Aimed limb movements in a hemimetabolous insect are intrinsically compensated for allometric wing growth by developmental mechanisms

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ABSTRACT
For aimed limb movements to remain functional, they must be adapted to developmental changes in body morphology and sensory-motor systems. Insects use their limbs to groom the body surface or to dislodge external stimuli, but they face the particular problem of adapting these movements to step-like changes in body morphology during metamorphosis or moulting. Locusts are hemimetabolous insects in which the imaginal moul to adulthood results in a sudden and dramatic allometric growth of the wings relative to the body and the legs. We show that, despite this, hind limb scratches aimed at mechanosensory stimuli on the wings remain targeted to appropriate locations after moulting. In juveniles, the tips of the wings extend less than halfway along the abdomen, but in adults they extend well beyond the posterior end. Kinematic analyses were used to examine the scratching responses of juveniles (fifth instars) and adults to touch of anterior (wing base) and posterior (distal abdomen) targets that develop isometrically, and to wing tip targets that are anterior in juveniles but posterior in adults. Juveniles reach the (anterior) wing tip with the distal tibia of the hind leg using anterior rotation of the thoraco-coxal and coxo-trochanteral (‘hip’) joints and flexion of the femoro-tibial (‘knee’) joint. Adults, however, reach the corresponding (but now posterior) wing tip using posterior rotation of the hip and extension of the knee, reflecting a different underlying motor pattern. This change in kinematics occurs immediately after the adult moult without learning, indicating that the switch is developmentally programmed.

KEY WORDS: Development, Scratching, Reaching, Locust, Schistocerca gregaria, Limb growth

INTRODUCTION
Fundamentally important behaviours such as reaching, stepping, grooming and prey capture all require accurately aimed limb movements. The neuronal networks underlying these movements must be adapted to changes in body morphology during development so that they remain functional throughout life. Sensory-motor plasticity is thus critical for tuning and maintaining behaviours during development (Easter, 1983). For example, human infants begin reaching towards targets at age 4–5 months, but the kinematics of their movements change over the first 3 years of life before assuming their adult-like patterns (Konczak and Dichgans, 1997). This reflects continual recalibration of the sensory-motor system as both the peripheral mechanics and the central nervous system develop (Konczak et al., 1997). Adult locusts spontaneously walk across larger gaps than juveniles, apparently judging the distance using visual cues. The change in maximum gap width crossed is reportedly not expressed immediately after the moult, but becomes apparent with some delay, and appears to be facilitated by a period of interaction with a complex external environment (Ben-Nun et al., 2013) – again suggesting age-dependent recalibration. Praying mantises that capture prey using visually triggered strikes of their raptorial front legs have longer strike distances as adults than as juveniles. Striking is triggered at age- and body-size-appropriate distances largely as a result of concomitant changes in the size and shape of the eyes and the size of the body and legs at each moult (Baladerrama and Maldonado, 1973; Maldonado et al., 1974; Kral and Poteser, 2009) – so here it seems that experience-dependent ‘recalibration’ is not required.

In the case of grooming movements aimed at a target elsewhere on the body, differential growth of body and limbs can cause a change in the spatial relationship between the target and the grooming limb. Changes in the sensory system, such as addition or loss of exteroceptors or changes in their properties, may lead to changes in the representation of both the body surface and features of the external environment (Murphey et al., 1980; Chiba et al., 1988). Growth leads to changes in limb lengths and musculoskeletal properties, which, given an unchanged motor pattern, would result in different and inappropriate movement trajectories. If behaviours are to be conserved during growth and development, then plasticity of the underlying neuronal pathways needs to match the biomechanical changes.

In locust scratching, deflection of tactile hairs on a wing elicits movement of the ipsilateral hind leg, with the distal end of the tibia and tarsus aimed toward the site of stimulation (Berkowitz and Laurent, 1996a; Matheson, 1997, 1998; Dürr and Matheson, 2003). Furthermore, stimulation of different locations along the wing surface elicits tarsal trajectories that are target specific from onset and accurately aimed to each stimulus site (Berkowitz and Laurent, 1996a; Matheson, 1997, 1998; Dürr and Matheson, 2003). To reach progressively more proximal targets requires increases in the elevation of the femur and increased flexion of the femoro-tibial joint (Fig. 1). These continuously graded changes in hind leg joint angles suggest that adult locusts use a single, systemically varied movement pattern for scratching, rather than using distinct motor strategies specific to each location within the wing receptive field (Dürr and Matheson, 2003). In addition, locusts compensate their
scratching movements against perturbations caused by loading the hind leg (Matheson and Dürr, 2003). This plastic behaviour provides an opportunity to investigate how aimed movements adapt to morphological changes that arise during development.

Hemimetabolous insects such as locusts maintain essentially the same body structure throughout life, enlarging abruptly at each successive moult (Uvarov, 1966). At the adult (imaginal) moult, however, wing length increases disproportionately more than total hind leg length. This allometric growth means that the wing tips lie relatively much further posterior in adults than in the last juvenile stage (fifth instar). To remain functional, scratching movements elicited by tactile stimulation of the wings must therefore adapt to the relative changes in limb and wing dimensions. To complicate matters further, there are also two morphological rotations of the wings during development (Roonwal, 1940, 1946; Ivanova, 1947; Burnett, 1951; Uvarov, 1966; Dirsh, 1967; Khatkar, 1972). In *Schistocerca gregaria*, wing pads develop in first instars as ventrally directed extensions of the dorsal cuticular plates. The fore wing pad lies anterior to the hind wing pad, and both have their dorsal surfaces outermost. The wing pads remain in the same position in the second and third instars (Fig. 1Aii,ii), but during the moult to fourth instar, both wing pads rotate: the ventral surfaces turn outermost, and the hind wing pad now largely covers the fore wing pad (Fig. 1Aii; Roonwal, 1940). In fifth instars, the wing pads remain in the same position (Fig. 1Aiv), but on moulting to adulthood, they rotate again so that the dorsal surfaces of both wings are outermost, and the fore wing now covers the hind wing at rest (Fig. 1Av; Roonwal, 1952). In adults, therefore, it is the dorsal surface of the fore wing that is outermost, whereas in fifth instars it is the ventral surface of the hind wing that is outermost (Fig. 1Av versus Aiv).

