Eight pairs of descending visual neurons in the dragonfly give wing motor centers accurate population vector of prey direction

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Intercepting a moving object requires prediction of its future location. This complex task has been solved by dragonflies, who intercept their prey in midair with a 95% success rate. In this study, we show that a group of 16 neurons, called target-selective descending neurons (TSDNs), code a population vector that reflects the direction of the target with high accuracy and reliability across 360°. The TSDN spatial (receptive field) and temporal (latency) properties matched the area of the retina where the prey is focused and the reaction time, respectively, during predatory flights. The directional tuning curves and morphological traits (3D tracings) for each TSDN type were consistent among animals, but spike rates were not. Our results emphasize that a successful neural circuit for target tracking and interception can be achieved with few neurons and that in dragonflies this information is relayed from the brain to the wing motor centers in population vector form.

Electrophysiology. We recorded from a total of 51 TSDNs from 38 different animals as follows: MDT1, 7 recordings; MDT2, 4 recordings; MDT3, 13 recordings; MDT4, 6 recordings; MDT5, 2 recordings; DIT1, 7 recordings; DIT2, 5 recordings; DIT3, 7 recordings. TSDNs respond only to small moving targets within the visual field (Fig. 1 A–C) and not at all to wide-field movement such as grating patterns (21). The identity of each recorded TSDN was ascertained from its target-direction preference and receptive field area. To sample TSDN responses in an unbiased manner and with high precision, we projected upon

We aimed to understand the information sent to the wings when a target moves across the dragonfly visual field. In particular, we have tested whether the population vector algorithm can successfully decode the directional component of the descending information. The population vector is the weighted sum activity of an ensemble of neurons with directional tuning. It was first shown to predict the direction of an upcoming arm movement in monkeys (7, 8). Since then, the population vector algorithm has described successfully the directional responses to mechanical stimulation in several invertebrate species (9–12). The dragonfly predatory flight provides a challenge for a reliable vector code. Not only are these animals highly maneuverable, with independent control of the fore and hind wings (13) and the ability to hover and even fly backward (14), but prey interception also demands a highly accurate representation of the prey’s direction vector (15). Nonetheless, computer modeling by Vaidyanathan et al. (15) suggested that this algorithm could drive an airborne target interception system successfully.

In the dragonfly brain, a group of neurons called small target movement detectors (STMDs) respond when a small target moves in a broad range of directions (16). Although there is no evidence showing that STMDs code a population vector, they are believed to be upstream of the target-selective descending neurons (TSDNs) (16, 17). TSDNs are thought to command the dragonfly trajectory during fast aerial attacks (5) for the following reasons: (i) each TSDN type, identifiable by its profile in the brain, behaves as a distinct feature detector, with a particular preference for target location, direction, and size (18, 19); (ii) TSDNs receive the information about the position and direction of a small target and relay it to all three (prothoracic, mesothoracic, and metathoracic) ganglia (18, 19); (iii) the axon diameters of the TSDNs are among the largest found in the ventral nerve cord (VNC) (18), affording a high conduction speed necessary for the behavior; and (iv) electrical stimulation of individual TSDNs alters the position/angle of the wings (20). Therefore, it is important to examine whether TSDNs use a population vector to accurately code prey direction and control wing movement.

In the CNS, there are only eight pairs of TSDNs with receptive fields facing dorsofrontally. We hypothesized that this small number of TSDNs is sufficient to transmit a finely tuned population vector that codes the direction of prey movement. To test this, we carried out intracellular recordings and confocal imaging of the TSDNs in the dragonfly Libellula luctuosa. Our results show that the population vector of the TSDNs codes with extreme accuracy the prey direction in all of the 360° tested.

Results

Electrophysiology. We recorded from a total of 51 TSDNs from 38 different animals as follows: MDT1, 7 recordings; MDT2, 4 recordings; MDT3, 13 recordings; MDT4, 6 recordings; MDT5, 2 recordings; DIT1, 7 recordings; DIT2, 5 recordings; DIT3, 7 recordings. TSDNs respond only to small moving targets within the dorsofrontal visual field (Fig. 1 A–C) and not at all to wide-field movement such as grating patterns (21). The identity of each recorded TSDN was ascertained from its target-direction preference and receptive field area. To sample TSDN responses in an unbiased manner and with high precision, we projected upon
a white screen a sequence of 3,497 trajectories, each composed of a moving dot that appeared in a random location and moved in a random direction, but always with a straight trajectory (Fig. S1A). To allow long recordings and prevent TSDN inhibition, the target size was small, subtending 1.48° (diameter) on the dragonfly’s eye, its speed fast (134°/s) and the trajectory time short (100 ms).

The latency between stimulus onset (first target movement) and the earliest TSDN response (Fig. S1B) was not significantly different among TSDN types (Fig. 1D). Thus, the timing of information arrival was similar among all TSDNs at the temperature tested (23–25 °C). This was also the case for the off latency (timing of last response – timing target movement cessation). After pooling, the TSDN onset latency (29.94 ± 5.75 ms, mean ± 1SD) and offset latency (28.82 ± 9.11 ms) were not significantly different (paired Student t test, P = 0.848, n = 51) (Fig. 1D).

