Ballistic Tongue Projection in Chameleons Maintains High Performance at Low Temperature

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Environmental temperature impacts the physical activity and ecology of ectothermic animals through its effects on muscle contractile physiology. Sprinting, swimming, and jumping performance of ectotherms decreases by at least 33% over a 10 °C drop, accompanied by a similar decline in muscle power. We propose that ballistic movements that are powered by recoil of elastic tissues are less thermally dependent than movements that rely on direct muscular power. We found that an elastically powered movement, ballistic tongue projection in chameleons, maintains high performance over a 20 °C range. Peak velocity and power decline by only 10%–19% with a 10 °C drop, compared to >42% for nonelastic, muscle-powered tongue retraction. These results indicate that the elastic recoil mechanism circumvents the constraints that low temperature imposes on muscle rate properties and thereby reduces the thermal dependence of tongue projection. We propose that organisms that use elastic recoil mechanisms for ecologically important movements such as feeding and locomotion may benefit from an expanded thermal niche.

Tongue projection and retraction in chameleons are powered by the tongue accelerator muscle (7). This mechanism is common to all chameleons and gives these slow, cryptic, sit-and-wait predators the element of surprise. Chameleons feed over a wide range of T\textsubscript{b}, including 0 °C during cold snaps (8). No significant effect on projection or retraction performance.

Ballistic tongue projection in chameleons achieves its extreme performance by rapid elastic recoil of collagen tissue within the tongue—tissue that is first stretched by slow contraction of the tongue accelerator muscle (7). This “bow and arrow” mechanism decouples muscle contraction temporally from tongue launch and thereby allows kinetic energy to be imparted to the tongue at a rate far exceeding that possible via direct muscle contraction (7). Once launched—at accelerations exceeding 400 ms\textsuperscript{−2} (41 g)—the tongue travels to the target on its momentum alone and then adheres to the prey. Tongue retraction relies on neither ballistic launch nor elastic recoil to bring prey to the mouth, but rather is driven by continuous contraction of the lengthy hyoglossus muscle (8).

The differing mechanisms of tongue projection and retraction in chameleons provide an opportunity to evaluate the hypothesis that the elastic-recoil mechanism confers low thermal dependence to tongue projection. We tested whether elastically powered tongue projection has a lower thermal dependence than nonelastic tongue retraction by examining the effects of temperature on performance parameters of these two movements. In addition, we propose that our findings can be generalized to explosive ballistic movements in other ectotherms, and that elastic-recoil mechanisms may serve to expand the thermal niche of ectotherms that use them for critical movements.

Results

Veiled chameleons (Chamaeleo calyptratus) were able to project the tongue and capture prey across the same range of distances regardless of temperature (15 °C–35 °C). Overall, projection distances ranged from 6.6 cm to 19.6 cm. Individual average projection distances ranged from 10.4 cm to 14.2 cm, with an overall average of 12.5 cm. No significant effect of temperature on prey distance, tongue projection distance, or tongue overshoot distance was found.

Inverse dynamic analysis of tongue movements revealed that as temperature increased, performance increased significantly (Table 1) for both tongue projection and retraction. Nonetheless, peak performance measures of ballistic tongue projection were maintained at a high level at all temperatures (Table 2). At the low end of our experimental T\textsubscript{b} range (15 °C), peak projection velocity averaged 3.4 ms\textsuperscript{−1}, peak acceleration averaged 357 ms\textsuperscript{−2}, and peak power averaged 1,892 W kg\textsuperscript{−1}. At 35 °C, values were somewhat higher: peak velocity averaged 4.4 ms\textsuperscript{−1}, peak acceleration averaged 433 ms\textsuperscript{−2}, and peak power averaged 2,900 W kg\textsuperscript{−1}. In contrast, performance parameters of retraction increased markedly at higher temperature. At 15 °C, peak velocity averaged 0.8 ms\textsuperscript{−1}, peak acceleration averaged 170.3 ms\textsuperscript{−2}, and peak power averaged 34.4 W kg\textsuperscript{−1}, whereas at 35 °C, peak velocity averaged 1.9 ms\textsuperscript{−1}, peak acceleration averaged 478 ms\textsuperscript{−2}, and peak power averaged 453 W kg\textsuperscript{−1} (Table 2). The average power of projection also was maintained at a high level, averaging 1,092 ± 78 W kg\textsuperscript{−1} at 15 °C (mean ± SE) and 1,911 ± 156 W kg\textsuperscript{−1} at 35 °C. The order of experimental temperatures experienced by an individual had no significant effect on projection or retraction performance.

