Mechanosensation and Adaptive Motor Control in Insects

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The ability of animals to flexibly navigate through complex environments depends on the integration of sensory information with motor commands. The sensory modality most tightly linked to motor control is mechanosensation. Adaptive motor control depends critically on an animal’s ability to respond to mechanical forces generated both within and outside the body. The compact neural circuits of insects provide appealing systems to investigate how mechanical cues guide locomotion in rugged environments. Here, we review our current understanding of mechanosensation in insects and its role in adaptive motor control. We first examine the detection and encoding of mechanical forces by primary mechanoreceptor neurons. We then discuss how central circuits integrate and transform mechanosensory information to guide locomotion. Because most studies in this field have been performed in locusts, cockroaches, crickets, and stick insects, the examples we cite here are drawn mainly from these ‘big insects’. However, we also pay particular attention to the tiny fruit fly, Drosophila, where new tools are creating new opportunities, particularly for understanding central circuits. Our aim is to show how studies of big insects have yielded fundamental insights relevant to mechanosensation in all animals, and also to point out how the Drosophila toolkit can contribute to future progress in understanding mechanosensory processing.

Introduction

The unpredictable structure of the natural world poses a problem for motor control systems. Because the environment and the body itself are always changing, a given motor command signal will not always result in an identical movement. A related problem is that internal frames of reference also shift during movement — for example, when limbs move relative to each other. In the terminology of control theory, flexible movements cannot be reliably executed in open-loop, but instead require closed-loop feedback from both external and internal sensors [1].

An important source of feedback is mechanosensation. Mechanical forces on the body are an inevitable consequence of self-movement. Mechanotransduction is also faster than phototransduction, and it is equally effective throughout the day, whatever the ambient light level. Perhaps for these reasons, most animals are absolutely reliant on mechanosensation for normal movement — for example, human patients who lack mechanosensory feedback can generate coarse limb movements but are unable to execute fine motor tasks [2].

Many robotic systems also use mechanical sensors to guide the control of mechanical actuators. Nonetheless, modern robots lack the motor flexibility of biological systems [3]. Although robots can excel at individual tasks, they frequently fail when placed in novel or unpredictable situations. One reason is that robots lack sensorimotor control circuits commensurate with those of animals. An important difference is that robots typically switch between explicit kinematic models to achieve different high-level behaviors, while biological systems appear to use modulation of low-level sensorimotor control loops [4]. As a result, it is difficult for a robot to find footholds on uneven terrain or to extricate itself when it becomes stuck. The fact that engineers have not yet succeeded in fully solving these problems is a powerful reminder that we do not fully understand how mechanosensory feedback interacts with top-down commands to control movement in animals.

Insects are particularly interesting exemplars of flexible sensorimotor control. Although early descriptions often portrayed insects as simple automatons, “denying them any portion of intellect”, careful observation has since revealed that insect motor patterns are, in fact, remarkably adaptable [5]. Insects execute a dizzying variety of complex behaviors, including running, hunting, flying, courting, fighting, foraging, building, and grooming. Moreover, each of these behaviors is robust to variability in the mechanical forces acting on the body. For example, cockroaches can react to an unexpected mechanical stimulus in less than 20 ms [6,7]. On a longer timescale, insects must be tolerant to the natural wear and tear of life. Foraging bumblebees collide with vegetation about 60 times a minute, sometimes resulting in a 5–10% loss of wing area in a single day [8]. Somewhat, bumblebees are able to compensate for these dramatic changes in the structure and aerodynamics of their wings, not to mention the added weight from the pollen and nectar they have collected. Even the most promising insect-inspired robots, such as the running VelociRoACH [9] and the flying RoboBee [10], can mimic only a small fraction of the motor behaviors executed by real insects.

In this Review, we will consider how specific design features of insect mechanosensation might contribute to the rapid and flexible control of movement. We will first examine the structure and function of mechanoreceptor organs, paying particular attention...
movements of one joint segment relative to the adjoining segment. (E) Chordotonal organs are stretch-sensitive mechanoreceptors that contain many individual sensory neurons with diverse mechanical sensitivities. They are found at leg joints, where they encode the angle and movement of the leg, as well as in Johnston’s organ in the fly antenna, where they encode auditory signals.

...to how sensory neurons achieve sensitivity and robustness. Next, we will discuss how central circuits integrate and transform mechanosensory signals in order to produce appropriate motor commands.

Our discussion will refer to many different insect species. The majority of the work in this field has been performed in what we will call ‘big insects’ — mainly locusts, cockroaches, crickets, and stick insects. However, there is also a small but growing body of work in the fruit fly, Drosophila melanogaster. Until recently, work on mechanosensation in Drosophila focused mainly on the development of peripheral organs [11,12] and the molecular genetics of mechanotransduction [13]. Recently, however, Drosophila has also become a major model for studying neural coding, central circuits, and the neural correlates of behavior. The Drosophila genetic toolkit now provides access to many specific cell types, and the major connectomics effort underway in Drosophila will soon tell us the synaptic connectivity of these cell types. To use these tools intelligently, we should consider the lessons already learned from studies of big insects. Because big insect studies have mainly been performed in adults rather than larvae, we have focused our discussion of Drosophila on the adult stage as well.

We hope this review will be useful for three types of readers. First, as Drosophila neuroscientists, we aim to provide a primer on big insect mechanosensation for the benefit of other Drosophilists. Second, in addressing readers familiar with big insects, we will indicate toward the end of the review how the Drosophila toolbox can help fill gaps in the existing literature. Third, for readers unfamiliar with insects of any kind, we hope to clarify how the study of insects is providing insights into the fundamental problems of mechanosensation and motor control that have relevance to all animals.

**Insect Mechanoreceptors**

Within a given region of the body — for example, the leg (Figure 1A) — one typically finds multiple mechanoreceptor types located in close proximity. Each mechanoreceptor type is sensitive to a particular range of mechanical stimuli, such that a naturalistic and complex mechanical stimulus will often co-activate multiple receptor types. Below we will summarize the distinctions between different mechanoreceptor types. Toward the end of this review, we will examine how signals from co-activated mechanoreceptor types are integrated in the central nervous system (CNS).

As in vertebrates, the different types of insect mechanoreceptors are traditionally divided into two functional groups: exteroceptors and proprioceptors. Exteroceptors directly detect mechanical forces generated in the external world, while proprioceptors detect the position or movement of body parts. By convention, we will generally treat mechanically isolated (i.e., not tightly clustered) tactile hairs on the external body surface as exteroceptors, and all other mechanoreceptors as proprioceptors. However, the distinction between exteroceptors and proprioceptors is murky. Exteroceptors can be stimulated during self-generated movement, and, conversely, proprioceptors can be stimulated when external stimuli cause body parts to move. Indeed, as we will see below, some neurons that are typically characterized as proprioceptors are in fact primarily dedicated to sensing external stimuli, because they monitor body parts that generate relatively large movements in response to small external forces.

