Food restriction affects maternal investment but not neonate phenotypes in a viviparous lizard

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ABSTRACT

Food availability significantly affects an animal’s energy metabolism, and thus its phenotype, survival, and reproduction. Maternal and offspring responses to food conditions are critical for understanding population dynamics and life-history evolution of a species. In this study, we conducted food manipulation experiments in field enclosures to identify the effect of food restriction on female reproductive traits and postpartum body condition, as well as on hatching phenotypes, in a lacertid viviparous lizard from the Inner Mongolian desert steppe of China. Females under low-food availability treatment (LFT) had poorer immune function and body condition compared with those under high-food availability treatment (HFT). The food availability treatments significantly affected the litter size and litter mass of the females, but not their gestation period in captivity or brood success, or the body size, sprint speed, and sex ratio of the neonates. Females from the LFT group had smaller litter sizes and, therefore, lower litter mass than those from the HFT group. These results suggest that female racerunners facing food restriction lay fewer offspring with unchanged body size and locomotor performance, and incur a cost in the form of poor postpartum body condition and immune function. The flexibility of maternal responses to variable food availability represents an important life strategy that could enhance the resistance of lizards to unpredictable environmental change.

Keywords: Eremias multiocellata; Food availability; Hatchling; Lizard; Reproductive output

INTRODUCTION

In nature, the food available to an animal exhibits spatio-temporal variation, significantly affecting its energy metabolism and consequently its phenotype, survival, and reproduction (Acheampong et al., 2011; Douhard et al., 2014; Du, 2006; Hoy et al., 2016). Food availability not only affects maternal fitness-related traits (e.g. body size, growth, and reproduction), but might also induce phenotypic variations in offspring (Ballinger & Congdon, 1980; Hillesheim & Stearns, 1991; Jones et al., 2015; Vaissi & Sharifi, 2016). For example, food availability can lead to variations in maternal reproduction (Ballinger, 1977) and offspring growth and survival in lizards (Dunham, 1978; Warner et al., 2015). Therefore, maternal and offspring responses to food conditions are critical for understanding ecological and evolutionary processes, such as population dynamics and life-history evolution.

Food availability can significantly affect female reproductive strategies, such as reproductive timing, investment, and output (offspring number and size) (Ballinger, 1977; Du, 2006; Ramírez-Pinilla, 2006). Two kinds of trade-offs regarding energy allocation are faced by a mother when the energy available to her is limited. First, a mother has to decide on energy allocation for multiple tasks, such as maintenance, growth, and reproduction, leading to important life-history trade-offs (e.g., maintenance vs. reproduction) (Hegemann et al., 2013; Rollinson & Rowe, 2016); for instance, the tropical house wren (Troglodytes aedon) decreases parental reproductive investment (i.e. nestling feeding frequency), but does not alter self-maintenance (metabolic rate and body condition) when the cost of activity increases during reproduction (Tieleman et al., 2008). Second, a mother needs to decide on the distribution of energy among offspring within a clutch, thus leading to a trade-off between clutch size and offspring size (Bleu et al., 2013; Du et al., 2005; Olsson et al., 2002). The “optimal egg size theory” assumes that one optimal egg size is appropriate under certain

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maternal and environmental conditions to maximize the fitness of offspring (Einum & Fleming, 1999; Smith & Fretwell, 1974; Williams, 1966). Accordingly, a mother might give priority of energy allocation to offspring size rather than to clutch size when faced with food restriction, especially in those species whose fitness is determined primarily by egg size (Styrsky et al., 2000). However, other studies have shown that offspring size is not always optimized, and can be highly condition-dependent in some species (Krist & Munchinger, 2015; Rollinson & Hutchings, 2013), with the shift in size being a function of maternal reproductive investment (Caley et al. 2001). In such cases, a mother might change both offspring and clutch sizes in response to food restriction. Despite extensive studies regarding the effect of food availability on maternal reproductive traits (e.g. Bonnet et al., 2001; Du, 2006; Hogstedt, 1981; Kitaysky et al., 1999; Warner & Lovern, 2014; Warner et al., 2015), how female reproduction responds to food availability might differ among species and deserves further investigation.

