

## RESEARCH ARTICLE

# Thermal effects on the dynamics and motor control of ballistic prey capture in toads: maintaining high performance at low temperature

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### SUMMARY

Temperature has a strong influence on biological rates, including the contractile rate properties of muscle and thereby the velocity, acceleration and power of muscle-powered movements. We hypothesized that the dynamics of movements powered by elastic recoil have a lower thermal dependence than muscle-powered movements. We examined the prey capture behavior of toads (*Bufo terrestris*) using high speed imaging and electromyography to compare the effects of body temperature (11–35°C) on the kinematics, dynamics and motor control of two types of movement: (1) ballistic mouth opening and tongue projection, which are powered by elastic recoil, and (2) non-ballistic prey transport, including tongue retraction and mouth closing, which are powered directly by muscle contraction. Over 11–25°C, temperature coefficients of ballistic mouth opening and tongue projection dynamics ( $Q_{10}$  of 0.99–1.25) were not significantly different from 1.00 and were consistently lower than those of prey transport movements ( $Q_{10}$  of 1.77–2.26), supporting our main hypothesis. The depressor mandibulae muscle, which is responsible for ballistic mouth opening and tongue projection *via* the recovery of elastic strain energy stored by the muscle prior to the onset of the movement, was activated earlier and for a longer duration at lower temperatures ( $Q_{10}$  of 2.29–2.41), consistent with a slowing of its contractile rates. Muscle recruitment was unaffected by temperature, as revealed by the lack of thermal dependence in the intensity of activity of both the jaw depressor and jaw levator muscles ( $Q_{10}$  of 0.754–1.12). Over the 20–35°C range, lower thermal dependence was found for the dynamics of non-elastic movements and the motor control of both elastic and non-elastic movements, in accord with a plateau of high performance found in other systems.

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Key words: feeding, tongue projection, *Bufo*, amphibian, elastic, electromyography.

### INTRODUCTION

A critical environmental challenge faced by organisms is the influence of temperature on physiological and biochemical processes (Cossins and Bowler, 1987; Hochachka and Somero, 2002; Huey and Kingsolver, 1989). Environmental temperature is especially significant for ectothermic animals because it directly affects body temperature and thereby physiological rate processes such as metabolic rate and muscle contraction velocity. The decline of these rates with decreasing temperature limits an organism's ability to effectively carry out critical behaviors at low environmental temperatures (Bennett, 1985; Hill et al., 2008).

Among the various physiological processes affected by temperature changes, the rate properties of muscle contraction have perhaps the most immediate impact upon an organism's ability to move and engage in behaviors such as foraging, predator evasion and courtship. Physiologists have long recognized that rate properties of muscle contraction – e.g. rate of tension development, maximum shortening velocity and peak power output – decline at low temperatures, dropping by at least half with each 10°C drop, i.e.  $Q_{10} > 2$  (where the temperature coefficient  $Q_{10}$  is the factor by which a rate increases with a 10°C increase in temperature). This marked decline has been found in a diversity of muscles from a vast array of animals, including vertebrates and invertebrates, endotherms and ectotherms (Putnam and Bennett, 1982; Bennett, 1984; Bennett,

1985; Hirano and Rome, 1984; Renaud and Stevens, 1984; Else and Bennett, 1987; John-Alder et al., 1989; Faulkner et al., 1990; Stevenson and Josephson, 1990; Barnes and Ingalls, 1991; Rome et al., 1992a; Rome et al., 1992b; Swoap et al., 1993; Asmussen et al., 1994; Sobol and Nasledov, 1994; Altringham and Block, 1997; Choi et al., 1998; Donley et al., 2007; Herrel et al., 2007).

The thermal dependence of these contractile rate properties strongly influences dynamic behaviors such as locomotion. The top speed at which a lizard can chase prey and the distance a frog can jump from a pursuing predator, for example, have been shown to be reduced by low body temperature (Huey and Stevenson, 1979; Bennett, 1984; Bennett, 1990). The reduction is not as great as that of the rate properties of isolated muscle, but it is nonetheless significant. For example, sprint velocity in the lizard *Trapelus* has a  $Q_{10}$  of 1.8 between 21 and 38°C (Herrel et al., 2007), and maximum jumping distance in *Rana* has a  $Q_{10}$  of 1.6 over 14–25°C (Hirano and Rome, 1984). Significant thermal dependence has been found in locomotor speed and frequency of oscillatory movements, including salamander locomotion, frog swimming, jumping and calling, lizard running and fish swimming, among others (Hirano and Rome, 1984; Marsh and Bennett, 1985; van Berkum, 1986; Else and Bennett, 1987; Huey and Bennett, 1987; John-Alder et al., 1988; John-Alder et al., 1989; Rome et al., 1990; Bauwens et al., 1995; Lutz and Rome, 1996; Altringham and Block, 1997;

Peplowski and Marsh, 1997; Navas et al., 1999; McLister, 2001; Marvin, 2003a; Marvin, 2003b; Herrel et al., 2007).

Many studies that demonstrate strong thermal dependence of contractile muscle dynamics also reveal minimal to absent thermal effects on static contractile properties such as maximum isometric twitch and tetanic tension, with  $Q_{10}$  values (i.e.  $R_{10}$  values, where  $R_{10}$  is equivalent to  $Q_{10}$  but is applied only to non-rate properties) of about 1.0 and sometimes lower. Tetanic tension remains approximately constant across a range of temperatures, while twitch tension may even increase slightly as temperature drops (Hill, 1951; Bennett, 1984). The thermal independence of static contractile properties of muscle suggests that behaviors for which performance relies upon the strength of isometric muscle contractions will themselves be relatively thermally independent. In fact, peak bite force, an ecologically relevant performance parameter in defense and feeding, has been shown in the lizard *Trapelus* to be largely independent of temperature between 20 and 38°C; peak tetanic force of a major mouth closing muscle in this species also shows almost complete thermal independence over this range (Herrel et al., 2007).

Ballistic movements such as jumping and tongue projection in many species do not rely on rapid muscle contraction or high muscle power output, but instead are powered by rapid recoil of elastic structures that are first loaded by much slower muscle contractions (Bennet-Clark, 1976a; Bennet-Clark, 1976b; Roberts and Azizi, 2011). In a classic example, jumping fleas load a resilin pad using the femoral depressor muscle during the preparatory phase prior to the jump, and allow the pad to recoil against isometrically contracted muscle to extend the legs and launch the flea into the air (Bennet-Clark and Lucey, 1967). Storing muscle energy in elastic structures enables the power of the jump to far exceed the power output of the muscle in a variety of animals including fleas, locusts, beetles and bushbabies, as the muscle energy is released more quickly than it was stored (Bennet-Clark, 1975; Aerts, 1998). For example, muscle power can be amplified 1000 times in the jump of the click beetle (Bennet-Clark, 1976a). Much like jumping in insects, ballistic tongue projection in chameleons and salamanders involves amplification of muscle power (10–100 times) *via* storage of energy in collagen fibers within the projector muscles (de Groot and van Leeuwen, 2004; Deban et al., 2007). Rapid release of energy from elastic structures allows ballistic movements to be among the most dynamically extreme of all animal movements in the acceleration and mechanical power that is achieved.

In addition to this high acceleration and power output, another potential benefit of elastic recoil mechanisms, which we examine in this study, is decreased thermal dependence of movement. Ballistic movements that are powered by rapid recoil of elastic structures loaded by slow or near-isometric muscle contraction may have low thermal dependence. By temporally decoupling muscle contraction from movement, ballistic systems like jumping and tongue projection may circumvent the influence of temperature on muscle performance that many locomotor systems experience. Through its effects on muscle dynamics such as time to peak tension, low temperature is expected to slow the rate of loading of elastic structures and to prolong muscle activity, but because peak isometric tension can be thermally independent, temperature may not alter the amount of energy that is stored in elastic structures. Further, stiffness of elastic tissues in animals – collagen, resilin and other animal protein rubbers – shows either very low thermal dependence ( $Q_{10}$  of 1.03–1.11) (Alexander, 1966; Denny and Miller, 2006) or complete thermal independence (Rigby et al., 1959). Therefore, temperature is not expected to affect the rate of recoil of elastic structures or, in turn, the performance of the movement. Consistent

with this reasoning, elastically powered ballistic tongue projection in chameleons displays relative thermal independence with  $Q_{10}$  values of less than 1.3, which contrasts with the strong thermal dependence of muscle-powered tongue retraction (Anderson and Deban, 2010).

Independently from chameleons, toads of the genus *Bufo* have evolved ballistic prey capture that makes use of elastic recoil to power rapid mouth opening and tongue projection (Nishikawa, 2000; Lappin et al., 2006). In a mechanism called inertial elongation, the tongue is whipped from the mouth by the accelerating mandible to which it is attached anteriorly; as the tongue rotates, or flips, over the mandible tip it also elongates to over 200% of resting length (Nishikawa, 2000). Activation of the bilaterally paired mouth opening muscle, m. depressor mandibulae (DM), for up to 250 ms prior to mouth opening and deactivation of the muscle at the start of mouth opening temporally separates muscle contraction from the ballistic mouth opening movement, and thus allows this movement to achieve extremely high acceleration and mechanical power. Power of ballistic mouth opening in *Bufo alvarius* reaches a peak of 9600 W kg<sup>-1</sup> of DM mass, far beyond what the muscles can produce directly. The tongue is then retracted by contraction of the hyoglossus muscle, transporting the prey into the mouth. Prey transport is accompanied by further mouth opening followed by mouth closing after the tongue is withdrawn. The prey capture behavior of *Bufo* is an ideal system for examining temperature effects because it comprises both elastically powered movements (mouth opening and tongue projection) and muscle-powered movements (tongue retraction and mouth closing).

We examined feeding in *Bufo terrestris* to test several hypotheses regarding temperature effects on the dynamics and motor control of both ballistic and non-ballistic movements. We hypothesized that the elastic recoil mechanism of ballistic mouth opening of toads liberates the movement from thermal effects on muscle contractile rates, and as a consequence ballistic prey capture (i.e. mouth opening and tongue projection) in *Bufo* should show thermal independence ( $Q_{10} \approx 1$ ). We also expected that the duration of activation of the mouth opening muscles would show thermal dependence ( $Q_{10} \approx 2$ ), commensurate with typical contractile rate properties and the assumption that the muscles achieve the same peak tension at all temperatures. We predicted, however, that the intensity of muscle activation would be maximal at all temperatures. Further, we hypothesized that non-ballistic movements associated with prey transport, such as tongue retraction and mouth closing, would show thermal dependence ( $Q_{10} \approx 2$ ). To test these hypotheses we performed kinematic and dynamics analyses of high speed image sequences, and corresponding electromyographic recordings of the main mouth opening and closing muscles, from prey capture events occurring across a range of body temperatures (11–35°C).

## MATERIALS AND METHODS

Four *B. terrestris* Bonnaterra 1789 (3.9–8.2 cm snout–vent length) were collected locally in Tampa, FL, USA, housed individually in plastic boxes at 22°C and maintained on a diet of crickets. All procedures in this study were approved by the Institutional Animal Care and Use Committee of the University of South Florida.

### Electromyography

Bipolar hook electrodes were constructed from Formvar-coated, 0.025 mm diameter nichrome wire (A-M Systems #7615, Sequim, WA, USA). Electrodes were made of two strands of wire 50–75 cm long, glued together at their ends with veterinary-grade

cyanoacrylate. Hook electrodes were made by threading the wire through a 30 gauge hypodermic needle, removing the insulation from the glued tips of the wires, and bending the two strands away from each other at the ends.

Prior to electrode implantation, toads were anesthetized by immersion in a  $1\text{g l}^{-1}$  buffered aqueous solution of MS-222 (3-aminobenzoic acid ethyl ester; Sigma, St Louis, MO, USA) for 10–30 min. For surgery each toad was positioned on its left side on moist paper towels on the stage of a dissecting microscope (Wild Heerbrugg M5 Stereomicroscope, Leica Microsystems GmbH, Wetzlar, Germany). Using 30 gauge hypodermic needles, electrodes were implanted through two incisions in the skin of the right side of the head into the DM, which originates on the posterior aspect of the cranium and inserts onto the retroarticular process of the lower jaw, and into the m. levator mandibulae posterior longus (LM), which originates on the dorsal and lateral aspects of the cranium and inserts onto the dorso-medial surface of the mandible rostral to the jaw joint. As in other non-mammalian tetrapods the DM functions as a first-order lever to open the mouth and the LM (i.e. adductor mandibulae) functions as a third-order lever to close the mouth. In toads the tympanum serves as an external landmark that widely separates the belly of the DM from that of the LM [see fig. 2 of Lappin et al. (Lappin et al., 2006)].

After electrode implantation the needles were withdrawn leaving the wire held in place by the hooks of the electrode tip. Incisions were closed with polypropylene monofilament suture with suture knotted around each electrode pair where it exited the incision, and the electrodes were bonded to the suture with veterinary-grade cyanoacrylate. Electrode wires from both recording sites were bundled together, sutured to the skin of the side of the head and at the dorsal midline, bonded to the sutures with cyanoacrylate, and all glued together along their length with modeling cement. The ends of the wires were stripped and soldered to a 40-pin plug, which mated with a socket on the amplifier probe.

Electromyographic (EMG) signals were amplified 2000 times using a 16-channel differential amplifier (A-M Systems 3500) and filtered to remove 60 Hz line noise. Conditioned signals were sampled at 4 kHz with a PowerLab 16/30 analog-to-digital converter coupled with LabChart software version 6 (AD Instruments Pty Ltd, Town, New South Wales, Australia) running on a Dell laptop PC. EMG recordings were synchronized with digital images *via* a trigger shared with the camera. Clean EMG signals were obtained from both muscles of all four individuals for all feedings.