Although tongue projection and retraction both showed effects of temperature, retraction showed a significantly stronger effect. For each 10 °C increment in temperature between 15 °C and 35 °C, a
Table 1. Results from repeated-measures ANCOVA examining the performance parameters peak velocity, peak acceleration, and peak power for effects of temperature, individual, feeding phase (projection vs. retraction), and projection distance (covariate)

<table>
<thead>
<tr>
<th></th>
<th>Peak velocity</th>
<th></th>
<th>Peak acceleration</th>
<th></th>
<th>Peak power</th>
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<tr>
<td></td>
<td>df</td>
<td>F value</td>
<td>P value</td>
<td>df</td>
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<td>P value</td>
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<tr>
<td>15 °C vs. 25 °C</td>
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<td>Individual</td>
<td>4</td>
<td>7.384</td>
<td>&lt;0.0001*</td>
<td>4</td>
<td>5.585</td>
<td>0.0003*</td>
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<tr>
<td>Temperature</td>
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<td>&lt;0.0001*</td>
<td>1</td>
<td>77.56</td>
<td>&lt;0.0001*</td>
</tr>
<tr>
<td>Projection distance</td>
<td>1</td>
<td>44.86</td>
<td>&lt;0.0001*</td>
<td>1</td>
<td>5.477</td>
<td>0.0205</td>
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<tr>
<td>Phase</td>
<td>1</td>
<td>8118</td>
<td>&lt;0.0001*</td>
<td>1</td>
<td>294.8</td>
<td>&lt;0.0001*</td>
</tr>
<tr>
<td>Individual × tempera</td>
<td>4</td>
<td>2.58</td>
<td>0.0395</td>
<td>4</td>
<td>2.439</td>
<td>0.0493</td>
</tr>
<tr>
<td>Individual × projec</td>
<td>4</td>
<td>1.665</td>
<td>0.1609</td>
<td>4</td>
<td>0.697</td>
<td>0.5952</td>
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<td>0.8633</td>
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<td>0.0157*</td>
<td>4</td>
<td>5.433</td>
<td>0.0004*</td>
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<tr>
<td>Temperature × phase</td>
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<td>1</td>
<td>73.86</td>
<td>&lt;0.0001*</td>
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<td>Projection distance × phase</td>
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<td>0.0126*</td>
<td>1</td>
<td>30.84</td>
<td>&lt;0.0001*</td>
</tr>
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</table>

Note the significant temperature × phase interaction effects, which indicate that tongue projection and tongue retraction are affected differently by changes in temperature.

*Significant difference in ANCOVA at Benferroni-corrected α = 0.017, indicating significant effect.

Discussion

Remarkably, C. calyptratus achieved extremely high-performance tongue projection even when cold. At a T<sub>b</sub> of 15 °C, time-averaged muscle-mass–specific power output averaged 1,092 W kg<sup>-1</sup>, and peak instantaneous muscle-mass–specific power output during projection averaged 1,892 W kg<sup>-1</sup>. This peak value is well in excess of peak power output of muscle tissue during active contraction as measured or estimated in other vertebrates operating at higher T<sub>b</sub>, including flying quail during vertical takeoff (1,121 W kg<sup>-1</sup>) (12), sprinting lizards (952 W kg<sup>-1</sup>) (13), and jumping frogs (373 W kg<sup>-1</sup>) (6). High power outputs for rapid movements using the elastic-recoil mechanism, including jumping in bushbabies (14) and insects (15, 16), predatory strikes of mantis shrimp (17), and tongue projection in salamanders (18) and chameleons (7), have been documented in numerous kinematic studies; little focus has been given to the maintenance of performance at low T<sub>b</sub>, however.