Tactile stimuli that elicit aimed scratching movements are detected by wing mechanosensory hairs (Page and Matheson, 2004). Developmental rotation of the wings at the imaginal moult means that those hairs which elicit scratching in resting fifth instars and adults are necessarily located on different wing surfaces. In fifth instars (where it is the ventral hind wing that is exposed), the sensory pathway originates from metathoracic wing hairs and passes to the metathoracic ganglion, where the interneurons and motor neurons driving hind leg movement are located (Matheson, 1997). In adults, the sensory pathway originates from mesothoracic wing hairs on the exposed dorsal fore wing surface (Page and Matheson, 2004) and passes via the mesothoracic ganglion (Campbell, 1961) and descending connectives to eventually drive the same metathoracic motor networks (Matheson, 2002). At the imaginal moult, locusts must therefore adapt their scratching movements to account not only for allometric elongation of the wing sensory surface, but also for a change from receptors on one surface to those on another.

Here, we describe the scratching movements of fifth instar locusts and show that for similar targets they have similar kinematics to those of adults. Both fifth instar and adult locusts accurately aim their hind limb movements at distal wing tip targets despite the profound developmental changes that occur at the imaginal moult. The compensation for wing allometry and developmental rotation is apparent immediately after the moult, indicating that the animal’s internal body image is adjusted by developmental rather than learned mechanisms.

**MATERIALS AND METHODS**

**Animals and experimental protocol**

**Movement kinematics**

Experiments were carried out on six fifth instar (3 male, 3 female) and six adult (3 male, 3 female) desert locusts (*Schistocerca gregaria*).
Forskål 1775) taken from a laboratory culture established at the University of Leicester, UK, with stock from Blades Biological Ltd (Kent, UK) and the Department of Zoology, University of Cambridge, UK. Animals were maintained in crowded cages under a 12 h:12 h light:dark photoperiod with a corresponding 36°C:25°C thermal regimen, and fed ad libitum with fresh wheat seedlings and organic wheat bran. For experiments, animals were tethered with a fine loop of wire that passed around their pronotum without obstructing movements of their legs. They were suspended above a foam ball on which they could stand or walk. The tether allowed each animal to adjust its body posture, with the exception of thorax height above the substratum, which was set to the height normally maintained by a walking locust. The hind leg was manipulated into a standardised start posture (black triangles in Fig. 2Ai,Bi) by placing the tarsus on a horizontal rod located at 67% of the distance between the anterior rim of the metathoracic coxal joint and the tip of the abdomen (Dürr and Matheson, 2003). The eyes and ocelli were covered with black acrylic paint (Cryla, Daler-Rowney, Bracknell, UK) to exclude visual cues. Experiments were carried out at 22–24°C, maintained by an infrared heat lamp suspended 50 cm above the animal. To facilitate video-based motion tracking, 1 mm diameter discs cut from reflective tape (3M Scotchlite) were attached to the body and the right hind leg with insect glue [Thorne (Beehives) Ltd, UK] (Fig. 2Aii,Bii). Movement of the unguis was ignored. For illustrative purposes, tarsal length was standardised to 6.75 mm. See Dürr and Matheson (2003) for additional details. All animals that responded with aimed limb movements to touch of all three sites were included in the analyses.

Scratching movements of the right hind leg ipsilateral to stimulus sites on a wing were analysed in the following way. All ipsilateral hind leg movements elicited by touch were included in the analyses, except for obvious struggling movements which typically involved several limbs, or movements in which the animal stepped from the start platform directly to the ground. The adult fore wing was notionally divided into five regions of equal length. The most anterior and posterior of these regions were used as ‘wing base’ and ‘wing tip’ stimulus sites, respectively, and the posterior fifth of the abdomen was used as a posterior ‘abdomen tip’ target (Fig. 2Ai). For fifth instar locusts, the wing pad was divided into anterior ‘wing base’ and posterior ‘wing tip’ regions (Fig. 2Bi). Stimulation was provided by gently touching the wing base, wing tip or abdomen tip with a fine paintbrush to elicit scratching behaviour (mean±s.e.m. stimulus positions are indicated by coloured squares in Fig. 2Ai,Bi). Wing base, wing tip and abdomen tip stimuli were given randomly until at least 10 scratches were recorded for each stimulus site. The main results are based on analyses of 793 scratches in 12 locusts. Sample sizes were guided by our published data (e.g. Dürr and Matheson, 2003). Power calculations were not carried out.

Locusts rarely touched their own body surface during the first few cycles of a scratching response, so responses were open-loop with respect to the stimulus. Responses in which the leg made contact with the body surface or with the stimulating brush were analysed up

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**Fig. 2. Body morphology and points of reference for kinematic analyses.** Coordinate frame of reference, start and stimulus (‘target’) positions for adults (Ai) and fifth instars (Bi). Squares indicate mean stimulus locations on the wing base (yellow), wing tip (red) or abdomen tip (black). Errors (±s.e.m.) lie within the symbol for all targets (N=6 fifth instars and 6 adults). The hind leg tarsus was placed on a rod at a single start position (black triangle, mean±s.e.m.). The body-centred coordinate frame of reference had its origin on the anterior rim of the hind leg coxa for adults (black axes in Ai) and was shifted so that wing base and abdomen tip stimuli in fifth instars were aligned with those of the adults (broken lines). Movements were tracked using reflective markers placed on the body and limbs (black circles in Ai,Bi) and fitted with a body model (black lines joining circles in Ai,Bi). For a single scratch it was thus possible to visualise in each video frame the locations of the hind leg segments (Ci, red lines; this example is fifth instar, wing base stimulation) and wing midline (blue line) relative to the thoracic frame of reference (short black lines).