### Feeding experiments

After recovery from surgery (12–24 h), toads were imaged at 6 kHz frame rate and 1/12,000 s shutter speed with a Fastcam 1024 PCI camera (Photron USA Inc., San Diego, CA, USA) as they captured crickets and beetles. All feeding trials and recordings were conducted within 3 days of surgery. Toads were positioned individually on a feeding stage of moistened foam rubber within a glass aquarium. Toads moved about unrestrained on the feeding stage and prey crickets and beetles were positioned at various distances in front of the toads to obtain lateral camera views of the toads during prey capture bouts. Because of movements of the toads within the imaging area, of the 109 feedings recorded, 65 yielded kinematic timing and EMG data, and a subset of 40 yielded the additional distance data necessary for the calculation of dynamics parameters. Only feedings in which toads were judged during the experiment to be oriented within approximately 15 deg of the focal plane of the camera were used for this latter subset. A cm scale was imaged in the same plane as the toad following each feeding to calibrate distances.

Feeding trials were conducted across a range of ambient temperatures (10–38°C) within an environmental chamber (Environmental Growth Chambers, Chagrin Falls, OH, USA). These temperatures were selected to encompass a range in which feedings could be reliably elicited, to be within the toads' thermal tolerance, and to span a range wide enough to detect potentially subtle effects of temperature. Each toad experienced each ambient temperature once as well as a unique sequence of ambient temperatures with regard to the direction and magnitude of temperature change. The temperature of the toad's head was measured immediately after each feeding using an infrared thermometer (Sixth Sense LT300, Williston, VT, USA;  $\pm 1^\circ\text{C}$  accuracy). Toad temperatures ranged from 11 to 35°C.

### Anatomy

Following feeding experiments, toads were killed with an overdose of MS-222 and electrode placement was confirmed by dissection. The mandible plus tongue and the paired DM were removed and masses were measured to  $\pm 0.001\text{ g}$  with a VB-302A digital balance (Virtual Measurements & Control, Santa Rosa, CA, USA). The mandible plus tongue specimen included all connective tissue and skin of the lower jaw. The distance from the jaw joint to the center of mass (CoM) of the mandible plus tongue was determined using a digital balance and a ruler as follows. A ruler was placed with one end on the digital balance and the other end on a fixed support, after which the balance was zeroed. The mandible plus tongue specimen was then placed in a resting position on the ruler with the mandibular symphysis facing the balance. The mass reported by the balance was divided by the mass of the specimen to yield the distance of the specimen's CoM from the fixed support as a percentage of the total distance between the balance and the fixed support. For example, if the CoM of the specimen was positioned on the ruler exactly halfway between the support and the balance, the balance would register 50% of the mass of the specimen, indicating that the CoM is positioned at 50% of the total distance, measured from the fixed support. Likewise, if the specimen's CoM was positioned 75% of the distance from the support to the balance, the balance would register 75% of the mass of the specimen. The distance between the CoM and the jaw joint was then measured with calipers, to be used along with the mass of the mandible plus tongue in dynamics calculations of ballistic mouth opening (below). Finally, the length of the mandible from the jaw joint center to the mandibular symphysis, perpendicular to the midline of the jaws, and the length of the retroarticular process of the mandible were measured with calipers.

### Kinematic and dynamic analyses

The timing and amplitude of movements of the mandible and tongue during prey capture, with respect to the maxilla as a fixed reference, were quantified from the digital image sequences. The positions of four anatomical landmarks were recorded from the image sequences using NIH ImageJ software running on an Apple iMac computer: (1) tip of the upper jaw, (2) tip of the mandible, (3) tip of the tongue and (4) jaw joint. Distances and angles were computed from these position data: (1) gape distance, or the distance between mandible and upper jaw tips, (2) gape angle in radians, computed as gape distance divided by anatomical mandible length, and (3) tongue reach, or the distance from the tongue tip to the mandible tip.

The times of eight events were measured relative to the start of ballistic mouth opening at time zero: (1) start of tongue projection, or the time the tongue crosses a plane connecting the upper and lower jaw tips, (2) end of ballistic mouth opening, (3)



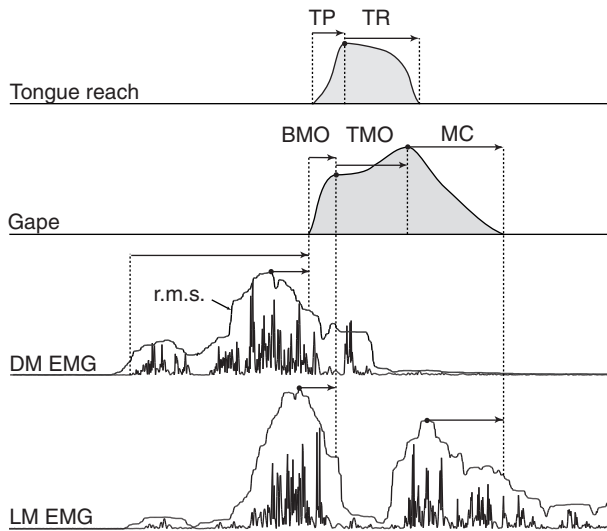


Fig. 1. Kinematic and motor control variables examined in this study as shown on a representative prey capture event. The two upper panels show tongue reach and gape profiles. For tongue reach, the horizontal arrows indicate the duration of tongue protraction (TP) and tongue retraction (TR) measured with respect to the maximum tongue projection distance (black dot). Gape variables include the durations of ballistic mouth opening (BMO), transport mouth opening (TMO) and mouth closing (MC), defined by the gape at the end of ballistic opening and transport opening (black dots). Vertical axes are in dimensions of distance and the horizontal axis is time. The lower panels illustrate rectified electromyographic (EMG) signals and the root mean square of the signals (r.m.s.) from the jaw depressor (DM) and levator (LM) muscles. Black dots indicate peak r.m.s. amplitudes. Horizontal arrows indicate latencies between the onset of DM activity and the start of ballistic mouth opening, peak DM amplitude and the start of ballistic mouth opening, peak LM amplitude (first burst) and the end of ballistic mouth opening, and peak LM amplitude (second burst) and the end of mouth closing. Vertical axes are in dimensions of voltage, and the horizontal axis is time. Additional variables are described in Materials and methods.

maximum tongue reach, (4) start of tongue retraction, (5) start of mouth opening associated with prey transport (i.e. 'transport mouth opening'), (6) end of mouth opening, (7) end of tongue retraction, or the time the tongue tip reaches the position of the jaw angle at the rear of the buccal cavity, and (8) end of mouth closing. Durations of movements were calculated from these timing variables (Fig. 1, Tables 1 and 2). The distance, angular and duration data were used to calculate mean linear and angular velocities of ballistic mouth opening, mouth opening associated with prey transport, and mouth closing, as well as mean linear velocities of tongue projection and retraction.

In addition to kinematics, the dynamics variables of ballistic mouth opening were calculated from the position data. Gape angle and gape distance were recorded for each image of ballistic mouth opening at 6kHz. To calculate the distance moved by the CoM of the mandible plus tongue (as measured in 'Anatomy' above), gape distance was multiplied by the percentage of mandible length that the CoM lies from the jaw joint. Because this distance represents changes in position of a point moving in an arc relative to another point and thus represents chord distance, the distance was converted to arc distance (i.e. displacement of the CoM along an arc) trigonometrically using gape angle and the radius of the arc (i.e. the distance from the jaw joint of the CoM of the mandible plus tongue). These arc distance data and gape angle data were

then smoothed using a quintic spline with the Pspline package in R statistical software ([www.r-project.org](http://www.r-project.org)). First and second derivatives of the spline function were computed to yield instantaneous velocity and acceleration, respectively, at a final interpolated rate of 10kHz.

Mass-specific instantaneous power of ballistic mouth opening in each prey capture event was calculated as the product of the acceleration and velocity of the CoM. Because ballistic mouth opening involves acceleration of the mandible, tongue and associated skin and connective tissue, absolute power output was computed by multiplying mass-specific power by the mass of the mandible plus tongue of the same individual toad (as measured in 'Anatomy' above). To calculate muscle mass-specific power output, this absolute power value was divided by the combined mass of the paired DM from the same individual. Maximum kinetic energy was calculated from the mass of the mandible plus tongue and the peak velocity achieved during ballistic mouth opening. Maximum values of velocity, acceleration, kinetic energy and power were used to examine the effects of temperature.

#### Analysis of electromyograms

The amplitudes of activity of the DM and LM and the timing of activity relative to kinematic events were quantified from the rectified EMG signals using AD Instruments LabChart software version 6 running on an Apple iMac computer. Bursts of activity with clear bounds were discernible only for the activity of the DM prior to mouth opening; the LM showed activity that was more dispersed in time, as did activity of the DM associated with prey transport. Therefore the duration of activity was measured only for the first burst of the DM. Onset of activity of the DM was defined as the signal reaching twice background noise level for at least 10ms; the end of activity was likewise defined as the signal dropping below twice the noise level for at least 10ms.

The duration of activity of the DM prior to ballistic mouth opening was hypothesized to increase with decreasing temperature because of the slowing of rate at which the muscle builds tension. Likewise, the latencies from the onset of DM activity and peak of DM activity to the start of ballistic mouth opening (Fig. 1) were expected to be greater at colder temperatures. The delay between activity of the DM and the start of ballistic mouth opening was quantified using three variables: (1) start of DM activity, (2) end of DM activity and (3) peak of DM activity (peak of root mean square, r.m.s.).

Unlike the duration of DM activity, the intensity of DM activity was expected to remain constant, under the assumption that muscle recruitment is maximized at all temperatures. Integrated area was measured for the DM as the sum of the values of the rectified signal between onset and end of activity, and intensity of the EMG burst was measured as (1) the r.m.s. within this period and (2) integrated area divided by duration of activity. Peak amplitude of muscle activity was measured as the maximum r.m.s. value using a 20ms time constant (i.e. the moving 20ms time window over which the r.m.s. was calculated).

Latencies between LM activity and mouth closing movements were expected to increase with decreasing temperature, because rates of muscle tension development and shortening slow at colder temperatures. The relationship between LM activity and the slowing of mandible depression (i.e. braking) at the end of ballistic mouth opening was measured as the time between the r.m.s. peak of the first burst of activity of LM and the end of ballistic mouth opening. The relationship between LM activity and mouth closing was measured as the time from peak r.m.s. activity of the second burst of the LM to the end of mouth closing (Fig. 1, bottom trace).

Table 1. Minimum and maximum values of kinematic, dynamic and electromyographic variables of prey capture in four *Bufo terrestris* from the 11–25°C temperature range

	Minimum	Maximum	Feedings
Muscle activity (s)			
DM duration	0.0498	0.3205	52 (6–22)
DM start to ballistic opening	0.0523	0.3488	52 (6–22)
DM end to ballistic opening	–0.0303	0.2990	52 (6–22)
DM max. amplitude to ballistic opening	–0.0015	0.1863	52 (6–22)
LM burst 1 max. amplitude to end ballistic opening	–0.0102	0.0394	52 (6–22)
LM burst 2 max. amplitude to mouth closed	0.0028	0.1937	53 (6–23)
Ballistic mouth opening			
Maximum gape distance (cm)	0.69	1.63	35 (4–13)
Maximum gape angle (rad)	0.70	1.44	35 (4–13)
Duration (s)	0.0036	0.019	56 (6–23)
Mean angular velocity (rad s <sup>–1</sup> )	53.36	303.49	35 (4–13)
Maximum angular velocity of jaw (rad s <sup>–1</sup> )	107.79	300.0	31 (4–12)
Maximum velocity of jaw CoM (m s <sup>–1</sup> )	0.66	1.17	31 (4–12)
Mean velocity of jaw CoM (m s <sup>–1</sup> )	0.36	1.18	35 (4–13)
Time of maximum velocity (s)	0.0012	0.0065	31 (4–12)
Maximum kinetic energy (J)	5.49E–05	6.71E–04	31 (4–12)
Maximum kinetic energy (J kg <sup>–1</sup> jaw mass)	0.2185	0.6845	31 (4–12)
Maximum kinetic energy (J kg <sup>–1</sup> depressor mass)	1.116	3.643	31 (4–12)
Maximum acceleration of CoM (m s <sup>–1</sup> s <sup>–1</sup> )	174.0	1440.0	31 (4–12)
Maximum power (W)	0.0414	0.3624	31 (4–12)
Maximum power (W kg <sup>–1</sup> jaw mass)	88.0	773.0	31 (4–12)
Maximum power (W kg <sup>–1</sup> depressor mass)	449.6	4114.4	31 (4–12)
Prey transport mouth opening and closing			
Maximum gape distance (cm)	1.11	2.31	35 (4–13)
Maximum gape angle (rad)	1.08	2.94	35 (4–13)
Duration of opening (s)	0.0070	0.0560	56 (6–23)
Mean angular velocity of opening (rad s <sup>–1</sup> )	22.09	244.60	35 (4–13)
Duration of closing (s)	0.0270	0.1970	56 (6–23)
Mean angular velocity of closing (rad s <sup>–1</sup> )	6.28	44.53	35 (4–13)
Tongue projection and retraction			
Maximum tongue reach (cm)	1.16	2.96	35 (4–13)
Maximum relative tongue reach (jaw lengths)	0.93	2.47	35 (4–13)
Duration of tongue projection (s)	0.002	0.014	55 (6–22)
Mean velocity of tongue projection (m s <sup>–1</sup> )	1.56	7.69	35 (4–13)
Mean velocity of tongue projection (jaw lengths s <sup>–1</sup> )	95.93	818.3	35 (4–13)
Duration of tongue retraction (s)	0.0134	0.065	55 (6–22)
Mean velocity of tongue retraction (m s <sup>–1</sup> )	0.178	1.73	35 (4–13)
Mean velocity of tongue retraction (jaw lengths s <sup>–1</sup> )	16.47	184.2	35 (4–13)

The total number of feedings is presented for each variable as well as the range of feedings for each individual (in parentheses). DM, m. depressor mandibulae; LM, m. levator mandibulae posterior longus; CoM, center of mass.