These results indicate that, within the 100 ms time window of each target trajectory, the latency between stimulus onset and cell response did not vary significantly.

The responses from 51 cells were grouped into eight TSDN types, whose electrophysiological properties (Fig. 2) and brain traces (Fig. S2) show homology to those described in the dragonfly Anax junius (18, 19). For example, DIT1, DIT3, and MDT1 all have receptive fields contralateral to their descending axons, whereas the rest (MDT5, MDT3, DIT2, MDT2, and MDT4) have receptive fields ipsilateral to their descending axons (Fig. 2 and Fig. S2).

Although all TSDNs were excited by a wide range of target directions and the total number of spikes for a TSDN type varied among animals (Fig. S3), the peak direction preferences were TSDN-type specific and consistent (Fig. 2A and B, polar plots). The tuning width of the directional tuning curves varied between 32° and 60° (Fig. S4A). The widest direction-tuning curve is that of DIT3, which may result from its sensitivity to an expanding stimulus (Fig. S4B). A similar range of tuning widths has been reported in the cells coding for direction in the monkey motor cortex (22). The basis for such cell-to-cell tuning-width differences remains to be clarified in both cases.

In addition to the consistency in the location and size of the receptive field, the zones of increased sensitivity (where a higher number of spikes were recorded in response to a moving target) also remained similar among replicates of each TSDN type (Fig. S5). To assess whether the preferred target direction is a function of a specific area within the receptive field, for each screen pixel the mean target direction was calculated from all of the targets that triggered a spike. For most cells, the preferred direction was homogeneous across the receptive field. However, DIT3 and MDT3 also displayed responses that were not directionally sensitive [Fig. 2A and B, direction receptive field (DRF) maps, area beyond the midline with highly variable coloration]. In the case of MDT3, such responses were tightly localized, and thus correlated with the location of the target stimulus. Hence, these responses are not likely to reflect intrinsic cellular noise. Instead, it is probable that these two cells receive additional, nondirectional, inputs from both eyes. This is important because only DIT3 responded significantly to an expanding target, a stimulus type that included moving edges in all directions (Fig. S4B).

Moreover, preliminary data from the Libellulid dragonfly Libellula pulchella indicates that, although MDT3 does not respond to a linearly expanding stimulus, it does so when an accelerated expansion, quasi-loom stimulus is presented in the middle of the view (DRF) maps show the directional preference of each recorded TSDN (red dots) and their mean direction tuning distribution (black bars). The red arrow indicates mean preferred direction. The color-coded direction receptive field (DRF) maps show the mean direction preference at each pixel, which was calculated by averaging the direction peak, at each pixel, for all recordings of each TSDN type. In addition, spike-triggered average (STA) maps are shown for each TSDN type. Note that, because number of spikes was normalized before and after computing the average, the same scale applies to all STA maps.

Fig. 2. Each TSDN type shows a unique direction tuning curve and receptive field consistent across animals. (A) Contralateral and (B) ipsilateral TSDN receptive fields whose axes were in the right connective of the ventral nerve cord (VNC). The polar plots show the directional preference of each recorded TSDN (red dots) and their mean direction tuning distribution (black bars). The red arrow indicates mean preferred direction. The color-coded direction receptive field (DRF) maps show the mean direction preference at each pixel, which was calculated by averaging the direction peak, at each pixel, for all recordings of each TSDN type. In addition, spike-triggered average (STA) maps are shown for each TSDN type. Note that, because number of spikes was normalized before and after computing the average, the same scale applies to all STA maps.
visual field (MDT3 responses per accelerated expansion stimulus, 4 spikes; \( n = 3 \)). We note here that receptive field laterality was inadvertantly switched for two TSDNs, MDT1 and MDT3, in the previous description of TSDN receptive fields (19). The corrected receptive field maps for Aeshnids agree with those reported here for Libellulids.

In all TSDNs, the “hot spots” of increased sensitivity are located near the visual midline. Although the size of our screen did not cover the receptive fields in the frontal-caudal axis entirely, the hot spots seemed to be placed dorsofrontal. This trend becomes more apparent when all of the spike-triggered average (STA) maps for each TSDN type are added and mirrored. This operation yields a sensitivity map for the space sampled by both eyes (Fig. 3A; see Fig. S6 for dragonfly head angles during the recordings). Our results show that dragonflies boast a matched detection system, with increased sensitivity to moving targets in the area of the retina where the prey image is focused during aerial hunts (1). In addition, plotting the directional tuning curve for each TSDN demonstrates that the directions pointing toward the back of the animal are sampled more intensely (Fig. 3B).

**Morphology.** The overlapping direction tuning curves and receptive fields of the different TSDNs suggest that the population vector from all TSDN responses could code a change in the prey’s bearing. However, to calculate a plausible population vector for each wing, knowledge of the TSDN arborization in wing motor centers is necessary. Moreover, to make morphological comparisons among the different TSDNs, the level of anatomical consistency of each TSDN type across different animals within the same species must be assessed first.

