# Imaging neural activity in the ventral nerve cord of behaving adult *Drosophila*

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## 1 Abstract

To understand neural circuits that control limbs, one must measure their activity during 2 behavior. Until now this goal has been challenging, because the portion of the nervous 3 system that contains limb premotor and motor circuits is largely inaccessible to large-4 scale recording techniques in intact, moving animals – a constraint that is true for both 5 vertebrate and invertebrate models. Here, we introduce a method for 2-photon 6 functional imaging from the ventral nerve cord of behaving adult Drosophila 7 melanogaster. We use this method to reveal patterns of activity across nerve cord 8 populations during grooming and walking and to uncover the functional encoding of 9 moonwalker ascending neurons (MANs), moonwalker descending neurons (MDNs), and 10 a novel class of locomotion-associated descending neurons. This new approach 11 enables the direct investigation of circuits associated with complex limb movements. 12

## 13 Introduction

14 Limbs allow animals to rapidly navigate complex terrain, groom, manipulate objects, and communicate. In vertebrates, neural circuits in the spinal cord coordinate the actions of 15 each arm or leq. Circuits within the thorax perform comparable tasks in insects. The 16 thoracic segments of the fruit fly, Drosophila melanogaster, house the ventral nerve 17 cord (VNC) which is a fusion of three thoracic and eight abdominal ganglia. The VNC 18 contains six spherical neuromeres, each controlling one leg, a flat dorsal neuropil 19 associated with the neck, wing, and halteres, and a set of intermediate neuropils 20 including the tectulum that may coordinate the action of the legs and wings. Also within 21 the thorax are descending and ascending axons that connect the VNC to the brain run 22 within a pair of neck or cervical connectives, which - like the VNC - are inaccessible in 23 most preparations. 24

In recent years, the VNC of adult *Drosophila* has gained attention as the site where some higher-order decisions are transformed into actions. Adult flies engage in complex limbed behaviors including walking<sup>1,2</sup>, reaching<sup>3</sup>, escape jumping<sup>4</sup>, courtship tapping<sup>5</sup>, aggressive boxing<sup>6</sup>, and grooming<sup>7</sup>. Our current understanding of how the VNC coordinates these actions is entirely based on behavioral genetics or recordings from a few neurons in tissue explants<sup>8</sup>, immobilized animals<sup>9-11</sup>, or sharp electrode studies in larger insects<sup>12,13</sup>.

To fully understand how VNC circuits orchestrate limb movements, it is necessary to record the activity of individual cells and populations of neurons during behavior. To date, these experiments have not been performed in *Drosophila* due to the difficulty of accessing the VNC in intact, behaving animals. Here we describe a preparation that overcomes this obstacle and makes it possible to record VNC population dynamics in adult animals during walking, grooming, and other actions involving limb movement.

## 39 **Results**

40 The VNC lies on the thoracic sternum - a cuticular structure that anchors the leg muscles and the proximal leg segments to the thorax (Fig. 1a). Consequently, it is 41 difficult to access the VNC by removing ventral thoracic cuticle without destroying 42 musculoskeletal elements required for limb movement. We chose instead to access the 43 VNC dorsally at the expense of flight-related behaviors<sup>14</sup>. This approach requires 44 removing the prescutum and scutum of the dorsal thoracic cuticle, the indirect flight 45 muscles (IFMs), and transecting the proventriculus, crop, and salivary glands of the gut 46 (Fig. 1a, see Methods). 47

Using this technique, it is possible to uncover the VNC for functional imaging in 48 flies that are still capable of exhibiting robust behavior, such as walking and grooming, 49 for at least 2-4 hours. To illustrate the extent of VNC access, we drove expression of the 50 genetically encoded calcium indicator, GCaMP6s<sup>15</sup>, together with a fiduciary 51 tdTomato<sup>16</sup>, throughout the 52 fluorophore, entire nervous system (GMR57C10>GCaMP6s; tdTomato)<sup>17</sup>, (Fig. 1b-c and Supplementary Video 1). To 53 perform 2-photon microscopy in semi-intact, behaving animals, we constructed a 54 customized fly holder and spherical treadmill (Supplementary Fig. 1a) that, in contrast 55 to previous methods used to record neural activity in the brain<sup>14,18,19</sup>, permits 56 unobscured optical access to the VNC and videography of leg movements 57 (Supplementary Fig. 1b). 58

