Facilitation of neural responses to targets moving against optic flow

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For the human observer, it can be difficult to follow the motion of small objects, especially when they move against background clutter. In contrast, insects efficiently do this, as evidenced by their ability to capture prey, pursue conspecifics, or defend territories, even in highly textured surroundings. We here recorded from target selective descending neurons (TSDNs), which likely subserve these impressive behaviors. To simulate the type of optic flow that would be generated by the pursuer’s own movements through the world, we used the motion of a perspective corrected sparse dot field. We show that hoverfly TSDN responses to target motion are suppressed when such optic flow moves syn-directional to the target. Indeed, neurone responses are strongly suppressed when targets move over either translational sideslip or rotational yaw. More strikingly, we show that TSDNs are facilitated by optic flow moving counterdirectional to the target, if the target moves horizontally. Furthermore, we show that a small, frontal spatial window of optic flow is enough to fully facilitate or suppress TSDN responses to target motion. We argue that such TSDN response facilitation could be beneficial in modulating corrective turns during target pursuit.

For the human observer, it can be difficult to follow the motion of small objects, especially when they move against background clutter. In contrast, insects efficiently do this, as evidenced by their ability to capture prey, pursue conspecifics, or defend territories, even in highly textured surroundings. We here recorded from target selective descending neurons (TSDNs), which likely subserve these impressive behaviors. To simulate the type of optic flow that would be generated by the pursuer’s own movements through the world, we used the motion of a perspective corrected sparse dot field. We show that hoverfly TSDN responses to target motion are suppressed when such optic flow moves syn-directional to the target. Indeed, neurone responses are strongly suppressed when targets move over either translational sideslip or rotational yaw. More strikingly, we show that TSDNs are facilitated by optic flow moving counterdirectional to the target, if the target moves horizontally. Furthermore, we show that a small, frontal spatial window of optic flow is enough to fully facilitate or suppress TSDN responses to target motion. We argue that such TSDN response facilitation could be beneficial in modulating corrective turns during target pursuit.

The ability of insects to successfully pursue targets in clutter (4, 19) is thus remarkable and suggests a high level of optimization, making the underlying neural mechanisms interesting to study. Indeed, insects that pursue targets, including predatory dragonflies and robberflies, as well as territorial hoverflies, have higher-order neurons in the optic lobes (20, 21) and the descending nerve cord (22, 23) that are sharply tuned to the motion of small, dark targets. Target-tuned neurons often have receptive fields (24–26) in the part of the compound eye that has the best optics (27, 28). Target selective descending neurons (TSDNs) project to the thoracic ganglia (25, 29) where wing and head movements are controlled (30), and electrically stimulating TSDNs leads to wing movements (31). Taken together, this suggests that TSDNs subserve target pursuit. However, how TSDNs respond to targets moving against translational and rotational optic flow is unknown.

Significance

Target detection in visual clutter is a difficult computational task that insects, with their poor spatial resolution compound eyes and small brains, do successfully and with extremely short behavioral delays. We here show that the responses of target selective descending neurons are attenuated by background motion in the same direction as target motion but facilitated by background motion in the opposite direction. This finding is important for understanding how target pursuit can occur in tandem with gaze stabilization. Indeed, the neural facilitation would come into effect if the hoverfly is subjected to background motion in one direction but the target is pursuing moves in the opposite direction and could therefore be used to override gaze stabilizing corrective turns.

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The authors declare no competing interest.

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We quantified the responses of *Eristalis* TSDNs to targets moving against six types of optic flow, three translations and three rotations. The optic flow stimulus was composed of thousands of dots. Thus, the target dot could not be discriminated by shape or contrast but only by its relative motion. We found that optic flow moving syn-directional to the target suppressed the TSDN response, regardless of whether the optic flow moved in a rotational or translational manner. We found that orthogonal optic flow attenuated the TSDN target response but to a lesser degree than syn-directional optic flow. This suggests that the vector divergence between the target and the optic flow is important. Most strikingly, we found that counterdirectional optic flow increased the TSDN response to target motion, if the target moved horizontally. We found that projecting optic flow to only a small frontal portion of the eye was sufficient to elicit both TSDN attenuation and facilitation. As descending neurons control behavioral output (30, 31), the response attenuation and facilitation could play a role in modulating optomotor, or gaze stabilizing corrective turns, as needed during target pursuit.

