in food-limited female tree lizards Urosaurus ornatus. Funct Ecol 21(6):1115–1122
- Gilman CA, Wolf BO (2007) Use of portable ultrasonography as a nondestructive method for estimating reproductive effort in lizards. J Exp Biol 210(11):1859–1867
- Grafen A (1988) On the uses of data on lifetime reproductive success. In: Clutton-Brock TH (ed) Reproductive Success: Studies of Individual Variation in Contrasting Breeding Systems. University of Chicago Press, pp 454–471
- Graham AL, Hayward AD, Watt KA et al (2010) Fitness correlates of heritable variation in antibody responsiveness in a wild mammal. Science 330(6004):662–665
- Hamilton WD, Zuk M (1982) Heritable true fitness and bright birds: a role for parasites? Science 218(4570):384–387
- Hines WGS, Hines ROH (2000) Increased power with modified forms of the Levene (Med) test for heterogeneity of variance. Biometrics 56(2):451–454
- Horváthová T, Cooney CR, Fitze PS et al (2013) Length of activity season drives geographic variation in body size of a widely distributed lizard. Ecol Evol 3(8):2424–2442
- Hudson SB, Kluever BM, Webb AC et al (2020) Steroid hormones, energetic state, and immunocompetence vary across reproductive contexts in a parthenogenetic lizard. Gen Comp Endocrinol 288:113372
- Hughes PW (2017) Between semelparity and iteroparity: Empirical evidence for a continuum of modes of parity. Ecol Evol 7(20):8232–8261
- Ilmonen P, Taarna T, Hasselquist D (2000) Experimentally activated immune defense in female pied flycatchers results in reduced breeding success. Proceed Royal Soc Lond Series b: Biol Sci 267(1444):665–670
- Jones MT, Gillham B (1988) Factors involved in the regulation of adrenocorticotropic hormone/betalipotropic hormone. Physiol Rev 68(3):743-818
- Knowles SCL, Nakagawa S, Sheldon BC (2009) Elevated reproductive effort increases blood parasitaemia and decreases immune function in birds: a meta-regression approach. Funct Ecol 23:405–415
- Kratochvíl L, Frynta D (2006) Body-size effect on egg size in eublepharid geckos (Squamata: Eublepharidae), lizards with invariant clutch size: negative allometry for egg size in ectotherms is not universal. Biol J Lin Soc 88(4):527–532

Krawchuk MA, Brooks RJ (1998) Basking behavior as a measure of reproductive cost and energy allocation in the painted turtle. Chrysemys Picta Herpetologica 54(1):112–121

Lack D (1947) The significance of clutch-size in the partridge (Perdix perdix). J Anim Ecol 16:19-25