Earlier reports were studies of the TSDNs of the large “hawking” dragonflies of the family Aeshnidae, but the species used in this study belongs to a family of smaller “perching” dragonflies, the Libellulidae. We have confirmed, by electrophysiology (Fig. 2) and brain morphology (Fig. S2), homology between TSDNs from Aeshnids and Libellulids, but with one striking difference. In the prothoracic ganglia of *L. lactuca*, MDT2, MDT3, and MDT4 travel through the dorsal intermediate tract (DIT) (Fig. S7).

In all but one preparation, only one TSDN was injected in each animal. Three-dimensionally imaged volumes of the complex formed by the mesothoracic and metathoracic ganglia and the first abdominal ganglion were registered and warped for comparison (Fig. 4A and B). The traces for each TSDN of the same type are remarkably consistent across all injected samples (Fig. 4C). All of the TSDNs targeted the dorsal part of mesothoracic and the metathoracic ganglia (Fig. 4C) dedicated to motor/afferent fibers in insects. Thus, if the TSDN branching pattern does not filter the electrical activity in a ganglion-specific manner, the information provided by each TSDN should be available nearly simultaneously to both the fore and hind wings of the targeted side. Once the TSDN morphology was confirmed across several animals, we chose the cell with the most extensive fill for comparisons among TSDN types. According to the complexity of their branching, we divided TSDNs into two categories: simple or complex.

Unilateral simple TSDNs, DIT1, DIT2, and MDT2, consistently target four ipsilateral locations in the lateral region of the mesothoracic and the metathoracic ganglia (Fig. 4C). DIT1 morphology is closely matched by DIT2 and MDT2 (Fig. 4D), although our best MDT2 fill is incomplete. MDT1 and MDT4 show bilateral asymmetry in their outputs (Fig. 4C), but a pairwise comparison with unilateral DIT1 reveals that the branches of MDT1 and MDT4 target the four stereotypical lateral locations in both sides of the ganglia (Fig. 4D). Indeed, when we injected the DIT2 cell on the left connective and the MDT1 on the right connective of the same animal, the contralateral branches of MDT1 extended to the same location as those of the unilateral DIT2 (Fig. 4B).

We have categorized DIT3, MDT5, and MDT3 as complex cells due to the additional number of short and highly variable branches found in the medial region of the ganglia (Fig. 4C). However, within the side of the connective in which they descend from the brain, the morphology of these complex cells also targets the four lateral locations mentioned previously (Fig. 4E).

**TSDN Population Vector Coding.** Because all of the TSDNs terminals arborize in similar areas, it is plausible that their outputs code a population vector that is converted into wing motor movements. However, a plausible population vector calculation must take into account the receptive fields and unilateral/bilateral TSDN branching morphology. Simmons (23) reported that the major motor neurons involved in flight control have extensive branching in the ipsilateral side, but rarely cross the midline. Therefore, we chose a parsimonious approach in which the branching side of the TSDNs indicated the input to the corresponding wing. To understand which TSDNs would form part of a population vector, we focused on the TSDN activity elicited by targets moving on the left side of the dragonfly's visual field. We looked at the TSDNs with left receptive fields (Fig. 2 plus the corresponding mirror plots that would account for the TSDNs present in the opposite connective) and noted whether their outputs feed the ipsilateral (left) and/or contralateral (right) wing (Fig. 4 plus the corresponding mirror traces that would account for the TSDNs present in the opposite connective). Hence, we obtained the population of TSDNs contributing to the vectors for the wings ipsilateral (left) and contralateral (right) to the target movement (Fig. 5A). We then binned the target directions (and corresponding cell activity) every 18° and calculated the population vector (Fig. 5B) (Materials and Methods and SI Text). The results show that both population vectors, for the left and the right wing, are strongly correlated with target movement in all 360 directions tested (Fig. 5 C, i and D, i). Moreover, the accuracy is such that the population vectors and target movements are not significantly different in any of the tested directions (Fig. 5 C, ii and D, ii). Thus, the impressive ability of the system to code the target motion direction accurately is unaffected by the fact that the TSDN tuning curves do not sample all possible directions equally.

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![Fig. 3](image-url)  
**Fig. 3.** TSDN sensitivity is matched to the predatory behavior. (A) The visual receptive fields of the TSDNs combine to create an area of increased sensitivity to target movement near midline. For more details on eye coordinates, see Fig. S6. (B) DIT1, DIT3, and MDT1 transmit motion information in the visual field contralateral to their axons in the VNC. MDT5, MDT3, DIT2, MDT2, and MDT4 transmit motion information in the visual field ipsilateral to their axons in the VNC. Together, all eight axons travel through the ventral connective and thoracic ganglia.
In the wing motor centers, all TSDNs share morphological features. (A) The mesothoracic, metathoracic, and first abdominal ganglia were imaged and warped to allow morphological comparison. (B) Unilateral DIT2 on the left and bilateral MDT1 on the right, injected in the same animal, target the same locations. (C) Traces within each TSDN type (grouped according to the electro-physiological results) are consistent, so the most complete fill from each TSDN type was used for comparison. TSDNs were categorized into “simple” or “complex” cells. A pairwise comparison (dorsal views) shows that unilateral simple cells, DIT1 (green), DIT2 (red), and MDT2 (magenta), are indistinguishable from each other (D, Upper), but the bilateral simple cells MDT1 (white) and MDT4 (yellow) display specific branching patterns (D, Lower). However, all simple TSDNs target the same location. In contrast, pairwise comparisons between the complex cells, DIT3 (red), MDT3 (green), and MDT5 (cyan) (E, all panels) are less informative because their additional intricate branching exhibits higher variability, particularly in the medial region of the ganglia (traces in C).