**Results**

**TSDNs Respond to Target Motion but Not to Optic Flow.** We simulated optic flow using the motion of a sparse dot field (24). By projecting the individual features in the sparse dot field onto a screen anterior of the hoverfly’s spatial location, and simulated depth, over time, provided the type of optic flow that would be generated by self-motion (24). In optic flow–sensitive descending neurons, this stimulus elicits strong direction-selective responses, similar to the responses to widefield sinusoidal gratings (24) or large moving images (22) with naturalistic statistics (32). We recorded from TSDNs (Fig. 1) in male *E. tenax* hoverflies and first confirmed that our optic flow stimulus did not generate a TSDN response (*SI Appendix, Fig. S1*). For example, translational sideslip to the left only generated a response in 18% of 222 repetitions across 12 TSDNs (SI Appendix, Fig. S1A). Furthermore, when leftward sideslip did generate a TSDN response, this was much less than the response to a target traversing a white background (*SI Appendix, Fig. S1B*). This lack of TSDN response confirms that the optic flow stimulus is qualitatively similar to other types of widefield background motion.

In contrast to the lack of response to the optic flow, TSDNs responded strongly to the motion of a small, dark target (22, 24) traversing a white background (compare spiking response during stimulation with the lack of activity before stimulation, Fig. 1B, *Left* and *Movies S1 and S2*), as well as when the target traversed the stationary sparse dot field (Fig. 1B, *Right* and *Movies S3 and S4*). Note that since the sparse dot field consisted of hundreds of “targets,” there were no spatial characteristics identifying the individual target when both were stationary (Fig. 1A, *Right*). For quantification across neurons, we calculated the mean spike frequency for the duration of target motion, except for the first and last 40 ms of each 500 ms target trajectory (dotted box, Fig. 1B and C). We found a small but significant response reduction to targets moving across a stationary dot field compared with a white background (Fig. 1D), as expected from the reduced motion energy when the target traversed pattern elements of the stationary dot field. As the response across neurons was variable (*N = 39, coefficient of variation 62 and 65%, respectively, Fig. 1D*), we normalized the response from each neuron to its own mean response to a target moving over a white background.

**Optic Flow Modulates the TSDN Target Response.** During translational optic flow, distant features move slower than closer ones, whereas during rotations, all features move at the same angular velocity (15). We asked whether optic flow rotations and translations would therefore have different effects on the TSDN response to target motion. To investigate this, we simulated different types of translations (Fig. 2A) at 50 cm/s and rotations (Fig. 2B) at 50 deg/s. We found that when the target moved horizontally across syn-directional sideslip, the TSDN response was strongly suppressed (mean inhibition 93%, Fig. 2C and D; “Left Sideslip” and *Movies S5 and S6*), compared with control where the target moved over a stationary dot field (gray, Fig. 2C and D and *Movies S3 and S4*). Similar effects of syn-directional background motion were previously seen when using panoramic images with naturalistic statistics (22). We also found that syn-directional yaw optic flow strongly attenuated the TSDN response to target motion (by

![Fig. 1. TSDN response to target motion.](https://doi.org/10.1073/pnas.2024966118)
We next asked whether counterdirectional optic flow would also attenuate the TSDN response to target motion but less than synchronodal optic flow. However, we found that when the target moved over counterdirectional sideslip the TSDN response was strongly enhanced (mean increase 71%, Fig. 2B) compared with the stationary control (gray, Fig. 3C). Similarly, yaw moving counterdirectional to the target also facilitated the TSDN response, by 85% (Fig. 2C and D, “Right Yaw”). Such response facilitation was not seen when displaying targets over counterdirectional background motion using panoramic images, in either TSDNs (22) or other target-tuned neurons in the fly optic lobes (21, 26, 33). This rejects the hypothesis that counterdirectional optic flow attenuates TSDNs.

**Frontal Optic Flow Is Required and Sufficient.** Our data show that yaw and sideslip have similar effects on TSDN responses to target motion (Fig. 2C and D). As both sideslip and yaw contain substantial local motion in the frontal visual field (34), we next asked if frontal optic flow is required. We investigated this by limiting the spatial extent of the sideslip to either cover the ipsilateral, dorsal, ventral, or contralateral position on the screen (Fig. 3A). Note that only the dorsal position covers the TSDN receptive field (Fig. 3A). In the other three positions, the sideslip optic flow was spatially separated from the receptive field (Fig. 3A) and thus the target trajectory.