**Tactile Hairs (Bristles)**

The most visible and abundant of the mechanosensory structures in the adult insect are the tactile hairs, which belong to a morphological class of sensory structures called trichoid sensilla. In the Drosophila literature, tactile hairs are referred to as bristles. A fruit fly has several thousand bristles, which fall into two discrete classes, macrochaetes and microchaetes, based on their size and developmental origins [14,15]. The larger bristles (macrochaetes, from the Greek macros [large] + khaitê [long hair]) are consistent in number and position across individuals within a species and even exhibit striking similarity across fly species that diverged 50 million years ago [16]. The smaller and more numerous microchaetes exhibit more variability in their distribution but are typically arranged in regularly spaced rows.
Each tactile hair is composed of a hollow hair shaft whose base is fixed to the dendritic tip of a single bipolar sensory neuron (Figure 1B). The hair acts as a lever arm that exerts forces on the tip of the sensory neuron dendrites, where mechanotransduction channels open to produce electrical currents [17]. Electrophysiological recordings from tactile hairs can be performed by clipping the hair shaft distal to the dendrite and simply placing an electrode in contact with the extracellular fluid that fills the shaft (Figure 2A) [18,19]. The recorded extracellular signal is dominated by the receptor potential, which appears as a downward deflection, and action potentials appear as small, superimposed transients (Figure 2B).

The ionic basis of mechanoreceptor transduction is probably similar for all insect mechanoreceptor types, but recordings from tactile hairs illustrate the relevant facts particularly clearly. In an extracellular recording from a tactile hair, the receptor potential reflects the inward flow of current into the mechanosensory cell dendrite from the endolymph that surrounds the dendrite (Figure 2C). The driving force for this current is due in part to the large K⁺ gradient that is maintained across the high-resistance layer of epithelial cells between the endolymph and the central hemolymph [20]. In Drosophila bristles, this ionic gradient gives rise to a standing transepithelial potential of approximately +30 mV [21]. Because only the dendrite is bathed in endolymph, the transepithelial potential and the transmembrane potential of the neuron act in series to drive K⁺ into the neuron when mechanotransduction channels open, thereby depolarizing the neuron [20]. This arrangement transfers much of the energetic burden of mechanotransduction to the pumps in the sheath cells that pump K⁺ into the endolymph. This arrangement also permits reliable, low-latency transduction, because the total driving force for mechanotransduction currents is very large; in Drosophila bristles, the latency of mechanotransduction is ~0.1 ms [22]. Interestingly, the same strategy is used by vertebrate epithelial mechanoreceptors, such as the hair cells of the cochlea [23].

Tactile hairs are directionally selective, and their preferred direction can be predicted from the bristle’s orientation in the cuticle. Fly bristles are typically positioned with their long axis oriented approximately 45° relative to the cuticle, and most bristle neurons are most sensitive to mechanical stimuli that push the bristle toward the cuticle [18,22]. Direction selectivity of tactile hairs is thought to be related to the morphology of the hair and socket [24,25], but other mechanisms may be involved, such as the identity and distribution of mechanotransduction channels.

In many insects, tactile hair neurons are of two physiological types: rapidly and slowly adapting. In the locust, slowly adapting hairs are more common and possess a lower mechanical threshold (10°), while rapidly adapting hairs have been found only on the tibia and have a higher mechanical threshold (~40°) [25]. Crickets [26] and cockroaches [27] also possess rapidly and slowly adapting tactile hairs with different mechanical thresholds. The same distinction pertains to cutaneous mechanoreceptors in mammals. In mammals, slowly adapting receptors, such as Merkel cells and Ruffini endings, respond best to sustained stimuli. By contrast, rapidly adapting receptors, such as Meissner’s corpuscles, respond only to the onset or offset of a stimulus [28]. In both insects and mammals, rapidly and slowly adapting mechanoreceptors are not spatially segregated but are instead intermingled across the body surface.

In comparison to what is known about tactile hairs in locusts, crickets, and cockroaches, far less is known about the physiological types of bristles found in the fly. Recordings from macrochaete bristles on the head of blowflies [24] and the notum of Drosophila [18,21,29,30] have identified only slowly adapting bristles with a very low mechanical threshold of ~1° (Figure 2). These bristles are probably not sensitive to wind or sound [24] but rather respond to transient mechanical deflections such as those created by contact with external objects or during grooming behavior. It is not known whether flies also possess high threshold, rapidly adapting bristles.

In addition to the purely tactile hairs, insects also possess sensory hairs that contain taste receptors. Each taste hair is innervated by multiple gustatory neurons (typically 4 in Drosophila) and one mechanosensory neuron. In the fly, taste hairs are found on the labial palps of the proboscis, the pharynx, the legs, wings, and genitals [31]. Whereas tactile hairs are thick, sharply pointed, and accompanied by a spine-like structure called a bract, the gustatory hairs are thin, have a blunt tip, and lack the bract cell.

Figure 2. Mechanosensitive properties of tactile hair neurons.

(A) Extracellular electrophysiological recordings from tactile hairs can be made by inserting the shaft of a clipped hair in a recording pipette which is mounted on a piezoelectric actuator so that the pipette can be used to move the hair. (B) An example extracellular recording from a fly femur bristle [168] shows a downward field potential deflection (the receptor potential) and a burst of superimposed action potentials. Repeated movement of the bristle leads to adaptation in the bristle response, depicted as a spike raster. (C) The equivalent circuit of the tactile hair epithelium recording (adapted from [20]). The hair neuron dendrite is bathed in endolymph (high K⁺), which is separated from the circulating hemolymph (low K⁺) by an electrically tight layer of epithelial cells. The K⁺ gradient is established by pumps in the apical membranes of epithelial cells, depicted here as a current source. The transepithelial potential (measured in the endolymph relative to the hemolymph) is about +30 mV. If the neuron rests at about ~60 mV, then the total driving force pushing K⁺ into the neuron’s dendrite through mechanotransduction channels (g_m) would be about 90 mV. It should be noted that the tactile hair neuron is separated from the hemolymph by a layer of glial cells (not depicted here). Membrane capacitance is also omitted for clarity.
taste hairs also belong to a different morphological class, known as basiconic sensilla. In the locust, mechanosensory neurons within taste hairs are directionally selective and rapidly adapting, with a lower mechanical threshold than purely tactile hairs [32]. Their response properties in *Drosophila* are not known.