The abdomen (long black line, partly obscured) was marked in the first frame of each sequence only, but did not move during a scratch. The corresponding trajectories of the hind leg tibio-tarsal joint (red dots and line) and femoro-tibial joint (purple dots and line) are shown in Cii (open triangle indicates stimulation site). Grey lines show the tarsus position in each frame (20 ms between frames), and the blue line indicates the initial movement vector for the first 200 ms of movement.
until the time of first contact. Contact was determined by visual inspection of the video.

**Responses to dorsal fore wing stimulation in fifth instars**

Ten female and two male fifth instar animals were used to examine responses to touch of the normally hidden dorsal fore wing surface. Animals were tethered as described above and their eyes were covered with black paint. They walked freely on the ball and the tibia was not placed onto a standard starting rod. The left hind wing bud and both of the wing buds on the right were ablated using fine scissors, leaving only short stumps of approximately one-quarter of the total length of each wing bud. The inner (morphologically dorsal) surface of the remaining intact left fore wing bud was then stimulated with a fine paintbrush. Each animal received 10 stimuli at 1 min intervals. All but one female animal responded to at least one stimulus with an aimed scratching movement (mean 3.4 responses per animal, median 3.5 responses per animal). The non-responding animal was omitted from further analysis. The responses were scored as: anterior, if the femur was strongly rotated anteriorly so that the tarsus passed anterior to the midpoint of the abdomen (which corresponds approximately to the tip of the wing bud, see Fig. 1Bi); middle, if the femur rotated anteriorly so that the tarsus passed between the tarsus and the midpoint of the abdomen; and posterior, if the femur was rotated posteriorly so that the tarsus passed posterior to the tip of the abdomen (as it would for a scratch aimed at the abdomen tip or adult wing tip (see Fig. 1Bi,ii)). Trials that elicited no scratching response were scored as ‘none’. Each animal was also tested with a touch to the tip of the abdomen, which in all cases led to robust posterior scratches.

**Development of aimed movements following the imaginal moult**

Nine animals were observed moulting from fifth instar to adult in their normal home cage. A fine paintbrush was used to touch the dorsal distal fore wing tip and other locations on the wings and body. Animals were observed continuously from the point at which the old exoskeleton split, and were stimulated periodically until the wings were expanded and rotated into their adult orientation. After emergence from the old exoskeleton, animals hung from a vertical support, head up, while the adult wings first expanded in their juvenile orientations (i.e. with the hind wing outmost, and both wings rotated so that their dorsal surfaces faced the body). During this period, animals remained in the same posture with little movement unless disturbed. In a brief movement, animals then rotated and adjusted the expanded wings so that they assumed their normal adult orientations. Responses to all stimulations were recorded.

A further 13 animals (6 male, 7 female) were also observed moulting, and had both hind legs restrained within 1 min after emergence from the old exoskeleton – before the animal had a chance to make any aimed scratching movements and before the cuticle had hardened. A loop of 370 μm diameter tinned copper wire was passed over the femoro-tibial joint of each fully flexed hind leg to restrain the tibia in the fully flexed position. The limb was free to move at the thoraco-coxal and coxo-trochanteral joints, but the animal could not extend the tibia or make aimed scratching movements. Animals were maintained together for 20–24 h in small holding cages (25×15×20 cm, length×width×height) in which they could walk, climb and feed. For testing, animals were suspended above a foam ball as described in Kinematic analyses above, with their eyes blacked out. They were not provided with a start rod. The wire loop was cut with fine wire-cutters to free the tibia and then the fore wing tip was immediately stimulated with a fine paintbrush. Responses were observed and recorded as either ‘walking/struggling’ or as ‘anterior’, ‘middle’ or ‘posterior’ scratches as defined in Responses to dorsal fore wing stimulation in fifth instars above. Subsequent stimulation of the fore wing base and tip permitted repeated observations.

**Video acquisition and analysis**

For kinematic analyses, animals were videotaped using a colour CCD camera (C1380, JVC, Yokohama, Japan) operated at a shutter speed of 1/500 s. The sVHS video signals were combined on a multiviewer (MV-40PS, For-A, Tokyo, Japan) and time stamped using a video timer (VTG-33, For-A). Images were taped on an sVHS video recorder (HR-S7500, JVC, Yokohama, Japan), displayed on a monitor (PVM-1450MD, Sony, Tokyo, Japan) and played back for capture by a personal computer video interface card (MiroVIDEO DC30plus, Pinnacle Systems, CA, USA) in PAL video format at a size of 720×540 pixels using MiroVIDEO Capture software (25 Hz frame rate for full interlaced video frames) and compressed using Microsoft ‘dvdv’ codec. A deinterlacing algorithm (Videotrack 2D written by J. Zakotnik, available on request from the corresponding author) then split each frame into even and odd frames, and linearly interpolated the missing lines to give a final frame rate of 50 Hz. Short clips containing individual scratches were produced in VirtualDub (A. Lee, http://www.virtualdub.org/index). A calibration target was filmed during every experiment.

Videotrack 2D was used to detect the co-ordinates of the eight markers within each frame that described the animal’s body position and leg movement (Fig. 2Ai,Bii; see Dürr and Matheson, 2003 for a discussion of the accuracy of this method). The line between the markers adjacent to the hind leg coxa and the front leg coxa defined the x-axis of the body-centred coordinate frame, with the marker 2 mm in front of thoraco-coxal joint of the hind leg set to the origin (Fig. 2Ai,Bii). The stimulus location and the start position of the tarsus were digitised on the first frame of each behavioural sequence. For the x-axis comparison of centres of density, the fifth instar coordinates were scaled by 140%, based on the mean difference in size for six fifth instar and six adult locusts, and wing base and abdomen tip stimulus locations were aligned by shifting the fifth instar frame of reference anteriorly by 2.7 mm. This standardisation permitted the direct comparison of movement trajectories between fifth instars and adults of different sizes.