We found that when sideslip moved syn-directional to the target, the TSDN response was attenuated if the optic flow covered the full, dorsal, or ventral position on the screen (Fig. 3B), compared with the stationary control (gray, Fig. 3B). However, when the optic flow only covered the ipsilateral or contralateral positions, there was no difference compared with control (Fig. 3B), suggesting that frontal optic flow is both required and sufficient. Previous work using moving images with naturalistic statistics showed that these did not have to spatially overlap with the target trajectory, nor have a large spatial extent, to attenuate TSDN responses (22).

We next found that when sideslip moved counterdirectional to the target, the TSDN response to target motion was facilitated if the optic flow covered the full, dorsal, or ventral positions of the screen (Fig. 3C). When the sideslip was limited to the ipsilateral or contralateral positions, there was no TSDN response facilitation (Fig. 3C). This suggests that the optic flow does not have to spatially overlap with the target trajectory. However, there has to be frontal, counterdirectional optic flow for facilitation to take place.

In summary, our results show that a small spatial window of optic flow in either the dorsal or ventral visual field is enough to strongly attenuate (Fig. 3B) or facilitate (Fig. 3C) the TSDN response to target motion.

**Vector Divergence between Target and Optic Flow Affects TSDNs Response.** Our data above (Figs. 2 and 3) show that syn-directional optic flow strongly attenuates TSDN responses to target motion, whereas counterdirectional optic flow facilitates the response. This suggests that the level of vector divergence between the target and the optic flow influences the TSDN responses, so that maximum
attenuation is generated at minimum vector divergence, whereas maximum facilitation is generated at maximum divergence. To explore this, we quantified the effect other types of optic flow had on TSDN responses to target motion.

We found that the TSDN target response was suppressed by lift, which moves orthogonal to the target (mean 44% suppression for downward lift, “Lift Down;” 34% for upwards lift, “Lift Up,” Fig. 4). When the target was displayed against pitch which also provides orthogonal motion in the frontal visual field, the response was also suppressed (mean 57% suppression, “Pitch Down;” 55% for “Pitch Up,” Fig. 4). The TSDN response to target motion was also attenuated when the target was displayed against thrust, which provides orthogonal motion along the animal’s anterior–posterior axis (mean 44% suppression for “Thrust Towards;” 58% for “Thrust Away,” Fig. 4). Thus, orthogonal optic flow attenuates the TSDN response (Fig. 4) but not as much as syn-directional optic flow (Fig. 2).

We next looked at roll optic flow, which creates no local motion straight ahead of the fly but opposite direction local motion in the dorsal and ventral visual fields. For example, during counterclockwise roll (“Roll CCW,” Fig. 4) the target and the optic flow move in the same direction through the dorsal receptive field but in opposite directions in the ventral visual field. Therefore, the TSDN would receive a combination of maximum and minimum vector divergence signals from the “dorsal” and “ventral” parts (as shown in Fig. 3A). We found that the response attenuation was stronger against counterclockwise roll (mean suppression 75% for “Roll CCW,” Fig. 4) than against clockwise roll (32%, “Roll CW,” Fig. 4).

Thus, our data (Figs. 2 and 4) support the notion that the level of vector divergence influences the TSDN response. To explore this in further detail, we recorded from TSDNs that respond robustly both to horizontal (gray, Fig. 5A and B) and vertical target motion (black, Fig. 5A and B). We found that when the target moved horizontally, the TSDN response was strongly attenuated against syn-directional sideslip (second column, Fig. 5C), strongly facilitated when displayed against counterdirectional sideslip (third column, Fig. 5C), and less attenuated against orthogonal lift in either direction (last two columns, Fig. 5C), consistent with previous results (Figs. 2 and 4). In the same eight TSDN neurons, we next moved the target vertically. We found that the TSDN response was completely attenuated against syn-directional lift (last column, Fig. 5D) and less attenuated against orthogonal sideslip in either direction (second and third column, Fig. 5D). This supports the suggestion that maximum TSDN response attenuation is generated at minimum vector divergence between the target and optic flow. However, when the vertical target was displayed against counterdirectional lift (fourth column, Fig. 5D), there was no response facilitation. This suggests that maximum vector divergence on its own is not enough to explain the TSDN response facilitation to target motion.