As the primary external mechanoreceptors in insects, tactile hairs contribute to a range of behaviors. For example, mechanical stimulation of small numbers of bristles on the *Drosophila* leg triggers appropriate postural changes, away from the site of stimulation [33]. In locusts, similar avoidance reflexes may contribute to object avoidance during walking [34].

Perhaps the most critical function of tactile hairs is to alert the animal to the presence of foreign objects, such as filth and parasites, on the body surface. In flies, stimulation of just one or two bristles is sufficient to trigger complex grooming sequences, in which the animal attempts to remove debris from its body [33,35,36]. These grooming behaviors involve precisely directed leg movements, which can be generated entirely by thoracic circuits, even in the absence of descending input from the brain [33,37]. Careful analysis of grooming behavior in locusts has shown that leg grooming movements are precisely targeted to the site of hair stimulation [38], irrespective of the initial position or mechanical loading of the leg [39].

Although most tactile hairs are excited by contact with solid objects, some specialized hairs are so sensitive that they can detect air particle fluctuations. In the locust, a small number of hairs on the head can respond to strong air currents and are thought to play a role in flight control [40,41]. Indeed, some hairs are almost entirely dedicated to detecting air currents — in crickets and cockroaches, bristles on specialized abdominal appendages, known as cerci, can detect wind, courtship song, and the wingbeats of a predator (reviewed in [42]). Depending on context and behavioral state, cercal stimulation can trigger rapid escape responses and jumping, as well as grooming and aggression [43]. Thus, tactile hairs can either drive or modulate many different motor programs in insects, in keeping with their role as the primary transducers of external mechanical stimuli.

**Hair Plates**

In addition to occurring as individual sensory hairs, tactile hairs also occur as tightly packed groups known as hair plates (Figure 1C). Proprioceptive hair plates are often positioned at folds in the cuticle, so that they will be deflected during joint movements [44]. However, they may also function as exteroceptors, as in the case of hair plates at the base of the cockroach antenna, which play an important role in active sensing and object localization [45]. In *Drosophila*, hair plates can be found at most leg joints [46,47] but have not been identified on the antenna.

Like a tactile hair, each individual sensillum within a hair plate is innervated by a single sensory neuron. Hair plate sensilla occur as two physiological types: rapidly adapting neurons that respond phasically to hair movements and slowly adapting neurons that respond tonically to maintained deflections [48,49].

Studies in big insects have shown that hair plates at leg joints provide sensory feedback to motor neurons that control walking. In the cockroach, a hair plate at the most proximal leg joint provides direct excitatory input to extensor motor neurons of the trochanter and indirect inhibitory input to the motor neurons that control flexion [50]. Ablation of this hair plate causes the leg to overstep and collide with the more anterior leg, indicating that proprioceptive signals from the hair plate limit the forward movement of the leg during the swing phase of the walking cycle [51].

In the fly, two hair plates located on the ventral surface of the neck form the prosternal organ, which encodes head rotations along all three axes and contributes to compensatory head movements [52]. Shaving the prosternal organ hairs on one side causes flies to compensate by rolling the head toward the operated side [53], supporting the hypothesis that the function of the prosternal organ is to monitor head position and provide feedback for the control of head posture.

**Campaniform Sensilla**

Campaniform sensilla are round or oval-shaped mechanosensory organs that respond to stress and strain within the cuticle [54]. Each sensillum consists of a socket spanned by a flexible cuticular dome that is innervated by the dendrite of a single bipolar sensory neuron (Figure 1D). The neuron is excited whenever the dome flattens [55], which occurs in response to compression and tension in the surrounding cuticle [56]. The elliptical shape of many campaniform sensilla endows them with directional selectivity — a fact first recognized by J.W.S. Pringle based on his experiments with rubber-and-paper models of the dome [57]. The dome is specifically elongated by tension along its long axis, or else compression along its short axis (Figure 3A).
The orientation of the sensillum within the cuticle therefore dictates which stimuli will preferentially excite it [58]. Approximately 1200 campaniform sensilla are distributed over the legs, wings, halteres, and antennae of the fly [59] and exhibit considerable structural diversity [60,61]. Campaniform sensory neurons in the fly can be classified into two physiological categories; rapidly and slowly adapting [62]. Groups of campaniform sensilla with similar directional sensitivities and response characteristics are often found clustered together in areas likely to encounter cuticular strains, such as joints (Figure 3A) [57].

Interestingly, the specific orientations of many campaniform sensilla position them to respond preferentially to self-generated force [58]. For example, campaniform sensilla on the stick insect trochanter are positioned to encode increases and decreases in mechanical load at the nearby leg joint [63]. Campaniform sensilla neurons are active when leg movements are resisted with a mechanical probe but do not fire during unresisted leg movements, indicating that they encode mechanical load as resistance to muscle contraction [63,64]. In this regard, the campaniform sensilla perform a function similar to that of vertebrate Golgi tendon organs [4]. However, unlike tendon organs, the coupling of campaniform sensilla to muscle tension is indirect and depends on joint position [65].

The functional contributions of insect campaniform sensilla have been studied in the context of both walking and flight control. In particular, the stick insect has served as an important model system for investigating the role of sensory feedback from campaniform sensilla in postural control and walking (reviewed in [66]). For example, campaniform sensilla on the trochanter, the second leg segment, contribute to joint coordination during walking by setting the firing phase of motor neurons that control movement of the first leg segment, the coxa [67]. Under normal conditions, input from trochanter campaniform sensilla terminates activity in the protractor motor neuron of the coxa and initiates activity in the corresponding retractor motor neuron. Accordingly, ablation of the trochanteral campaniform sensilla substantially decreases the magnitude of bursts in coxal motoneurons [67]. Campaniform sensilla can also contribute to inter-leg coordination by producing muscle bursts at appropriate phases of the step cycle, after a leg is placed on the substrate [68]. These studies have shown that sensory feedback from campaniform sensilla plays an important role in coordinating joint movement both within a leg and across legs and that natural walking gaits are shaped by sensory feedback to motor circuits.

In flies, campaniform sensilla also play an important role in flight control through feedback from both the halteres and wings. Halteres are small dumb-bell shaped organs derived from the hindwings, which beat back and forth in antiphase to the wings. When the body rotates during flight, arrays of campaniform sensilla at the base of the halteres detect inertial (Coriolis) forces that are linearly proportional to the angular velocity of the body (reviewed in [69]). Campaniform sensilla on the wings of locusts [70], moths [71], and flies [72] also encode wing-bending forces.