By focusing on dorsoventral horizontal image planes in animals expressing 59 GCaMP6s and tdTomato pan-neuronally (GMR57C10>GCaMP6s; tdTomato), we could 60 record the detailed time course of neural activity in the right and left prothoracic leg 61 neuromeres during walking and grooming (Fig. 2a-b,e and Supplementary Video 2). 62 We identified two regions-of-interest (ROIs) in the right prothoracic neuromere that 63 correlated with spontaneous prothoracic leg grooming and walking, respectively (Fig. 64 **2e**). Alternatively, we could use a piezo-driven microscope objective to acquire coronal 65 x-z image planes. These coronal sections allowed us to simultaneously record activity at 66 different depths of the VNC corresponding to layers housing sensory neuron axons<sup>20</sup>, 67 interneurons<sup>11</sup>, and motor neuron dendrites<sup>21</sup> (Fig. 2a,c; Supplementary Video 3), or 68

monitor activity patterns across populations of descending<sup>22,23</sup> and ascending
fibers<sup>8,20,23</sup> within the cervical connective (Fig. 2a,d and Supplementary Video 4).
Thus, we confirmed that our new preparation provides optical access to previously
inaccessible thoracic neural populations in behaving animals.

Using *Drosophila*, it is possible to repeatedly and systematically investigate the 73 functional properties of sparse sets of genetically-identifiable neurons. In one recent 74 study, a thermogenetic activation screen was used to identify a pair of descending 75 76 neurons – Moonwalker Descending Neurons (MDNs) – which cause flies to walk backwards<sup>23</sup>. Additionally, concurrent thermogenetic activation of a set of ascending 77 neurons that project from the VNC to the brain – Moonwalker Ascending Neurons 78 (MANs) - resulted in even more sustained backwards walking, perhaps by arresting 79 forward walking<sup>23</sup>. While these activation experiments show that these neurons play an 80 81 important role in the control of backwards walking, their native activity patterns and the means by which they regulate and report limb movements remain unknown. 82

Because MAN and MDN axons terminate in the gnathal ganglia (GNG) and the 83 84 VNC, which are both relatively inaccessible regions of the nervous system, it is difficult 85 to record the activity of these cells during any behavior. We used our functional imaging approach to overcome this challenge and recorded the activity of this set of ascending 86 and descending interneurons within the VNC. Because of the vertical movement 87 artifacts associated with walking, we imaged the activity of MAN axons through coronal 88 sections within the cervical connective (Fig. 3a). With this approach, axons are visible 89 as small ellipses (Fig. 3b). The MAN split-GAL4 line we used drives expression in a pair 90 of dorsal and a pair of ventral neurons. We focused our analysis on the dorsal pair of 91 neurons - hereafter referred to as dMANs - because they showed conspicuous 92 93 changes in activity (Fig. 3c). The activity of left and right dMANs were strongly correlated (**Supplementary Fig. 2a**; Pearson's  $r = 0.96 \pm 0.01$ , n = 5 flies), allowing us 94 to study their collective response properties. Specifically, we automatically identified the 95 occurrence of transient increases in dMAN fluorescence - referred to as 'events' - and 96 examined the corresponding behavioral changes reflected in the spherical treadmill data 97 (see Methods). Our analysis revealed that dMAN events were associated with rapid 98 bimodal anterior-posterior rotations of the spherical treadmill (Fig. 3d, n = 746 left and 99

748 right dMAN events from 9773 s of data from 5 flies). By close inspection of the
 video data, we observed that these rotations occur when flies extend all six legs to push
 down on the ball (Supplementary Videos 5 and 6).