**Data analysis**

Analysis of individual scratching responses started with the frame before the movement began and ended if the tarsus touched the ground, the tarsus hit the brush, the leg completed three cycles of movement (‘loops’, see below), or the leg stopped moving for longer than 120 ms. The two-dimensional coordinates of hind leg joints throughout a scratch were output by Videotrack 2D and subsequently analysed by Scratch Analysis (written by V. Dürr, available on request from the corresponding author; Fig. 2Ci,ii). The position of the hind leg relative to the body is shown as an example for one movement toward a wing base target in a fifth instar locust (Fig. 2Ci). The hind leg tarsus moved from the start position (small blue circle) anteriorly and dorsally toward the stimulus position (triangle) on the wing (blue line) driven by anterior rotation of the thoraco-coxal joint and flexion of the femoro-tibial joint. Fig. 2Cii shows the sequential locations of the femoro-tibial joint (purple), the tibia-tarsal joint (red), and the wing bud tip (blue) for the same movement. Scratch Analysis computed joint angles and angular velocities for all four hind leg joints in each frame of the sequence. Statistical tests were performed using SPSS Version 12 (SPSS Inc., Chicago, IL, USA). Contrasts between stimulus sites were examined using repeated-measures ANOVAs (or non-parametric
repeated-measures Friedman tests where distributions were non-normal as noted in the Results. Contrasts between fifth instars and adults were conducted using one-way ANOVAs (or non-parametric Mann–Whitney U-tests where appropriate) as noted in the Results. Where necessary, post hoc two-tailed t-tests were used to contrast the three stimulus sites. The outcomes of statistical tests are described using the nuanced terminology recommended by Colquhoun (2014, 2015): *P* > 0.1, ‘no evidence for a real effect’; 0.05 < *P* < 0.1, ‘very little evidence for a real effect’; 0.01 < *P* < 0.05, ‘weak evidence for a real effect’; 0.001 < *P* < 0.01, ‘moderate evidence for a real effect’; and *P* < 0.001, ‘strong evidence for real effect’. Where data were pooled, means were first calculated for each animal and then averaged across animals and presented together with the standard error of the mean.

To determine which part of the hind leg was being aimed at the target, the hind leg was divided into 24 “units” of equal length (see Dürr and Matheson, 2003). The femur was divided into 10 equal units beginning proximally (units 1–10). The tibia was likewise divided into 10 equal units (units 11–20), and the tarsus was divided into four equal units (units 21–24). The shortest distance between each unit and the target during the scratch was calculated. Having determined the point on the leg that on average most effectively reached all three target sites, the movement trajectory of this effector point (approximately 4 mm proximal to the tibio-tarsal joint) was used to define the closest point of approach to the target in each scratch, thus providing a measure of the accuracy of each movement. A mean accuracy was established for each animal for movements towards wing base, wing tip and abdomen targets.

Two components of each scratching response were analysed: a short (200 ms) initial vector component (Fig. 2Ci) and a second cyclic component (Dürr and Matheson, 2003). The first 200 ms of movement was used to calculate the initial speed and angle of movement of the effector (scaling was not applied to these values). The cyclic component was quantified by first determining the number of movements with 0, 1, 2 or 3 loops for fifth instars and adults for each target. A loop was defined by both extension and flexion of the femoro-tibial joint, and ended with the transition of the femoro-tibial joint from flexion to extension. The cyclic component was further characterised by calculating a probability distribution of the location of the distal tibia across scratches (see Dürr and Matheson, 2003 for details). The centre of density of the probability distribution was used as a single measure of the location of the scratch. To distinguish between pairs of movement distributions, the probability distributions were treated as empirical likelihood functions, and Bayes’ rule was applied to decide which one of the two sites had most likely been stimulated to cause the leg to move across point X during a single video frame (see Dürr and Matheson, 2003 for derivation and details; these computations were carried out in Microsoft Excel). The likelihood of the probability estimate being wrong for a single observation (frame) was used to determine a critical number of frames required to permit two maps to be distinguished. If the average number of frames of the two maps being compared was more than the critical number of frames needed to distinguish between them, then the two distributions were considered statistically different. In other words, the distributions were considered significantly different if observation of a single scratch of average duration would permit a correct prediction of the observed probability distribution in 95% of cases (see Dürr and Matheson, 2003). Probability distributions were compared for pairs of target sites to determine: (1) the percentage volume overlap of the two distributions; and (2) the minimum number of observations needed to obtain a 95% chance of making a correct decision (>50% correct single assignments).

The point of closest approach was shown in joint angle space for comparison with data from Dürr and Matheson (2003). Measurements of the thoraco-coxal and the coxo-trochanteral joint angles of the hind leg were smoothed with a sliding window having a width of five frames, weighted by binomial coefficients.

**RESULTS**

**Wings grow allometrically at the imaginal moult**

Body, leg and wing lengths were measured in six animals prior to and six different animals following the imaginal moult (Fig. 1Bi,ii, C). Adult bodies were 135% longer than those of fifth instars, while hind leg segments were 122–161% longer (Table 1). In contrast, adult wings were 447% longer than the wing buds of fifth instars (Fig. 1C, Table 1). Relative to the increase in body length, all hind limb segments thus increased isometrically by a factor of 0.9–1.2, whereas wing length increased allometrically by a factor of 3.3 (Table 1). The tips of the fifth instar wing pads lay approximately 15 mm anterior to the tip of the abdomen, whereas the tips of the adult wings lay approximately 12 mm posterior to the tip of the abdomen (Fig. 1Bi,ii).