Fast sensory feedback from haltere campaniform sensilla is thought to mediate compensatory flight control reflexes in flies [73]. For example, one haltere campaniform field (dF2) provides direct synaptic input to a motor neuron (mnb1) that controls a flight steering muscle [74]. The mnb1 motor neuron can also be entrained by direct input from wing campaniform sensilla [75]. Wing and haltere campaniform sensilla each fire a single action potential at unique phases within a wing stroke cycle, which in Drosophila lasts about 4–5 ms. Together, the relative phase difference between wing and haltere feedback may contribute to maintaining flight equilibrium by tuning the firing phase of steering muscles during flight [75]. Haltere campaniform sensilla exhibit diverse phase sensitivity [76], suggesting that sensory feedback is capable of fine-tuning flight control throughout the wingstroke.

**Proprioceptive Chordotonal Organs**

Chordotonal organs are internal mechanoreceptors found at nearly every joint and between joints within individual limb and body segments. The fundamental unit of a chordotonal organ is the scoloparium, which is composed of 1–3 bipolar mechano-sensory neurons and 2 accessory cell types, the scolopale cells and cap cells, which envelop and anchor the sensory neurons, respectively (Figure 1E). The chordotonal organ is attached to the cuticle wall or muscles by connective tissue or can be linked to a joint by an outgrowth of cuticle called an apodeme. Within a given chordotonal organ, there is diversity in the structure and attachment of individual scolopidia, which may be related to the diverse mechanical tuning of the chordotonal sensory neurons [77].

Most chordotonal organs are proprioceptors, but some (discussed separately below) function as exteroceptors. In flies, proprioceptive chordotonal organs are associated with the legs, wings, halteres, and mouthparts. Genetic experiments in the fly have demonstrated the important proprioceptive role of chordotonal neurons in many motor functions. Perturbation of the chordotonal organs in adult Drosophila produces deficits in locomotion and posture [78–82].

Although relatively little is known about the physiology of proprioceptive chordotonal organs in adult Drosophila, there is an extensive literature about these neurons in big insects (reviewed in [83]). In particular, the femoral chordotonal organ (fCO) of the locust and stick insect has been investigated in great detail. The fCO is comprised of several hundred chordotonal neurons that are located in the femur and mechanically coupled to the tibia [77,84,85]. Neurons of the fCO are organized into two distinct clusters; the larger cluster (~400 neurons) is more distal and ventral within the femur and the smaller (~80 neurons) is more proximal and dorsal [77,86]. This organization is similar to that found in Drosophila, though the fly fCO contains fewer neurons [87].

The dorsal and ventral clusters of the fCO are thought to detect distinct mechanical stimuli and contribute to different behaviors [77,86]. Neurons in the larger, dorsal cluster encode high-frequency (200–800 Hz) vibration of the tibia [88] and may play a role in sensing substrate vibration [89]. In contrast, neurons in the smaller, ventral cluster are sensitive to the position of the tibia; individual cells in this group are typically classified as either slowly adapting (tonically firing) neurons or rapidly adapting (phasically firing) neurons (Figure 4A,B). Tonic neurons encode position of the joint between the femur and the tibia, while phasic neurons encode velocity and acceleration of joint movement [84,90,91]. Interestingly, individual neurons within the ventral cluster respond to rather specific features of the leg’s position...
and movement. Single neurons may strictly encode the leg's position but only over a narrow range, or else may encode velocity invariant to position, or else position-and-velocity, or velocity-and-acceleration, etc. [84,92–95].

A notable property of chordotonal neuron encoding is hysteresis — the relationship between firing rate and joint position can depend on the preceding direction of movement (Figure 4C) [84,90–92]. From a decoding perspective, hysteresis introduces ambiguity about joint position. However, it is also possible that hysteresis in sensory neurons may actually compensate for nonlinear dynamics in other parts of the circuit, such as muscles [96].

An important function of the fCO is to mediate basic resistance reflexes. For example, movement of the joint between the femur and the tibia is detected by chordotonal neurons within the ventral cluster, which directly excite flexor motor neurons and inhibit extensor motor neurons through polysynaptic pathways, leading to changes in muscle activity to oppose the movement [77,86]. This reflex serves to stabilize the position of the joint and maintain the posture of the animal. Importantly, the reflex is reliable across a wide range of initial joint positions, movement directions, and velocities [97,98]. The sign and efficacy of the femoral resistance reflex is not fixed but depends on behavioral context (see below). Given that individual chordotonal neurons are often narrowly tuned to particular combinations of position, velocity, and acceleration, it would seem that many chordotonal neurons must contribute to the control of these seemingly simple leg reflexes. However, the function of chordotonal neuron population coding remains relatively unexplored.

Studies in the locust have shown that the presynaptic terminals of fCO neurons receive rhythmic inhibitory synaptic input during walking [99]. This presynaptic inhibition can be driven by activity of chordotonal neurons [100] or other proprioceptors within the same leg [101]. This inhibition does not arise from direct interactions between mechanosensory neurons but more likely through GABAergic or glutamatergic interneurons whose identity is currently unknown [102]. Presynaptic inhibition may serve to maximize dynamic range by controlling the output gain of the chordotonal neuron population [103]. Alternatively, it may suppress expected mechanosensory input at particular phases within a motor rhythm in order to increase sensitivity to unpredicted sensory signals [99]. Presynaptic modulation of neurotransmitter release is also a feature of vertebrate mechanosensory afferents, though its function remains poorly understood [104].

The activity of chordotonal neurons can be strongly influenced by the presence of neuromodulators. For example, neurons in the fCO of the stick insect [105] and the locust [106] increase their firing in the presence of octopamine, the insect analog of noradrenaline. Interestingly, this gain increase is specific to tonically firing neurons that encode joint position [105]. Octopamine can act by directly increasing the excitability of chordotonal neurons in the leg or by modulating presynaptic inhibition of chordotonal neuron axon terminals [106].

**Exteroceptive Chordotonal Organs**

In addition to the proprioceptive chordotonal organs described above, many insects possess specialized chordotonal organs that are used to detect external mechanical signals. Although these organs are formally still proprioceptors — because they are located inside the body and because they detect the position or movement of body parts — they function primarily as exteroceptors. This is because they monitor body parts that generate relatively large movements in response to small external forces.

In *Drosophila*, the most prominent example is Johnston’s organ, a chordotonal organ that resides inside the second-most distal segment of the antenna. It senses movements of the most distal antennal segment, which rotates easily about its long axis and therefore generates relatively large movements in response to small fluctuations in air particle velocity. In *Drosophila*, Johnston’s organ is comprised of ~500 neurons [107]. This organ detects sound, including male courtship song, which is produced when the male fly extends and vibrates his wing while pursuing a female (reviewed in [108,109]). Johnston’s organ also detects wind [110], and it even responds to the fly’s own wingbeats in flight [111]. Finally, there is some evidence that Johnston’s organ is involved in encoding gravity, because the distal antennal segment can move as much as 1 μm in response to changes in gravitational forces [112]. Johnston’s organ neurons seem to be divided into rapidly adapting types and slowly adapting types, some of which are also selective for the direction of antennal rotation (toward the head or away from the head) [110,112,113].