The Q<sub>10</sub> values for tongue projection (1.1–1.3; Fig. 1) are well below the Q<sub>10</sub> values of contractile rate properties of isolated muscles and of other dynamic behaviors, which generally exceed 1.5 (1–6). This degree of temperature independence is similar to that of static contractile muscle properties, such as maximum isometric tetanic tension (3, 6), and of static behaviors, such as exertion of peak bite force (1); however, the extent of temperature dependence on tongue retraction (Q<sub>10</sub> = 1.7–2.9; Figs. 1 and 2) resembles that of contractile rate properties of isolated muscles and of dynamic behaviors, such as sprinting (1–6). Jump distance in frogs, for example, exhibits a Q<sub>10</sub> value of 1.6 over 14 °C–25 °C, and the power generated by the muscles activated during jumping has a Q<sub>10</sub> value of 2.7 (5). Similarly, sprint speed in lizards has an average Q<sub>10</sub> value of 1.5 at temperatures below the estimated optimal temperature (2).

The contrasting thermal dependence of tongue projection and retraction (Fig. 3 and Movie S1) supports the hypothesis that the low thermal dependence of tongue projection in chameleons is due to the elastic-recoil mechanism, in which temperature-dependent muscle shortening occurs during the loading phase before tongue launch, and is temporally decoupled from the temperature-independent elastic recoil of connective tissue that powers ballistic tongue projection. This mechanism not only endows chameleons with spectacular performance, but also liberates projection from the
constraints on muscle rate properties imposed by low temperature. Thus, the thermal dependence of the contractile rate properties of the tongue accelerator muscle need not be unusually low to maintain high performance at low temperature. In contrast, tongue retraction declines at low temperature, because it relies on direct muscle power output, which is thermally dependent. Projection performance depends instead on peak muscle tension and the elastic modulus of collagen, both of which show low thermal dependence or complete thermal independence (3, 5, 6, 19). Peak isometric muscle tension typically exhibits $Q_{10}$ values of 1.0–1.2 (5), and the load–strain relationship of collagenous tendon exhibits a $Q_{10}$ of 1 across the large physiological temperature range of 0 °C–37 °C (19). Studies of other animal systems that use elastic structures to power movements lend additional support to the conclusion that elastic-recoil mechanisms confer relative thermal independence compared with movements that rely on muscle rate properties. Among ballistic systems, jumping in frogs is powered partially by recoil of in-series elastic elements that supplements muscle power output (20). Frog jumping appears to show a reduced effect of temperature on performance (5), but it is not liberated to the same extent as tongue projection in chameleons, probably because elastic recoil and muscle contraction overlap temporally (20). Among cyclical systems, wingbeat frequency of beetles shows very low temperature sensitivity, apparently because frequency is determined by the resonant frequency of the flight system, which is dictated by its physical properties rather than by its muscle rate properties (21).

Because the mechanical properties of elastic tissues are known to have low thermal sensitivity (19, 22, 23), temperature manipulation may be a valuable methodological approach to test for the presence or prevalence of elastic recoil in powering movements. Elastic recoil is implicated if performance of a movement is maintained at a high level over a wide range of body temperatures. Our findings on chameleons thus serve as independent validation for the presence of an elastic-recoil mechanism in tongue projection.

Finally, chameleons have increased the thermal breadth of their feeding mechanism by decreasing the temperature effects on performance of ballistic tongue projection and thus are able to feed at very low $T_b$ (9–11, 24, 25). This ability likely grants them an expanded thermal niche, allowing them to feed early in the morning when effective thermoregulation is not possible (10) and enabling them to be active over a wider temperature range than other sympatric lizard species (11). The ability of chameleons to forage at low temperatures also may reduce thermoregulatory behavior and its ecological costs (26). Other ectothermic organisms that use explosive, ballistic movements for prey capture or locomotion across a range of temperatures may similarly benefit from the relative thermal independence of elastic recoil mechanisms.

Materials and Methods

Five Chamaeleo calyptratus (12.5–14.0 cm snout–vent length) were imaged at 3 kHz at a $T_b$ of 15 °C, 25 °C, and 35 °C while feeding on crickets at a range of distances, using a Photron Fastcam high-speed digital camera. Crickets were placed on a square of insect screen suspended vertically from above by thread. This “cricket trapeze” allowed the chameleon’s tongue to complete its trajectory naturally without being stopped by an immovable target, and thus permitted examination of performance and physiological parameters at a range of actual tongue projection distances.