Next, we asked to what extent MDNs are active during periods of backwards 103 walking, a possibility suggested by behavioral responses to thermogenetic<sup>23</sup> or 104 optogenetic<sup>24</sup> MDN stimulation. To address this guestion, we applied our approach of 105 imaging coronal sections of the thoracic cervical connective using MDN driver line flies 106 107 expressing GCaMP6s and tdTomato (*MDN-1>GCaMP6s; tdTomato*)(Fig. 4a-b). As for dMANs, the activity of pairs of MDNs were strongly correlated (Fig 4c), allowing us to 108 focus on their collective response properties (Supplementary Fig. 2b; Pearson's r = 109  $0.93 \pm 0.001$ , n = 3 flies). As predicted, MDNs were active prior to anterior rotations of 110 111 the spherical treadmill, corresponding to brief episodes of backward walking (Fig. 4c-d, n = 900 left and 900 right MDN events from 3 flies and 7790 s of data; **Supplementary** 112 Videos 7 and 8). 113

In addition to resolving the functional properties of previously identified neurons, 114 our method facilitates the discovery of novel cell classes that are active during walking, 115 grooming, and other behaviors involving the limbs or abdomen. As a proof-of-concept, 116 we selected four split-GAL4 lines<sup>25</sup> that drive sparse expression in pairs of descending 117 neurons<sup>22</sup> whose axons project to leg neuromeres in the VNC (classes DNa01, DNb06, 118 119 DNg10, and DNg13). Among these, we found that DNa01 neurons – hereon referred to 120 as A1 cells – were active in a manner that was clearly linked to locomotor state (A1>GCaMP6s; tdTomato)(Fig. 5a-b and Supplementary Video 9). The activity of left 121 and right A1 neurons were not highly correlated (Fig. 5c and Supplementary Fig. 2c: 122 Pearson's  $r = 0.53 \pm 0.17$ , n = 4 flies). Therefore, we investigated the response 123 properties of the left and right cells separately. We found that although the activities of 124 both cells are linked to forward walking, events associated only with left A1 activity were 125 correlated with positive medial-lateral and yaw rotations, or rightward turning by the fly 126 (Fig. 5d and Supplementary Video 10; n = 1644 events from 4 flies and 8784 s of 127 data). As expected from bilateral symmetry, activity in the right A1 neuron coincided 128 with negative medial-lateral and yaw rotations, or leftward turning (Fig. 5e and 129 **Supplementary Video 11**; n = 1651 events from 4 flies and 8784 s of data). 130

This approach for recording neural activity in the VNC of behaving Drosophila 131 opens up many new avenues for studying premotor and motor circuits. Nevertheless, 132 we can imagine further improvements that will accelerate the study of the thoracic 133 nervous system. For example, in our preparation we found it challenging and time-134 consuming to remove the indirect flight muscles (IFMs) which fill most of the thorax. 135 Although large, these muscles are quite fragile and tend to disintegrate over the course 136 of an hour after the cuticle of the notum is removed. To increase the efficiency of our 137 dissection, we devised a transgenic strategy to selectively ablate IFMs. We drove the 138 expression of Reaper – a protein promoting apoptosis<sup>26</sup> – in the IFMs by using a 5' 139 Act88F promotor sequence<sup>27</sup>. Act88F:Rpr animals show a nearly complete loss of the 140 IFMs after 7 days post-eclosion (dpe) when raised at 25°C (Fig. 6a-b). This loss results 141 in highly elevated or slightly depressed wings, phenotypes identical to those seen in 142 IFM developmental mutants<sup>28</sup>. The heterozygous *Act88F:Rpr* transgenic background 143 greatly accelerated the dorsal thoracic dissection. Although the imaging data in this 144 manuscript were performed without the Act88F:Rpr transgene, we envisage that this 145 genetic reagent will greatly simplify and accelerate use of this method in the 146 neuroscience community. 147

Here we have described a new preparation that enables the visualization of genetically identified neurons in the VNC and cervical connective of *Drosophila* during behavior. We can record the activity of entire neural populations (**Fig. 2**) or measure the encoding of known (**Figs. 3** and **4**) and novel sparse cell classes (**Fig. 5**). This method fills a critical gap in the study of sensory-motor pathways and serves as a complement to ongoing genetic behavioral screens<sup>23,29,30</sup> aimed at elucidating how populations of neurons coordinate limb movements and orchestrate a variety of legged behaviors.