Many insects (including crickets, bees, locusts, and some flies) possess tympanal organs, specialized hearing structures that consist of a thin cuticular membrane stretched across an
air-filled sac. Chordotonal organs attached to the tympanal membrane detect high frequency (2–100 kHz) mechanical vibrations, acting much like a vertebrate eardrum (reviewed in [114]. Drosophila do not have tympanal organs and therefore rely on the Johnston’s organ for audition, limiting their hearing to lower frequencies (< 1 kHz).

Another example of a specialized chordotonal organ is the subgenual organ, a fan-shaped array of scolopidia in the tibia of most insects that is sensitive to vibrations of the substrate [115,116]. Subgenual organs have not been reported in flies. However, Drosophila females appear to detect substrate-borne vibrations during courtship [117], possibly through the fCO or campaniform sensilla.

**Multipolar Receptor Neurons**

Multipolar receptors are non-ciliated neurons with multiple dendritic branches that terminate on a variety of internal organs and tissues. In Drosophila larvae, such multipolar neurons are commonly referred to as multidendritic neurons [118,119], which are distinct from ciliated neurons with a single dendrite, such as those associated with tactile hairs, campaniform sensilla, and chordotonal organs. Multidendritic neurons in the larval body wall are active during peristaltic muscle contraction, indicating that they respond to self-generated forces during locomotion [81]. Some multidendritic neurons persist through metamorphosis and are found in the abdomen of the adult fly [120].

Multipolar receptors are found throughout the insect body, typically embedded within an accessory structure such as a strand of tissue or muscle, or slung between protuberances of cuticle called sclerites. The proprioceptive role of multipolar receptors has been studied in great detail in larger insects. For example, in the locust, feedback from wing hinge multipolar receptors directly modulates the motor patterns that control flight [121,122]. Multipolar stretch receptors in the moth wing signal the position of the wing during flight by encoding the amplitude, rather than the velocity, of stretch [123]. Ablation of wing hinge stretch receptors decreases the amplitude of visually evoked flight responses, indicating that multipolar receptor feedback is integrated with descending motor commands to control wing movement [124].

Multipolar receptors are also found in the viscer. For example, in the blowfly, internal stretch receptor neurons monitor the distension of the fly gut and provide a signal that terminates feeding behavior [125]. In adult Drosophila, multipolar receptors have been identified in the proboscis [126] and at leg joints [127–129], though their function in the fly remains poorly understood.

**Other Mechanoreceptors**

There are several mechanoreceptors found in insect legs whose physiology and function remain enigmatic — these include the strand receptors, muscle receptor organs, and tension receptors. None of these mystery mechanoreceptors has yet been described in Drosophila, but given the overall homology of mechanoreceptor structure and location in the legs of insects [130,131], genetic tools for labeling specific neuron types may soon enable their identification in the fly.

Strand receptors are unique among mechanoreceptor neurons, in that their cell bodies are located within the CNS and they project dendrites into the periphery [132]. Like multipolar stretch receptors, they monitor stretch across joints or muscles, most notably at joints in the locust and cockroach legs [133].

Muscle receptor organs (MROs) are single multipolar receptor neurons associated with a modified muscle, which is itself innervated by a specialized motor neuron. MROs are thought to monitor changes in muscle position in a manner similar to vertebrate muscle spindles; for example, MROs that encode joint position have been identified in the locust [134]. However, in addition to detecting mechanical movements of the joint, the MRO motor neuron receives direct input from other proprioceptive organs, so that responses of muscle receptor organs result from both afferent and efferent signals from other mechanoreceptors [135].

Finally, some multipolar proprioceptors, referred to as tension receptors, can be found directly embedded in muscles. For example, in the locust, a single tension receptor is associated with the flexor tibiae muscle in the leg [136], while 200 tension receptors are located within a single ovipositor muscle that controls egg-laying behavior [137]. Tension receptors detect the forces generated by a muscle — they are most sensitive to active muscle contraction, rather than passive movement [136–138].

**Central Projections of Peripheral Mechanoreceptors**

Peripheral mechanoreceptors on the insect body send direct axonal projections into the central nervous system. Detailed anatomical studies in a number of insect species have shown that these axons are systematically organized depending on mechanoreceptor location, physiological tuning, and receptor type.

A well-studied example of axonal mapping is the somatotopic organization of afferents from leg bristle neurons. In Drosophila bristles on each of the three pairs of legs project to a corresponding compartment, or neuronomere, within the ventral nerve cord (VNC). Axons from bristle sensory neurons are then topographically organized within each neuronomere — bristle neurons on the anterior surface of the front leg arborize along the anterior edge of the prothoracic neuronomere, while posterior bristles arborize posteriorly (Figure 5A) [139]. This topographic map of the leg surface within each neuronomere would allow central neurons to more easily sample inputs from groups of nearby bristles. This is relevant because, due to the structure of the natural world, nearby bristles are most likely to exhibit correlated activity.

Similar somatotopic maps of tactile hair axons have been described in big insect species, including the locust [140], cockroach [141], and cricket [142]. A topographic map of mechanoreceptor afferents also exists in the dorsal horn of the mammalian spinal cord [143] and the trigeminal nuclei of the brainstem [144]. The existence of such maps across distantly related species suggests that topographic organization of primary sensory afferents plays an important functional role in central processing of somatosensory signals.

Roughly orthogonal to the somatotopic map, there is another axis of organization in the fly VNC. Namely, different sensory modalities arborize in different layers along the dorsal-ventral axis of the VNC (Figure 5B). Axons from bristle neurons arborize along the ventral edge of the neuropil, while campaniform sensilla and hair plate axons terminate in a more dorsal region termed the intermediate layer [47]. Leg chordotonal neurons arborize along the medial border of this intermediate layer [128]. The
most dorsal region of the VNC is dominated by motor neurons [145,146]. Thus, proprioceptive afferents are positioned close to dorsal motor neuron dendrites, while tactile signals from hairs are segregated in more ventral layers. Again, the organization described here for Drosophila is roughly similar to that in other insects, such as the locust (reviewed in [147]).

As a general rule, mechanoreceptors are not thought to make synapses outside the CNS. An exception is the fCO of Drosophila. Although the majority of their synaptic output is probably in the VNC, some femoral chordotonal neurons also form a peripheral ‘glomerulus’ of synapses within the leg nerve [87].