### Statistical analyses

All kinematic, dynamic and EMG data were log<sub>10</sub> transformed prior to statistical analysis. This transformation was performed because both the body temperature at which feedings occurred and the body size of the individual toads were expected to have an exponential relationship with the kinematic, dynamic and EMG variables. The dataset was divided into two overlapping subsets based on the temperature at which prey capture occurred, 11–25°C and 20–35°C, to examine whether the same thermal relationship held across the entire range. The lower range was expected to capture strong effects of temperature, whereas the upper range was expected to capture weaker effects, based on published results from other ectotherms (van Berkum, 1986; Huey and Kingsolver, 1993; Bauwens et al., 1995; Huey and Kingsolver, 1989). The ranges (i.e. subsets) were overlapping by 5°C because many feedings were recorded in the 20–25°C range and this increased the number of feedings within each subset, and because we had no *a priori* expectation of an exact temperature at which thermal effects would transition from strong to weak. A separate analysis of covariance (ANCOVA) was conducted on each subset of the data to examine three effects on

the variables: (1) temperature, (2) individual and (3) prey distance (Tables 3 and 4 and supplementary material Tables S1 and S2). Temperature effects were included as a continuous variable to examine how elastically powered and non-elastic movements and their motor control responded to changes in body temperature. An individual effect was included to account for differences in body size and other random individual differences, and for characteristics of the EMG electrodes and resulting variation in signal strength. Prey distance was included because it has been found in another study to influence prey capture kinematics (Lappin et al., 2006); it was dropped from the model when non-significant for a given variable to increase sample size and statistical power. Prey distance was consequently retained in the model for only three of the 27 variables (see Results), but in none of these cases did its inclusion change the significance of the temperature effect. A simultaneous Bonferroni correction for experiment-wise error adjusted the significance level from  $\alpha$  of  $\leq 0.05$  to 0.0019 (for 27 tests corresponding to the 27 variables examined).

Temperature coefficients ( $Q_{10}$ ) for each performance variable were computed across each temperature range (11–25°C and

Table 2. Minimum and maximum values of kinematic, dynamic and electromyographic variables of prey capture in four *B. terrestris* from the 20–35°C temperature range

	Minimum	Maximum	Feedings
Muscle activity (s)			
DM duration	0.0315	0.1760	37 (7–11)
DM start to ballistic opening	0.0290	0.3488	37 (7–11)
DM end to ballistic opening	–0.0119	0.2990	37 (7–11)
DM max. amplitude to ballistic opening	–0.0015	0.0505	37 (7–11)
LM burst 1 max. amplitude to end ballistic opening	–0.0102	0.0324	37 (7–11)
LM burst 2 max. amplitude to mouth closed	–0.0009	0.0969	37 (7–11)
Ballistic mouth opening			
Maximum gape distance (cm)	0.71	1.63	24 (4–8)
Maximum gape angle (rad)	0.70	1.43	24 (4–8)
Duration (s)	0.0045	0.0130	38 (7–12)
Mean angular velocity (rad s <sup>–1</sup> )	58.51	303.49	24 (4–8)
Maximum angular velocity of jaw (rad s <sup>–1</sup> )	99.60	300.00	23 (4–7)
Maximum velocity of jaw CoM (m s <sup>–1</sup> )	0.70	1.17	23 (4–7)
Mean velocity of jaw CoM (m s <sup>–1</sup> )	0.44	1.18	24 (4–8)
Time of maximum velocity (s)	0.0012	0.0065	23 (4–7)
Maximum kinetic energy (J)	6.98E–05	6.71E–04	23 (4–7)
Maximum kinetic energy (J kg <sup>–1</sup> jaw mass)	0.2429	0.6845	23 (4–7)
Maximum kinetic energy (J kg <sup>–1</sup> depressor mass)	1.241	3.643	23 (4–7)
Maximum acceleration of CoM (m s <sup>–1</sup> s <sup>–1</sup> )	174.0	1070.0	23 (4–7)
Maximum power (W)	0.0465	0.3624	23 (4–7)
Maximum power (W kg <sup>–1</sup> jaw mass)	99.0	817.0	23 (4–7)
Maximum power (W kg <sup>–1</sup> depressor mass)	505.8	4348.5	23 (4–7)
Prey transport mouth opening and closing			
Maximum gape distance (cm)	1.01	2.30	24 (4–8)
Maximum gape angle (rad)	0.75	2.94	24 (4–8)
Duration of opening (s)	0.0041	0.0375	38 (7–12)
Mean angular velocity of opening (rad s <sup>–1</sup> )	38.49	244.60	24 (4–8)
Duration of closing (s)	0.0255	0.1005	38 (7–12)
Mean angular velocity of closing (rad s <sup>–1</sup> )	11.29	44.53	24 (4–8)
Tongue projection and retraction			
Maximum tongue reach (cm)	1.19	2.79	24 (4–8)
Maximum relative tongue reach (jaw lengths)	0.89	2.47	24 (4–8)
Duration of tongue projection (s)	0.002	0.0113	38 (7–12)
Mean velocity of tongue projection (m s <sup>–1</sup> )	1.96	7.69	24 (4–8)
Mean velocity of tongue projection (jaw lengths s <sup>–1</sup> )	116.85	818.3	24 (4–8)
Duration of tongue retraction (s)	0.0105	0.047	38 (7–12)
Mean velocity of tongue retraction (m s <sup>–1</sup> )	0.530	1.73	24 (4–8)
Mean velocity of tongue retraction (jaw lengths s <sup>–1</sup> )	27.59	184.2	24 (4–8)

Total number of feedings is presented for each variable as well as the range of feedings for each individual (in parentheses). DM, m. depressor mandibulae; LM, m. levator mandibulae posterior longus; CoM, center of mass.

20–35°C) from the partial regression coefficients of the temperature effect in the ANCOVAs, because the ANCOVA model includes effects of individual toad (and in some cases prey distance) that influence the estimate of the relationship between temperature and the performance variable. The  $Q_{10}$  value for each variable was calculated as the base 10 antilogarithm of the partial regression coefficient of the temperature effect (PRC) multiplied by 10:

$$Q_{10} = 10^{(\text{PRC} \times 10)} \quad (1)$$

Temperature coefficients of duration variables are reported as inverse  $Q_{10}$  values (i.e.  $1/Q_{10}$ ) to express them as rates. Statistical analyses were performed on an Apple iMac computer using JMP 5.1 software (SAS Institute, Cary, NC, USA).

## RESULTS

### Prey capture kinematics

Toads captured prey by ballistic tongue projection, sometimes combined with a forward lunge of the body. The mouth was opened in two phases; ballistic mouth opening associated with tongue projection followed by a second phase during tongue retraction and prey transport. The two phases were separated by a pause when the

tongue elongated to maximum length and adhered to the prey. The tongue was then retracted, simultaneous with the second phase of mouth opening that accommodated the prey, immediately after which the mouth was closed, typically with the prey delivered to the rear of the buccal cavity and clear of the jaws (Fig. 2 and supplementary material Movie 1).

Considering feedings examined across the entire 11–35°C range, ballistic mouth opening required 0.0036–0.019 s, mouth opening during prey transport took 0.0041–0.056 s and mouth closing took 0.026–0.197 s, based upon 70 total feedings and 10–24 feedings per individual. Gape angle after ballistic mouth opening was 0.7–1.4 rad, and this was increased to 0.75–2.9 rad in the second phase of mouth opening based upon 44 total feedings and 8–13 feedings per individual. Tongue projection required 0.002–0.014 s and tongue retraction took longer, at 0.011–0.065 s (69 feedings, 10–23 per individual). The tongue was projected 0.89–2.5 mandible lengths (44 feedings, 8–13 per individual) beyond the tip of the mandible. Summary results from the two temperature ranges are shown in Tables 1 and 2.

Temperature had a significant effect on three kinematic variables across the 11–25°C range (Fig. 3, Table 3 and supplementary

Table 3. Results of ANCOVA examining effects on kinematic, dynamic and electromyographic variables over the 11–25°C range

	Individual <i>P</i> -value	Temp. <i>P</i> -value	Prey distance <i>P</i> -value	Slope	$Q_{10}$	$1/Q_{10}$
<b>Kinematic variables</b>						
Duration of ballistic mouth opening	0.6973	0.1449	0.1119	-0.0127	0.75	<b>1.34</b>
Duration of tongue projection	0.0044	0.1668	0.3659	-0.0105	0.78	1.27
Gape distance at end of ballistic opening	0.0020	0.8688	0.0110	-0.0002	1.00	1.00
Maximum tongue reach	0.0167	0.1464	<b>&lt;0.0001</b>	0.0028	1.07	0.94
Duration of mouth opening during transport	0.5418	<b>0.0003</b>	0.6559	-0.0214	0.61	<b>1.64</b>
Final gape distance	<b>&lt;0.0001</b>	0.2725	0.0068	0.0012	1.03	0.97
Duration of tongue retraction	0.0256	<b>0.0012</b>	0.0331	-0.0196	0.64	<b>1.57</b>
Duration of mouth closing	0.3021	<b>&lt;0.0001</b>	0.5633	-0.0339	0.46	<b>2.18</b>
<b>Dynamic variables</b>						
Mean velocity of ballistic mouth opening	<b>&lt;0.0001</b>	0.0121	0.0660	0.0053	1.13	0.89
Maximum velocity of ballistic opening	<b>&lt;0.0001</b>	0.1425	0.0262	0.0025	1.06	0.94
Maximum acceleration of ballistic opening	<b>&lt;0.0001</b>	0.8950	0.6371	-0.0006	0.99	1.01
Maximum power of ballistic opening	<b>&lt;0.0001</b>	0.4343	0.0647	0.0042	1.10	0.91
Mean velocity of tongue projection	<b>&lt;0.0001</b>	0.0857	0.2727	0.0097	1.25	0.80
Mean velocity of mouth opening during transport	0.6133	<b>&lt;0.0001</b>	0.0887	0.0302	<b>2.00</b>	0.50
Mean velocity of tongue retraction	0.0619	<b>&lt;0.0001</b>	<b>&lt;0.0001</b>	0.0247	<b>1.77</b>	0.57
Mean velocity of mouth closing	<b>0.0010</b>	<b>&lt;0.0001</b>	0.9841	0.0354	<b>2.26</b>	0.44
<b>Electromyographic (EMG) variables</b>						
Depressor EMG integrated area	<b>&lt;0.0001</b>	<b>&lt;0.0001</b>	<b>&lt;0.0001</b>	-0.0343	0.45	<b>2.20</b>
Depressor EMG duration	<b>&lt;0.0001</b>	<b>&lt;0.0001</b>	0.1449	-0.0381	0.42	<b>2.41</b>
Depressor EMG r.m.s.	0.0035	0.0254	0.0059	0.0081	1.20	0.83
Depressor EMG integrated area/duration	0.1850	0.1661	0.0032	0.0049	1.12	0.89
Depressor EMG r.m.s. max. amplitude	0.7349	0.1280	0.0545	0.0050	1.12	0.89
Levator EMG (second burst) r.m.s. max. amplitude	<b>&lt;0.0001</b>	0.0749	0.3593	-0.0123	0.75	1.33
Depressor EMG start to start of ballistic opening	0.0045	<b>&lt;0.0001</b>	0.0070	-0.0360	0.44	<b>2.29</b>
Depressor EMG end to start of ballistic opening	0.0033	0.2954	0.9636	0.0090	1.23	0.81
Depressor EMG max. amplitude to start of ballistic opening	0.9674	<b>&lt;0.0001</b>	0.4487	-0.0563	0.27	<b>3.66</b>
Levator EMG (first burst) max. amplitude to end ballistic opening	0.2043	0.0253	0.5845	-0.0239	0.58	1.73
Levator EMG (second burst) max. amplitude to mouth closed	<b>0.0010</b>	<b>0.0013</b>	0.9162	-0.0265	0.54	<b>1.84</b>

*P*-values are shown for individual, temperature (Temp.) and prey distance, as is the partial regression coefficient for the temperature effect (i.e. Slope) from the model from which  $Q_{10}$  values were calculated. Prey distance was included as a covariate only when it showed a significant effect for that variable.

Bold *P*-values indicate significant effects at the Bonferroni-corrected  $\alpha$  (0.0019 for 27 variables). Bold  $Q_{10}$  values indicate significant temperature effects.

material Table S1): duration of mouth opening during prey transport ( $1/Q_{10}=1.64$ ;  $P=0.0003$ ), duration of mouth closing ( $1/Q_{10}=2.18$ ;  $P<0.0001$ ) and duration of tongue retraction ( $1/Q_{10}=1.57$ ;  $P=0.0012$ ). Temperature did not significantly affect the duration of ballistic mouth opening ( $1/Q_{10}=1.34$ ;  $P=0.1449$ ), the gape at the end of ballistic opening ( $Q_{10}=1.00$ ;  $P=0.8688$ ), the duration of tongue projection ( $1/Q_{10}=1.27$ ;  $P=0.1668$ ), the final gape distance ( $Q_{10}=1.03$ ;  $P=0.2725$ ) or maximum tongue reach ( $Q_{10}=1.07$ ;  $P=0.1464$ ; Fig. 2).

