

INTRODUCTION

The environment that organisms experience during development can have a considerable influence on phenotypic expression. For example, individuals are primed to adjust their phenotype to developmental cues through epigenetic changes (Weaver et al. 2004; Ledon-Rettig et al. 2012) which can have long-term consequences on morphology, physiology, and behavior (West-Eberhard 2003; Kasumovic 2013; Snell-Rood 2013). Understanding the contribution of the developmental environment to phenotypic expression is therefore fundamental to understanding the generation of stable individual differences in behavior (i.e., animal personality and behavioral syndromes, Sih et al. 2004a, 2004b; Réale et al. 2007).

Personality variation arises from the combined influence of genetic and environmental factors (Dingemanse and Dochtermann 2014, Han and Dingemanse 2015) with each accounting for around 50% of observed personality variation (Dochtermann et al. 2015). A large part of personality research has focused its attention on quantifying the genetic inheritance of behavioral variation (Taylor et al. 2012), its fitness consequences (Bergeron et al. 2013), and implications for a population’s evolutionary trajectories (Dochtermann and Dingemanse 2013). However, these areas of attention largely deal with ultimate causes and consequences of personality variation and behavioral syndromes and as such are fundamentally multigenerational topics. In contrast, variation in the environment that an individual experiences during development—such as food abundance, food quality, conspecific density, or presence of predator cues—typically operate within the life-time of individuals.
Focusing on how within-lifetime processes affect the expression of behavioral variation is necessary because differences in early experiences may place individuals along different life-history trajectories in interaction with their behavioral types (Buss and Greiling 1999; Carere et al. 2005). As changes in the expression of a behavioral phenotype are often associated with costs (reviewed by Snell-Rood [2013]), such changes might be restricted to sensitive periods during early development (Groothuis and Trillmich 2011). Manipulative studies have identified many of the developmental conditions that influence the expression of behaviors (diet quality: Noguera et al. 2013; immune challenge: Butler et al. 2012; predation pressure: Niemela et al. 2012b; access to shelter: Bengston and Jandt 2014; conspecific cues: DiRienzo et al. 2012; Niemela et al. 2012a; physical and social environment: Liedtke et al. 2015). Most of these studies, however, focus strictly on changes in the population average while effects on behavioral variation in and of itself remain largely ignored (but see DiRienzo et al. [2013]). This is a missed opportunity because determining the effects of developmental conditions on behavioral variation can shed light on the set of factors that promote the generation of stable individual differences (Duckworth 2010; Stamps and Groothuis 2010; Jandt et al. 2014; Han and Dingemanse 2015).

The developmental environment can also create conditions by which morphological and behavioral variation dynamically interact with each other via feedback loops. For example, diet quality impacts growth rates (Hunt et al. 2005) and resulting morphological differences can in turn affect behavioral and life history strategies (; Metcalfe and Monaghan 2001; Lee et al. 2015). Feedback loops connecting behavior with an individual’s morphology (e.g., condition or other state variables, sensu Houston and McNamara [1999]) can then affect the expression of behavioral variation within a population. Under a negative feedback scenario, if individuals with access to higher quality or more food are larger or have positive energy balances, they are expected to engage in asset protection (Clark 1994; Clark and Mangel 2000) and limit the amount of risk-taking behaviors over time. This behavioral change is then expected to reduce the amount of personality variation observed over time (Dall et al. 2004; Luttbeg and Sih 2010). In a positive feedback scenario, such as state-dependent safety (Luttbeg and Sih 2010), individuals with higher body sizes or more positive energy balances are less likely to be captured by predators and can increase their foraging effort under predation risk, in turn increasing their state value. Such a feedback loop is again expected to increase the amount of among-individual variation present within a population and lead to stable behavioral differences (Luttbeg and Sih 2010). Although state-dependent models of personality have been extensively investigated through theoretical and conceptual models (Dall et al. 2004; Wolf et al. 2007; Luttbeg and Sih 2010; Montigio and Royauté 2014; Sih et al. 2015), they remain poorly studied empirically (but see Mathot et al. [2011]). In the absence of direct tests, state-dependent feedback can be indicated by among-individual state-based differences in behavior (Sih et al. 2015).