Sensory axons from a single receptor type may also form maps that correspond to their physiological tuning. This organization may facilitate the integration of correlated signals in downstream neurons. For example, the neurons that make up the cricket tympanal organ arborsize in a tonotopic pattern related to their frequency sensitivity [148–150]. There is also evidence for a correlation between the physiological tuning of neurons in the locust fCO and their axonal morphology in the CNS [151]. Similarly, in the Drosophila brain, the axonal projections of chordotonal neurons from Johnston’s organ are organized according to their tuning [110,152].

Compared with mechanoreceptors on the legs, less is known about the central projections of mechanoreceptor neurons from other regions of the insect body. One well-characterized set of central projections are those of the large filiform hairs of the locust head and thorax, whose axonal projections into the VNC have been used to study activity-dependent circuit formation (reviewed in [153]). Similarly, the central projections of Drosophila thorax bristles have been characterized in genetic studies of axon targeting [154] and synaptogenesis [155]. However, the central organization of head mechanoreceptors remains poorly characterized. In the few cases that have been examined, dye fills have shown that the arborizations of individual head mechanoreceptor neurons can be extensive. For example, tactile hairs on the cricket eye [156], filiform hairs on the locust head [157], and campaniform sensilla on the blowfly antennae [158] send axonal projections to both the brain and the VNC. In these examples, arborizations in the brain appear to be concentrated in ventral regions, including the subesophageal ganglion and the antennal mechanosensory and motor center.

From Mechanosensation to Action: the Problems Faced by the Central Nervous System

In turning to the topic of central mechanosensory circuits, it is useful to consider the universal ‘problems’ that the mechanosensory systems of all animals must solve. Below, we list some of these basic problems, and we briefly describe recent work that illustrates how insect nervous systems can solve these problems. We then focus on two of these problems in greater detail in the following sections. Readers wanting more detail on any of these topics should consult Malcolm Burrows’ book [147] and other classic reviews [159,160], as well as some more recent perspectives [4,42,66,161–164].

Problem 1: Mechanoreceptor spikes often must be processed rapidly. Some mechanical signals demand an immediate behavioral response. The fastest motor responses to mechanical perturbations are mediated by the intrinsic biomechanical properties of the body, prior to the involvement of any neurons at all (e.g., [165,166]). But there is also evidence that neural mechanisms are often adapted for speed. The extreme diversity and specialization of mechanoreceptor cells is one such adaptation: in essence, the peripheral nervous system performs much of the computational work of the mechanosensory processing system, and in doing so, it eliminates the need for additional layers of CNS processing. Another adaptation for speed can be seen at the very first stage of processing in the CNS, where single mechanoreceptor axons can diverge to synapse onto central neurons that perform distinct computations in parallel [167,168]. By parallelizing central computations in this manner, the CNS can extract information as rapidly as possible. Speed is also achieved in part by making computations very local, thus minimizing the number of long axons and synaptic relays in any given circuit. For example, in the extreme case, many proprioceptive mechanoreceptors synapse directly on motor neurons [147,169,170]. Finally, in some cases, neural processing delays can be avoided altogether by taking advantage of passive mechanical conduction. For example, a mechanical stimulus applied to the locust tarsus propagates through the leg cuticle to be detected by campaniform sensilla on the proximal femur and trochanter [171]. This mechanical conduction occurs in less than 1 ms, compared to the 8 ms delay for conduction in the axons of distal campaniform sensilla neurons.
**Problem 2:** The implications of a mechanoreceptor spike train for motor control can depend on the position of the body. An intuitive example of this problem can be seen whenever an insect makes searching movement with its forelegs or antennae. To accurately locate the position of an object, the nervous system must take into account the leg’s position when an object is contacted. A recent study shed light on this problem in stick insects. When a leg makes transient contact with an object, a stick insect responds by searching in a tight local pattern around the site of contact [172]. Accurate local searching behavior does not require visual input but does require an intact hair plate at the coxa-trochanter joint, suggesting that this proprioceptive organ provides key information about the joint’s position at the moment of contact. Intriguingly, searching behavior can be evoked by depolarizing current injection into a single identified ‘command interneuron’ in the appropriate neuroromere [173]. However, if the foot is in contact with the substrate, then depolarization of this command interneuron does not produce searching. It will be interesting to determine where foot touch receptor signals act to gate searching behavior, and to learn how proprioceptors determine the coordinate frame of searching patterns. The coordinate frame problem also applies to the relationships between different appendages. For example, when a stick insect is exploring a gap with its antenna, a contact between the antenna and the far edge of the gap is sufficient to initiate a directed movement of the forelegs to the far side of the gap, implying that the antenna’s position at the moment of contact is being relayed to the leg control system [174].

**Problem 3:** Mechanosensory cues must be integrated with cues from other sensory modalities. In many cases, motor actions are guided by signals from multiple sensory modalities, including mechanosensation, vision, and olfaction. The integration of cues from different sensors is a hard problem because different sensory modalities operate on different timescales and are also formatted within different coordinate frames (e.g., retinacentric versus leg-centric coordinates). The solution thus requires both temporal integration and a spatial coordinate transformation. Recent studies in insects illustrate specific instances where mechanosensory cues are integrated with cues from other modalities. For example, cockroaches respond to an obstacle with a range of different climbing strategies depending on both antennal mechanoreceptor signals and visual cues [175]. Similarly, the wings of flying flies are controlled by both visual and mechanosensory cues operating on different timescales [176–178]. Multi-sensory integration may occur at multiple sites within the central nervous system. In flies, for example, visual commands from the brain descend to the VNC, but campaniform sensilla neurons on the wings and halteres also project directly to the brain [179].

**Problem 4:** Mechanoreceptors must modify the activity of synergistic muscle groups. Most movements involve the synergistic action of multiple muscles, with different motor patterns engaging different synergies (Figure 6A). This situation creates a constraint on mechanosensory feedback. Specifically, it means that mechanosensory feedback will be most effective when it targets the muscles that participate synergistically in the same locomotor pattern. Below we will discuss the topic in more detail in a section dedicated to muscle synergies.

**Problem 5:** There is a fundamental tradeoff between stability and maneuverability. For example, when an animal is standing still, postural reflexes counteract disturbances in order to ensure a robust stance. However, when the organism needs to execute a motor pattern such as walking, postural reflexes may actively oppose the movements required for swinging the leg. The solution to this problem is to suppress (or even reverse) postural reflexes during voluntary movement. In essence, reflexes are managed by the CNS to promote stability in some cases and maneuverability in others. Below we will examine this topic in more detail in a section dedicated to ‘reflex reversal’.