To control $T_b$ after an acclimation period of at least 1 h, imaging trials were conducted in an environmental chamber set to the experimental $T_b$. Supplemental lighting was switched on immediately before tongue projection and turned off immediately after tongue retraction to prevent elevation of body temperature through light source radiation. During the prey reduction phase, immediately after tongue retraction, $T_b$ was verified orally using a calibrated Sixth Sense LT300 infrared thermometer (± 1 °C accuracy). Only feeding sequences with a postfeeding $T_b$ of the target experimental temperature ± 1 °C were included in the analysis.

Ten feeding sequences were collected from each of four individuals at each experimental $T_b$, for a total of 120 feedings. Five feeding sequences from a fifth individual were collected at each experimental $T_b$ before this animal was removed from the experiment due to illness. Between one and five feeding events were collected per individual at each feeding session. The sequence of experimental $T_b$ for each individual was selected randomly, and no two animals were exposed to an identical $T_b$ sequence. To account for natural variation in the distance between the prey and the chameleon’s snout because of the distance that the chameleon leaned its body forward off the perch for any given feeding event, distance to the “cricket trapeze”...
was varied within a normal range of projection distances. Thus, feedings were collected over an 8-20 cm range of tongue projection distances. Distance to the prey was adjusted to elicit maximal tongue projection length for each individual at each experimental Tp. Effects of temperature on preprojection distance to the target and overshoot distance of the tongue beyond the target were examined using repeated-measures ANOVA.

An inverse-dynamics approach was used to compute the instantaneous velocity, acceleration, and power of tongue projection and retraction. Using National Institutes of Health Image J software (http://rsb.info.nih.gov), the distance of tongue projection for each scale-calibrated feeding sequence was recorded. Image J software was used to record the x,y coordinates of the tip of the tongue on each frame throughout the tongue projection sequence. Using a custom script for the P-Spline package of R statistical software (R Project for Statistical Computing), a quintic spline was fitted to the position trace of the tongue and smoothed to remove secondary oscillation artifacts from the first and second derivatives of position. From these smoothed position data, instantaneous velocity (m s⁻¹) and acceleration (m s⁻²) (i.e., first and second derivatives of the position) were calculated. For tongue retraction, coordinates of four positions along the length of the retractor muscle were recorded on each frame of the retraction sequence. These coordinate data were used to quantify the length of the retractor muscle in each frame, and this length was then used to compute the length change through the retraction sequence. These length data were then smoothed and subjected to the same inverse dynamics analysis as the tongue projection position data. Mass-specific power (in W kg⁻¹) was calculated as the product of velocity and acceleration (7) and corrected for the mass of the active muscle in each phase. As in other species (7), dissection and mass measurements of the tongue apparatus of seven C. calyptratus (12.0–15.5 cm snout–vent length) determined that the circular portion of the accelerator muscle accounts for ~50% (mean, 48.2% ± 2.9%) of the mass of the accelerator muscle complex and tongue pad, whereas the retractor muscle accounts for ~25% (mean, 25.8% ± 1.7%) of the mass of the accelerating muscle complex, tongue pad, and retractor muscle. Thus, mass-specific power is multiplied by a factor of 2 for projection and by a factor of 4 for retraction (7).

To examine the effects of temperature on performance, tongue projection distance, peak velocity (m s⁻¹), peak acceleration (m s⁻²) and peak mass specific power (W kg⁻¹) for both tongue projection and retraction were computed for each feeding sequence. Performance was log-transformed and examined for effects of temperature (fixed effect), phase of feeding (fixed effect), and individual (random effect) using repeated-measures ANCOVA with projection distance as a covariate. The temperature × phase interaction term of the model allowed us to examine whether tongue projection and tongue retraction responded differently to temperature changes. In addition, the influence of experimental temperature sequence on performance was assessed using repeated-measures ANOVA to test for an effect of previous temperature on the projection distance residuals of each performance parameter.

Least squares regression of performance parameters during both projection and retraction, with projection distance as the independent variable, was performed for each individual at each temperature. The interpolated value of each performance parameter at the overall average projection distance (12.5 cm) was calculated for each individual and used to calculate temperature coefficient (Q10) values using the equation $Q_{10} = (R_t/R_d)(10^{(t_1-t_d)/10})$, where $R_t$ and $R_d$ are the interpolated performance values at temperatures $t_1$ and $t_d$, respectively, and $t_1$ is greater than $t_d$. The $Q_{10}$ values for each individual were then used to calculate an average $Q_{10}$ value with SE.

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