**Scratching movement trajectories of fifth instar and adult locusts**

In resting adult locusts, the dorsal fore wing surface is outermost and thus exposed. As a consequence of wing bud rotation during development, however, it is the ventral hind wing surface that is exposed in resting fifth instars. Both fifth instar and adult locusts performed scratching movements in response to touches on the exposed wing surfaces or abdomen (Fig. 3). Following stimulation of an anterior target at the base of the wing (yellow squares in Fig. 3Ai,Bi), fifth instars, like adults, moved the distal end of the ipsilateral hind leg towards the target in an anteriorly directed arc, and performed cyclical movements near to the target (black trajectories in Fig. 3Ai,Bi). Following stimulation of a posterior target on the tip of the abdomen (black squares in Fig. 3Aii,Bii), fifth instars, again like adults, moved the distal end of the leg towards the target in posteriorly directed movements and made cyclical movements near the target (black trajectories in Fig. 3Aii,Bii).

The hind wing tip in fifth instars is an anterior target and its stimulation elicited anteriorly aimed movements (Fig. 3Aiii) that were similar in form to those aimed toward the hind wing base (Fig. 3Ai). In adults, however, the fore wing tip is a posterior target, and its stimulation elicited movements that were directed posteriorly (Fig. 3Biii). They were similar in form to those aimed towards nearby abdomen tip targets (Fig. 3Bii).

In fifth instars, tactile stimulation of the normally hidden dorsal fore wing tip (see Materials and Methods) elicited anteriorly directed scratches. Of 110 trials in 11 locusts in which the fore wing

![Table 1. Comparison of body and limb dimensions for the fifth instar and adult Schistocerca gregaria used in subsequent experiments](image-url)
tip was stimulated, 33 responses were aimed anterior to the midpoint of the abdomen, 19 were aimed anterior to the tip of the abdomen, and none were aimed posterior to the tip of the abdomen. The remaining 58 stimuli did not elicit scratching responses. Touching the dorsal fore wing tip therefore elicits anteriorly directed scratches in fifth instars but posteriorly directed scratches in adults.

Rapid change in behaviour at the imaginal moult
The switch from making anteriorly directed to posteriorly directed scratches in response to touch of the wing tip occurs instantaneously following the imaginal moult, and does not require a period of calibration or learning. In nine of nine animals observed continuously during moulting, the first touch of the newly expanded adult fore wing tip led immediately to posterior scratches, prior to any other significant hind limb movements, including walking or scratching. During the early part of moulting, animals hang with their head down, and make small wriggling movements to pull themselves out of their old exoskeleton with the help of gravity. As soon as the adult emerges from the exuvium, it rotates to a head-up posture and remains essentially motionless while haemolymph is pumped through the wings to expand them, again assisted by gravity. These newly moulted animals responded reliably to tactile stimulation. To confirm that these posterior movements were retained beyond the moulting period, the hind legs of a further 13 animals – also observed continuously during moulting – were restrained immediately following the moult to prevent tibial extension that is required to reach the adult wing tip. On removal of the restraint 20–24 h after the moult, all 13 of these animals responded immediately to touch of the adult fore wing tip with posterior scratches. In no case did the initial touch of the adult fore wing tip lead to an anterior scratch.

Both fifth instars and adults aim the distal tibia at wing and distal abdomen targets
For movements toward anterior wing base targets in both fifth instars and adults, two different regions of the hind leg approach the target closely, resulting in a W-shaped distribution of points of closest approach (Fig. 4Ai,Bi). Unit 4 at the centre of the femur (see Materials and Methods for a description of the division of the leg into analysis ‘units’) was closest for both fifth instars and adults (‡ in Fig. 4Ai,Bi). A second minimum was centred on unit 18 of the distal tibia (* in Fig. 4Ai,Bi). The physical constraints of hind leg movement mean that for the distal tibia or tarsus to reach the anterior targets (wing base and wing tip in fifth instars; wing base in adults), the femur must rotate far anteriorly, itself crossing the target site (see Fig. 2Ci). The femur typically crossed the target once, and was then held anteriorly for the rest of the movement. In contrast, the distal tibia approached the target several times during the cyclic component of a scratch (Fig. 3). The local minimum at unit 18 thus reflected the aimed effector (distal tibia), whereas that at unit 4 reflected a mechanical consequence of that aimed movement. In movements toward (posterior) abdomen tip targets in both fifth instars and adults, units 18 or 19 of the distal tibia most closely approached the target (* in Fig. 4Aii,Bii). The wing tip is an anterior target in fifth instars, but a posterior target in adults, and this difference in target position is reflected in the parts of the hind leg that approach the target. In fifth instars, both unit 5 of the femur and unit 19 of the distal tibia approached the target as described above for wing base targets (‡ and *, respectively, in Fig. 4Aiii). In adults, only unit 23 of the tarsus reached the very posterior position of the wing tip target (* in Fig. 4Biii). The distal tibia and tarsus are thus used as effectors during scratching movements in both fifth instars and adults, for both anterior and posterior targets.
Targeting accuracy is similar for all sites

The minimal distance from the distal effector point to the target during each trajectory provided a measure of accuracy. For fifth instars, the mean minimal distance was similar for movements toward the wing base, abdomen tip and wing tip (repeated-measures ANOVA, Greenhouse–Geisser correction, $F_{1.084,5.421}=0.448$, $P=0.548$). The mean minimal distances were 5.1±0.8, 5.1±1.4 and 4.3±0.6 mm (means±s.e.m.) for wing base, abdomen tip and wing tip targets, respectively.

For adults, there was weak evidence that mean minimal distance may be different for different targets (repeated-measures ANOVA, $F_{2,10}=4.167$, $P=0.048$). There was no evidence for a difference in mean minimal distance for adult wing base and wing tip targets (post hoc two-tailed t-test, $P=0.537$), and only weak evidence for differences between the abdomen tip target and either the wing base target ($P=0.028$) or the wing tip target ($P=0.036$). The mean minimal distances were 7.4±1.7, 3.5±0.7 and 6.2±0.7 mm for wing base, abdomen tip and wing tip targets, respectively.