**Solving the Muscle Coordination Problem: Feedback Loops Targeting Muscle Synergies**

Movements are executed by the coordinated action of multiple muscles. For example, when a flexor is activated, its corresponding extensor is often relaxed. These ‘muscle synergies’ may extend across multiple muscle groups that control different segments of a limb (reviewed in [180,181]). For example, to pull an object toward our body, we synergistically activate flexors in both our arm and hand. Because mechanosensation is important for fine-tuning motor control, mechanosensory feedback signals might be expected to engage these same muscle synergies.

A recent study approached this question by studying how mechanosensory feedback engages muscle synergies in both the cockroach and the stick insect. Zill et al. [182] focused on a muscle synergy involving three muscles of the same leg — namely, the depressor of the trochanter, the tibial flexor within the femur, and the retractor unguis within the tibia (Figure 6A). In order to exert a strong inward grip on the substrate underneath the foot, these three muscles must contract together.

This study found that when specific groups of leg campaniform sensilla were stimulated, the motor neurons innervating all three of the relevant muscles were co-activated (Figure 6A). For example, stimulation of campaniform sensilla on the foot had this effect, as did stimulation of campaniform sensilla on the proximal trochanter. Thus, distinct peripheral mechanoreceptors can generate the same pattern of muscle co-contraction — that is, the same muscle synergy.

Importantly, the campaniform sensilla that had these effects are normally stimulated when the animal grips the substrate. The sensilla on the foot detect resisted forces exerted by the retractor muscle [183,184]. Similarly, the sensilla on the proximal trochanter detect resisted forces exerted by the trochanteral depressor. Thus, this circuit consists of multiple positive feedback loops, which operate in parallel to increase grip.

Compared with vertebrates, insects have far fewer motor neurons per muscle. For example, Drosophila leg muscles are typically innervated by less than five motor neurons and sometimes as few as two [145,146]. Invertebrates also make extensive use of inhibition and neuromodulation to fine tune muscle activity (reviewed in [185]). In the future, it will be interesting to explore whether the organizing principle of muscle synergies extends to these inhibitory and neuromodulatory motor neurons.

**Solving the Stability–Mobility Tradeoff Problem: Reflex Reversal**

We define a reflex as a behavior mediated by signaling from sensory neurons to motor neurons through a direct or nearly direct
pathway (i.e., with few intervening synapses). Naively, we might imagine that reflex behaviors are immutable and that the neurons that mediate them have fixed relationships. In reality, however, most reflexes are flexible, and can be dramatically modified depending on behavioral context.

For example, stretch reflexes, or resistance reflexes, serve to stabilize posture in both invertebrates and vertebrates. These resistance reflexes are mediated by feedback circuits. The most basic implementation requires just three components: a sensory neuron, a motor neuron, and a muscle (Figure 6B). When the muscle is stretched, the stretch is detected by the sensory neuron, which provides excitation to the motor neuron, leading to contraction of the muscle, thereby opposing stretch. The function of this reflex is to help maintain posture during changes in mechanical load.

The circuits that underlie resistance reflexes can be remarkably flexible, exhibiting both short-term modulation as well as long-term plasticity (reviewed in [186]). One of the best examples of reflex flexibility is that of the stick insect tibia, originally described by Bässler [187]. In a stationary stick insect, stretching of the fCO excites tibia extensor motor neurons and inhibits tibia flexor motor neurons (Figure 6C). During contralateral stepping, turning to the right, or backward walking, fCO stretch excited the tibial extensor (top row), but during stepping of the ipsilateral front leg, leftward turning, or straight forward walking, fCO stretch produced a distinct pattern of tibial muscle activity (bottom row).

Figure 6. Mechanosensory processing in reflex circuits of insects.
(A) Left: schematic illustrating the concept that a muscle synergy involves multiple co-activated muscles, and different behaviors recruit different muscle synergies. Middle: a specific muscle synergy in the insect leg. Muscles indicated in green are co-activated when leg grips the substrate. (Antagonist muscles are shown in gray.) Right: campaniform sensilla on the leg provide positive force feedback to motor neurons that participate in this muscle synergy. Specific clusters of campaniform sensilla on the tarsus and trochanter (green arrows) are activated when the leg grips the substrate, and their activation leads to increased grip (adapted from [182]). (B) Schematic connectivity diagram of mechanosensory reflex circuits in the insect VNC. The basic circuit supporting the resistance reflex is outlined in gray, and inputs that modulate the sign and efficacy of the reflex are outlined in red. (C) Reflex reversal in the stick insect is specific to behavioral context (adapted from [188]). Extracellular recordings were made from the tibial flexor and extensor muscles and motor neurons while mechanically stimulating the femoral chordotonal organ (fCO) by pulling on the receptor apodeme. During contralateral stepping, turning to the right, or backward walking, fCO stretch excited the tibial extensor (top row), but during stepping of the ipsilateral front leg, leftward turning, or straight forward walking, fCO stretch produced a distinct pattern of tibial muscle activity (bottom row).
phenomenon is known as reflex reversal (reviewed in [159]). Reflex reversal is an important element of maneuverability: without it, the resistance reflex would oppose voluntary movement. Because the reversed reflex now reinforces rather than opposes movement, it is sometimes called an ‘assistance reflex’.

A recent study revealed the contextual specificity of the stick insect reflex reversal. Hellekes and colleagues [188] recorded activity from tibia motor neurons of the stick insect’s middle leg. They then mechanically stretched the ICO of the middle leg while the animal was walking. They found that reflex reversal did not occur during all active movements but depended on the particular task being executed. For example, they observed reflex reversal during stepping of the ipsilateral front leg but not stepping of the contralateral legs (Figure 6C). For the front leg, reflex reversal only occurred during forward walking; when the insect walked backward, this particular reflex did not reverse. This is likely related to the fact that the femur–tibia joint is flexed during stance in forward walking but is extended during stance in backward walking. In other words, during forward and backward walking, the leg muscles work together differently to create propulsion, and so the nature of reflex modification would naturally also be different.

Finally, Hellekes et al. [188] observed reflex reversal when the animal was attempting to turn to the same side as the manipulated leg but not when the animal was turning in the opposite direction (Figure 6C). Again, the sign of the reflex is matched to the requirements of the behavior: to execute a turn, the middle leg on the inside must pull by flexing the femur–tibia joint, whereas the middle leg on the outside must push. Overall, these experiments show that the resistance reflex is not simply reversed whenever the animal is active but can be specifically modulated during particular locomotor patterns. Indeed, it has been suggested that reflex modification is not simply a corollary of switching locomotor patterns but is the essence of the switch. For example, in a walking insect, a command to turn may consist simply of the modification of leg mechanosensory reflexes such that turning is the natural result [189,190].