State-dependent feedback loops and their effects on behavioral variation are further complicated by the fact that behavioral responses are frequently integrated within behavioral syndromes (Sih et al. 2004a, 2004b). Because they are generated in part through genetic correlations (Dochtermann and Roff 2010; Dochtermann 2011; Han and Dingemanse 2015), behavioral syndromes may constrain evolutionary responses (Dochtermann and Dingemanse 2013) and may prevent populations from reaching adaptive peaks if behavioral syndromes are robust to environmental variability. There is, however, considerable variation among organisms and within populations in the behaviors associated with a syndrome, often depending on environmental conditions (Bell 2005; Bell and Sih 2007; Dingemanse et al. 2007; Royauté et al. 2014, 2015a). Unfortunately despite the stability of behavioral syndromes over multiple life-stages having been well studied (Sinn et al. 2008; Brodin 2009; Wilson and Krause 2012), the developmental characteristics that shape the emergence of syndromes are poorly understood (Han and Dingemanse 2015).

Here we investigated how variation in a single component of the developmental environment, diet quality, influenced subsequent variation in and correlations among body mass, activity, and response to cues of predator presence in juvenile house crickets (Acheta domestica). Specifically, we aimed to answer the following questions: (1) Does variation in diet quality generate different developmental trajectories in growth rates, activity, and response to cues of predator presence? (2) Does diet quality increase or decrease body mass and behavioral variation? (3) Does diet quality affect the strength of correlations between morphological and behavioral traits? At the population average level, we expected a low quality diet to decrease growth rate and, as a result, increase an individual’s propensity to exhibit risky behaviors in order to meet energetic demands (state-dependent safety and starvation avoidance principles, Luttbeg and Sih 2010; Figure 1a). Here, this would translate into higher activity levels and a higher propensity to ignore the presence of predatory cues. In contrast, we expected individuals fed a high quality diet to increase their growth rate and avoid risky situations through the asset protection principle (Clark 1994; Luttbeg and Sih 2010; Figure 1b). For effects on trait covariance, we predicted a negative correlation between activity and predator cue response in both treatments as the patterns of effect of growth rates on these traits are opposite and symmetrical across diet quality treatments (Figure 1). We also predicted that due to the cost of plasticity, individuals fed a low quality diet would decrease the amount of behavioral plasticity (measured here as the residual within-individual variance $V_R$ — sensu Westneat et al. [2015]—although here, this measure conflates measurement error, organismal error, and plasticity). Finally, since individuals would be more constrained in terms of energy allocation we also expected a “ceiling effect” on among-individual variation ($V_M$) such that $V_M$ in body mass would be lower in the low quality diet treatment.

**MATERIALS AND METHODS**

**House cricket rearing and diet preparation**

We purchased 1-week old *A. domestica* from Fluker’s Farm (Port Allen, LA). Crickets were housed individually in 0.71-L containers and provided with *ad libitum* food and water, as well as pieces of egg carton for shelter. All individuals were maintained under a 12:12 h light:dark photoperiod. Individuals were fed a standard laboratory diet and allowed to habituate to their containers for a minimum of 7 days before being randomly assigned to either a high or low quality diet (Table 1) and beginning behavioral trials.

We varied diet quality by producing food pellets baked from high protein fish flakes (high quality diet, TetraMin Plus) and cellulose mixed with fish flakes (low quality diet, 2 parts cellulose to 1 part fish flakes). Thus, the nutritive quality of the high quality diet was 3 times higher than that of the low quality diet (Table 1). The use of non-nutritive cellulose also imposed gut limitations on
intake, preventing individuals in the low quality diet treatment from circumventing diet quality differences. Food pellets were produced by mixing ground fish flakes with water or cellulose and water and cooking the mixture in a Plexiglas mold at 60 °C for 12 h following Hunt et al. (2005). A preliminary study completed on a subset of 40 crickets (20 low quality and 20 high quality) showed substantial differences in growth rates between the 2 diets after 10 days of treatment (diet × day interaction, \( P < 0.05 \)) (Supplementary Figure 1). Because these crickets originate from captive populations, a direct comparison with their wild diet is difficult. However, the low quality diet treatment provided considerably lower energy than our standard laboratory diet (ground Purina chick starter chow, 2.88 cal/g, Purina Mills, St Louis, MO; Table 1).