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### 156 Methods

#### 157 Drosophila lines

Several lines (*GMR57C10-Gal4*, *UAS-GCaMP6s*, *UAS-GCaMP6f*, *UAS-CD4:tdGFP*,
and *UAS-tdTomato*) were obtained from the Bloomington Stock Center. *MAN-Gal4*(*VT50660-AD*; *VT14014-DBD*) and *MDN-1-Gal4* (*VT44845-DBD*; *VT50660-AD*) were
provided by B. Dickson (Janelia Research Campus). *DNa01-Gal4* (SS00731:

GMR22C05-AD; GMR56G08-DBD), DNb06-Gal4 (SS02631: GMR20C04-AD; 162 BJD113E03-DBD), DNg13-Gal4 (SS02538: BJD118A10-AD; BJD123E03-DBD), and 163 DNg16-Gal4 (SS01543: BJD104A07-AD; BJD107F12-DBD) were provided by G. Rubin 164 (Janelia Research Campus). Transgenic Actin88F:Rpr strains (Act88F:Rpr flies) were 165 generated using an Actin88F:eGFP construct described previously<sup>27</sup> and injected 166 (BestGene, Chino Hills, CA, USA) with the phiC31-based integration system using the 167 attP18 (X chromosome) landing site<sup>31</sup>. 168

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### 170 Fluorescence imaging of indirect flight muscles

Fluorescent microscopy of hemi-thoraces was performed as described previously<sup>32,33</sup>. 171 Briefly, flies were anesthetized and their heads and abdomens were then removed. 172 Thoraces were fixed overnight in 4% paraformaldehyde at 4°C and rinsed in 1x 173 phosphate buffered saline (PBS) the following day. The specimens were arranged on a 174 glass slide, snap frozen in liquid nitrogen and bisected down the midsagittal plane using 175 a razor blade. IFMs were stained with Alexa-Fluor 568 Phalloidin (1:100 in PBS with 176 0.1% Triton-X (PBST)) overnight at 4°C, rinsed with PBS and visualized using EVOS® 177 FL Cell Imaging System (Life Technologies) at 4X magnification. For whole-mount 178 imaging of IFM myofibrils, flies were prepared and thoraces bisected as described 179 above. Hemi-thoraces were stained with Alexa-Fluor 568 phalloidin (1:100 in PBST) 180 181 overnight at 4°C. Samples were rinsed in PBS, mounted with Vectashield (Vector Laboratories) and visualized using a Leica TCS SPE RGBV confocal microscope (Leica 182 183 Microsystems) at 100X magnification.

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#### 185 Immunofluorescence imaging of whole-mount brains and ventral nerve cords

Brains and ventral nerve cords (VNCs) were dissected out of 2-3 dpe female flies in PBS. Tissues were then fixed for 20 minutes in 4% paraformaldehyde in PBS at room temperature. After fixation, brains and VNCs were washed 2-3 times in PBS with 1% Triton-X-100 (PBST) and then incubated at 4°C overnight in PBST. Samples were then placed in PBST with 5% normal goat serum (PBSTS) for 20 min at room temperature. They were then incubated with primary antibodies (rabbit anti-GFP at 1:500, Thermofisher; mouse anti-Bruchpilot/nc82 at 1:20, Developmental Studies Hybridoma

Bank) diluted in PBSTS for 48 h at 4°C. Brains and VNCs were rinsed 2-3 times in PBST for 10 min each before incubation with secondary antibodies (goat anti-rabbit secondary antibody conjugated with Alexa 488 at 1:500; Thermofisher; goat anti-mouse secondary antibody conjugated with Alexa 633 at 1:500; Thermofisher) diluted in PBSTS for 48 h at 4 °C. Finally, brains and VNCs were rinsed 2-3 times for 10 min each in PBST and mounted onto slides with bridge coverslips in Slowfade mounting-media (Thermofisher).