Across the 20–35°C range (Fig. 3, Table 4 and supplementary material Table S2), temperature had a significant effect on the duration of ballistic mouth opening ( $1/Q_{10}=1.34$ ;  $P<0.0001$ ), duration of mouth opening during prey transport ( $1/Q_{10}=1.82$ ;  $P=0.0002$ ) and final gape distance ( $Q_{10}=0.72$ ;  $P<0.0001$ ). No significant effect of temperature was found for the gape at the end of ballistic opening ( $Q_{10}=0.95$ ;  $P=0.1061$ ), duration of mouth closing ( $1/Q_{10}=1.12$ ;  $P=0.2846$ ), duration of tongue projection ( $1/Q_{10}=0.79$ ;  $P=0.2582$ ), duration of tongue retraction ( $1/Q_{10}=1.25$ ;  $P=0.0224$ ) or maximum tongue reach ( $Q_{10}=0.88$ ;  $P=0.0295$ ). Across both temperature ranges, prey distance affected only maximum tongue reach ( $P<0.0001$ ), which was shorter for nearer prey (Tables 3 and 4).

#### Prey capture dynamics

Considering feedings examined across the entire 11–35°C range, ballistic mouth opening achieved a peak angular velocity of 100–300 rad s<sup>-1</sup> and an mean angular velocity of 53–303 rad s<sup>-1</sup> based

on 44 total feedings and 8–13 feedings per individual (Fig. 4). The CoM of the mandible plus tongue reached a maximum linear velocity of 0.66–1.17 m s<sup>-1</sup> and a maximum linear acceleration of 174–1440 m s<sup>-2</sup>. Peak velocity and kinetic energy were achieved 1.2–6.5 ms after the start of mandible movement. Maximum kinetic energy was 0.05–0.67 mJ; mass-specific kinetic energy was 1.1–3.6 J kg<sup>-1</sup> of depressor muscle mass; peak power reached 0.04–0.36 W, and mass-specific peak power was 450–4349 W kg<sup>-1</sup> depressor muscle mass. These last six dynamics calculations were based on a total of 40 feedings, 8–12 per individual (Fig. 4). The second phase of mouth opening was slower, at 22–245 rad s<sup>-1</sup>, as was mouth closing at 6.3–45 rad s<sup>-1</sup>. Tongue projection velocity was 1.6–7.7 m s<sup>-1</sup> or 96–818 mandible lengths s<sup>-1</sup>, while tongue retraction was slower, at 0.17–1.7 m s<sup>-1</sup> or 16–184 mandible lengths s<sup>-1</sup>. These final four calculations were based on 44 total feedings, 8–13 feedings per individual.

Temperature significantly affected only three dynamics variables across the 11–25°C range (Figs 3–5 and Table 3): the mean velocity of transport mouth opening ( $Q_{10}=2.00$ ;  $P<0.0001$ ), mean velocity of mouth closing ( $Q_{10}=2.26$ ;  $P<0.0001$ ) and mean velocity of tongue retraction ( $Q_{10}=1.77$ ;  $P<0.0001$ ). Dynamics variables that were not significantly affected by temperature included the mean velocity of ballistic mouth opening ( $Q_{10}=1.13$ ;  $P=0.0121$ ), maximum velocity of ballistic mouth opening ( $Q_{10}=1.06$ ;  $P=0.1425$ ), maximum acceleration of ballistic mouth opening ( $Q_{10}=0.99$ ;  $P=0.8950$ ), maximum power of ballistic mouth opening ( $Q_{10}=1.10$ ;  $P=0.4343$ ) and mean velocity of



Table 4. Results of ANCOVA examining effects on kinematic, dynamic and electromyographic variables over the 20–35°C range

	Individual <i>P</i> -value	Prey Temp. <i>P</i> -value	Prey distance <i>P</i> -value	Slope	$Q_{10}$	$1/Q_{10}$
<b>Kinematic variables</b>						
Duration of ballistic mouth opening	<b>&lt;0.0001</b>	<b>&lt;0.0001</b>	0.0765	-0.0126	0.75	<b>1.34</b>
Duration of tongue projection	<b>&lt;0.0001</b>	0.2582	0.8717	0.0104	1.27	0.79
Gape distance at end of ballistic opening	0.0353	0.1061	0.2323	-0.0024	0.95	1.06
Maximum tongue reach	0.0019	0.0295	<b>&lt;0.0001</b>	-0.0057	0.88	1.14
Duration of mouth opening during transport	0.0162	<b>0.0002</b>	0.5201	-0.0261	0.55	<b>1.82</b>
Final gape distance	0.0216	<b>&lt;0.0001</b>	0.2870	-0.0140	0.72	<b>1.38</b>
Duration of tongue retraction	<b>&lt;0.0001</b>	0.0224	0.2962	-0.0097	0.80	1.25
Duration of mouth closing	<b>&lt;0.0001</b>	0.2846	0.0664	-0.0048	0.90	1.12
<b>Dynamic variables</b>						
Mean velocity of ballistic mouth opening	0.0064	0.7305	0.0261	0.0018	1.04	0.96
Maximum velocity of ballistic opening	<b>&lt;0.0001</b>	0.1261	0.1253	0.0042	1.10	0.91
Maximum acceleration of ballistic opening	<b>&lt;0.0001</b>	0.3219	0.6880	0.0065	1.16	0.86
Maximum power of ballistic opening	<b>&lt;0.0001</b>	0.1702	0.5394	0.0112	1.29	0.77
Mean velocity of tongue projection	0.0027	0.0160	0.2348	-0.0357	0.44	2.28
Mean velocity of mouth opening during transport	0.7525	0.9304	0.7919	-0.0007	0.98	1.02
Mean velocity of tongue retraction	0.0207	0.8545	0.1292	-0.0011	0.97	1.03
Mean velocity of mouth closing	<b>&lt;0.0001</b>	0.1025	0.0991	-0.0092	0.81	1.24
<b>Electromyographic (EMG) variables</b>						
Depressor EMG integrated area	<b>&lt;0.0001</b>	<b>0.0002</b>	0.1762	-0.0159	0.69	<b>1.44</b>
Depressor EMG duration	<b>0.0007</b>	0.1119	0.0303	-0.0072	0.85	1.18
Depressor EMG r.m.s.	<b>&lt;0.0001</b>	0.0086	0.6499	-0.0099	0.80	1.26
Depressor EMG integrated area/duration	<b>&lt;0.0001</b>	0.0179	0.2593	-0.0087	0.82	1.22
Depressor EMG r.m.s. max. amplitude	<b>&lt;0.0001</b>	0.0100	0.6670	-0.0083	0.83	1.21
Levator EMG (second burst) r.m.s. max. amplitude	<b>&lt;0.0001</b>	0.0369	0.2768	-0.0174	0.67	1.49
Depressor EMG start to start of ballistic opening	0.1970	0.0265	0.0020	-0.0158	0.69	1.44
Depressor EMG end to start of ballistic opening	0.0175	0.0861	0.1334	-0.0136	0.73	1.37
Depressor EMG max. amplitude to start of ballistic opening	0.8851	0.5968	0.2322	-0.0100	0.79	1.26
Levator EMG (first burst) max. amplitude to end ballistic opening	0.0749	0.7538	0.4303	0.0054	1.13	0.88
Levator EMG (second burst) max. amplitude to mouth closed	0.0131	0.8417	0.0657	-0.0042	0.91	1.10

*P*-values are shown for individual, temperature (Temp.) and prey distance, as is the partial regression coefficient for the temperature effect (i.e. Slope) from the model from which  $Q_{10}$  values were calculated. Prey distance was included as a covariate only when it showed a significant effect for that variable.

Bold *P*-values indicate significant effects at the Bonferroni-corrected  $\alpha$  (0.0019 for 27 variables). Bold  $Q_{10}$  values indicate significant temperature effects.

tongue projection ( $Q_{10}=1.25$ ;  $P=0.0857$ ). Prey distance influenced only the mean velocity of tongue retraction ( $P<0.0001$ ), which was faster for more distant prey.

Across the 20–35°C range, all dynamics variables were unaffected by temperature (Fig. 3, Table 4 and supplementary material Table S2): mean velocity of ballistic mouth opening ( $Q_{10}=1.04$ ;  $P=0.7305$ ), maximum velocity of ballistic opening ( $Q_{10}=1.10$ ;  $P=0.1261$ ), maximum acceleration of ballistic opening ( $Q_{10}=1.16$ ;  $P=0.3219$ ), maximum power of ballistic opening ( $Q_{10}=1.29$ ;  $P=0.1702$ ), mean velocity of transport mouth opening ( $Q_{10}=0.98$ ;  $P=0.9304$ ), mean velocity of mouth closing ( $Q_{10}=0.81$ ;  $P=0.1025$ ), mean velocity of tongue projection ( $Q_{10}=0.44$ ;  $P=0.0160$ ) and mean velocity of tongue retraction ( $Q_{10}=0.97$ ;  $P=0.8545$ ). Prey distance influenced none of the dynamics variables significantly across the 20–35°C range.

#### Motor control of prey capture

In feedings across the entire 11–35°C range (65 total feedings, 10–23 per individual), the DM was active for 31.5–320 ms, and the start of activity preceded mouth opening by 29–349 ms (Figs 6 and 7). The time of end of activity of the DM ranged from 299 ms before to 30 ms after the start of mouth opening. The peak of activity of the DM occurred from 186 ms before to 1.5 ms after mouth opening.

The LM was active at two times during the gape cycle; the first period of activity was associated with braking the mandible at the end of ballistic mouth opening, and the second was responsible for closing the mouth (Figs 1 and 7). The first burst of LM activity

reached its peak amplitude 39 ms before to 10 ms after the end of ballistic opening. The second peak of LM activity occurred 194 ms before to 1 ms after the mouth was closed.

Temperature significantly affected five motor control variables across the 11–25°C range (Table 3 and supplementary material Table S1): integrated area of DM ( $Q_{10}=0.45$ ;  $1/Q_{10}=2.20$ ;  $P<0.0001$ ), duration of activity of DM ( $1/Q_{10}=2.41$ ;  $P<0.0001$ ) (Fig. 6A), latency between onset of DM activity and start of mouth opening ( $1/Q_{10}=2.29$ ;  $P<0.0001$ ), time from peak amplitude of DM to start of mouth opening ( $1/Q_{10}=3.66$ ;  $P<0.0001$ ) and time from peak amplitude of LM activity (second burst) to end of mouth closing ( $1/Q_{10}=1.84$ ;  $P=0.0013$ ). In all cases, values of variables increased with decreasing temperature. Prey distance influenced only the integrated area of the DM significantly across the 11–25°C range ( $P<0.0001$ ), which was greater in feedings on more distant prey.

The remaining six variables showed no significant effect of temperature across the 11–25°C range: r.m.s. of DM ( $Q_{10}=1.20$ ;  $1/Q_{10}=0.83$ ;  $P=0.0254$ ) (Fig. 6B), integrated area of DM divided by duration of activity of DM ( $Q_{10}=1.12$ ;  $P=0.1661$ ), peak amplitude of DM ( $Q_{10}=1.12$ ;  $P=0.1280$ ), peak amplitude of the second burst of LM ( $Q_{10}=0.75$ ;  $P=0.0749$ ), time from end of DM activity to start of mouth opening ( $1/Q_{10}=0.81$ ;  $P=0.2954$ ) and time from peak LM activity (first burst) to end of ballistic opening ( $1/Q_{10}=1.73$ ;  $P=0.0253$ ).