### Table 1
**Nutritional composition of the low and high quality diets**

<table>
<thead>
<tr>
<th>Nutrient content</th>
<th>Low quality diet</th>
<th>High quality diet</th>
<th>Standard laboratory diet</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total energy (cal/g)</td>
<td>1.12</td>
<td>3.35</td>
<td>2.88</td>
</tr>
<tr>
<td>Protein (%)</td>
<td>15.33</td>
<td>45.00</td>
<td>18.00</td>
</tr>
<tr>
<td>Lipid (%)</td>
<td>3.66</td>
<td>11.00</td>
<td>3.00</td>
</tr>
<tr>
<td>Carbohydrate (%)</td>
<td>7.66</td>
<td>23.00</td>
<td>56.90</td>
</tr>
<tr>
<td>Non-nutritive fiber (%)</td>
<td>63.00</td>
<td>3.00</td>
<td>5.50</td>
</tr>
</tbody>
</table>

Behavioral tests

To test whether diet quality affected patterns of behavioral variance and covariance, we recorded individuals’ activity levels in a modified open-field arena (obstacle course assay) and response to diluted gecko excreta (response to a cue of predator response assay). Behavioral testing began within 1 week of arrival and testing occurred between 1 October 2014 and 3 March 2015. Due to logistical constraints, crickets were assayed in batches of 20 individuals with 10 individuals randomly assigned to either the high or low diet quality treatments. Five separate batches were sequentially reared and assayed and “batch” was included in all analyses (see below) to statistically account for any potential effects on average behavior. Behavioral trials for batches 1–5 began on 1 October 2014, 11 November 2014, 12 January 2015, 27 January 2015, and 6 February 2015, respectively. In order to minimize potential carry-over effects from exposure to cues of predator presence, the obstacle course assay was always conducted first followed by testing an individual’s predator cue response. We thoroughly cleaned each arena in between trials with 70% ethanol wipes to avoid chemical trace of conspecifics from influencing the behavior of subsequently tested individuals. We recorded mass to the nearest mg immediately after the predator cue response assay. This procedure was repeated twice every week over a 3-week period for a maximum of 6 behavioral measures per individual per assay. In total, we were able to record the behavioral responses of 98 individuals (low quality diet: \( n = 45 \), high quality diet: \( n = 53 \)) with a total of 446 individual observations (low quality diet: \( n = 198 \), high quality diet: \( n = 248 \); Supplementary Table 1). Due to mortality during the course of development, particularly in the low quality diet, only 35% of individuals survived until sexual characteristics were noticeable. We thus did not include sex in analyses.
Behavioral Ecology

Obstacle course activity
Activity was measured in a modified open field test, in which individuals had to navigate around multiple obstacles to explore the entire arena. The arena was 60 cm × 60 cm × 15 cm high, constructed of sealed plastic, and had a Plexiglas lid. This obstacle course behavioral protocol has previously been used with A. domestica to evaluate exploratory behavior (Doehrmann and Nelson 2014) and behavior-physiology correlations (Royauté et al. 2015). Individuals were left to rest for 30 s in a 5-cm-diameter container introduced into the lower right section of the arena (Z1, Supplementary Figure 2a). We then allowed the cricket to move freely through the arena for 220 s. We calculated the total distance travelled through the arena (in cm) with Ethovision X (Noldus Information Technology).

Response to cues of predator presence
Considerable research with a variety of cricket species has examined latency to emerge from shelter following disturbance under the assumption that this latency relates to antipredator behavior (e.g., Hedrick 2000; Kortet and Hedrick 2007; Hedrick and Kortet 2012; Niemela et al. 2015). Such latencies are ecologically reasonable measures of antipredator behavior, particularly for males, but are not the only means by which crickets assess or respond to potential predator presence. For example, over the last decade it has been repeatedly demonstrated that Gryllidae crickets respond to chemical cues of predators via either escape or avoidance behaviors (Kortet and Hedrick 2004; Storm and Lima 2008; Storm and Lima 2010). Response to chemical cues has likewise been used to measure response to potential predation threat in a variety of invertebrates and vertebrates (e.g., Herman and Valone 2000; Doehrmann et al. 2012; Nelson et al. 2013; Paterson et al. 2013).