In the absence of a moving target, the basal activity of TSDNs is zero. Therefore, a target moving in the antiprefered direction does not reduce the spiking rate. For such a system to code target direction equally well in all 360°, a minimum of four cells with cosine directional tuning are needed (24). We investigated whether all eight TSDN types are necessary for an accurate code of the target direction. We removed various numbers of TSDN types, from one (i.e., seven TSDN types left) to seven (i.e., one TSDN type left), from the population vector calculation and assessed the resulting performance. We found that, on average, if the population vector is calculated with six or more TSDN types, the vector is not significantly different from the target direction (r test, mean difference left wing, 1.23 ± 3.72° SD, P = 0.089; right wing, 1.19469 ± 3.49° SD, P = 0.07624; Fig. 5C, iii and D, iii), and thus its accuracy is not affected. The two exceptions arise when DIT2+MDT3 are removed from the ipsilateral wing calculation and MDT3+MDT4 are removed from the contralateral wing calculation; these changes result in large coding errors (population vector direction – target direction = 20.06 and 19.28°, respectively). When only two TSDN types were used in the calculations, the constant bias in the population vector outcome, also called constant error, was not far from the target direction (mean population vector direction – target direction, left wing, 9.67°, and right wing, 14.50°). However, the variable error, which can be regarded as a plus-or-minus confidence interval on the constant error, was large (mean error, left wing, 29.52°, and right wing, 40.12°), making the code unreliable. On average, at least three types of TSDNs are needed to keep the coding error with a constant bias below the 10° of the presented target direction. To describe the accuracy of this system in more detail, the activity of all TSDNs will have to be recorded with extracellular probes in the same animal. This is because although the preferred direction for a type of TSDN is highly constant between animals, the magnitude (the spike rate) of the cell response is not (Fig. S3). The properties acquired in this study would allow appropriate TSDN type identification on extracellular traces. This information is necessary to investigate simultaneous responses from several TSDNs within one animal.

Discussion
We have shown that TSDN responses are suited to act as matched filters for the retinal slip of the prey during the predatory behavior. TSDNs are most sensitive to targets moving in the frontal-dorsal region of the eye, where dragonflies fixate the prey image during interception flights (1). It would be informative to study if the other insects known to visually track and fixate on targets, such as mantis (25) and hover flies (26), also have a group of neurons most sensitive to target motion away from the retinal fovea. Moreover, although a descending looming neuron has been recorded extracellularly in a mantis species (27), TSDNs in this taxa are yet to be reported.

Our results show that the TSDN properties make these cells excellent information channels for a prey-direction population vector that steers wing movements during predatory flights. These properties include the following: (i) overlapping receptive fields, (ii) wide and overlapping direction tuning curves, (iii) latencies of cell activation and contraction, and (iv) shared dorsal branching organization and location at the wing motor centers.

We have confirmed homology between the TSDNs in representatives of two dragonfly families with markedly different foraging behavior. The subject of this study was L. luctuosa (family Libellulidae, most of whose members forage from a perch). TSDNs were originally characterized in dragonflies of family Aeshnidae, which detect and pursue prey during continuous flight. We showed that, in L. luctuosa, three of the TSDNs travel in a different tract of the thoracic ganglion. We have also found that MDT3 travels through the DIT in another libellulid species (Libellula hydra; Fig. S8). Given that such tracts are laid down early in development and are highly conserved among insect groups (28), it is possible that this variation in pathway reflects a change in the neural “wiring” of the wing control circuitry. For example, a homologous neuron involved in swimming pattern generation has been found to carry out different functions in two closely related sea slugs (nudibranchs) that exhibit similar swimming modalities (29). To address such evolutionary questions about the dichotomy between the TSDNs in hawker and percher dragonflies, the mesothoracic and metathoracic branching patterns of the Aeshnid TSDNs also need to be investigated.

It is interesting that all TSDNs target the mesothoracic and metathoracic ganglia with four identifiable locations in the most lateral region of each ganglion. These results raise two questions. First, as the dragonfly fore and hind wings are controlled...
independently (30), what additional information alters the TSDN motor inputs to yield a specific movement for the rear and the fore wings? Further studies on the central pattern generators underlying the dragonfly flight motor are necessary to answer this question. Second, because the projection from leg sensory neurons into the thoracic ganglia of locusts follows a 3D somatotopic map (31), do the four locations of TSDN branching reveal a pattern of signal integration with other localized sensory inputs? Each dragonfly wing has four sensory structures called crevice organs. Crevice organs resemble the campaniform sensilla of insects and are believed to act as mechanoreceptors for the degree of wing twist (32). In the locust, campaniform sensilla measure lift and twisting amplitude of the hind wings (33). Because TSDN activation alters the attack angle of the wing (20), it seems likely that information from the TSDNs and input from the crevice organs is combined in some manner, but a follow-up investigation on the information flow from the TSDNs to the wing motor neurons is needed.