There was no evidence for a difference between fifth instars and adults in mean minimal distance for movements towards wing base targets (one-way ANOVA, $F_{1,10}=1.498$, $P=0.249$), abdomen tip targets ($F_{1,10}=1.242$, $P=0.291$) or wing tip targets ($F_{1,10}=3.815$, $P=0.079$). Overall, there is therefore little evidence for any difference in accuracy between sites or between ages.

Initial direction of movement and speed is influenced by target site

The first 200 ms of each scratch was used to calculate a movement vector that was used to determine whether movements were target specific from their onset and thus predetermined. For fifth instars, the initial directions of movement towards all three targets were directed dorsally and anteriorly (yellow, red and black vectors in Fig. 5A). There was no detectable effect of target position on the initial direction of movement, despite the fact that the abdomen tip target was posterior to the initial tarsus position while the two wing targets were anterior (non-parametric repeated-measures ANOVA (Friedman test): $\chi^2_{2,6}=0.333$, $P=0.846$). Nevertheless, the movement vectors were all correctly biased towards the appropriate targets (corresponding yellow, red and black squares in Fig. 5A inset).

For adults, initial movements toward wing tip targets (red in Fig. 5B) were directed more posteriorly than those for either wing base (yellow) or abdomen tip (black) targets (Fig. 5B). Target position influenced the initial movement direction (Friedman $\chi^2_{2,6}=9.000$, $P=0.011$).

For adults, initial movements toward wing tip targets (red in Fig. 5B) were directed more posteriorly than those for either wing base (yellow) or abdomen tip (black) targets (Fig. 5B). Target position influenced the initial movement direction (Friedman $\chi^2_{2,6}=9.000$, $P=0.011$). There was no evidence for a difference between fifth instars and adults in initial direction of movement towards wing base targets (yellow in Fig. 5A,B; Mann–Whitney $U=18$, $P=1.000$), or towards abdomen tip targets (black in Fig. 5A,B; Mann–Whitney $U=15$, $P=0.631$). The initial movement towards wing tip targets, however,
The initial movement vector (direction and speed) of the distal tibia was measured over the first 200 ms for movements in fifth instars (A) and adults (B), in response to touches at wing base (yellow), wing tip (red) and abdomen tip (black) targets. The initial movement vector angle for each target is indicated by the colour-coded circle on the perimeter of the polar plot. The start and target locations are also indicated in the inset diagrams of fifth instar and adult locusts. Movements toward wing tip targets were directed more posteriorly in adults than in fifth instars. Speed was similar for fifth instars and adults for movements towards the wing base and abdomen tip, but different for movements towards the wing tip.

Cyclic component of scratching is influenced by target site

Most scratches consisted of an initial outward trajectory followed by several cyclical ‘loops’ near the target. Relatively few scratches (9.6±1.3%, mean±s.e.m.) had no cyclical component (‘loops’). There was no difference in the number of loops between fifth instars and adults for movements towards wing base stimuli ($\chi^2=1.201$, d.f.=3, $P=0.867$), abdomen tip stimuli ($\chi^2=0.671$, d.f.=3, $P=1.000$) or wing tip stimuli ($\chi^2=2.237$, d.f.=3, $P=0.588$).

The pattern of movement during the cyclic part of scratching was summarised as an effector probability distribution (Fig. 6). Regions of highest probability are darkest in Fig. 5 and the weighted centre of the distribution (‘centre of density’) is indicated by a circle (mean±s.e.m.).

In fifth instars, the probability distributions for movements aimed at targets at the base of the wing or the tip of the wing both formed arcs (shaded regions in Fig. 6Ai,iii) that curved from near the start positions (black triangles), dorsally and anteriorly, then ventrally and further anteriorly to near the targets (squares). The mean centres of density yellow and red circles, respectively) were dorsal and posterior to the targets in both cases, so that the targets lay near the anterior ventral margin of the probability distributions. The probability distribution for movements aimed at targets at the tip of the abdomen formed a compact distribution over the target site (Fig. 6Aii). The centre of density (white circle) was close to the target (white square).

Probability distributions for movements aimed at the wing base and abdomen tip in fifth instars overlapped by 40% but could be reliably distinguished (Fig. 6Aii), as could those for movements toward wing tip and abdomen tip targets (47% overlap; Fig. 6Aiii). These movement distributions were thus target specific. Probability distributions for movements aimed at the fifth instar wing base and wing tip targets (which are separated by only 4 mm) overlapped by 76% and could not be reliably distinguished (Fig. 6Aiii). The distribution for the more anterior wing base target was nevertheless biased anteriorly compared with that for the more posterior wing tip target (Fig. 6Aiv).

Target position (wing base, wing tip or abdomen tip) had a strong effect on the anterior–posterior (x-axis) location of the centre of density of the probability distribution (repeated-measures ANOVA, $F_{2,10}=31.13$, $P<0.001$). There was no evidence for a difference between the x-axis locations of the centres of density for movements aimed at wing base and wing tip targets (post hoc two-tailed t-test, $P=0.110$). However, both were more anterior than the centre of density for movements towards abdomen tip targets (wing base, $P=0.004$; wing tip, $P=0.004$).

In adults, the probability distribution for movements aimed at targets at the base of the wing formed an arc that curved from near the start position (black triangle in Fig. 6Bi), dorsally and anteriorly, then ventrally and further anteriorly to near the target (square). The mean centre of density (circle) was posterior to the target. In adults, the probability distributions for movements aimed at targets at the abdomen tip or wing tip formed compact clusters close to the corresponding target sites (squares in Fig. 6Bii,iii). The centre of density (circle) was ventral and anterior to the target (square). These movement distributions were thus target specific. Probability distributions for movements towards targets at the abdomen tip or wing tip formed compact clusters close to the corresponding target sites (squares in Fig. 6Bii,iii). The centre of density (circle) was ventral and anterior to the target (square). Probability distributions for movements aimed at targets at the abdomen tip or wing tip formed compact clusters close to the corresponding target sites (squares in Fig. 6Bii,iii). The centre of density (circle) was ventral and anterior to the target (square). Probability distributions for movements aimed at targets at the abdomen tip or wing tip formed compact clusters close to the corresponding target sites (squares in Fig. 6Bii,iii). The centre of density (circle) was ventral and anterior to the target (square).