Across the 20–35°C range (Table 4 and supplementary material Table S2), only DM integrated area was significantly increased by decreasing temperature ( $Q_{10}=0.69$ ;  $1/Q_{10}=1.44$ ;  $P=0.0002$ ), while the



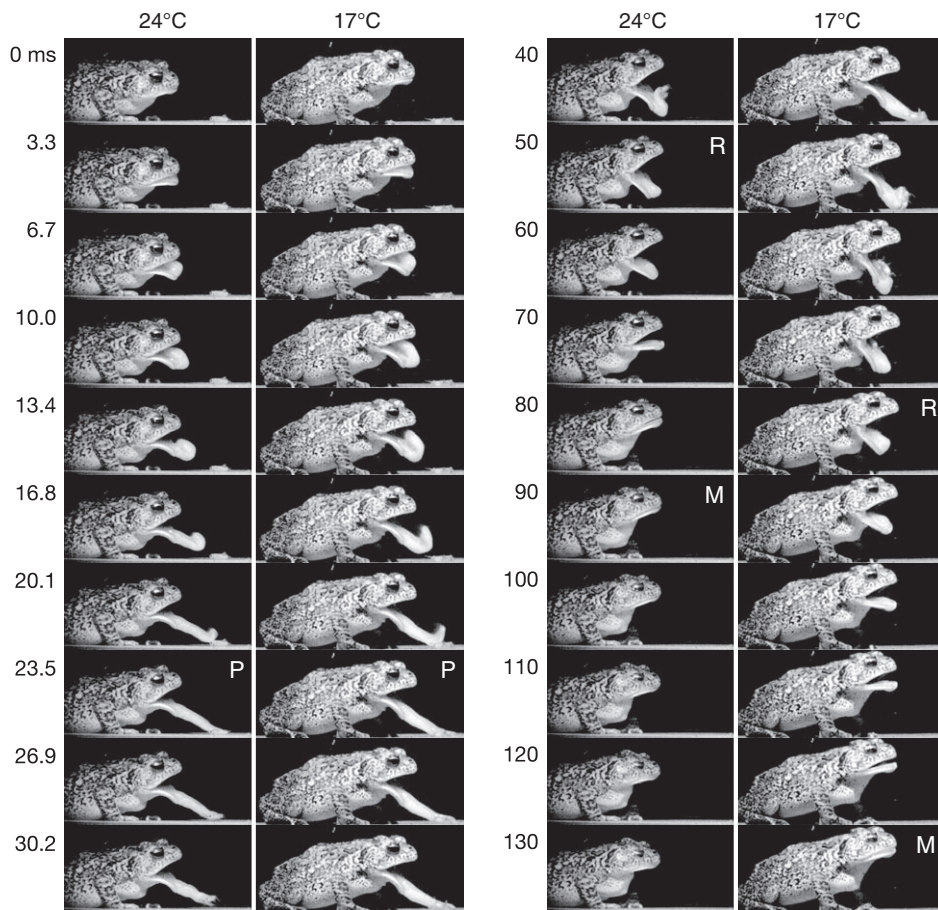


Fig. 2. Image sequences of one individual of *Bufo terrestris* feeding at 24 and 17°C showing no difference in the durations of ballistic mouth opening and tongue projection (P indicates peak projection) yet pronounced differences in the durations of non-ballistic tongue retraction (R) and mouth closing (M). Sequences progress downward beginning at the top of the left column and continue at the top of the right column, and begin at the start of ballistic mouth opening at time 0. The time step is 3.3 ms in the left column and 10 ms in the right column. EMG signals obtained from these feedings are shown in Fig. 7.

remaining 10 variables showed no significant temperature effect: duration of activity of DM ( $1/Q_{10}=1.26$ ;  $P=0.1119$ ) (Fig. 6A), r.m.s. of DM ( $Q_{10}=0.80$ ;  $1/Q_{10}=1.18$ ;  $P=0.0086$ ) (Fig. 6B), integrated area of DM divided by duration of activity of DM ( $Q_{10}=0.82$ ;  $P=0.0179$ ), peak amplitude of DM ( $Q_{10}=0.83$ ;  $P=0.0100$ ), peak amplitude of the second burst of LM ( $Q_{10}=0.67$ ;  $P=0.0369$ ), latency between onset of DM activity and start of mouth opening ( $1/Q_{10}=1.44$ ;  $P=0.0265$ ), time from end of DM activity to start of mouth opening ( $1/Q_{10}=1.37$ ;  $P=0.0175$ ), latency between peak amplitude of DM and start of mouth opening ( $1/Q_{10}=1.26$ ;  $P=0.5968$ ), time from peak LM activity (first burst) to end of ballistic opening ( $1/Q_{10}=0.88$ ;  $P=0.7538$ ) and time from peak amplitude of LM activity (second burst) to end of mouth closing ( $1/Q_{10}=1.10$ ;  $P=0.8417$ ). Prey distance influenced none of the motor control variables significantly across the 20–35°C range.

#### Anatomy of the mandible

Mandible length from the jaw to the mandible tip was 0.94–1.92 cm ( $1.46\pm 0.21$  cm) and retroarticular process length was 0.13–0.28 cm ( $0.18\pm 0.03$  cm). Combined mass of the paired DM muscles was 0.031–0.250 g ( $0.13\pm 0.05$  g) and the mass of the mandible plus tongue was 0.165–1.129 g ( $0.65\pm 0.21$  g). Combined DM mass was 17–22% ( $19\pm 1\%$ ) of mandible plus tongue mass, and the CoM of the mandible plus tongue was positioned 39–45% ( $42\pm 1.2\%$ ) of the distance from the jaw joint to the mandible tip.

### DISCUSSION

#### Prey capture kinematics

The toads in this study captured prey by ballistic tongue projection, in which the mouth was opened rapidly and the tongue flipped over

the mandible and underwent ‘inertial elongation’ (*sensu* Nishikawa, 2000), as momentum was transferred from the accelerated mandible (Mallett et al., 2001). The tongue was then retracted into the mouth, transporting the prey into the buccal cavity. Prey transport included a second phase of mouth opening followed by mouth closing after the tongue was withdrawn. Although prey transport was slower at the lower temperatures, the toads were capable of ballistic feeding at all temperatures (11–35°C), and tongue projection distance and gape distance were unaffected by temperature.

The effects of temperature on the durations of movements differed across the two temperature ranges, however, indicating that the muscle physiology underlying the movements of the mandible and tongue experienced different thermal effects. In the lower temperature range (11–25°C), ballistic movements – ballistic mouth opening and tongue projection – showed no effect of temperature in their durations or excursions and exhibited low  $Q_{10}$  values of 1.0–1.3. The durations of non-ballistic movements during prey transport – mouth opening, tongue retraction and mouth closing – were affected by temperature and showed higher  $Q_{10}$  values of 1.6–2.2. In the higher temperature range (20–35°C), temperature had a significant effect on duration of ballistic mouth opening, yet the  $Q_{10}$  was still low at 1.3, and among the non-ballistic movements only duration of transport mouth opening and final gape distance ( $Q_{10}$  values of 0.72–1.82) were affected by temperature (Table 4). These results combined with the fact that the durations of mouth closing and tongue retraction showed no effect of temperature and low  $Q_{10}$  values (1.1–1.3) indicate a plateau of high performance and low thermal sensitivity at the higher end of the temperature range.

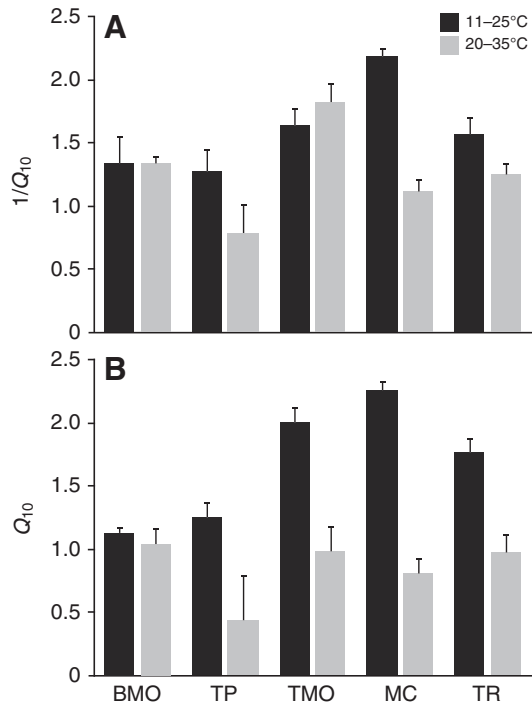


Fig. 3. Bar graphs of mean temperature coefficients of duration of prey capture phases (A) and the mean velocities of movements (B) across the 11–25°C range (dark gray bars) and the 20–35°C range (light gray bars). Note that the effects of temperature are greater across the lower temperature range (dark gray bars), and that ballistic mouth opening (BMO) and tongue projection (TP) show values below 1.5 across both ranges. Non-ballistic movements – transport mouth opening (TMO), mouth closing (MC) and tongue retraction (TR) – show stronger temperature effects across the lower range. Inverse  $Q_{10}$  values ( $1/Q_{10}$ ) are shown for durations.  $Q_{10}$  values and standard errors (error bars) are calculated from partial regression coefficients of the temperature effect in the ANCOVA (see Materials and methods for details).

A pattern of relatively high thermal dependence at lower temperatures and lower thermal dependence at moderate and high temperatures has also been observed in the muscle-powered movements of other ectotherms including swimming speed in tadpoles (Wilson and Franklin, 1999), swimming speed and jumping distance in frogs (John-Alder et al., 1989; Wilson and Franklin, 2000; Knowles and Weigl, 1990), tongue projection velocity of chameleons (Anderson and Deban, 2010) and sprint speed of lizards (Mautz et al., 1992; Marsh and Bennett, 1986; Marsh and Bennett, 1985; Bennett, 1990), as well as in the contractile rates of isolated frog skeletal muscle (Barnes and Ingalls, 1991). This pattern indicates a plateau of high performance at higher temperature (but below the abrupt decline observed near the critical thermal maximum) and often corresponds to the temperature range over which the animals thermoregulate when active (van Berkum, 1986; Huey and Kingsolver, 1993; Bauwens et al., 1995; Huey and Kingsolver, 1989). Species of *Bufo* have been found to select temperatures of approximately 26°C when placed in a thermal gradient (Lillywhite et al., 1973), which is within the thermal performance breadth of sprinting locomotion (Tracy et al., 1993), and within the upper temperature range in which we recorded low thermal dependence of feeding movements in *B. terrestris*.

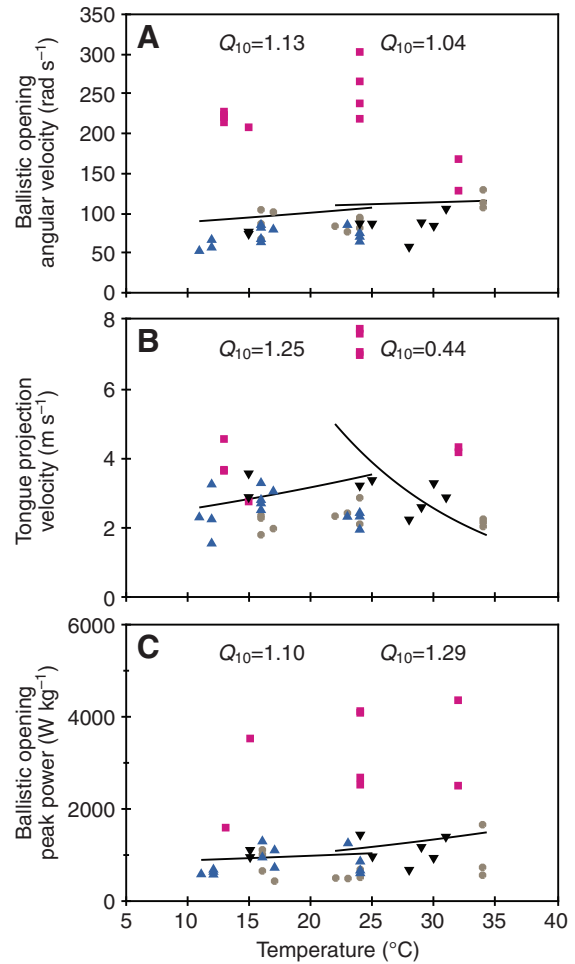


Fig. 4. Scatterplots of dynamics variables from the ballistic movements of all prey capture events *versus* temperature. Mean angular velocity of ballistic mouth opening (A), mean velocity of tongue projection (B) and peak power of ballistic mouth opening (C) show weak relationships with temperature across both temperature ranges. Exponential regressions representing  $Q_{10}$  values are overlaid on the data points across the 11–25°C and the 20–35°C ranges.  $Q_{10}$  values are derived from partial regression coefficients of the temperature effect in the ANCOVA (see Materials and methods for details). Individual toads are shown as different symbols.

### Prey capture dynamics

Tongue projection in *B. terrestris* was explosively dynamic at all temperatures. A toad feeding at 11°C, for example, achieved a peak acceleration of the mandible plus tongue of 393 m s<sup>-2</sup> and peak power of 598 W kg<sup>-1</sup> muscle mass. Our lowest value for peak power of ballistic opening of 450 W kg<sup>-1</sup> muscle mass (which occurred at a body temperature of 17°C) exceeded the maximum direct muscle power estimated for frog limb muscles during jumping (373 W kg<sup>-1</sup> at 25°C) (Lutz and Rome, 1994). These high values and the high mean values of velocity, acceleration and power confirm that *B. terrestris* relies on elastic power enhancement during ballistic mouth opening, as do other toads (Lappin et al., 2006).

In contrast to ballistic mouth opening and tongue projection, mouth opening during prey transport as well as tongue retraction and mouth closing showed lower peak and mean values for the dynamics parameters. For example, tongue retraction had a mean peak velocity of 56 mandible lengths s<sup>-1</sup> compared with 255 mandible lengths s<sup>-1</sup> for tongue projection.

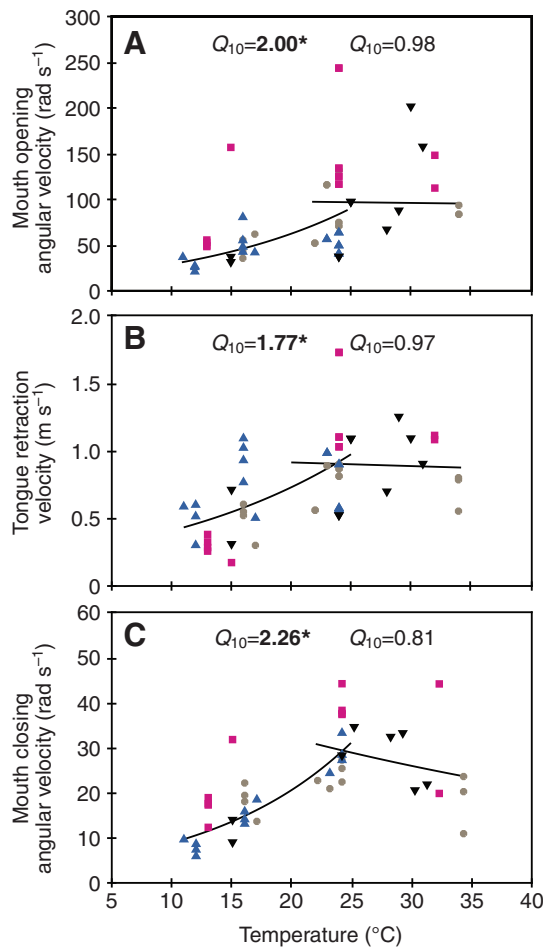


Fig. 5. Scatterplots of dynamics variables from the non-ballistic movements of all prey capture events *versus* temperature. Mean angular velocity of transport mouth opening (A), mean velocity of tongue retraction (B) and mean angular velocity of mouth closing (C) show strong effects of temperature across the lower range. Exponential regressions representing  $Q_{10}$  values are overlaid on the data points across the 11–25°C and the 20–35°C ranges.  $Q_{10}$  values are derived from partial regression coefficients of the temperature effect in the ANCOVA (see Materials and methods for details). Individual toads are shown as different symbols.