Previous reports have shown that the underlying principles used by dragonflies to calculate motion (17) and orientation (16) are similar to those found in vertebrate systems. Here, we have shown that population vector code could successfully direct target interception in these ancient animals. The population vector algorithm has been coined the “geometrical universal actuator” because sensory information coded in vector form can be used directly for appropriate motor actuation, without a need for a time-consuming transformation (12). Our data further support this view. Even though the visual information flows through the brain, the TSDN population vector carries the information about prey direction, and the appropriate compensatory forces are implemented in the wing motor centers. Thus, the population vector could allow this steering system to act as an “autopilot” (see ref. 34 for a discussion on the feasibility of automated steering in insects), providing the fast reaction speed displayed by dragonflies during predatory behavior (~30 ms from stimulus onset to compensatory wing strokes (1)).

The dragonfly system shows a significant level of robustness; even when only six TSDN types are used for the calculation, the population vector is still not significantly different from the target direction. Unlike the population vector coding in the leech (9), TSDNs do not sample the visual space and target directions homogeneously. Bias in direction sampling has also been reported in the cells of the monkey motor cortex, which code population vectors for arm movements (35). Moreover, in insects, the visual performance of motion detection is matched to the species-specific flight movements (36), a topic recently reviewed by Taylor and Krapp (37). Regardless of such bias, for the 2D plane tested in this study, TSDN population vector coding performs equally well across all directions. It is possible that the bias in the preferred direction of the different TSDNs becomes significant once the animal is presented with a more realistic target showing freedom of movement in all three dimensions (i.e., including contraction or expansion of the prey image).

Although we have focused on prey pursuit, dragonflies also pursue larger targets, such as conspecifics in male/male territorial chases. In such nonpredatory flight trajectories, collision may not be desirable, and an interception strategy may not be beneficial. It remains to be elucidated whether the TSDNs are used in an exclusively predatory context, or whether they play a part in flights with different strategies.

In summary, the fast and highly precise prey interception flight system of dragonflies is particularly amenable to investigation at many levels: electrophysiological, morphological, and behavioral. The ultimate goal is to elucidate the neural underpinnings from molecular basis to behavior, something already achieved for the swimming movement of the lamprey (38). The electrophysiological and morphological evidence presented here support the notion that TSDNs provide population vector information about target movement that can be used to direct wing movement.
during a predatory flight. This system relies on a low number of cells performing with high accuracy. The functional significance of this result is that, even in an insect that displays a behavior as complex as interception, the integration of signals and calculation of motor output seem to occur at the thoracic ganglia, and not in the brain. A process as seemingly complicated as interception appears automated to yield the high speed and performance necessary for insect predation. This study is of particular importance because dragonflies are the only insect group that exclusively sport direct wing muscles, allowing independent control of both sets of wings. With the view to developing bio-inspired robots, recent studies have investigated dragonfly vision (39) and wing dynamics (40). Thus, understanding these unique neurological control of the dragonfly wings may prove invaluable for future technological applications. The 3D atlas presented here forms a basis for further understanding of this circuit.

Materials and Methods

The ventral nerve cord of adult L. fuctuosa was impaled with a sharp electrode while the animal viewed a visual display (360 Hz). TSDNs were recognized initially by robust responses to a vertical or horizontal small moving target of fixed size, increased response for certain target directions, and lack of responses to a wide-field stimulus. A stimulus composed of 3,497 consecutive small moving targets was presented, and the resultant cell activity was used to calculate its latency, directional tuning (see refs. 41 and 42) and receptive field. Looming sensitivity was tested with a linearly expanding stimulus. Using negative current (~2 nA), we injected 32 of the TSDNs with Lucifer yellow. The preparations were processed and imaged with a confocal microscope (Zeiss 710; see ref. 43). To visualize the cell morphology, we used Vaa3D (www.vaa3d.org) (44, 45) and BrainAligner (46). The population vector was calculated with a bootstrap method, with turning curves calculated by binning target directions every 18° and averaging cell activity for those bins. The population vector correlation was based on weight 12 in appendix 1 of ref. 47. See Fig. 59 and SI Materials and Methods for more information.

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The right connective.

the midsagittal plane, so all responses are shown for the TSDNs in properties: robust responses to small moving targets of neurons (TSDNs) were recognized initially by the following a black-and-white grating in any orientation and direction). The increased response for certain target directions (i.e., directional selectivity), and lack of responses to a wide-

trace was calculated. The time from movement onset to the weighted

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cord was calculated. The time from movement onset to the
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cord [for a diagram, see Adelman et al. (1)]. The ventral nerve cord was impaled between the prothoracic and mesothoracic legs, and a metal support was placed underneath the ventral nerve cord [for a diagram, see Adelman et al. (1)]. The ventral nerve cord was impaled between the prothoracic and mesothoracic ganglia with borosilicate electrodes (Sutter; 10 cm; o.d., 1; i.d., 0.5; #BF-100-50-10; pulled with a Sutter P2000) filled with 1.5% Lucifer Yellow (In-vitrogen; #L-453) in 100 mM LiCl and backfilled with 1 M LiCl (100-180 MΩ). The intracellular signal was amplified (Axoclamp; Axon Instruments), digitized (40 kHz), displayed, and recorded (Spike GL; A. Leonardo, Janelia Farm Research Campus, Ashburn, VA) for off-line analysis, which was carried out in MATLAB.