**Optic Flow at Different Dot Densities Modulates TSDN Responses.** The experiments above (Figs. 1–5) used optic flow with dot densities of 100 per m$^2$. Is it possible that the optic flow’s effect on TSDN responses depends on the dot density? To investigate this, we used six different dot densities, ranging from 10/m$^2$ to 500/m$^2$ (SI Appendix, Fig. S2A and B). All dot densities had naturalistic Fourier spectra (SI Appendix, Fig. S2C), and the contrasts ranged from below naturalistic (35) for the sparsest, to above naturalistic for the densest (SI Appendix, Fig. S2D). We verified that optic flow with these dot densities stimulated optic flow–sensitive descending neurons (SI Appendix, Fig. S2E).

We first quantified the TSDN response to a target moving over stationary optic flow and found that this remained robust across dot densities (gray data, Fig. 6). We next quantified the TSDN response to targets moving over syn-directional sideslip and found that this was significantly inhibited compared with stationary
control at all dot densities (two-way ANOVA, $P < 0.0001$, Fig. 6). In contrast, when the target moved over counterdirectional side-slip, there was no facilitation at the lowest dot density (Fig. 6A). However, the facilitation was significant for dot densities of 50/m$^3$ to 500/m$^3$ (two-way ANOVA, $P < 0.0001$, Fig. 6B–F). Our data thus suggest that the effect optic flow has on TSDN target responses does not depend on dot density, at least not for the densities tested here (two-way ANOVA, $P = 0.2399$, Fig. 6).

Fig. 4. The TSDN response to target motion is attenuated by orthogonal optic flow. (A) The response from one TSDN to a target traversing a stationary sparse dot field (Left, gray, replotted from Fig. 2C) or different types of optic flow (red pictograms). All histograms shown as mean ± SEM ($n = 18$) at 1-ms resolution, after smoothing with a 20-ms square-wave filter. (B) The mean response across TSDNs ($N = 12$, same neurons as Fig. 2D) to a target traversing a stationary sparse dot field (Left, gray, replotted from Fig. 2D) or different types of optic flow. Stars indicate significance, one-way ANOVA followed by Dunnett’s multiple comparisons test (**$P < 0.01$, ***$P < 0.001$, and ****$P < 0.0001$), done together with the data shown in Fig. 2D. Translations were simulated at 50 cm/s and rotations at 50°/s. The lines show the mean, and the filled data points correspond to the neuron in A.

Fig. 5. Vector divergence between optic flow and target. (A) The TSDN response to a target traversing a white background horizontally (Left, gray) or vertically (Right, black). Histograms shown as mean ± SEM ($n = 18$) at 1-ms resolution, after smoothing with a 20-ms square-wave filter. (B) The mean spiking response of eight TSDNs to targets moving horizontally (gray, Left) or vertically (black, Right) over a white background. The filled data points correspond to the neuron in A. (C) TSDN responses to targets moving horizontally across sideslip or lift, syn-directional, counterdirectional, or orthogonal to the target, as indicated by the pictograms. Significance shown using one-way ANOVA followed by Dunnett’s multiple comparisons test, with *$P = 0.0232$, **$P = 0.006$, ***$P = 0.0003$, and ****$P < 0.0001$. (D) TSDN responses to targets moving vertically across sideslip or lift, orthogonal, counterdirectional, or syn-directional to the target, as indicated by the pictograms. Significance shown using one-way ANOVA followed by Dunnett’s multiple comparisons test, with *$P = 0.0198$, **$P = 0.0021$, ****$P < 0.0001$, and NS for $P > 0.05$. Same $N = 8$ in B–D.
Our data suggest that there are differences between the TSDN response suppression and facilitation (Figs. 5 and 6). To determine whether there are other differences, we investigated if the inhibition or facilitation was affected by preceding optic flow (green, SI Appendix, Fig. S3A). During target motion (blue, SI Appendix, Fig. S3A), the concurrent optic flow (red, SI Appendix, Fig. S3A) was either stationary or moving. We found that preceding optic flow affected the TSDN response if it was syn-directional (SI Appendix, Fig. S3B) but that counterdirectional preceding optic flow did not (SI Appendix, Fig. S3D and E). This supports the suggestion that there are different circuit mechanisms driving TSDN response suppression and facilitation.