In addition to maternal reproductive traits, food availability can also profoundly affect offspring phenotypes and fitness. Previous studies have shown that a variety of phenotypic traits of offspring (e.g., body size, locomotor performance, and growth rate) can be affected by maternal food availability (Du, 2006; Hafer et al., 2011; Raveh et al., 2016; Warner et al., 2015), likely through a maternal effect—a phenomenon in which environmental information passes through generations by means of plasticity rather than direct genetic transmission (Fox & Mousseau, 1998; Hsu et al., 2016); for example, low food availability compromises the snout-vent length (SVL) of offspring, as well as the mass, performance ability, and fat reserves of the viviparous lizard *Pseudemoia entrecasteauxii* (Itonaga et al., 2012b). Offspring sex is one of the most interesting phenotypes affected by maternal food availability. The “sex allocation hypothesis” suggests that females can manipulate the sex ratio of their offspring, contingent upon local conditions, to maximize offspring fitness (Rosenfeld & Roberts, 2004; Trivers & Willard, 1973). Food availability can affect the offspring sex ratio in several avian species; for example, when provided with supplementary food, the kakapo (*Strigops habroptilus*) female produces an excess of males (Clout et al., 2002). However, such studies are scarce for reptilian species (Wapstra & Warner, 2010; Warner et al., 2007).

Viviparous lizards make an excellent model for studying maternal and offspring responses to food availability because many lizards experience fluctuating food resources (Meserve et al., 2016; Zhu et al., 2014), and viviparous species retain their eggs for a longer period than oviiparous species do, with higher locomotor and thermoregulatory costs (Le Galliard et al., 2003; Shine, 2003). Moreover, most ectothermic vertebrates are considered capital breeders, in which reproduction is financed from stored energetic capital (Bonnet et al., 1998), which makes the direct causal relationship between food availability and reproduction harder to detect. Conversely, many viviparous reptiles are considered income breeders because they can use maternal nutrients (matrotrophy) to supplement or replace yolk nutrients (lecithotrophy) for embryonic development (Bonnet et al., 2001; Ramirez-Pinilla, 2006; Winne et al., 2006). As a result, viviparous lizards may confront more severe trade-offs between reproduction and self-maintenance when experiencing limited resources during gestation, and thus provide a unique system to increase our understanding of the effects of food on life history strategies in animals. In this study, we conducted food manipulation experiments in field enclosures to identify the effect of food restriction on female reproductive traits (such as gestation period in captivity and litter size) and postpartum body condition, as well as hatching phenotypes (body size and locomotor performance and immune response) in a lacertid viviparous lizard from the Inner Mongolian desert steppe of China. We aimed to address the following questions: (1) How does female reproductive investment respond to food restriction? (2) Does food restriction affect postpartum body condition of females? (3) How does maternal food restriction influence neonate phenotypes?

**MATERIALS AND METHODS**

**Study species**

The multi-ocellated racerunner (*Eremias multiocellata*) is a small viviparous lizard that inhabits desert or semiarid areas, with the SVL of adults ranging from 58 to 73 mm. This species is the main lizard fauna in our field study site at the Shierlancheng Field Station, Institute of Grassland Research of the Chinese Academy of Agricultural Sciences (Zeng et al., 2016), located in Ordos, Inner Mongolia, China (N40°12′17″; E111°07′43″; elevation 1 036 m). Thermal and hydric environments significantly affect the reproductive traits and offspring phenotypes in different populations of this species, with the offspring sex ratio biased towards males when gravid females are kept at high temperatures (Tang et al., 2012; Wang et al., 2016).

**Maternal food treatment in field enclosures**

Adult *E. multiocellata* (16 males and 48 females) were collected from our field study site and were housed in round enclosures (high×diameter=40 cm×180 cm) built at the field site (one male and three females in each enclosure) from May 20 to May 30, 2014, which is the beginning of the reproductive season for this species. All females had blemishes on their belly, suggesting that they had mated in the field. The enclosures were covered with plastic nets to avoid predation by birds.