The fact that crickets respond to chemical cues of predator presence should not be surprising as chemosensory cues are known to be important and often sufficient and/or necessary to illicit a variety of behavioral responses. The importance of chemical cues to crickets has been demonstrated, for example, in male–male agonistic interactions (Iwasaki and Katagiri 2008), in female assessment of male quality (Kortet and Hedrick 2005), self-reference to prevent repeated matings with the same individuals (Weddle et al. 2013), and, as mentioned above, in eliciting response to potential predator presence.

To measure response to cues of potential predator presence, here, we collected excreta from 2 subadult and 1 adult leopard gecko (Eublepharis macularius) that were fed a mixed diet of crickets (A. domestica) and mealworms (Tenebrio molitor). Leopard geckos were housed according to North Dakota State University IACUC standards (Protocol number: A14006). Collected excreta was ground weekly and diluted with deionized water (1 ml H2O:5 mg excreta). This solution was then applied to 15-cm-diameter Whatman filter paper discs (Supplementary Figure 2b) with a 5-cm-diameter central cutout that allowed crickets to be left to rest unexposed to the predatory cue. Each predatory disc was left to dry for a minimum of 2 h and was stored at 4 °C before trials. We inserted the predatory cue disc at the bottom of a 15-cm-diameter arena and left the cricket to rest for 30 s under a 5-cm-diameter cup in the nontreated central cutout. We then allowed the cricket to move freely for 220 s and estimated the distance travelled (in cm) through Ethovision. Previous experiments with this protocol showed that crickets had increased activity levels in presence of diluted gecko excreta compared to a water control (t = 2.05, P = 0.046; Supplementary Figure 3), thus greater activity during predator cue response trials was interpreted as greater responsiveness to predator cues.

Data analysis
All analyses were conducted in R 3.2.2 (R Development Core Team 2013) with the package MCMCglmm for Bayesian mixed models (Hadfield 2010).

Effect of diet quality on average trait value and trait repeatability
We estimated the effects of diet type and development on activity, response to cues of predator presence, and mass using Bayesian univariate mixed models. Mass was log10 transformed and all response variables were expressed as standard deviation units. The third batch of crickets had stronger mortality due to winter shipment and we included a 3-way interaction for diet type, day (centered) since placed on either the low or high quality diet, and batch number. Additional fixed effects included condition (injured or not), time of testing, temperature (expressed as among and within individual values; van de Pol and Wright 2009), and whether the cricket crawled under the filter paper during the predator cue response assay to control for potential confounds and “pseudo-repeatability” (Nakagawa and Schielzeth 2010; Westneat et al. 2011). Individual identity was included as a random effect and variance components were estimated separately by diet treatment to allow comparison of trait repeatabilities (Royauté et al. 2015a). We specified a Markov chain Monte Carlo (MCMC) chain with 1.3 × 10^6 iterations, 300,000 burn-in period and a thinning interval of 1000. We tested the influence of prior type on our results by comparing results from an inverse-Wishart prior to those estimated with maximum likelihood. Prior type had very little influence on our estimates and we present the results based on the inverse-Wishart prior.

MCMC approaches to fitting mixed models, like used above, do not allow classic tests of “significance.” Thus, to test for the “significance” of the diet × day × batch effects on body size and behaviors we employed 2 approaches. First, we specified 7 a priori models (Supplementary Table 3) of different ways in which our predictor variables might have affected body size or behaviors. The models we evaluated differed in the complexity of interactions included, ranging from a full model including the 3-way interaction of diet × day × batch, all 2-way interactions, and all main effects to reduced models that included single main effects. For these different models, we then compared their “Deviance Information Criterion values” (DIC). Models with lower DIC have better support and models with ADIC < 4 were considered as statistically equivalent. This procedure is analogous to the Likelihood Ratio tests for fixed effects performed with frequentist mixed models (Zuur 2009). Second, to assess the “significance” of particular terms we examined whether individual model terms as estimated had 95% credibility intervals that overlapped zero.