Responses to ON-OFF Edges Consistent with 1-Point Correlation. TSDNs have been proposed to get their input from small target motion detectors (STMDs, see ref. 36). As opposed to our TSDN data, hoverfly STMD responses are rarely suppressed by syn-directional background motion and not facilitated by counter-directional background motion (21). As there are many other target-tuned neurons in the fly optic lobes (e.g., refs. 33 and 37–40), it is possible that TSDNs do not get their input from STMDs. We can investigate the potential input using the underlying target tuning mechanisms, which can be distilled down into three fundamentally different concepts. For example, visual neurons can become target tuned by receiving inhibitory feedback from the widefield system (40–42) or by using center-surround antagonism together with rapid adaptation (39, 43). Alternatively, they can use an elementary STMD model, which is tuned to the unique spatiotemporal profile of a moving target, with a dark-contrast change (OFF) from the leading edge followed by a bright-contrast change (ON) by the trailing edge. Importantly, while the first two mechanisms rely on comparisons from neighboring points in space, the elementary STMD compares input from one point in space (39, 44, 45). Therefore, the first two models will respond similarly to the motion of a target, to the motion of a leading OFF edge, and the motion of a trailing ON edge (black, Fig. 6, redrawn from refs. 39 and 46). In contrast, the elementary STMD model only responds strongly to the target (gray, Fig. 7, redrawn from ref. 46).

Our results show that TSDNs do not respond well to a leading OFF edge or to a trailing ON edge (white, Fig. 7). However, a complete target, where the leading edge is rapidly followed by a trailing edge, gives a robust TSDN response (white, Fig. 7). Indeed, the physiological responses (white, Fig. 7) match the elementary STMD model output (gray, Fig. 7). Since STMD physiology also matches the elementary STMD model output (46, 47), this suggests that TSDNs receive input from STMDs.

Discussion

We show that TSDNs do not get their input from small target motion detectors (STMDs, see ref. 36). As opposed to our TSDN data, hoverfly STMD responses are rarely suppressed by syn-directional background motion and not facilitated by counter-directional background motion (21). As there are many other target-tuned neurons in the fly optic lobes (e.g., refs. 33 and 37–40), it is possible that TSDNs do not get their input from STMDs. We can investigate the potential input using the underlying target tuning mechanisms, which can be distilled down into three fundamentally different concepts. For example, visual neurons can become target tuned by receiving inhibitory feedback from the widefield system (40–42) or by using center-surround antagonism together with rapid adaptation (39, 43). Alternatively, they can use an elementary STMD model, which is tuned to the unique spatiotemporal profile of a moving target, with a dark-contrast change (OFF) from the leading edge followed by a bright-contrast change (ON) by the trailing edge. Importantly, while the first two mechanisms rely on comparisons from neighboring points in space, the elementary STMD compares input from one point in space (39, 44, 45). Therefore, the first two models will respond similarly to the motion of a target, to the motion of a leading OFF edge, and the motion of a trailing ON edge (black, Fig. 6, redrawn from refs. 39 and 46). In contrast, the elementary STMD model only responds strongly to the target (gray, Fig. 7, redrawn from ref. 46).

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Discussion

We found that optic flow moving syn-directional to a target almost completely attenuated the TSDN target response (Fig. 2), even when the optic flow only covered a small part of the frontal visual field (Fig. 3). More strikingly, we found that optic flow counter-directional to target motion increased the TSDN response (Fig. 2), if the target moved horizontally (Fig. 5), across a range of dot densities (Fig. 6). We also found that orthogonal optic flow attenuated the TSDN target response (Figs. 4 and 5), but less than syn-directional motion, suggesting that the vector divergence between the target and the optic flow is important for response suppression. However, we found that vector divergence was not enough to explain the response facilitation (Fig. 5D).

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might have to be more sensitive to horizontal motion than to vertical motion. Since some STMDs might respond to features within the optic flow used here (47), and furthermore, since there are some STMDs that respond both to target motion and to features within background motion (21), these could potentially play a role in facilitation. Future investigation of STMD and TSDN responses are clearly required to explore the underlying mechanisms.