Samples were imaged using a Carl Zeiss LSM 700 Laser Scanning Confocal Microscope with the following settings: 20X magnification, 8-bit dynamic range, 2x image averaging,  $0.52 \times 0.52 \mu m$  pixel size,  $0.57 \mu m$  z-axis interval. Standard deviation z-projections of imaging volumes were made using Fiji<sup>34</sup>.

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#### 205 Thoracic dissection for ventral nerve cord imaging

Custom holders used to mount flies during imaging were fabricated as described previously<sup>35</sup>. For VNC imaging, these stages were modified to have (i) flat rather than folded frames, and (ii) chamfered vertices to make the spherical treadmill visible to optic flow sensors (Shapeways, https://www.shapeways.com/model/upload-andbuy/5963553).

All experiments were performed on 1-3 dpe female flies raised at 25°C on 211 212 standard cornmeal food on a 12 h light: 12 h dark cycle. Flies were anaesthetized at 4°C. A female fly was selected and, in some cases, its wings were clipped to simplify the 213 214 mounting process. The fly's dorsal thorax was then pushed through a hole in the steel shim of a custom imaging stage. The stage was then flipped over, UV-curing glue 215 216 (Bondic, Aurora, ON Canada) was carefully applied around the perimeter of the thorax and hardened through UV illumination (LED-200, Electro-Lite Co. Bethel, CT USA). UV 217 glue was then similarly applied to fix the head and abdomen to the underside of the 218 stage. The stage was then filled with extracellular saline as described previously<sup>14</sup>. 219 Under a high-magnification dissection microscope (Leica M165C), a hypodermic needle 220 (30G, BD PrecisionGlide, Franklin Lakes, NJ USA) was used to slice and lift the cuticle 221 off of the dorsal thorax<sup>36</sup>, being careful not to sever the neck connective. Subsequently, 222 in non-Act88F:Rpr animals, a pair of dull forceps was used to remove IFMs, 223

predominantly from the anteriomedial region of the thorax overlying the proventriculus 224 (this step is unnecessary in aged Act88F:Rpr animals). This process exposes the dorsal 225 surface of the proventriculus – a large bulbous structure in the gut. With great care, a 226 pair of super-fine forceps was used to grasp and lift the proventriculus to displace much 227 of the gut (including the crop and salivary glands) from the more ventrally located 228 nervous system. With the gut thus elevated, ultra-fine clipper scissors (Fine Science 229 Tools, Foster City, CA USA) were used to transect it at its anterior-most section. The 230 proventriculus was then peeled back and a posterior incision was made to completely 231 remove these portions of the gut, fully revealing the underlying nervous tissue. In some 232 cases, we observed that gut or muscle tissue would begin to obscure the VNC during 233 imaging. Therefore, loose tissue should be removed at this stage while taking great care 234 not to sever the VNC. After each dissection, we examined the extent to which the 235 animal moved its legs in response to a puff of air. This proved to be an accurate 236 predictor of the success of the preparation. 237

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#### 239 **2-photon microscopy during behavior**

Experiments were performed in the evening zeitgeber time (Z.T.) and animals were typically imaged 30-60 min following dissection. We found that individual animals provided useful data for 2-4 h. Fly holders were secured to a platform raised over the spherical treadmill (**Supplementary Fig. 2a**). The VNC was located using microscope oculars and then aligned to the center of the field-of-view using 2-photon microscopy.

The spherical treadmill was an aluminum rod with a ball-shaped hole milled near 245 its tip<sup>18</sup>. We fabricated 10 mm diameter foam balls (Last-A-Foam FR-7106, General 246 247 Plastics, Burlington Way, WA USA) and manually spotted them using a Rapidograph pen (Koh-I-Noor, Leeds, MA USA) to provide high-contrast features for optic flow 248 measurements. A 500-600 mL/min stream of filtered and humidified air was passed 249 through the holder using a digital flow controller (Sierra Instruments, Monterey, CA 250 USA). Movements of the ball were measured using optical flow sensors (ADNS3080) 251 outfitted with zoom lenses (Computar MLM3X-MP, Cary, NC USA). The ball and fly 252 were illuminated using a pair of IR LEDs (850-nm peak wavelength) coupled to optic 253 fibers and collimator lenses (ThorLabs, Newton, NJ USA). Optic flow measurements 254

were passed to a microcontroller board (Arduino Mega2560) to be recorded using custom-written Python code. Simultaneously, videography of behaviors on the ball were made using an IR-sensitive firewire camera (Basler, Ahrensburg, Germany) at approximately 30 frames per second.