As was the case for kinematics, the greatest effect of temperature on dynamics was across the 11–25°C range, in which three dynamics variables were significantly affected – velocities of mouth opening during transport, mouth closing and tongue retraction – showing  $Q_{10}$  values of 1.8–2.3 (Table 3). The lack of an effect of temperature across the 20–35°C range (Table 4) on dynamics variables ( $Q_{10}$  values of 0.44–1.3) also points to a plateau of temperature sensitivity at the higher end of the temperatures examined, probably the result of a plateau in the thermal sensitivity of muscle dynamics.

These results provide evidence that the elastic recoil mechanism that powers ballistic mouth opening and tongue projection confers low thermal sensitivity to these movements –  $Q_{10}$  of 1.1 for velocity and power – over a range of temperatures that strongly influence muscle contractile dynamics. In contrast, prey transport movements that rely upon direct muscular power do not escape limitations imposed by thermal effects on muscle contractile dynamics afforded by an elastic mechanism and therefore show significant thermal sensitivity.

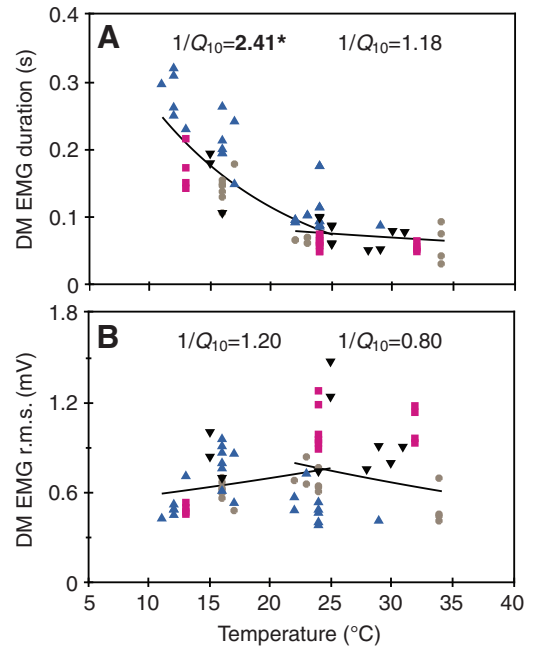


Fig. 6. Scatterplots of EMG variables from all prey capture events *versus* temperature. Duration of jaw depressor muscle activity (A) shows a strong increase at lower temperature and a plateau at higher temperature. Exponential regressions representing  $Q_{10}$  values are overlaid on the data points across the 11–25°C and the 20–35°C ranges. Intensity of jaw depressor muscle activity (B) is nearly constant at all temperatures, indicating consistent recruitment.  $Q_{10}$  values are derived from partial regression coefficients of the temperature effect in the ANCOVA (see Materials and methods for details). Individual toads are shown as different symbols.

High power output of ballistic movements based on elastic mechanisms is widespread among animals (Roberts and Azizi, 2011) and has been shown not only in the prey capture of toads (Lappin et al., 2006) but also in the tongue projection of chameleons (Anderson and Deban, 2010; de Groot and van Leeuwen, 2004; Van Leeuwen et al., 2000) and plethodontid salamanders (Deban et al., 1997; Deban et al., 2007), the striking of mantis shrimp (Zack et al., 2009; Patek et al., 2004; Patek et al., 2007), jaw closure of trap-jaw ants (Gronenberg, 1995; Gronenberg, 1996; Patek et al., 2006), jumping of frogs (Roberts and Marsh, 2003) and bushbabies (Aerts, 1998), feeding of pipefish (Van Wassenbergh et al., 2008), and claw closure of snapping shrimp (Versluis et al., 2000). Numerous jumping insects also rely on high power output afforded by elastic recoil of springy tissues (Burrows, 2006; Burrows, 2009; Rothschild et al., 1975; Bennet-Clark and Lucey, 1967; Bennet-Clark, 1976a; Bennet-Clark, 1976b).

The mechanical properties of the tissues used to store strain energy in many of these systems have not been thoroughly studied, and the potential for thermal dependence of these properties is less well known. Nonetheless, some elastic proteins that serve as energy stores – collagen, resilin and abductin – have been found to have very low thermal dependence or complete independence (Alexander, 1966; Denny and Miller, 2006; Rigby et al., 1959). Movements that rely on recoil of these tissues to achieve their high performance may show low thermal dependence, such as jumping in fleas, which use recoil of resilin pads, in part, to launch themselves (Burrows, 2009), and suction feeding in pipefish, which rely on tendon recoil to snap the head upward (Van Wassenbergh et al., 2008). The thermal



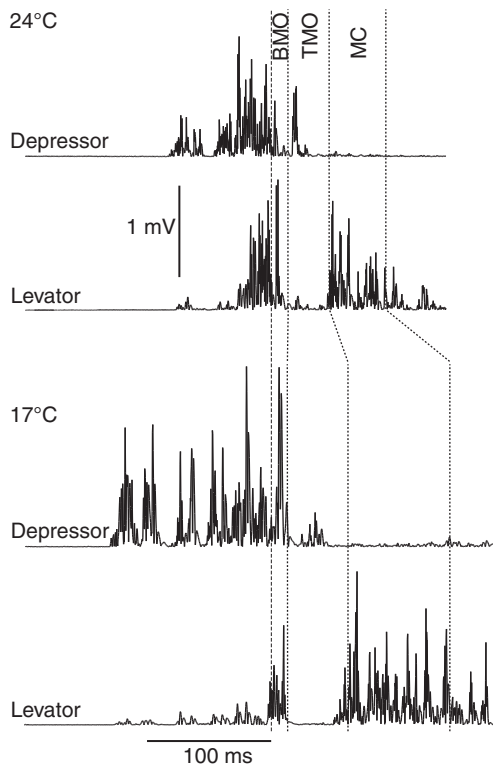


Fig. 7. Rectified EMG signals from the jaw depressor and levator muscles in the same individual of *B. terrestris* feeding at 24°C (top) and 17°C (bottom). Traces are aligned at the start of ballistic mouth opening (first dashed line) and the phases of the gape cycle are shown: ballistic mouth opening (BMO), transport mouth opening (TMO) and mouth closing (MC). Note the activation of the depressor muscle prior to mouth opening and the extended activation of both muscles at 17°C compared with 24°C. Amplitude of the muscles is similar at the two temperatures. All signals are shown on the same scale. Image sequences of these feedings are shown in Fig. 2.

dependence of the spring properties may be similarly low for the cuticular structures of arthropods, in which case the accompanying explosive movements are likely to maintain high performance across a broad range of temperatures. Some evidence exists in support of this; locusts have been reported to jump immediately upon removal from a 4°C cold room (Gabriel, 1985), and fleas have been found to retain jumping ability when cold and may use resilin as a temperature-independent energy store that allows them to jump from the cold ground to a warm host (Rothschild et al., 1975). Further research aimed at determining the temperature sensitivity of insect jumping performance and explosive actions of other arthropods would complement the current study.

#### Motor control of prey capture

Muscle activation patterns matched the elastic recoil mechanism of ballistic mouth opening. The mean duration of activity of the DM prior to mouth opening was 127 ms, which was 13 times the mean duration of ballistic mouth opening and sufficient time for the muscles to load elastic structures with strain energy. This duration of activity was nearly identical to the latency between the start of DM activity and the start of mouth opening, at 128 ms, and the range of durations (32–320 ms) encompassed the range of durations of DM activity (49–247 ms) recorded in *B. alvarius* (Lappin et al., 2006). The fact that DM activity ended 0.3 ms before mouth opening, on average, and rarely ended after the start of mouth opening (Tables 1 and 2)

is further evidence that ballistic opening is powered by recoil of elastic structures loaded prior to mouth opening.

Activation of the DM in *B. terrestris* for up to 320 ms prior to mandible movement is consistent with a ‘bow and arrow’ mechanism of elastic recoil, as has been found or implicated in many vertebrate and invertebrate systems that show activation of muscles well in advance of ballistic movement, including mantis shrimp and trap-jaw ants (Patek et al., 2004; Patek et al., 2006; Patek et al., 2007), various jumping insects (Burrows, 2006; Burrows, 2009), pipefish (Van Wassenbergh et al., 2008) and chameleons (Anderson and Deban, 2010; de Groot and van Leeuwen, 2004; Wainwright and Bennett, 1992a; Wainwright and Bennett, 1992b).

Activity of the LM was consistent with two functions: braking the mandible at the end of ballistic mouth opening, and closing the mouth. The latency between the peak of LM activity (first burst) and the end of ballistic opening was brief, averaging 16 ms, and covered a narrow range, whereas the activity associated with mouth closing (second burst) reached its peak a mean of 55 ms before the mouth closed.

The character of the LM activity was quite different from that of the DM, with activity of the LM dispersed in small bursts of varying durations and intervals, compared with the well-defined large burst of the DM prior to mouth opening. These differences match the different mechanisms of movement; the DM activates maximally to load elastic structures and ceases after the appropriate strain (or stress) is achieved, while the LM is active to drive movement of the mandible against the variable gravitational load imposed by the prey.

Measures of intensity of the DM EMG signal such as integrated area divided by duration, r.m.s. and peak r.m.s. amplitude showed no significant effect of temperature across either temperature range, with  $Q_{10}$  values of 0.8–1.2. This provides evidence that the muscle was recruited to the same degree at all temperatures, and thus excludes the possibility that toads were holding muscle fibers in reserve when warm and recruiting them when cold. The peak r.m.s. amplitude of the LM during mouth closing was also devoid of temperature effects. The thermal independence of the intensity of muscle activation found in the jaw muscles of *Bufo* contrasts with findings in muscle-powered movements such as swimming in fish (e.g. scup and carp), in which more fast, white muscle fibers are recruited when fish are cold (Rome et al., 1992a; Rome et al., 1984; Rome et al., 1990) to compensate for the loss of power at low temperatures.

Changes in activity of the DM and LM of *Bufo* associated with temperature were mainly changes in duration, and were more pronounced across the lower temperature range. Across the 11–25°C range, durations and latencies were longer at lower temperature ( $1/Q_{10}$  of 1.8–3.7) and integrated area was higher ( $1/Q_{10}$  of 2.2), consistent with the muscle fibers requiring longer to achieve tension or to shorten. Activity durations and latencies showed a plateau in the higher temperature range ( $1/Q_{10}$  of 0.9–1.5), which is consistent with a thermal plateau in muscle dynamics. A pattern of higher thermal dependence of muscle contractile rates at lower temperatures has frequently been found in muscles of other organisms (Bennett, 1984; Bennett, 1985; Putnam and Bennett, 1982; Hirano and Rome, 1984; John-Alder et al., 1989; Swoap et al., 1993; Stevenson and Josephson, 1990). The EMG data here thus indicate indirectly that toad jaw muscles are typical with regard to thermal effects on their contractile properties.

#### CONCLUSIONS

Ballistic prey capture in *B. terrestris* is accomplished by activation of jaw depressor muscles up to 320 ms prior to mouth opening



followed by inactivation as the mouth is rapidly opened and the tongue is projected. This early muscle activation combined with the high muscle mass-specific power output of ballistic mouth opening confirm that ballistic prey mouth opening is elastically powered (Lappin et al., 2006). As hypothesized, ballistic mouth opening shows virtual thermal independence, likely as a consequence of the low thermal dependence of both the isometric force developed by the depressor muscles and the elastic properties of the energy storage tissue. Further, the low thermal dependence of tongue projection dynamics confirms previous studies showing that the energy for tongue projection comes from ballistic mouth opening (Mallett et al., 2001). Non-ballistic prey transport movements, in contrast, are accomplished by direct muscle power with little contribution from elastic recoil, as evidenced by their higher thermal dependence at cold temperatures, which matches the thermal dependence that has been well established for the contractile rates of muscle (Bennett, 1984).

The motor patterns of the jaw muscles reveal constant and presumably maximum recruitment at all temperatures. The increase in duration of activity as temperature drops indicates a typical thermal response of slowing contractile properties at lower temperatures, which requires the muscles to remain active for longer during both ballistic and non-ballistic movements.

Our results reveal that the predatory strike of toads – ballistic mouth opening and tongue projection – is thermally independent across a broad temperature range. The elastic recoil mechanism expands the thermal breadth over which high performance prey capture can be accomplished. This in turn may allow toads to reduce thermoregulatory behavior and thus avoid associated costs in energy expenditure or water loss (Huey, 1974; Feder, 1982). *Bufo* has been shown in preference experiments to thermoregulate behaviorally near a temperature that maximizes sprinting performance when hydration is not an issue but to choose lower temperatures when water is limiting, revealing a potential conflict between performance and hydration (Lillywhite et al., 1973; Tracy et al., 1993). The elastic mechanism of prey capture may allow toads to partially circumvent this tradeoff by remaining cool and allowing muscle contractile performance to drop in circumstances where thermoregulating would cause undue water loss, while nonetheless maintaining high prey capture performance.