We calculated repeatability (τ) as the posterior mode of the between individual variance and τ within-individual (or residual) variance. We also calculated the MCMC posterior distribution of differences in repeatability between the (Δτ = τHighQual − τLowQual). This metric provides an estimate of the effect size for the difference in repeatability (Royauté et al. 2014, 2015a). Here positive values of Δτ indicate greater repeatability in the high quality diet treatment. “Significance” and inference of this difference was based on the proportion of posterior estimates that excluded zero. We repeated this procedure on each variance component (τHigh and τLow) to determine whether changes in repeatability were linked to changes in any specific variance component. Importantly, estimates of values
like Δr may differ from a comparison of r_{HighQual} − r_{LowQual} because the former is calculated directly from each MCMC iteration.

**Effect of diet quality on behavioral correlations**

We tested whether diet quality affected correlations among activity, predator cue response, and mass by specifying a multivariate mixed model, which was fit separately for each diet type. We included all 3 traits as response variables and used individuals as random effects. All fixed effects and model conditions were as described above. This procedure allowed us to estimate and compare among- and within-individual (Δr) correlation matrices between diet types (following Dingemanse and Dochtermann 2013). As above, we compared the magnitude of the difference in posterior estimates of correlation coefficients between the treatments (Δr = r_{HighQual} − r_{LowQual}) and base our inferences on the proportion of estimates excluding zero.

**RESULTS**

**Effect of diet quality on average trait value and trait variation**

Diet quality positively affected growth rate (diet × day interaction, estimate ± CI; β = 0.04 ± [0.03:0.05]) but had no effect on average activity (β = 0.00 ± [−0.05:0.05]) or response to cues of predator presence (β = 0.02 ± [−0.03:0.08]) (Figure 2, Supplementary Table 2). These inferences are consistent with our model comparison results which indicated similar patterns as none of the best models had substantially higher support than the null model for behavioral data (ΔDIC < 4, Supplementary Table 3). We did, however, detect a significant 3-way interaction between diet, day, and batch number on body mass, indicating that the positive effect of the high quality diet on growth rate varied among batches (Supplementary Table 3, Supplementary Figure 4).

The repeatability of response to cues of predator presence was lower in the high quality diet (low quality: r = 0.36 ± [0.21:0.53], high quality: τ = 0.16 ± [0.07:0.27], Δτ = −0.24, P = 0.01; Figure 3), but no significant differences were detected in repeatability for activity or mass (activity: Δτ = −0.02, P = 0.43; mass: Δτ = 0.03, P = 0.27; Figure 3, Supplementary Table 2). The observed changes in the repeatability of predator cue response were more strongly influenced by an increase in within-individual variation in the high quality diet than a decrease in among-individual variance (among-individual variance: ΔG_m = −0.18, P = 0.07; within-individual variance: ΔG_r = 0.27, P = 0.008; Figure 3 and Supplementary Figure 5). This suggests that individuals provided with a high quality diet were more inconsistent across repeated testing in their response to predator cues whereas individuals provided with a poor diet quality remained relatively consistent over the duration of the experiment.

**Effect of diet quality on behavioral correlations**

We found no evidence for a change in behavioral syndrome structure mediated by diet quality. Instead, activity and predator cue response were positively correlated with similar effect sizes in both treatments (low quality: r_0 = 0.39 ± [0.05:0.71], r_0 = 0.31 ± [0.18:0.47]; high quality: r_0 = 0.31 ± [0.02:0.64], r_0 = 0.40 ± [0.27:0.51], Figure 4). In the high quality diet, larger individuals had a greater predator cue response while this association was not detected in the low quality diet (i.e., body mass × predator cue correlation, low quality: r_0 = 0.01 ± [−0.41:0.33]; high quality: r_0 = 0.39 ± [0.02:0.62]; Δr_0 = 0.38, P = 0.04) (Figure 4, Supplementary Table 4).