We found some notable differences between attenuation and facilitation. For example, as opposed to the response attenuation, there was not facilitation to the lowest dot density (Fig. 6A), and whereas attenuation was affected by preceding optic flow (SI Appendix, Fig. S3B), facilitation was not (SI Appendix, Figs. S3 D and E). In addition, only horizontally moving targets could be facilitated (Fig. 5D). Taken together, this suggests that the underlying circuit mechanisms differ. The response suppression could be driven by active inhibition from widefield motion detectors, as previously suggested (22), or inherited from upstream processes. We previously showed that TSDN responses to target motion are suppressed by background motion consisting of an image with naturalistic statistics and that background stimuli that drive optic flow–sensitive neurons suppress TSDN responses to target motion, whether spatially overlapping with the target trajectory or not (22). Indeed, we here confirmed that stimuli that drive optic flow–sensitive neurons (SI Appendix, Fig. S2E) suppress TSDN responses to target motion (Fig. 6), whether overlapping or not (Fig. 3F). Thus, TSDNs appear to be suppressed by widefield motion in general.

Nevertheless, our findings make behavioral sense (SI Appendix, Fig. S4). Prior to initiating target pursuit, male Eristalis hoverflies predict the flight course required to successfully intercept the target, based predominantly on the target’s angular velocity (49). To successfully execute an interception flight, the hoverfly turns in the direction that the target is moving (49). In doing so, the hoverfly creates self-generated optic flow counterdirectional to the target’s motion. In this case, the TSDNs would be facilitated (Fig. 2), which could be beneficial. Importantly, the facilitation would take place across a range of dot densities (Fig. 6 B–F), suggesting that even relatively sparse background textures (SI Appendix, Fig. S2B) would affect the TSDN response.

Once pursuit is initiated, if the pursuer drifts rightward due to, for example, a gust of wind, this induces lefthward optic flow across the frontal retina, which would evoke a lefthward optomotor response and/or gaze stabilizing turn (red, SI Appendix, Fig. S4 B, Left). If the target was also moving lefthward, in the direction of the corrective maneuver, this maneuver would also pursue the target, and no TSDN signal would be required (SI Appendix, Fig. S4 C, Left). Indeed, we found that the TSDNs were quiet under such conditions (“Left,” Fig. 2). However, if the optic flow instead moved rightward, counterdirectional to the target, the TSDNs would be strongly facilitated (“Right,” Fig. 2 and SI Appendix, Fig. S4 C, Right). Considering that TSDNs project to motor command centers in the thoracic ganglia (25, 31), such TSDN facilitation could then potentially override corrective turns. The TSDN response modulation by optic flow shown here could thus be beneficial for controlling behavioral output.

We only saw response facilitation if the target moved horizontally and not vertically (Fig. 5). This facilitation difference could be affected by the fact that the TSDNs that we recorded from responded better to horizontal motion than to vertical target motion (Fig. 5 A and B). In addition, it is currently unknown whether hoverfly TSDNs control head movements, wing movements, or maybe both. Indeed, recent work suggests that gaze stabilizing turns by the head, and the wing optomotor response, are controlled independently and by different visual components (13, 50, 51). Furthermore, the difference could be explained by different behavioral strategies used for vertical and horizontal target deviations (52). For example, whereas horizontal deviations...
Materials and Methods

Electrophysiology. *E. tenax* hoverflies were reared and maintained as previously described (53). For electrophysiology, a male hoverfly was immobilized ventral side up using a beeswax and resin mixture. A small hole was cut at the anterior end of the thorax to expose the cervical connective, which was then raised slightly and supported using a small wire hook, for insertion of a sharp polyimide-insulated tungsten microelectrode (2 MΩm, Microprobes). The animal was grounded via a silver wire inserted into the ventral part of the hole.

Extracellular signals were amplified at 100× gain and filtered through a 10- to 3,000-Hz bandwidth filter on a DAM50 differential amplifier (World Precision Instruments), with 50 Hz noise removed with a HumBug (Quest Scientific). The data were digitized via Powerlab 4/30 (ADInstruments) and acquired at 40 kHz with the Labchart 7 Pro software (ADInstruments). Single units were discriminated by amplitude and half-width using Spike Histogram software (ADInstruments).