Stimulus. The animal was supported ventral-side-up by a clamp and the head tilted so that the dorsal area of the eye looked at the backprojected screen (see Fig. S6 for more details). Custom-written software (A. Leonardo) controlled the display of an image. The stimuli output were rendered by a graphics card with two display ports. Each display port was connected to a custom-

ized DLP projector (Light Speed; model: DepthQ) running at 360 Hz. The projectors were aligned without a gap between the images and the stimuli backprojected on a single piece of paper to create a continuous screen. This arrangement allowed stimuli to run seamlessly between the two displays. The full screen size (in pixels) was 1,600 wide by 600 high, placed 12 cm away from the head of the animal. The image projection size was 12.8 cm (height) × 33 cm (width). Therefore, the subtended screen size was 108° wide by 56° high. The target diameter subtended 1.48° (0.45 cm, 22 pixels), and the distance between target locations was 108° wide by 56° high. The target diameter subtended 1.48° (0.45 cm, 22 pixels), and the distance between target locations within a trajectory was 0.43° (0.13212 mm, 6.4 pixels). Hence, the targets between two consecutive locations overlapped by two-thirds of the target size. This assured that the target was seen as an object in continuous motion. A second signal from a photodiode monitoring the screen frame rate and timing was also recorded, allowing synchronization of the recording with the stimulus.

Electrophysiological Data Analysis. Target-selective descending neurons (TSDNs) were recognized initially by the following properties: robust responses to small moving targets of fixed size, increased response for certain target directions (i.e., directional selectivity), and lack of responses to a wide-field stimulus (such as a black-and-white grating in any orientation and direction). The data from cells impaled in the left connective were flipped along the midsgittal plane, so all responses are shown for the TSDNs in the right connective.

Latency Calculation. We obtained the minimum latency between stimulus onset/offset and cell first/last responses by creating a peristimulus histogram of spike timing for each recording (Fig. S1). The data were smoothed with an adjacent average weighted filter (3 ms). The second derivative of the smoothed trace was calculated. The time from movement onset to the first local maximum of the second derivative (corresponding to the moving target response) was taken as the latency. To avoid mistaking noise for spikes, the second derivative and the smoothed trace were plotted together. Visual inspection confirmed that the local maxima of the second derivative corresponded to the start of the responses to a moving target, and not to background spiking. The same procedure was carried out for the last responses.

Directional Tuning Calculation. For every spike recorded, the direction of the moving target that elicited it was noted. The data were grouped in 10° bins and plotted in polar form (Fig. 2A). Note that the spikes due to the “flash” response caused by the appearing target were not taken into account. The preferred cell direction (Fig. 2A, red dot) was obtained by using the CircStat MATLAB toolbox developed by Berens (2), which fits a Gaussian to the direction preference distribution of each cell. To obtain the tuning width of each cell accurately, the MATLAB toolbox published by Cronin et al. (3) was used. This toolbox uses a Bayesian method to estimate the tuning parameters.

Receptive Field Calculation. The onset latency obtained for each cell was used to calculate the target locations that elicited each spike. The average direction of the targets was calculated for each pixel [Fig. 2, direction receptive field (DRF) plots]. In addition, the spike-triggered average (STA) was calculated to yield the receptive fields shown in Fig. 2 (STA plots). Note that, in DRF plots, a pixel with a single spike is always marked with a bright color, because color denotes direction. However, in the STA maps, only the pixels with proportionally larger number of spikes look bright, because brightness codes number of spikes.

Looming Sensitivity. Libellula luctuosa TSDNs were tested with a linearly looming stimuli. The linear looming stimulus presented consisted of a target that grew in diameter from 1.35 to 31.86° in 158 ms and was presented in three different locations of the screen (left, center, and right) consecutively (Fig. S4B). Thus, the rate of diameter expansion was 19° per 100 ms. For comparative purposes, the reference system for the three locations of the expanding stimulus has been changed to ipsilateral, center, and contralateral. For example, an expanding stimulus was ipsilateral if it was presented on the right side of the screen and the cell was impaled in the right side of the connective. We counted the number of spikes elicited by each stimulus expansion. The mean spike number and the SE were calculated for each type of TSDN. In addition, Libellula pulchella TSDNs were tested with an angular expansion (1 m/s) of a 1-cm target.

Recording Temperature. Caution should be used when interpreting the electrophysiological results from this study. The animal and the room temperature were set at 23–25 °C during the recordings. Although this is above 19 °C, the minimum flying temperature in a Libellulid species, it is drastically lower than the Libellulid body temperature measured in specimens that were flying at the time of capture (28–36 °C) (4).