What are the underlying sites and mechanisms of reflex reversal? As described above, there is some evidence that the output of mechanoreceptor neurons is directly modulated during behavior — for example, through presynaptic inhibition of the chordotonal neuron axon terminals by GABAergic feedback neurons [102]. However, it is also likely that the effect of sensory input upon motor neurons is influenced by VNC interneurons. One population of non-spiking interneurons in the stick insect has been shown to receive input from chordotonal neurons and provide input to tibia motor neurons [191,192]. It is thought that the balance of excitation and inhibition within this population regulates the sign of chordotonal feedback within leg motor neurons [193].

Descending signals from the brain may also be involved in modulating reflex reversals during locomotion. For example, in the cockroach, removing the descending projections from the brain alters certain mechanosensory reflexes [189]. A recent study extended this finding to show that electrical stimulation of neurons in the cockroach central complex can alter the tibial resistance reflex [194]. An appealing hypothesis is that diverse motor patterns can be generated by top-down modulation of basic sensorimotor loops like the resistance reflex. This is a very different strategy than that implemented in robots, in which high-level behaviors are typically achieved with explicit kinematic models [4].

The ‘Hard Problems’ of Sensorimotor Control and the Value of Insect Models

One way to identify compelling scientific problems is to ask which features of biological systems are most difficult to replicate in artificial systems. Thirty years ago, a reasonable answer might have been the problem of visual object recognition, which motivated many neuropsychological and computational studies of the primate visual cortex [195]. However, recent advances in artificial neural networks have made it possible for a computer to automatically classify natural images with accuracies that match, and sometimes surpass, human performance [196]. A plausible argument can be made that the efforts of both neuroscientists and computer scientists contributed to these advances in object recognition technology.

Today, flexible sensorimotor control represents a set of problems of a similar scale. Studies of insects are particularly well-suited to understanding the solutions to these problems, because insect nervous systems are compact, relatively stereotyped, and amenable to in vivo recordings — even intracellular recordings from targeted cell types in alert, behaving animals. As detailed above, there are some explicit and fundamental ‘hard problems’ of sensorimotor control that are faced by insects as well as other organisms. Already, neuropsychological and behavioral studies have taught us a great deal about how insects solve these problems. These biological insights are now providing inspiration for a new generation of artificial systems [4].

The Role of Drosophila in the Study of Mechanosensory Processing

There are many outstanding questions in insect mechanosensory processing. One of the most critical concerns the role of CNS circuits. Certainly, there is already a literature on CNS circuitry, and we have noted highlights from this literature. However, we are far from understanding how central circuits select and control complex motor behaviors in the face of external mechanical perturbations. One difficulty is the complexity of central circuits. In an insect, each thoracic neuromere — the central circuitry controlling a single leg — contains >100 motor neurons and several times as many local interneurons [4]. To understand central circuits of this complexity, we should aim to (1) identify neurons that are active during a particular behavior, (2) map their connections, (3) inject signals at particular locations in the circuit, and finally (4) break connections (and especially loops) in the circuit. Given this mission, Drosophila offers some powerful emerging tools.

Chief among these is the availability of many cell-type specific ‘driver’ lines that allow transgenes of interest to be expressed in specific neurons. Thousands of these lines are publicly available [197]. For example, a recent study identified driver lines targeting each of the major mechanoreceptor types in the Drosophila leg [168]. These sorts of driver lines are already being used to determine how gait is affected by silencing specific mechanoreceptor neurons [78–82]. Driver lines are also being identified that allow genetic access to specific central neurons of the VNC [168,198,199]. Meanwhile, another effort is currently underway...
to generate genetic driver lines that individually target many of the ~350 pairs of descending neurons that project from the brain to the VNC [200–202]. Descending signals from the brain are known to modulate the sign and amplitude of mechanosensory reflexes [189], for example, but the format of these signals is not well understood in any insect species. A collection of driver lines targeting descending neurons will be an important tool for answering this question.

Driver lines have two broad applications. One application is to perturb specific circuit elements — using optogenetic tools, or other genetic perturbations (reviewed in [203]). For example, driver lines targeting descending neurons will make it possible to inject signals directly into descending pathways, thereby showing whether these signals are sufficient to reverse sensory-motor loops like the leg resistance reflex, and whether this modulation is specific to particular reflexes and behaviors. The other application is equally important: driver lines allow neuroscientists to target electrodes and fluorescent activity reporters to almost any cell type of interest. Historically, insect neurophysiology was limited to recording from cell types that could be targeted using sharp glass microelectrodes — generally neurons with large-diameter dendrites or axons that could be stably penetrated with a sharp electrode. Now, in Drosophila, it is possible to label almost any cell type with a fluorescent protein, and thus make targeted patch-clamp recordings from virtually any central neuron of interest (reviewed in [204]). Another powerful tool in Drosophila is the ability to image neural activity in vivo using genetically encoded calcium or voltage indicators.

Studies in Drosophila will also benefit from a major ongoing effort to map the synaptic connections in the adult fruit fly. This effort uses serial-section electron microscopy combined with cell-type specific genetic tools to map synaptic connections between morphologically identifiable cell types [205,206]. Cell types identified morphologically can then be matched with driver lines using new bioinformatic tools [207,208]. This effort to map synaptic connections using large-scale electron microscopy has focused on the brain thus far, but it is currently being extended to the VNC as well.

The challenge, then, is to apply these tools to determine how central circuits integrate mechanosensory information to select different actions, such as grooming, walking, and take-off. To identify neurons that initiate changes in behavioral state, one could use pan-neuronal imaging to record activity while tracking behavior. Once relevant neurons have been identified, targeted in vivo electrophysiological recordings could then be used to understand how mechanosensory signals are integrated with other sensory cues and behavioral state changes to select appropriate behavioral responses. To test specific models of action selection, one could then bias behavioral transitions through targeted optogenetic stimulation in freely behaving flies. In order to make wise use of these tools, it is important to consider the lessons learned from work in big insects.

Indeed, although Drosophila presents many experimental advantages, there are also many questions that are still more amenable to investigation in other insect species. For example, some experiments are simply more feasible in larger insects. Ultimately, the identification of general principles for sensorimotor control will require investigation of multiple species and circuits. In the future, genome-editing tools like CRISPR/Cas9 may permit the use of genetic tools for targeted circuit manipulation in larger insects like the locust and cockroach. All of these considerations emphasize the broader point that insects are likely to be a source of future insights into mechanosensory processing, as well as a source of inspiration for roboticists and engineers.

REFERENCES