**DISCUSSION**

Our aim here was to test whether the developmental environment experienced by individuals affects behavioral means, behavioral variation, and the covariance between activity and response to cues of predator presence. We did find evidence of diet manipulation affecting multiple levels of trait variation; however, those patterns often contradicted our theoretical predictions (Figure 1). Besides greater growth (and thus mass) for the high versus low quality diet, we predicted decreased activity and increased response to predator cues in the high versus low quality diets. We also had initially predicted that among-individual variation in mass would be lower in the low quality diet versus the high quality diet and that within-individual variation in behaviors would be higher in the high quality diet than the low, resulting in greater behavioral repeatability in the low quality diet. Although a higher diet quality increased cricket growth rate, we found surprisingly little effect on either

![Figure 2](image-url)

**Figure 2**

Effect of diet quality on behaviors and mass. Diet quality affected growth rate but not behavioral response. Values are presented on their original scales; mass data were fitted with loess smoothed curves. Black squares and dashed line: low quality diet; white circles and solid line: high quality diet.
average activity or response to predator cues. Further, we found decreased within-individual variation—evidence of lower behavioral plasticity in the low quality diet—but only for the response to cues of predator presence. Finally, the activity-predator cue response behavioral syndrome was not affected by diet quality but the sign of the correlation was opposite to that expected. This suggests that the developmental diet, as manipulated here, can have non-intuitive consequences on trait expression when hierarchical patterns of variation are taken into account.

As expected, increasing diet quality resulted in faster growth rates and thus greater body mass. However, this change in population growth rate did not have consequences on population average behaviors. This led us to generally reject our prediction that individuals in a low quality diet would follow the state-dependent safety principle while individuals in the high quality diet would follow an asset protection model (but see below). Instead, the influence of diet quality had stronger consequences on trait repeatability and covariance. That diet affected the magnitude of behavioral variation in the absence of a population-level behavioral change is particularly intriguing. Because individuals were maintained in their treatments for only 30 days, it is possible that the effects of diet quality manifest themselves quickly on individual variation but require longer exposure before population shifts are detected. In such a case, maintaining individuals on the different diet regimes over the entirety of development and maturity would be necessary to determine the long-term consequences of diet type on activity and response to cues of predator presence.

Crickets reared on a lower quality diet demonstrated a higher repeatability of their response to cues of predator presence but not their activity. As expected, plasticity in responses to cues of predator presence were favored only when individuals had access to sufficient nutritive resources (i.e., the high quality diet) and individuals under nutritive stress exhibited higher individual consistency in their responses to cues of predator presence, causing higher repeatability. In addition, the largest crickets from the high quality diet treatment reduced risk-taking by expressing a stronger response to cues of predator presence (Figure 4). This indicates partial support...
for the asset protection principle—wherein individuals that accumulate more assets over time (e.g., increase in body mass) preserve assets by reducing risky behaviors—despite a lack of general support for the principle herein.

These observations lead to 2 important insights on the influence of diet quality on behavioral variation: First, a higher diet quality and higher caloric content diet may alleviate the costs to behavioral plasticity by attenuating individual trade-offs. In contrast, individuals experiencing nutritive stress may face stronger allocation trade-offs due to limits on energy acquisition and a higher cost of switching behavioral responses. Second, changes in diet quality may have the potential to change the magnitude of state-dependent feedback loops (e.g., in this case the correlation between body mass and response to cues of predator presence between the 2 treatments), which in turn can have a profound influence on the maintenance of among-individual variation. Unfortunately, our experimental design only allowed us to test for the direct effect of growth rate on behavior through diet manipulation and not for recursive effects of behavior on growth rates. Recursive effects are expected to reduce the amount of personality differences over time, because individual protecting assets would acquire resources at a lower rate (Luttbeg and Sih 2010; Montiglio and Royauté 2014).