Recording Duration and STA Maps. Of the total number of intracellular recordings, 17 did not complete the full sequence of trajectories (only four records had less than 2,500 trajectories and none of those were below 1,500 trajectories). Although a lower number of presented trajectories also lowered the total number of recorded spikes, it did not affect the direction or receptive field results (Fig. S5). The spikes from each recording were used to create the STA (Fig. S5). All of the STAs from the same type of
TSDNs were normalized, averaged, and normalized again, shown as STA maps in Fig. 2.

**Population Vector Computation.** In some trials, the target position presented a “wrapping” effect as they hit the edge of the screen. The 2,776 trajectories that did not exhibit target wrapping were used to compute the population vectors.

In the ensuing part, the raw electrophysiology data were reanalyzed. This time, target directions were binned every 18°, cell activity was averaged per bin, and preferred directions were recalculated from these binned data. To obtain the cell activity for each target presented, we used the following bootstrap method. In a single bootstrap, a new sample of \( n = 2,776 \) target motion directions was formed for each cell by resampling with replacement the existing 2,776 trials, and so on for 100 bootstraps.

To calculate a population vector from the activity of a neural ensemble, it is first necessary to obtain a tuning curve function that describes the directional properties of each cell. A tuning curve function with a good fit yields the cell’s activity \( d(\theta) \) in response to a particular stimulus \( \theta \). For example, Eq. S1 is a suitable tuning function when the frequency discharge of a cell is sinusoidal and depends on stimulus direction (5) as follows:

\[
d(\theta) = b_0 + b_1 \sin \theta + b_2 \cos \theta + e,
\]

where \( d(\theta) \) is the rate of cell activity, \( b_0, b_1, b_2 \) are regression coefficients, and \( e \) is an error term.

For TSDNs, we have found that, in addition to the main bell-shaped cell activity curve, the cells also display a second smaller peak. This is taken into account by Eq. S2 as follows:

\[
d(\theta) = b_0 + b_1 \sin \theta + b_2 \cos \theta + b_3 \sin 2\theta + b_4 \cos 2\theta + e.
\]

We confirmed the good fit of this function by calculating the multiple correlation coefficients \( R \). This function yielded highly statistically significant fits \((P < 0.01; P < 0.001)\) for 47 of 50 cells and high \( R^2 \) (range: 0.65–0.978; mean, 0.918; median, 0.938). Such a good fit allows us to estimate correctly the preferred direction of the cell \((\theta_0)\), given by Eq. S3 as follows:

\[
\theta_0 = \tan^{-1} \frac{b_1}{b_2}.
\]

Before calculating the population vector, the activity of each \( i \)th cell for stimulus \( \theta, C_i(\theta) \), was weighted using Eq. S4, shown as weight 12 in appendix 1 of ref. 6, as follows:

\[
w_{i\theta} = \frac{d(\theta) - b_0}{b_0}.
\]

Finally, the population vector for each target presented, \( P(\theta) \), was calculated by summing weighted vectorial contributions from each of \( N \) cells in the population (6) as follows:

\[
P(\theta) = \sum_{i=1}^{N} w_{i\theta} \cdot \theta_0.
\]

Within the mesothoracic and metathoracic ganglia, the vast majority of motor neurons are ipsilateral. Therefore, the information is not likely to cross over from one side of the ganglia to the other side. For this reason and in absence of additional information, it is parsimonious to assume that the movement of each wing can only be formed by a population vector arising from the inputs into that specific side of the ganglia. Due to the receptive field location and to the anatomy of the TSDNs, a target moving in the left side of the visual field generates a different set of inputs for each side of the ganglia. This is to say that the population vector of the right and left wings arise from different sets of TSDNs inputs. This is the reason behind dividing TSDNs into groups that form right and left population vectors.

Correlation coefficients for circular data were calculated as shown by ref. 7.

For a graphical explanation of population vector coding, see Fig. S9.

**Dye Injection, Sample Processing, and Confocal Imaging.** Following electrophysiological characterization, the injection period was typically between 30 min and 2 h. To visualize injection of the dye, a Kramer quad illumination system with an X-cite fluorescence light source was used. Immunohistochemistry and streptavidin binding were used to tag the injected Lucifer yellow with DyLight 633, shifting the signal to the red spectrum. Each sample was imaged with three different channels; the background fluorescence, the tra chea fluorescence, and the injected dye signal (Fig. S7). The first two were used to place landmarks for warping.

**Image Reconstruction, Warping, and Tracing.** To visualize the cell morphology, Vaa3D software was used (www.vaa3d.org) (8). The Vaa3D-Stitch plug in was used to stitch the images (9). To register the images, one sample was used as a model, and all subsequent samples were registered based on 48 markers manually determined using Vaa3D. The BrainAligner tool (10) was used to warp all samples to the model. To obtain the morphology of these cells, we developed a neuron-tracing method based on distance-transform of images.
Fig. S1. Visual stimulus. (A) Each presented trajectory was composed of a target (1.48°, solid black circle on white background) that appeared in a random location and waited for 50 ms before moving for 100 ms and traveling 13.4° (4.1 cm) along a straight path in a random direction with a constant speed of 134°/s (41 cm/s). The target then disappeared for 150 ms before appearing again. The same random sequence was presented to all impaled cells. (B) We created a peristimulus histogram of spike timing for each recording (only one shown). This histogram provided the minimum latency between stimulus onset/offset and cell first/last responses.