Visual Stimuli. *Eristalis* males were placed ventral side up, centered and perpendicular to an Asus liquid-crystal display (LCD) at 6.5 cm distance. The screen had a refresh rate of 165 Hz, a linearized contrast with a mean illuminance of 200 Lux, and a spatial resolution of 2,560 × 1,440 pixels, giving a projected screen size of 153° × 138°. Visual stimuli were displayed using custom written software based on the Psychophysics toolbox (54, 55) in Matlab (Mathworks).

TSDNs were identified as described (22, 24). In short, we mapped the receptive field of each neuron by scanning a target horizontally and vertically at 20 evenly spaced elevations and azimuths (24) to calculate the local motion sensitivity and local preferred direction. We then scanned targets of varying size through the small, dorso-frontal receptive fields (Fig. 3A) to confirm that each neuron was sharply size tuned with a peak response to targets subtending 3° to 6°, with no response to larger bars to looming or to widefield stimuli (22, 24).

Unless otherwise mentioned, targets were black and round and had a diameter of 15 mm, moving at a velocity of 900 pixels/s for 0.48 s. When converted to angular values and taking the small frontal receptive fields of TSDNs into account, this corresponds to an average diameter of 3° and a velocity of 130°/s (22). Unless otherwise stated, each target traveled in each neuron’s preferred horizontal direction (i.e., left or right) and across the center of its receptive field. Between repetitions, we varied the target elevation slightly, to minimize habituation (22). There was a minimum 4 s between stimulus presentations. Stimulus order was randomized.

Optic flow was generated as previously described (24). Briefly, the optic flow consisted of a simulated cube with 4-m sides, filled with 2-cm-diameter spheres at a density of 100/m³, unless otherwise stated, with the hoverfly placed in the center. The motion of these randomly placed circa 6,400 spheres around the hoverfly was used to simulate self-generated optic flow. The circa 1,200 spheres anterior to the hoverfly were projected onto the screen, with their size indicating the distance from the hoverfly. Circles closer than 6 cm were not displayed. Six types of optic flow were simulated: three translations at 50 cm/s (sidelisp, lift, and thrust) and three rotations at 50°/s (yaw, pitch, and roll). Unless otherwise stated, optic flow was displayed for 0.48 s prior to the target. Both target motion and optic flow disappeared simultaneously.

In most experiments, the optic flow covered the entire visual display. In some experiments, we limited the spatial extent of the optic flow into four spatial positions. TSDN receptive fields tend to be located slightly offset from the visual midline, with preferred direction of motion away from the midline (Fig. 3A). We defined the lateral parts of the display as either ipsilateral or contralateral based on the preferred direction of each TSDN.

Data Analysis and Statistics. We recorded from 39 TSDNs in 39 male hoverflies. We kept data from all TSDNs that showed a robust response to a target moving over a white background (Fig. 1 and Movies S1 and S2). We repeated this control throughout the recording and only kept data from neurons that responded consistently. We only kept data from experiments with a minimum of nine repetitions. The data from repetitions within a neuron were averaged and shown as spike histograms (mean ± SEM) with 1-ms resolution, after smoothing with a 20-ms square-wave filter. For quantification across neurons, we calculated the mean spike rate for each neuron from the spike histogram for the duration of target motion, after excluding the first and last 40 ms of each 0.48-s target trajectory (dotted boxes, Fig. 1 B and C), unless otherwise indicated. We normalized the responses to each neuron’s own mean response to a target moving over a white background. Percentage change was defined as (Response_{experimental condition} – Response_{stationary control}) / Response_{stationary control}.

Data analysis was performed in Matlab and statistical analysis in Prism 7.0c for Mac OS X (GraphPad Software). Throughout the paper, n refers to the number of repetitions within one neuron and N to the number of neurons. The sample size, type of test (paired t tests or one-way ANOVAs, followed by Dunnett’s post hoc test for multiple comparisons), and P value are indicated in each figure legend. All data have been deposited to DataDryad (https://doi.org/10.5061/dryad.rn8pk0p6z).

Data Availability. Prism files data have been deposited in DataDryad (https://doi.org/10.5061/dryad.rn8pk0p6z).

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