We performed 2-photon microscopy using a Bergamo II microscope (ThorLabs) 259 outfitted with two GaAsP PMT detectors for GCaMP6 and tdTomato imaging, 260 respectively, and coupled to a Ti:Sapphire laser (MaiTai DeepSee, Newport Spectra-261 Physics, Santa Clara, CA USA) tuned to 930 nm. We used an Olympus 20X objective 262 water-immersion lens with 1.0 NA (Olympus, Center Valley, PA USA). The microscope 263 was controlled using Thorlmage software (ThorLabs). Occasionally, a puff of air was 264 used to elicit walking behaviors. These puffs were digitally encoded (Honeywell AWM 265 3300V, Morris Plains, NJ USA). Custom ROS software interfaced through an analog 266 output device (Phidgets, Calgary, Canada) to ThorSync software (ThorLabs) was used 267 to synchronize optic flow measurements, behavior videography, air puff measurements, 268 and 2-photon image acquisition. For coronal section imaging, a piezo collar (Physik 269 Instrumente, Karlsruhe, Germany) was used for rapid z-axis movements of the 270 microscope objective lens. 271

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#### 273 Data analysis

We analyzed all data using custom scripts written in Python. Because the data acquisition frequency differed for optic flow, behavior videography, and 2-photon imaging we interpolated signals to that of the highest frequency. Subsequently, optic flow data were smoothed using a running average and then translated into rotations s<sup>-1</sup> for the anterior-posterior, medial-lateral, and yaw axes as described in<sup>18</sup>.

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Pan-neuronal image registration, ROI identification, and fluorescence processing
 (related to Fig. 2)

We observed that large tissue deformations could occur during behavior. Therefore, we performed post-hoc registration of pan-neuronal imaging data. To do this, we registered all frames of an imaging experiment with one reference image. Because the complexity of the deformations could not be captured by simple parametric motion models (e.g.,

affine transformations), we used a non-parametric, variational approach, designed to model arbitrarily complex deformations. We computed the motion field w between the reference image, denoted  $I_r$ , and the image at time t, denoted  $I_t$ , by solving the minimization problem

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$$\widehat{\mathbf{w}} = \arg\min_{\mathbf{w}} D(\mathbf{w}) + \lambda \sum_{\mathbf{x} \in \Omega} \|\nabla \mathbf{w}(\mathbf{x})\|_2^2, \tag{1}$$

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where  $D(\mathbf{w})$  is a data fitting term, the second term is a regularization promoting smoothness of w by penalizing its gradient  $\nabla \mathbf{w}^{37}$ ,  $\Omega$  is the discrete image domain, and the parameter  $\lambda$  weights the contributions of the two terms.

The sequence tagged with GCaMP6s images is characterized by two main difficulties for motion estimation. First, fast motion of the fly induces very large deformations. Second, the activation of neurons produces large local intensity changes between corresponding pixels in  $I_r$  and  $I_t$ . To address these issues, we defined a data term of the form

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$$D(\mathbf{w}) = \rho(\mathbf{w}, I_r, I_t) + \gamma \phi(\mathbf{w}, J_r, J_t).$$
<sup>(2)</sup>

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The first term models the standard assumption of conservation of intensity along the trajectory of each pixel. It is defined by

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$$\rho(\mathbf{w}, I_r, I_t) = \sum_{\mathbf{x} \in \Omega} |I_t(\mathbf{x} + \mathbf{w}(\mathbf{x})) - I_r(\mathbf{x})|,$$
(3)