Fig. S2. The traces of TSDNs* from brains of the hawker dragonfly Anax junius (Aeshnid) and the percher dragonfly L. luctuosa (Libellulid) show homology. (Only TSDNs to the right of the midsagittal plane are shown. Trace for MDT4 is not available. The trace for DIT1 does not display a cell body due to an incomplete fill.) Traces were acquired from maximum-intensity projections of the brains. Aeshnid data were reproduced from Frye and Olberg (1).

Fig. S3. Number of spikes recorded for each TSDN type differs among animals. This variation in number of spikes is not likely to be due to recording quality because we have observed recordings with large spikes, stable over prolonged periods but few in number. Although we have found that poor animal health is correlated with a low success in recording TSDNs for prolonged periods, we have also observed strong healthy animals responding with low number of spikes. Thus, on its own, animal health cannot explain spike number variance. It is most likely that the variance in spike numbers arise from differences in the state of the animal. In addition, because these animals are immobilized for the experiment, the lack of appropriate movement and feedback signals may impact some specimens more strongly.

Fig. S4. TSDNs tuning width and looming sensitivity. (A) The tuning width of the TSDNs tuning curves, i.e., one-half the maximum width at one-half the maximum height (\(I\)), were calculated using the MATLAB toolbox published by Cronin et al. (1). This toolbox uses a Bayesian method to fit the data and estimate the tuning parameters. (ii) Analysis output example showing data points (grouped into 10° bins) from one MDT1 cell (black dots), mean fit (red), and the posterior samples (gray). (iii) The results show that the tuning widths varied significantly between TSDN types. (B) A linearly expanding stimulus consisting of a target which grew in diameter from 1.35 to 31.86° in 158 ms was presented in three different locations of the screen. This expanding stimulus only elicited a significant number of spikes at the contralateral (\(P = 0.006, n = 5\)) and midline (\(P = 0.004, n = 7\)) locations of DIT3 (asterisk). One-sample \(t\) test was used to test statistical significance.

For two TSDN types, DIT1 and MDT1, the characteristics of five of the contributing cells are shown. Number of spikes are plotted according to the direction of the target that elicited them (polar plot) and to the location where the target was, after taking into account the appropriate latency (spike-triggered average map). The receptive fields and direction preferences of the cells are constant across specimens, but their spike numbers are not. The scale for each STA notes the number of spikes at each pixel.

The dragonfly was immobilized, with the eye pointing at the backprojected screen with the angles shown in the graph.
Fig. S7. In Libellulids, three TSDN types travel through a different track in the thoracic ganglia. (A) After injecting Lucifer yellow into the TSDN just above the mesothoracic ganglion, the central nervous system was dissected out and the ganglia imaged (prothoracic ganglion shown inside box). (B–D) (Top) Dorsal maximum-intensity projection showing TSDNs branching in the prothoracic ganglion. The eight middle and bottom panels show optical cross-sections through the prothoracic ganglion, each containing the fluorescence profile of a different TSDN type. Although we found homologous TSDNs between *L. luctuosa* and *A. junius*, we also found consistent differences between them. In the prothoracic ganglion of *L. luctuosa*, the axons of MDT3, MDT2, and MDT4 are found in the DIT track (B). Thus, MDT1 and MDT5 are the only two TSDNs found in the MDT track (C). All three DIT cells travel in the DIT track (D). The white arrows point at TSDN axons in cross-section. Due to the certainty of homology between Libellulid and Aeshnid TSDNs, and to avoid potential confusion in future comparisons, we followed the original naming system (1).

Fig. S8. MDT3 in the Libellulid *L. lydia* also travels through an alternative track. (A) Tracing of the brain fill showing the T shape characteristic of the MDT3 profile. (B) MIP projection of the neuron in the mesothoracic and metathoracic ganglia, showing the morphology typical of MDT3. (C) Cross-section of the prothoracic ganglion showing the axon traveling in the DIT track.
Fig. S9. Several cells are needed for a population vector to code equally well in all directions. The polar plots (Left) and linear plots (Right) both show the same information. (A) A single cell with a 45° tuning width does not provide full coverage for the detection of a target’s direction (shown as areas of no information). In addition, only when the target travels in the preferred cell direction (100% cell activity, which equals to the maximum number of spikes) is the direction of the target coded without ambiguity. For every other direction covered by the tuning curve, the cell activity would code for two possible, opposing, directions. (B) Adding a second cell that displays a preferred direction shifted by 90°, decreases the number of directions where a target movement is not coded. In addition, ambiguity in the coding is now reduced. For example, both cells would need to spike at 50% of their maximum activity to code for a target moving with a 225° direction. If only cell 1 spikes at 50% and cell 2 remains silent, the target would be moving at 135°. The opposite ratio (cell 1 = 0%; cell 2 = 100%) would code for a target direction at 315°. Adding more cells would increase the ability of the system to code target direction more precisely.