To test whether movements were target specific in adults, the overlap between pairs of probability distributions was compared in the same way as for fifth instars. The probability distributions for movements aimed at wing base and wing tip targets overlapped by only 37% and could be distinguished reliably (Fig. 6Bi,iii), as could movements aimed at the wing base or abdomen tip (34% overlap, Fig. 6Bi,ii). Probability distributions for movements aimed at wing tip and abdomen tip targets overlapped by 66% and could
not be distinguished reliably (Fig. 6Bii,iii). The distribution for the more dorsal wing tip target was nevertheless biased dorsally compared with that for the more ventral abdomen tip target (Fig. 6Biv).

As for fifth instars, target position had a strong effect on the x-axis location of the centre of density (repeated-measures ANOVA, $F_{2,10}=61.05, P<0.001$). In adults, movements towards the wing base had a more anterior centre of density than movements towards either the abdomen tip (post hoc two-tailed t-test, $P=0.002$) or the wing tip ($P=0.001$). In contrast, movements towards the abdomen tip and wing tip had centres of density in a similar location along the x-axis ($P=1.000$). Overall, these results indicate that both fifth instars and adults accurately aim the cyclical part of their scratching movements at the correct target sites.

**Direct contrast between fifth instar and adult scratching movements**

For movements towards an anterior target such as the base of the wing, the probability distributions for fifth instars and adults both had a similar shape and similar centres of density (Fig. 6Ai,Bi). The
adult distribution was less dense because the trajectories covered a larger area (note scale bars in Fig. 6). For movements towards a posterior target, such as the tip of the abdomen, the probability distributions for fifth instars and adults also had similar shapes and centres of density (Fig. 6Aii,Bii). The location of the wing tip changes from an anterior position in fifth instar locusts to a posterior position in adults, and this was accompanied by a clear change in the behavioural response. In response to stimulation of the wing tip, fifth instars made an anteriorly directed movement (Fig. 6Aiii) that had a similar shape to the probability distribution for movements aimed at the (anterior) base of the wing (Fig. 6Ai), whereas adults made a posteriorly directed movement (Fig. 6Biii), that had a similar shape to the probability distribution for movements aimed at the (posterior) tip of the abdomen (Fig. 6Bii).

**Joint kinematics are similar for corresponding targets in fifth instars and adults**

To determine whether fifth instars and adults use similar patterns of inter-joint coordination to reach anterior or posterior targets, the combined thoraco-coxal+coxo-trochanteral joint angle was plotted against the femoro-tibial joint angle at the closest point of approach to the target (Fig. 7). Mean joint angles for each animal (coloured circles in Fig. 7) were superimposed on a large dataset of individual adult scratches from Dürr and Matheson (2003) (grey circles in Fig. 7). Dürr and Matheson (2003) stimulated multiple sites along the length of the adult wing surface and showed that combinations of joint angles were gradually shifted as the target position was moved along the wing’s length. Adult data in the present study correspond well with this pattern: for anterior wing base targets, the thoraco-coxal+coxo-trochanteral joint angle was large and the femoro-tibial joint angle small (yellow circles in Fig. 7A), reflecting elevation of the femur and flexion of the tibia at the point of closest approach to the target. For posterior wing tip targets, the thoraco-coxal+coxo-trochanteral joint angle was large, and the femoro-tibial joint angle small (red circles in Fig. 7A), reflecting depression of the femur and extension of the tibia at the point of closest approach to the target. Joint angle combinations used to reach the abdomen tip lay on the same curve and were near those used to reach the wing tip (black circles in Fig. 7A).

For fifth instars, movements towards the anterior wing base target used joint angle combinations that coincided with those used to reach the same anterior target in adults (yellow circles in Fig. 7B cf. A). The combinations of joint angles used to reach the posterior abdomen tip target in fifth instars similarly coincided with those for the abdomen tip adults (black circles in Fig. 7B cf. A). In contrast, the combinations of joint angles for movements aimed towards the wing tip were different in fifth instars and adults (red circles in Fig. 7B cf. A), while still lying on the same curve. Specifically, the combinations of joint angles used to reach the wing tip target in fifth instars were similar to those used to reach the wing base targets in both adults and fifth instars, whereas those used to reach the wing tip target in adults were similar to those used to reach the abdomen tip targets of both adults and fifth instars. Fifth instars used systematically larger femoro-tibial joint angles for a given thoraco-coxal+coxo-trochanteral angle to reach wing targets than they did for wing base targets (red circles in Fig. 7B lie above yellow circles).

**DISCUSSION**

We show that gentle tactile stimulation of the wings elicits aimed hind leg movements in juvenile locusts that closely resemble those of adults. Despite marked allometric growth and radical changes in wing morphology during imaginal development, these aimed movements are immediately retargeted to appropriate locations, while the characteristic form of the movements remains the same.

**Fifth instar and adult locusts use the same effector and movement patterns**

Fifth instar locusts, like adults (Berkowitz and Laurent, 1996; Matheson, 1997, 1998; Dürr and Matheson, 2003), perform aimed hind leg scratching movements in response to, and accurately aimed at, stimuli on the wings and abdomen. First, both fifth instars and adults use the distal tibia as the effector. Second, both move the effector at similar speeds for the initial 200 ms of movement. Third, both achieve a similar accuracy, as determined by the minimum distance between the effector and the target. Fourth, both use comparable combinations of hind leg joint angles to reach a range of targets. A single movement pattern that can be shifted towards anterior or posterior targets on the wings (Dürr and Matheson, 2003) thus appears to be conserved across the imaginal moult. Combinations of joint angles lie on a continuum in joint angle
space, allowing the effector to reach anterior targets using joint angle combinations at one end of the continuum (thoraco-coxal+coxo-trochanteral joint rotated anteriorly, and the femoro-tibial joint fully flexed), whereas posterior targets are reached using joint angle combinations at the other extreme of the continuum (thoraco-coxal+coxo-trochanteral joints rotated posteriorly, and the femoro-tibial joint extended). Targets between these two extremes are reached by intermediate joint angle combinations in both fifth instars and adults. We therefore hypothesise that essentially the same central neuronal network, with properties conserved across the imaginal moult, generates scratching in fifth instars and adults, but that the sensory inputs eliciting the behaviour must be modified.