On June 1, 2014, the SVL and body mass of all females were measured, after which each female was randomly assigned to a treatment group with either high (*n=24*) or low (*n=24*) prey availability. Lizards in the high-food availability treatment (HFT) group were fed every other day with 0.05 g of mealworms per gram mass of female per day (amounting to 120% of the average consumption of a gravid female, as measured prior to the experiment), whereas those in the low-food availability treatment (LFT) group were fed at the same intervals with 0.025 g of mealworms per gram mass of female per day (amounting to 60% of the average consumption of a gravid female). Each treatment was replicated eight times (for a total of 16 enclosures). Given that all viviparous amniotes have a placenta that transfers maternal nutrients to the embryo (Flemming et al.,
Female reproductive traits and phenotypes of neonates

On July 1, 2014, females (16 females from HFT and 15 females from LFT) in the field enclosures were retrieved and transferred to the laboratory for measurement of reproductive traits. Females were maintained in small cages (long×wide×high=310×210×180 mm) with a substrate of sand and two small pieces of brick as shelter. The cages were exposed to the natural light regime of the field station, and a 60W incandescent light bulb was suspended 5 cm above each cage for thermoregulation from 0800 to 1200h. Food (mealworms and crickets dusted with additional vitamins and minerals) was provided daily ad libitum. Each cage contained two females and was checked once per day for neonates, and four times per day following first parturition.

Immediately after a female produced a litter of neonates, the mother was measured and weighed. The SVL and body mass of neonates were then measured to 0.01 mm and 0.001 g accuracy after the absorption of the yolks one day later. Litter size was determined as the number of neonates and litter mass was calculated as the total mass of neonates produced by a female (Li et al., 2006; Ramírez-Bautista et al., 2000). The gestation period in captivity was calculated as the days between the initiation of treatment and female parturition, which did not include the gestation period in the field before the females were collected. To measure the locomotor performance, each neonate was made to run on a racetrack (80 cm long, marked at 20-cm intervals), within 2–3 days of birth, by stimulation with a soft paintbrush (Irschick & Losos, 1998). This procedure was repeated twice at 30±1 °C allowing a break of 1 h between each test. For quantitating locomotor performance, the sprint speed was determined by averaging the fastest speeds for covering a distance of 20 cm in each race. The sex of neonates was identified by observing the preanal scales—males have large, square, regularly-distributed preanal scales, whereas females have small, round, and scattered preanal scales (Wang et al., 2016).

A total of 12 and 14 females produced offspring from the LFT and HFT groups, respectively. Offspring from four clutches in the LFT group and one clutch in the HFT group were stillborn and excluded from further analysis of locomotor performance and offspring sex. The brood success of females was calculated as the number of females producing live neonates / total number of females.

Cellular immune response of postpartum females

Cellular immune response was assessed by administering an injection (20 µL) of 50 mg of phytohemagglutinin (PHA) in the right foot of postpartum E. multocellata females. The thickness of the right and left feet were measured every 12 h from 0 to 36 h following the PHA injection. The difference in foot thickness was considered as an index of immune response (Brown et al., 2011; Vinkler et al., 2010). The largest difference in thickness (i.e., peak immune response) was found at 12 h. Thus, PHA-induced skin swelling at 12 h was used to identify the effect of food availability on the immune response of females.

Statistical analysis

All analyses were performed with SPSS Statistics software (ver. 22; IBM Corp. 2014). Data were normalized by log-transformation when necessary. Maternal body condition was calculated as a residual score from the regression of body mass on SVL (Jakob et al., 1996). Treatment effects on maternal body condition, PHA response, reproductive output, and neonate traits were evaluated by one-way analysis of variance (ANOVA). Clutch means were calculated for neonate traits to avoid pseudo-replication. Pearson Chi-Square test was used for comparing between-treatment differences in brood success.