According to our predictions, we expected a negative among-individual correlation between activity and response to cues of predator presence. However, the traits were positively correlated. Because activity and response to cues of predator presence were both measured as distance moved through an arena, it is possible that a positive structural correlation will be present between measurements, over-riding more biologically relevant and interesting relationships. However, we view this explanation as insufficient given past experiments with this species. For example, movement rates of A. domestica measured in different arenas can be uncorrelated even if each behavior demonstrates repeatability (Dochtermann and Nelson 2014). Specifically, Dochtermann and Nelson (2014) found that activity (distance moved) measured in an obstacle course arena and activity (distance moved) in maze exploration trials were not correlated. A more plausible explanation for our results, therefore, is that the activity-predator cue response syndrome is robust to changes in the permanent environment as the among-individual correlations did not vary between low and high quality diet treatments (Figure 4). This general inference holds even if we were assaying activity in different contexts (e.g., open field trials: “exploration of a control environment,” predator cue trials: “exploration of an altered environment”; sensu Dingemanse et al. 2007). That is, even if considering activity in the presence and absence of predators as separate contexts, we still demonstrated that the among-individual correlation was robust to permanent environmental differences.

Among-individual correlations, that is, behavioral syndromes (Dingemanse et al. 2012), are generated by both additive genetic correlations and permanent environment correlations (Dingemanse and Dochtermann 2013; Dingemanse and Dochtermann 2014). In our study, additive genetic correlations should have been the same between the 2 treatments as individuals were randomly assigned to the 2 diet types. Any differences in among-individual correlations between the 2 treatments are therefore attributable to changes in permanent environmental correlations or $G \times E$ effects (Han and Dingemanse 2015). The fact that the activity-predator cue response syndrome was unchanged suggests 2 alternative explanations: First, this syndrome might be primarily underpinned by genetic correlations. Second, diet quality may have little contribution to the permanent environment correlation and other environmental cues (e.g., temperature, predator presence, and maternal effects) could be more influential. We view this latter explanation as less likely because of the strong influence of diet quality on life-history trajectories (Tatar and Carey 1995; Houslay et al. 2015), morphological (e.g., Braendle et al. 2006), physiological (e.g., Cruz et al. 2004), and behavioral traits (Akman et al. 2004; Wilder and Rypstra 2008; Tremmel and Muller 2013) that has been demonstrated in a considerable number of species, including crickets (Hunt et al. 2004; Hunt et al. 2005; Hunt et al. 2006; Zajitschek et al. 2009; Zajitschek et al. 2012).

As mentioned, there has been extensive research demonstrating the effects of diet and condition on behavior in crickets (see above). Moreover, beyond just general condition or energy availability, attention to particular stoichiometric relationships has highlighted subtle differences in the effects of diet on behavior (e.g., Bertram et al. 2009; Han et al. 2016; reviewed by Han and Dingemanse [2015]). Our results are particularly interesting even with this existing literature for several reasons. For example, despite a major difference in dietary value (Table 1), no mean effects on behavior were statistically detectable. This lack of detectable effects is surprising not only because of the existing literature but also given the clear predictions available from conceptual and theoretical frameworks (e.g., the asset protection principle; Luttbeg and Sih 2010). Moreover, our approach of partitioning variation and explicitly estimating among-versus within-individual variation and correlations revealed that, despite a lack of mean effects, within-individual variation in response to cues of predator presence differed between the 2 diet treatments. Most previous manipulations of diet, condition, and state have largely failed to explicitly estimate these types of effects and thus have failed to detect the effect noted here (but see Han et al. [2016] for a notable exception). More generally, our study shows that the outcomes of state manipulation on behavioral variation and behavioral syndromes are not obvious and may fail to confirm theoretical predictions. Indeed, while we found partial support for the asset protection principle regarding predator cue response in the high quality diet, the changes in behavioral variance were more due to an increase in behavioral plasticity than to a decrease in among-individual variance contrary to our predictions. It remains unclear whether manipulating state generally has stronger effects on behavioral averages, behavioral variances, or on behavioral syndromes.

SUPPLEMENTARY MATERIAL

Supplementary material can be found at http://www.beheco.oxfordjournals.org/

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Data accessibility: Analyses reported in this article can be reproduced using the data provided by Royauté and Dochtermann (2016).

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REFERENCES


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