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where we use an  $\ell_1$  norm to gain partial robustness to intensity changes<sup>38</sup>. The second term in (2) is a feature matching constraint inspired by Revaud and co-workers<sup>39</sup>, written as

$$\phi(\mathbf{w}, J_r, J_t) = \sum_{\mathbf{x} \in \Omega} \|\mathbf{w}(\mathbf{x}) - \mathbf{m}(\mathbf{x}, J_r, J_t)\|_1,$$
(4)

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where the images  $J_r$  and  $J_r$  are the analog of  $I_r$  and  $I_t$  in the second channel tagged 311 with tdTomato, for which we do not expect activity-dependent intensity changes. 312 Minimizing the function  $\phi$  favors motion vectors w(x) to be close to feature 313 correspondences  $m(x, J_r, J_t)$ , computed on a sparse set of relevant keypoints. We obtain 314 m with the feature matching algorithm proposed by Revaud and co-workers<sup>39</sup>, which is 315 specifically designed to handle large image deformations. We compute m using  $I_r$  and 316  $J_t$ , such that the correspondences are also insensitive to the intensity changes between 317  $I_r$  and  $I_t$ . As a result, the estimation is guided by reliable feature matches. We found 318 that it is necessary to keep a standard data term (3) defined on the GCaMP6s imaging 319 channel, because the tdTomato channel may not provide information in some regions of 320 the image. The parameter  $\gamma$  balances the two terms in (2). 321

We solved the optimization problem (1) with an alternated direction method of multiplier (ADMM) algorithm<sup>40</sup>. We introduced two splitting variables, associated with the regularization and the feature matching terms, respectively. Each sub-problem of the algorithm was solved analytically. We used parts of the inverse problems library described in <sup>41</sup>. A post processing based on weighted median filtering was applied with the method of <sup>42</sup>.

From these registered imaging data, regions-of-interest (ROIs) were manually selected.  $\Delta F/F$  values were then calculated from fluorescence signals averaged within the ROIs.  $\Delta F = F_t - F$ , where  $F_t$  is the average fluorescence within an ROI at time, *t*. F is a baseline fluorescence signal that was calculated as the average pixel value for the first ten sequential GCaMP6s images for which no cellular activity was observed (i.e., minimal and unchanging GCaMP6s fluorescence).

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335 Sparse neuron ROI identification, and fluorescence processing (related to Figs. 3-5)

For single-neuron fluorescence data, ROIs were selected using custom Python scripts relying on OpenCV and Numpy libraries. First, a reference frame was selected for which software identified all potential ROIs. To do this, the GCaMP6s channel image was smoothed to reduce background noise and then an Otsu filter threshold was applied to the image. An erosion factor was then applied on all objects detected within the image.

Contours of all detected objects were then presented to the user for manual selection. 341 Once these reference ROIs were selected for left and right neurons, we used a cross-342 correlation-based image registration algorithm<sup>43</sup> to identify the most likely left and right 343 ROIs for each image frame based on those manually selected on the reference frame. 344 A second script was used to manually verify automatically selected ROIs and, if 345 incorrect, to display all potential ROIs within the frame for manual selection. If chosen 346 erosion values yielded malformed ROI shapes, another script was used to manually 347 position elliptical ROIs with arbitrary orientations on a given frame. Finally, binary ROI 348 images were used as an image mask to extract mean fluorescence intensities from the 349 original GCaMP6s or tdTomato images. These signals were reported as % AR/R as in <sup>44</sup> 350 to reduce the effects of motion artifacts on fluorescence signals. Due to the absence of 351 stimuli for eliciting behaviors, the baseline R was calculated as the minimum ratio of 352 GCaMP6s / tdTomato within a 2.5 s bin. 353