RESULTS

At the beginning of the experiment, both male and female body sizes did not differ between food treatments (Table 1). However, the postpartum body conditions were worse for females in the LFT group than for those in the HFT group ($F_{1,24}=7.928$, $P=0.010$) (Figure 1A), although initial body condition did not differ between the two food treatment groups ($F_{1,24}=1.162$, $P=0.292$). In addition, females from the LFT group had poorer immune function than those from the HFT group, as indicated by the lower PHA response of LFT group females ($F_{1,23}=7.214$, $P=0.014$) (Figure 1B).

Food availability treatment did not affect brood success or gestation period in captivity of females, but did significantly affect both litter size and litter mass (Table 1). The females from the LFT group had smaller litter sizes and, therefore, lower litter mass compared with those for females from the HFT group (Table 1). Moreover, maternal food availability did not affect the body size (SVL and body mass), sprint speed, or sex ratio of the neonates (Table 1).

DISCUSSION

When facing food restriction, female racerunners lay fewer offspring with unchanged body size and locomotor performance, but at a cost in terms of poor postpartum body condition and immune function. These maternal and offspring responses to food restriction have interesting implications for our understanding of the reproductive strategies of lizards under temporally fluctuating food abundance in nature.

A female must decide how to allocate limited resources to the processes of self-maintenance and reproduction (Du, 2006; Hegemann et al., 2013; Itonaga et al., 2012a). Obviously, female racerunners from the LFT group produced neonates with a similar body size to those produced by females from the HFT group, but at the cost of poor body condition and immune function. Understandably, lizards adopt a strategy of keeping reproductive investment relatively constant and sacrificing other requirements like those of immune function and energy storage under constrained food conditions, because maintaining a
Table 1  Reproductive traits of *Eremias multiocellata* under high- and low-food availability treatment

| Variable                          | Treatment          |                | P          |
|----------------------------------|--------------------|----------------|------------|
|                                  | Low food           | High food      | F, df, P   |
| Adult body sizes                 |                    |                |            |
| Male SVL (mm)                    | 63.19±0.65 (n=8)   | 62.94±0.65 (n=8) | F₁, 1₄=0.740, P=0.789 |
| Male mass (g)                    | 6.94±0.25 (n=8)    | 7.18±0.25 (n=8) | F₁, 1₄=0.435, P=0.520 |
| Female SVL (mm)                  | 66.08±0.38 (n=24)  | 66.16±0.38 (n=24) | F₁, 4₆=0.020, P=0.889 |
| Female mass (g)                  | 7.24±0.16 (n=24)   | 7.28±0.16 (n=24) | F₁, 4₆=0.041, P=0.841 |
| Reproductive parameters          |                    |                |            |
| Brood success                    | 53.33% (n=15)      | 81.25% (n=16)  | χ²=1.631, P=0.202 |
| Gestation period in captivity (days) | 41.58±1.28 (n=12) | 42.00±1.19 (n=14) | F₁, 2₄=0.057, P=0.814 |
| Litter size (n)                  | 2.67±0.27 (n=12)   | 3.71±0.25 (n=14) | F₁, 2₄=7.907, P=0.010 |
| Litter mass (g)                  | 1.56±0.17 (n=12)   | 2.16±0.16 (n=14) | F₁, 2₄=6.650, P=0.016 |
| Neonate traits                   |                    |                |            |
| Neonate SVL (mm)                 | 30.15±0.33 (n=12)  | 29.52±0.30 (n=14) | F₁, 2₄=1.991, P=0.171 |
| Neonate mass (g)                 | 0.59±0.02 (n=12)   | 0.58±0.02 (n=14) | F₁, 2₄=0.008, P=0.930 |
| Sprint speed (m/s)               | 0.85±0.13 (n=8)    | 0.95±0.10 (n=13) | F₁, 1₉=0.355, P=0.558 |
| Sex ratio (male%)                | 42.10±8.50 (n=8)   | 51.10±11.30 (n=13) | F₁, 1₉=0.397, P=0.536 |

Values are expressed as means±SE. One-way ANOVA and Fisher’s Exact Test were used to compare the between-treatment differences in reproductive traits. Brood success=number of females producing live neonates/total number of females. Neonate traits (except for sex ratio) were calculated as clutch means to avoid pseudo-replication. SVL, snout-vent length.