Finally, our results demonstrate that elastic recoil mechanisms can confer not just high performance but also the maintenance of high performance at temperatures that would normally hinder muscular performance. The uncoupling of muscle power from movement power afforded by elastic storage and release mechanisms frees the ballistic movement from factors such as temperature that influence muscle contractile rates. A similar effect has been demonstrated in the elastically powered tongue projection of chameleons (Anderson and Deban, 2010), which evolved independently from that of toads. We expect that the phenomenon of low thermal sensitivity of ballistic movements is widespread among organisms that use elastic recoil mechanisms.

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#### REFERENCES

- Aerts, P. (1998). Vertical jumping in *Galago senegalensis*: the quest for an obligate mechanical power amplifier. *Philos. Trans. R. Soc. Lond. B.* **353**, 1607-1620.
- Alexander, R. M. (1966). Rubber-like properties of the inner hinge-ligament of Pectinidae. *J. Exp. Biol.* **44**, 119-130.
- Altringham, J. D. and Block, B. A. (1997). Why do tuna maintain elevated slow muscle temperatures? Power output of muscle isolated from endothermic and ectothermic fish. *J. Exp. Biol.* **200**, 2617-2627.
- Anderson, C. V. and Deban, S. M. (2010). Ballistic tongue projection in chameleons maintains high performance at low temperature. *Proc. Natl. Acad. Sci. USA* **107**, 5495-5499.
- Asmussen, G., Beckers-Bleukx, G. and Marechal, G. (1994). The force-velocity relation of the rabbit inferior oblique muscle; influence of temperature. *Pflugers Arch.* **426**, 542-547.
- Barnes, W. S. and Ingalls, C. P. (1991). Differential effects of temperature on contractile behavior in isolated frog skeletal muscle. *Comp. Biochem. Physiol.* **100A**, 575-580.
- Bauwens, D., Garland, T. J., Castilla, A. M. and Van Damme, R. (1995). Evolution of sprint speed in lacertid lizards: morphological, physiological and behavioral covariation. *Evolution* **49**, 848-863.
- Bennet-Clark, H. C. (1975). The energetics of the jump of the locust *Schistocerca gregaria*. *J. Exp. Biol.* **63**, 53-83.
- Bennet-Clark, H. C. (1976a). Energy storage in jumping animals. *Perspect. Exp. Biol.* **1**, 467-479.
- Bennet-Clark, H. C. (1976b). Energy storage in jumping insects. In *The Insect Integument* (ed. H. R. Hepburn), pp. 421-443. London: Elsevier Scientific Publishing Company.
- Bennet-Clark, H. C. and Lucey, E. C. (1967). The jump of the flea: a study of the energetics and a model of the mechanism. *J. Exp. Biol.* **47**, 59-67.
- Bennett, A. F. (1984). Thermal dependence of muscle function. *Am. J. Physiol.* **247**, R217-R229.
- Bennett, A. F. (1985). Temperature and muscle. *J. Exp. Biol.* **115**, 333-344.
- Bennett, A. F. (1990). Thermal dependence of locomotor capacity. *Am. J. Physiol.* **259**, R253-R258.
- Burrows, M. (2006). Jumping performance of frog hopper insects. *J. Exp. Biol.* **209**, 4607-4621.
- Burrows, M. (2009). How fleas jump. *J. Exp. Biol.* **212**, 2881-2883.
- Choi, I. H., Cho, Y., Oh, Y. K., Jung, N. P. and Shin, H. C. (1998). Behavior and muscle performance in heterothermic bats. *Physiol. Zool.* **71**, 257-266.
- Cossins, A. R. and Bowler, K. (1987). *Temperature Biology of Animals*. London, New York: Chapman and Hall.
- de Groot, J. H. and van Leeuwen, J. L. (2004). Evidence for an elastic projection mechanism in the chameleon tongue. *Proc. R. Soc. B* **271**, 761-770.
- Deban, S. M., Wake, D. B. and Roth, G. (1997). Salamander with a ballistic tongue. *Nature* **389**, 27-28.
- Deban, S. M., O'Reilly, J. C., Dicke, U. and van Leeuwen, J. L. (2007). Extremely high-power tongue projection in plethodontid salamanders. *J. Exp. Biol.* **210**, 655-667.
- Denny, M. and Miller, L. (2006). Jet propulsion in the cold: mechanics of swimming in the Antarctic scallop *Adamussium colbecki*. *J. Exp. Biol.* **209**, 4503-4514.
- Donley, J. M., Shadwick, R. E., Sepulveda, C. A. and Syme, D. A. (2007). Thermal dependence of contractile properties of the aerobic locomotor muscle in the leopard shark and shortfin mako shark. *J. Exp. Biol.* **210**, 1194-1203.
- Eise, P. L. and Bennett, A. F. (1987). The thermal dependence of locomotor performance and muscle contractile function in the salamander *Ambystoma tigrinum nebulosum*. *J. Exp. Biol.* **128**, 219-233.
- Faulkner, J. A., Zerba, E. and Brooks, S. V. (1990). Muscle temperature of mammals: cooling impairs most functional properties. *Am. J. Physiol.* **259**, R259-R265.
- Feder, M. E. (1982). Thermal ecology of neotropical lungless salamanders (Amphibia: Plethodontidae): environmental temperatures and behavioral responses. *Ecology* **63**, 1665-1674.
- Gabriel, J. M. (1985). The development of the locust jumping mechanism: II. Energy storage and muscle mechanics. *J. Exp. Biol.* **118**, 327-340.
- Gronenberg, W. (1995). The fast mandible strike in the trap-jaw ant *Odontomachus*. *J. Comp. Physiol. A* **176**, 399-408.
- Gronenberg, W. (1996). The trap-jaw mechanism in the dacetine ants *Daceton armigerum* and *Strumigenys* sp. *J. Exp. Biol.* **199**, 2021-2033.
- Herrel, A., James, R. S. and Van Damme, R. (2007). Fight versus flight: physiological basis for temperature-dependent behavioral shifts in lizards. *J. Exp. Biol.* **210**, 1762-1767.
- Hill, A. V. (1951). The influence of temperature on the tension developed in an isometric twitch. *Proc. R. Soc. B* **1951**, 349-354.
- Hill, R. W., Wyse, G. A. and Anderson, M. (2008). *Animal Physiology*. Sunderland, MA: Sinauer Associates.
- Hirano, M. and Rome, L. C. (1984). Jumping performance of frogs (*Rana pipiens*) as a function of muscle temperature. *J. Exp. Biol.* **108**, 429-439.
- Hochachka, P. W. and Somero, G. N. (2002). *Biochemical Adaptation: Mechanism and Process in Physiological Evolution*. New York: Oxford University Press.
- Huey, R. B. (1974). Behavioral thermoregulation in lizards: importance of associated costs. *Science* **184**, 1001-1003.
- Huey, R. B. and Bennett, A. F. (1987). Phylogenetic studies of coadaptation preferred temperatures versus optimal performance temperatures of lizards. *Evolution* **41**, 1098-1115.
- Huey, R. B. and Kingsolver, J. G. (1989). Evolution of thermal sensitivity of ectotherm performance. *Trends Ecol. Evol.* **4**, 131-135.
- Huey, R. B. and Kingsolver, J. G. (1993). Evolution of resistance to high temperature in ectotherms. *Am. Nat.* **142**, S21-S46.
- Huey, R. B. and Stevenson, R. D. (1979). Integrating thermal physiology and ecology of ectotherms: a discussion of approaches. *Am. Zool.* **19**, 357-366.
- John-Alder, H. B., Morin, P. J. and Sharon, L. (1988). Thermal physiology, phenology, and distribution of tree frogs. *Am. Nat.* **132**, 506-520.
- John-Alder, H. B., Barnhart, M. C. and Bennett, A. F. (1989). Thermal sensitivity of swimming performance and muscle contraction in northern and southern populations of tree frogs (*Hyla crucifer*). *J. Exp. Biol.* **142**, 357-372.

- Knowles, T. W. and Weigl, P. D.** (1990). Thermal dependence of anuran burst locomotor performance. *Copeia* **1990**, 796-802.
- Lappin, A. K., Monroy, J. A., Pilarski, J. Q., Zepnewski, E. D., Pierotti, D. J. and Nishikawa, K. C.** (2006). Storage and recovery of elastic potential energy powers ballistic prey capture in toads. *J. Exp. Biol.* **209**, 2535-2553.
- Lillywhite, H. B., Licht, P. and Chelgren, P.** (1973). The role of behavioral thermoregulation in the growth energetics of the toad, *Bufo boreas*. *Ecology* **54**, 375-383.
- Lutz, G. J. and Rome, L. C.** (1994). Built for jumping: the design of the frog muscular system. *Science* **263**, 370-372.
- Lutz, G. J. and Rome, L. C.** (1996). Muscle function during jumping in frogs. II. Mechanical properties of muscle: implications for system design. *Am. J. Physiol.* **271**, C571-C578.
- Mallett, E. S., Yamaguchi, G. T., Birch, J. M. and Nishikawa, K.** (2001). Feeding motor patterns in anurans: insights from biomechanical modeling. *Am. Zool.* **41**, 1364-1374.
- Marsh, R. L. and Bennett, A. F.** (1985). Thermal dependence of isotonic contractile properties of skeletal muscle and sprint performance of the lizard *Dipsosaurus dorsalis*. *J. Comp. Physiol. B* **155**, 541-551.
- Marsh, R. L. and Bennett, A. F.** (1986). Thermal dependence of sprint performance of the lizard *Sceloporus occidentalis*. *J. Exp. Biol.* **126**, 79-87.
- Marvin, G. A.** (2003a). Aquatic and terrestrial locomotor performance in a semiaquatic plethodontid salamander (*Pseudotriton ruber*): Influence of acute temperature, thermal acclimation, and body size. *Copeia* **2003**, 704-713.
- Marvin, G. A.** (2003b). Effects of acute temperature and thermal acclimation on aquatic and terrestrial locomotor performance of the three-lined salamander, *Eurycea guttolineata*. *J. Therm. Biol.* **28**, 251-259.
- Mautz, W. J., Daniels, C. B. and Bennett, A. F.** (1992). Thermal dependence of locomotion and aggression in a xantusiid lizard. *Herpetologica* **48**, 271-279.
- McLister, J. D.** (2001). Physical factors affecting the cost and efficiency of sound production in the treefrog *Hyla versicolor*. *J. Exp. Biol.* **204**, 69-80.
- Navas, C. A., James, R. S., Wakeling, J. M., Kemp, K. M. and Johnston, I. A.** (1999). An integrative study of the temperature dependence of whole animal and muscle performance during jumping and swimming in the frog *Rana temporaria*. *J. Comp. Physiol. B* **169**, 588-596.
- Nishikawa, K. C.** (2000). Feeding in frogs. In *Feeding: Form, Function, and Evolution in Tetrapod Vertebrates* (ed. K. Schwenk), pp. 117-147. San Diego: Academic Press.
- Patek, S. N., Korff, W. L. and Caldwell, R. L.** (2004). Deadly strike mechanism of a mantis shrimp. *Nature* **428**, 819-820.
- Patek, S. N., Baio, J. E., Fisher, B. L. and Suarez, A. V.** (2006). Multifunctionality and mechanical origins: ballistic jaw propulsion in trap-jaw ants. *Proc. Natl. Acad. Sci. USA* **103**, 12787-12792.
- Patek, S. N., Nowroozi, B. N., Baio, J. E., Caldwell, R. L. and Summers, A. P.** (2007). Linkage mechanics and power amplification of the mantis shrimp's strike. *J. Exp. Biol.* **210**, 3677-3688.
- Peplowski, M. M. and Marsh, R. L.** (1997). Work and power output in the hindlimb muscles of Cuban tree frogs *Osteopilus septentrionalis* during jumping. *J. Exp. Biol.* **200**, 2861-2870.
- Putnam, R. W. and Bennett, A. F.** (1982). Thermal dependence of isometric contractile properties of lizard muscle. *J. Comp. Physiol. B* **147**, 11-20.
- Renaud, J. M. and Stevens, E. D.** (1984). The extent of short-term and long-term compensation to temperature shown by frog and toad sartorius muscle. *J. Exp. Biol.* **108**, 57-75.
- Rigby, B. J., Hirai, N., Spikes, J. D. and Eyring, H.** (1959). The mechanical properties of rat tail tendon. *J. Gen. Physiol.* **43**, 265-283.
- Roberts, T. J. and Marsh, R. L.** (2003). Probing the limits to muscle-powered accelerations: lessons from jumping bullfrogs. *J. Exp. Biol.* **206**, 2567-2580.
- Rome, L. C., Loughna, P. T. and Goldspink, G.** (1984). Muscle fiber activity in carp as a function of swimming speed and muscle temperature. *Am. J. Physiol.* **247**, R272-R279.
- Rome, L. C., Funke, R. P. and Alexander, R. M.** (1990). The influence of temperature on muscle velocity and sustained performance in swimming carp. *J. Exp. Biol.* **154**, 163-178.
- Rome, L. C., Choi, I. H., Lutz, G. and Sosnicki, A.** (1992a). The influence of temperature on muscle function in the fast swimming scup. I. Shortening velocity and muscle recruitment during swimming. *J. Exp. Biol.* **163**, 259-279.
- Rome, L. C., Stevens, D. and John-Alder, H. B.** (1992b). The influence of temperature and thermal acclimation on physiological function. In *Environmental Physiology of the Amphibians* (ed. M. E. Feder and W. W. Burggren), pp. 183-205. Chicago: University of Chicago Press.
- Rothschild, M., Schlein, J., Parker, K., Neville, C. and Sternberg, S.** (1975). The jumping mechanism of *Xenopsylla cheopis*. III. Execution of the jump and activity. *Philos. Trans. R. Soc. Lond. B Biol. Sci.* **271**, 499-515.
- Sobol, C. V. and Nasedlov, G. A.** (1994). Thermal dependence of force-velocity relation of lamprey live striated muscle fibres. *Gen. Physiol. Biophys.* **13**, 215-224.
- Stevenson, R. D. and Josephson, R. K.** (1990). Effects of operating frequency and temperature on mechanical power output from moth flight muscle. *J. Exp. Biol.* **149**, 61-78.
- Swoap, S. J., Johnson, T. P., Josephson, R. K. and Bennett, A. F.** (1993). Temperature, muscle power output and limitations on burst locomotor performance of the lizard *Dipsosaurus dorsalis*. *J. Exp. Biol.* **174**, 185-197.
- Tracy, C. R., Christian, K. A. and O'Connor, M. P.** (1993). Behavioral thermoregulation by *Bufo americanus*: the importance of the hydric environment. *Herpetologica* **49**, 375-382.
- van Berkum, F. H.** (1986). Evolutionary patterns of the thermal sensitivity of sprint speed in anolis lizards. *Evolution* **40**, 594-604.
- Van Leeuwen, J. L., de Groot, J. H. and Kier, W. M.** (2000). Evolutionary mechanics of protrusible tentacles and tongues. *Neth. J. Zool.* **50**, 113-139.
- Van Wassenbergh, S., Strother, J. A., Flammang, B. E., Ferry-Graham, L. A. and Aerts, P.** (2008). Extremely fast prey capture in pipefish is powered by elastic recoil. *J. R. Soc. Interface* **5**, 285-296.
- Versluis, M., Schmitz, B., von der Heydt, A. and Lohse, D.** (2000). How snapping shrimp snap: through cavitating bubbles. *Science* **289**, 2114.
- Wainwright, P. C. and Bennett, A. F.** (1992a). The mechanism of tongue projection in chameleons. 1. Electromyographic tests of functional hypotheses. *J. Exp. Biol.* **168**, 1-21.
- Wainwright, P. C. and Bennett, A. F.** (1992b). The mechanism of tongue projection in chameleons. 2. Role of shape change in a muscular hydrostat. *J. Exp. Biol.* **168**, 23-40.
- Wilson, R. S. and Franklin, C. E.** (1999). Thermal acclimation of locomotor performance in tadpoles of the frog *Limnodynastes peronii*. *J. Comp. Physiol. B* **169**, 445-451.
- Wilson, R. S. and Franklin, C. E.** (2000). Inability of adult *Limnodynastes peronii* (Amphibia: Anura) to thermally acclimate locomotor performance. *Comp. Biochem. Physiol.* **127A**, 21-28.
- Zack, T. I., Claverie, T. and Patek, S. N.** (2009). Elastic energy storage in the mantis shrimp's fast predatory strike. *J. Exp. Biol.* **212**, 4002.