- Lessels CM (1991) The evolution of life histories. In: Krebs JR, Davies NB (eds) Behavioural ecology: an evolutionary approach. Blackwell, pp 32–68
- Ljungström G, Stjernstedt M, Wapstra E et al (2016) Selection and constraints on offspring size-number trade-offs in sand lizards (Lacerta agilis). J Evol Biol 29(5):979–990
- Lochmiller RL, Deerenberg C (2000) Trade-offs in evolutionary immunology: just what is the cost of immunity? Oikos 88(1):87–98
- Lucas LD, French SS (2012) Stress-induced tradeoffs in a free-living lizard across a variable landscape: consequences for individuals and populations. PLoS ONE 7(11):e49895
- MacLean GS, Lee AK (1973) Wilson KJ (1973) A simple method of obtaining blood from lizards. Copeia 2:338–339
- Martin LB, Weil ZM, Nelson RJ (2008) Seasonal changes in vertebrate immune activity: mediation by physiological trade-offs. Phil Trans Royal Soc B: Biol Sci 363(1490):321–339
- Metcalfe NB, Alonso-Alvarez C (2010) Oxidative stress as a life-history constraint: the role of reactive oxygen species in shaping phenotypes from conception to death. Funct Ecol 24(5):984–996
- Olsson M, Shine R (1997) The limits to reproductive output: offspring size versus number in the sand lizard (Lacerta agilis). Am Nat 149(1):179–188
- Parker GA, Begon M (1986) Optimal egg size and clutch size: effects of environment and maternal phenotype. Am Nat 128(4):573–592
- Pianka ER (1970) On r and K selection. Am Nat 104(940):592-597
- Pinheiro J, Bates D, DebRoy S et al (2021). R Core Team, nlme: Linear and nonlinear mixed effects models. R package version 3.1–152
- Qualls CP, Shine R (1998) Costs of reproduction in conspecific oviparous and viviparous lizards. Lerista Bougainvillii Oikos 82(3):539–551
- Recknagel H, Elmer KR (2019) Differential reproductive investment in co-occurring oviparous and viviparous common lizards (Zootoca vivipara) and implications for life-history trade-offs with viviparity. Oecologia 190(1):85–98
- Roff DA (2002) Life history evolution. Sinauer Associates, Sunderland, MA
- Rohwer FC (1988) Inter and intraspecific relationships between egg size and clutch size in waterfowl. Auk 105(1):161–176
- Romero LM, Reed JM (2005) Collecting baseline corticosterone samples in the field: is under 3 min good enough? Comp Biochem Physiol A: Mol Integr Physiol 140(1):73–79
- Romero LM, Wingfield JC (2001) Regulation of the hypothalamic-pituitary-adrenal axis in free-living pigeons. J Comp Physiol B 171(3):231–235
- Rowe JW (1994) Reproductive variation and the egg size-clutch size trade-off within and among populations of painted turtles (Chrysemys picta bellii). Oecologia 99:35–44
- Royston JP (1982) An extension of Shapiro and Wilk's W test for normality to large samples. J Roy Stat Soc: Ser C (appl Stat) 31(2):115–124
- Sheldon BC, Verhulst S (1996) Ecological immunology: costly parasite defences and trade-offs in evolutionary ecology. Trends Ecol Evol 11(8):317–321
- Sheriff MJ, Dantzer B, Delehanty B et al (2011) Measuring stress in wildlife: techniques for quantifying glucocorticoids. Oecologia 166(4):869–887
- Sinervo B (1990) The evolution of maternal investment in lizards: an experimental and comparative analysis of egg size and its effects on offspring performance. Evolution 44:279–294
- Smith GD, French SS (2017) Physiological trade-offs in lizards: costs for individuals and populations. Integr Comp Biol 57(2):344–351
- Smith CC, Fretwell SD (1974) The optimal balance between size and number of offspring. Am Nat 108:499–506
- Stearns SC (1992) The evolution of life histories. Oxford University Press, New York, p 249
- Sullivan BK, Kwiatkowski MA (2007) Courtship displays in anurans and lizards: theoretical and empirical contributions to our understanding of costs and selection on males due to female choice. Funct Ecol 21(4):666–675
- Sonosite Turbo ultrasound, FUJIFILM SonoSite Inc., Bothell, Washington, USA
- Uller T, Olsson M (2009) Offspring size-number trade-off in a lizard with small clutch sizes: tests of invariants and potential implications. Evol Ecol 23:363–372

Wickham H, Wickham MH (2020) plyr package https://cran.rproject.org/web/packages/dplyr/dplyr.pdf Wickham H (2016) Programming with ggplot2. Springer, pp 241–253

- Williams GC (1966) Natural selection, the costs of reproduction, and a refinement of Lack's principle. Am Nat 100(916):687–690
- Williams TD (2001) Experimental manipulation of female reproduction reveals an intraspecific egg size clutch size trade-off. Proceed Royal Soc Lond Series b: Biol Sci 268(1465):423–428

Publisher's Note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Authors and Affiliations

Hannah E. Caracalas¹ · S. S. French^{3,4} · S. B. Hudson^{3,4} · B. M. Kluever⁵ · A. C. Webb^{3,4} · D. Eifler⁶ · A. J. Lehmicke⁷ · L. M. Aubry^{1,2}

- ¹ Department of Fish, Wildlife, and Conservation Biology, Colorado State University, Fort Collins, CO 80523-1474, USA
- ² Graduate Degree Program in Ecology, Colorado State University, 102 Johnson Hall Fort Collins, Fort Collins, CO 80523-1021, USA
- ³ Department of Biology, Utah State University, Logan, UT 84322-5305, USA
- ⁴ Ecology Center, Utah State University, Logan, UT 84322-5205, USA
- ⁵ Department of Agriculture, National Wildlife Research Center, Florida Field Station, Wildlife Services, Gainesville, FL 32641-6033, USA
- ⁶ The Erell Institute, Lawrence, KS 66047, USA
- ⁷ DPW Environmental Division, 1626 Evans St., Bldg 1219, Fort Carson, CO 80913, USA