Figure 1  Effects of food availability treatment on female postpartum body condition (A) and immune function (B)

Data are summarized as means±SE. Immune function represents cellular immune response assessed by phytohemagglutinin (PHA)-induced skin swelling within 12 h of PHA application. LFT, low-food availability treatment; HFT, high-food availability treatment. A competent immune system is a nutritionally demanding process that necessitates trade-off decisions with competing nutrient demands, such as those of growth, reproduction, and thermal regulation (Sheldon & Verhulst, 1996; Uller et al., 2006; Zamora-Camacho et al., 2016). This strategy of females giving priority to reproduction rather than to self-maintenance has also been reported in other reptiles and vertebrate species; for example, the body condition of female jacky dragons (*Amphibolurus muricatus*) fed a low-quality diet declined dramatically throughout the reproductive season, whereas their hatchlings were larger and in better body condition than those produced from females fed a high-quality diet (Warner et al., 2007). Such a life strategy helps the reproduction and recruitment of populations, but may significantly weaken the immune response and, therefore, the survival of females via elevated risks of predation or starvation in their natural habitat (Iglesias-Carrasco et al., 2016; Neuman-Lee et al., 2015).

The pre-breeding nutritional condition of females is closely related to the production of neonates in oviparous vertebrates, with reduced fecundity under conditions of food limitation (Du, 2006; Donelson et al., 2008; Johnson et al., 2014; Lehman & Smith, 1988; Warner & Lovern, 2014). In response to decreased resources, the females of multiple-clutch species can reduce their reproductive output by decreasing litter size (Sun et al., 2002), offspring size (Abell, 1999), or reproductive frequency (Du, 2006). The viviparous racersrunners in our study produced one litter per reproductive season, and decreased their litter size rather than offspring size in response to food restriction. This reproductive strategy reflects the highly critical
reproductive investment for each hatchling, suggesting that an optimal neonate body size could exist independent of nutritional conditions. This is consistent with the optimal offspring size theory in which females reproducing in a given environment divide available resources into optimally-sized offspring (Sinervo & Licht, 1991; Smith & Fretwell, 1974). Nonetheless, further studies on the relationship between initial body size and fitness of neonates are needed to verify this hypothesis. At the same time, the reduction in clutch size under conditions of low food availability could be an effective strategy to improve offspring fitness. Fewer offspring means more abundant food per neonate in an environment with finite resources, which will reduce competition and maximize the survival and reproduction of descendants (Bartlett, 1988).

The traditional sex allocation theory generates two conflicting predictions on how maternal nutrition affects the offspring sex ratio (Hamilton, 1967; Trivers & Willard, 1973). Females may produce higher numbers of male offspring under low-food conditions because males might have higher fitness or are more likely to disperse under stressful food conditions (Komdeur et al., 1997; Warner et al., 2007). Alternatively, the opposite pattern may occur, because producing more daughters under low-food conditions will gain greater fitness return given that offspring fitness is less dependent on body size (hence, reproductive energy input) in daughters than in sons (Krist & Munclinger, 2015; Trivers & Willard, 1973). In the current case, however, females did not adjust the sex ratio of their offspring, giving no support to either prediction above.

Overall, female viviparous lizards may sacrifice their own health to produce high-quality offspring and maintain the sex ratio, which could maximize their reproductive success and individual fitness when experiencing low food availability. In addition to maternal and offspring responses to environmental factors like temperature and precipitation (Ma et al., 2014; Wang et al., 2016), the flexibility of maternal responses to variable food availability represents an important life strategy that might enhance the resistance of lizards to unpredictable environmental change. Understanding maternal and offspring responses to the combined impact of these biotic and abiotic factors is a considerable challenge and should be of great interest for future investigations.

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