Table S1. Results of ANCOVA examining effects on kinematic, dynamic and electromyographic variables over the 11-25°C range in four *Bufo terrestris*. Prey distance was included as a covariate only when it showed a significant effect for that variable. Bold values with asterisks indicate significant effects at the Bonferroni-corrected alpha (0.0019 for 27 variables). Data from four individual toads were included in all analyses.

	Individual				Temperature				Prey Distance			
	# Trials	F-ratio	df	P-value	# Trials	F-ratio	df	P-value	# Trials	F-ratio	df	P-value
<b>Kinematic variables</b>												
Duration of ballistic mouth opening	56	0.481	3,51	0.6973	56	2.192	1,51	0.1449	35	2.689	1,29	0.1119
Duration of tongue projection	55	4.947	3,50	0.0044	55	1.969	1,50	0.1668	35	0.844	1,29	0.3659
Gape distance at end of ballistic opening	31	6.485	3,26	0.0020	31	0.028	1,26	0.8688	28	7.712	1,22	0.0110
Maximum tongue reach	34	4.037	3,28	0.0167	34	2.232	1,28	0.1464	34	40.302	1,28	<b>&lt;0.0001*</b>
Duration of mouth opening during transport	56	0.725	3,51	0.5418	56	15.203	1,51	<b>0.0003*</b>	35	0.203	1,29	0.6559
Final gape distance	34	19.786	3,29	<b>&lt;0.0001*</b>	34	1.251	1,29	0.2725	34	8.539	1,28	0.0068
Duration of tongue retraction	55	3.369	3,50	0.0256	55	11.834	1,50	<b>0.0012*</b>	35	5.004	1,29	0.0331
Duration of mouth closing	56	1.248	3,51	0.3021	56	114.627	1,51	<b>&lt;0.0001*</b>	35	0.342	1,29	0.5633
<b>Dynamic variables</b>												
Average velocity of ballistic mouth opening	34	50.055	3,29	<b>&lt;0.0001*</b>	34	7.157	1,29	0.0121	34	3.660	1,28	0.0660
Maximum velocity of ballistic opening	31	85.455	3,26	<b>&lt;0.0001*</b>	31	2.287	1,26	0.1425	28	5.683	1,22	0.0262
Maximum acceleration of ballistic opening	31	34.044	3,26	<b>&lt;0.0001*</b>	31	0.018	1,26	0.8950	28	0.229	1,22	0.6371
Maximum power of ballistic opening	31	32.061	3,26	<b>&lt;0.0001*</b>	31	0.631	1,26	0.4343	28	3.782	1,22	0.0647
Average velocity of tongue projection	34	12.927	3,29	<b>&lt;0.0001*</b>	34	3.164	1,29	0.0857	34	1.252	1,28	0.2727
Average velocity of mouth opening during transport	34	0.611	3,29	0.6133	34	35.580	1,29	<b>&lt;0.0001*</b>	34	3.110	1,28	0.0887
Average velocity of tongue retraction	34	2.742	3,28	0.0619	34	28.270	1,28	<b>&lt;0.0001*</b>	34	20.899	1,28	<b>&lt;0.0001*</b>
Average velocity of mouth closing	34	7.098	3,29	<b>0.0010*</b>	34	122.424	1,29	<b>&lt;0.0001*</b>	34	0.000	1,28	0.9841
<b>Electromyographic variables</b>												
Depressor EMG integrated area	35	14.494	3,29	<b>&lt;0.0001*</b>	35	125.225	1,29	<b>&lt;0.0001*</b>	35	125.225	1,29	<b>&lt;0.0001*</b>
Depressor EMG duration	52	18.795	3,47	<b>&lt;0.0001*</b>	52	244.604	1,47	<b>&lt;0.0001*</b>	35	2.245	1,29	0.1449
Depressor EMG RMS	52	5.189	3,47	0.0035	52	5.328	1,47	0.0254	35	8.836	1,29	0.0059
Depressor EMG integrated area/duration	52	1.676	3,47	0.1850	52	1.979	1,47	0.1661	35	10.332	1,29	0.0032
Depressor EMG RMS max. amplitude	52	0.427	3,47	0.7349	52	2.400	1,47	0.1280	35	4.014	1,29	0.0545
Levator EMG (second burst) RMS max. amplitude	53	29.468	3,48	<b>&lt;0.0001*</b>	53	3.315	1,48	0.0749	35	0.868	1,29	0.3593
Depressor EMG start to start of ballistic opening	52	4.964	3,47	0.0045	52	91.734	1,47	<b>&lt;0.0001*</b>	35	8.439	1,29	0.0070
Depressor EMG end to start of ballistic opening	52	5.243	3,47	0.0033	52	1.120	1,47	0.2954	35	0.002	1,29	0.9636
Depressor EMG max. amp. to start of ballistic opening	52	0.086	3,47	0.9674	52	20.159	1,47	<b>&lt;0.0001*</b>	35	0.590	1,29	0.4487
Levator EMG (first burst) max. amp to end ballistic open	52	1.590	3,47	0.2043	52	5.338	1,47	0.0253	35	0.306	1,29	0.5845
Levator EMG (second burst) max. amp to mouth closed	53	6.371	3,48	<b>0.0010*</b>	53	11.640	1,48	<b>0.0013*</b>	35	0.011	1,29	0.9162

Table S2. Results of ANCOVA examining effects on kinematic, dynamic and electromyographic variables over the 20-35°C range in four *Bufo terrestris*. Prey distance was included as a covariate only when it showed a significant effect for that variable. Bold values with asterisks indicate significant effects at the Bonferroni-corrected alpha (0.0019 for 27 variables). Data from four individual toads were included in all analyses.

	Individual				Temperature				Prey Distance			
	# Trials	F-ratio	df	P-value	# Trials	F-ratio	df	P-value	# Trials	F-ratio	df	P-value
<b>Kinematic variables</b>												
Duration of ballistic mouth opening	38	45.012	3,33	<b>&lt;0.0001*</b>	38	24.722	1,33	<b>&lt;0.0001*</b>	23	3.556	1,17	0.0765
Duration of tongue projection	38	10.853	3,33	<b>&lt;0.0001*</b>	38	1.324	1,33	0.2582	23	0.027	1,17	0.8717
Gape distance at end of ballistic opening	23	3.552	3,18	0.0353	23	2.895	1,18	0.1061	22	1.541	1,16	0.2323
Maximum tongue reach	23	7.660	3,17	0.0019	23	5.649	1,17	0.0295	23	42.591	1,17	<b>&lt;0.0001*</b>
Duration of mouth opening during transport	38	3.959	3,33	0.0162	38	17.192	1,33	<b>0.0002*</b>	23	0.432	1,17	0.5201
Final gape distance	24	4.073	3,19	0.0216	24	28.035	1,19	<b>&lt;0.0001*</b>	24	1.204	1,18	0.287
Duration of tongue retraction	38	12.449	3,33	<b>&lt;0.0001*</b>	38	5.739	1,33	0.0224	23	1.161	1,17	0.2962
Duration of mouth closing	38	11.509	3,33	<b>&lt;0.0001*</b>	38	1.183	1,33	0.2846	23	3.847	1,17	0.0664
<b>Dynamic variables</b>												
Average velocity of ballistic mouth opening	24	5.586	3,19	0.0064	24	0.122	1,19	0.7305	23	5.939	1,17	0.0261
Maximum velocity of ballistic opening	23	75.977	3,18	<b>&lt;0.0001*</b>	23	2.573	1,18	0.1261	22	2.617	1,16	0.1253
Maximum acceleration of ballistic opening	23	25.565	3,18	<b>&lt;0.0001*</b>	23	1.038	1,18	0.3219	22	0.167	1,16	0.688
Maximum power of ballistic opening	23	27.777	3,18	<b>&lt;0.0001*</b>	23	2.041	1,18	0.1702	22	0.393	1,16	0.5394
Average velocity of tongue projection	24	6.753	3,19	0.0027	24	6.985	1,19	0.0160	23	1.517	1,17	0.2348
Average velocity of mouth opening during transport	22	0.403	3,17	0.7525	22	0.008	1,17	0.9304	22	0.072	1,16	0.7919
Average velocity of tongue retraction	24	4.122	3,19	0.0207	24	0.035	1,19	0.8545	23	2.544	1,17	0.1292
Average velocity of mouth closing	24	13.190	3,19	<b>&lt;0.0001*</b>	24	2.944	1,19	0.1025	24	3.025	1,18	0.0991
<b>Electromyographic variables</b>												
Depressor EMG integrated area	37	12.311	3,32	<b>&lt;0.0001*</b>	37	17.048	1,32	<b>0.0002*</b>	22	2.002	1,16	0.1762
Depressor EMG duration	37	7.316	3,32	<b>0.0007*</b>	37	2.673	1,32	0.1119	22	5.646	1,16	0.0303
Depressor EMG RMS	37	35.935	3,32	<b>&lt;0.0001*</b>	37	7.825	1,32	0.0086	22	0.214	1,16	0.6499
Depressor EMG integrated area/duration	37	30.517	3,32	<b>&lt;0.0001*</b>	37	6.227	1,32	0.0179	22	1.368	1,16	0.2593
Depressor EMG RMS max. amplitude	37	26.974	3,32	<b>&lt;0.0001*</b>	37	7.505	1,32	0.0100	22	0.192	1,16	0.667
Levator EMG (second burst) RMS max. amplitude	37	56.100	3,32	<b>&lt;0.0001*</b>	37	4.743	1,32	0.0369	22	1.268	1,16	0.2768
Depressor EMG start to start of ballistic opening	37	1.652	3,32	0.1970	37	5.408	1,32	0.0265	22	13.542	1,16	0.0020
Depressor EMG end to start of ballistic opening	37	3.902	3,32	0.0175	37	3.136	1,32	0.0861	22	2.500	1,16	0.1334
Depressor EMG max. amp. to start of ballistic opening	37	0.215	3,32	0.8851	37	0.286	1,32	0.5968	22	1.542	1,16	0.2322
Levator EMG (first burst) max. amp to end ballistic open	37	2.527	3,32	0.0749	37	0.100	1,32	0.7538	22	0.655	1,16	0.4303
Levator EMG (second burst) max. amp to mouth closed	37	4.188	3,32	0.0131	37	0.041	1,32	0.8417	22	3.904	1,16	0.0657