**Isometric growth can explain the maintenance of aiming accuracy for targets on the abdomen**

Isometric growth of the hind legs means that combinations of joint angles used by fifth instars to reach locations on the abdomen will continue to direct the larger adult hind leg to the same region of the larger adult abdomen. Nevertheless, growth must also lead to changes in muscle mass and strength, leg mass, cuticular properties and joint mechanics (Andersen, 1973, 1974; Norman, 1995), so it remains possible that leg motor patterns used by fifth instars differ from those used by adults to achieve the same kinematics.

Praying mantises use rapid aimed lunges of their fore legs and body to grasp moving prey. Striking distance is precisely estimated by a binocular triangulation mechanism of the visual system (Maldonado and Levin, 1967), with the longest distance that elicits a strike being related to the length of the fore legs in each developmental stage (Balderrama and Maldonado, 1973). The growth of the head and the eyes is such that throughout development, the fovea registers a stimulus at the maximum catching distance. This triggers a striking movement at a distance that is appropriate for the length of the limbs at each developmental stage (Maldonado et al., 1974). In an analogous way, as the locust abdominal body wall elongates during development, distal targets on it move posteriorly, but it is likely that little or no remapping of sensory neurons is required, because unchanged patterns of connectivity with motor networks would ‘automatically’ drive the isometrically larger hind leg to the new target location, at least within the levels of accuracy seen for these movements. Such automatic compensation is not possible for wing tip targets, however, because of allometric wing growth and wing rotation.

**Developmental wing rotation at the imaginal moult does not explain changed responses to wing tip stimulation**

Touching the exposed fore or hind wing tips of fifth instar locusts elicits anterior movements, whereas touching the wing tips of adults elicits posterior movements. We have ruled out the possibility that fifth instars fail to make posterior movements simply because the receptors are hidden under the (rotated) wing buds; stimulating the normally hidden dorsal fore wing receptors in fifth instar locusts leads to anterior movements. The central circuits underlying posterior scratching are present in fifth instars – because distal abdominal stimulation elicits posteriorly directed movements – but they do not produce posterior movements in response to wing tip stimulation until adulthood.

**Remodelling of sensory inputs may explain changed responses to wing tip stimulation**

In contrast to the apparently learned ability of adult locusts to spontaneously walk across larger gaps than juveniles (Ben-Nun et al., 2013), appropriately aimed scratching movements did not require prior experience of either stimulation or movement following the adult moult. The most parsimonious explanation of the observed change in aiming is that the adult fore wing develops new tactile hair receptors that drive the existing central motor circuits to produce more posteriorly directed movements than those driven by comparable receptors on the juvenile wing buds. Aimed scratches are elicited by activation of touch-sensitive hairs distributed across the wing surface, with a dorsally directed row on the adult forewing postcubitus vein being particularly efficacious (Page and Matheson, 2004). The somatosensory representation of the wing surface is unknown for any insect – in part because the afferent neurons appear to have small diameters, yet long axons, which makes staining difficult. It is reasonable to assume, however, that the afferents form a somatotopic map in the local ganglion, as is the case for locust leg and body wall receptors (Newland, 1991; Newland et al., 2000). We propose that the juvenile wing bud somatosensory map, and its connections to leg motor networks, is extended ‘posteriorly’ through the addition of new receptors born on the adult wing during the final larval instar (Altman et al., 1978). In the cricket, clavate mechanosensory hairs and their receptors are likewise added at each moult. The central projections of later-born receptor project less far, and less extensively, into the terminal abdominal ganglion, so that the population of afferents forms a topographic representation of the surface of the cercus depending on both birth date and hair location (Murphey et al., 1980; Murphey, 1981). The number of wind-sensitive hairs on the head of a locust similarly increases with each instar (Sviderskii, 1969), and this is accompanied by a progressive development of the flight motor pattern – which they drive – in juveniles and adults (Altman, 1975; Kutsch, 1971, 1985; reviewed by Pfüger and Wolf, 2013). Immobilising the wings at the imaginal moult does not prevent expression of the adult flight motor pattern later in adult life (Altman, 1975), and the development of the final adult wingbeat frequency does not require flight experience, suggesting that, as for aimed scratching, changes are ‘hard-wired’ rather than learned. Although the flight motor pattern begins to develop during larval stages, the full adult pattern is not expressed until the day of (following) the imaginal moult. We have shown that although the motor system of fifth instar locusts can produce posteriorly aimed hind limb movements (e.g. in response to touch of the abdomen), such posterior movements are not elicited by touch of the wings until the adult wings are fully expanded following the imaginal moult, when the new stimulus–response relationship is apparent even on the very first touch of the new sensory surface. Locusts thus possess a mechanism to automatically compensate their aimed limb movements for allometric growth of the wing sensory surface.

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**Competing interests**

The authors declare no competing or financial interests.

**Author contributions**

Conceptualization: A.J.P., T.M.; Methodology: A.J.P., T.M.; Software: T.M.; Validation: T.M.; Formal analysis: A.J.P., T.M.; Investigation: A.J.P., T.M.; Resources: T.M.; Data curation: A.J.P., T.M.; Writing - original draft: A.J.P., T.M.; Writing - review & editing: A.J.P., T.M.; Visualization: A.J.P., T.M.; Supervision: T.M.; Project administration: T.M.; Funding acquisition: T.M.

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