To detect transient increases in activity above the  $\%\Delta R/R$  baseline, we 354 developed an algorithm based partly on <sup>45</sup>. We first determined when the first order 355 derivative of the  $\%\Delta R/R$  signal crossed an arbitrary threshold, which was calculated as 356 a percentile determined by examining all derivatives values for a given neuron class 357 (i.e., MDN, MAN, or A1). We reasoned that threshold values should be characteristic 358 and potentially different for each neuron class because fluorescence dynamics are 359 360 related to intrinsic physiological properties that can differ across neuron classes but not across experiments investigating a single class. We set this threshold for the derivative 361 value as the 97.5<sup>th</sup> percentile for MDNs and dMANs and 90<sup>th</sup> percentile for A1 neurons. 362 A lower threshold value was selected for A1 neurons because many more fluorescence 363 364 transients were observed in A1 % AR/R fluorescence traces. These transients would be overlooked using the 97.5<sup>th</sup> percentile. To identify the onset of fluorescence increases 365 we found the nearest preceding time point where the derivative crossed zero (i.e., 366 typically an inflection between decreases and increases in fluorescence). This zero-367 crossing is considered the time-point of an 'event' associated with the identified 368 fluorescence increase. Events detected close to one another with no intervening 369 derivative zero-crossing were compressed into one event associated with the first time 370 point. There were ~10 separate experiments per animal. Events in the first and last 10 s 371

of each experiment were not considered since the window for data presentation encompassed 10 s before and 10 s after each event.

Because left and right MDN and dMAN cells strongly covaried (**Supplementary Fig. 2**), an additional step was performed for event detection: if events were detected in both left and right neurons within 2 s of one another, both events were retained; otherwise, an event identified for neuron A (e.g., left MDN) and not neuron B (e.g., right MDN) was also added to neuron B's event library.

By contrast, left and right A1 neural activities did not strongly covary. Therefore, analyzed events associated uniquely to one and not the other neuron. To accomplish this, if an event was detected in both left and right A1 neurons within a time window of 0.25 s, neither of the events were logged for analysis.

 $\%\Delta R/R$  and optic flow traces linked to each event were aligned by setting the 383 event time points to 0 s. We then computed the mean and bootstrapped 95% 384 confidence intervals for these aligned traces using the Python Seaborn library. Optic 385 flow and  $\Delta R/R$  measurements were downsampled to 500 values/s for this analysis. To 386 increase clarity, %AR/R traces were baseline-subtracted to make them zero at the time 387 of the event in the summary **Figs. 3d**, **4d**, and **5d-e**. Control shuffled data (gray traces) 388 389 were computed by instead assigning random time-points in place of real, identified events. These random time points were treated as real events and their mean and 390 391 bootstrapped 95% confidence intervals were computed and plotted for comparison.

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393 Covariance analysis (related to Supplementary Figure 2)

Covariance analysis was performed with a custom Python script using Matplotlib and Numpy libraries. Scatter plots were computed comparing left and right neuron  $\%\Delta R/R$ values from all experiments for each fly separately. All data were included for analysis in these scatter plots with the exception of two MDN flies that exhibited unusually low fluorescence values. Pearson's r values are reported as mean ± standard deviation.

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Event-related behaviors (related to Supplementary Videos 6, 8, 10, and 11)

Events chosen for behavioral summary videos were chosen from automatically detected

402 events as described above. For dMANs, events were manually selected from those that

403 maximized the difference in anterior-posterior ball rotations between 1 s before the 404 event and 2 s after the event. For MDNs, events were manually selected from among 405 those that minimized anterior-posterior ball rotations up to 2 s after the event. For A1 406 neurons, events were manually selected from among those that maximized the average 407 yaw ball rotations (positive for left A1 neuron examples and negative for right A1 neuron 408 examples) for up to 2 s after the events.

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417

## 418 **Author contributions**

- 419 C.L.C. generated strains; performed experiments; analyzed data
- 420 L.H. analyzed data
- 421 M.C.V. performed experiments; analyzed data
- 422 D.F. wrote analysis code
- 423 M.U. supervised the project
- 424 A.C. designed and supervised the project
- 425 M.H.D. designed and supervised the project
- 426 P.R. conceived of, designed, and supervised the project; performed experiments;
- 427 analyzed data
- 428 All authors contributed to writing the paper

429

## 430 **Competing financial interests**

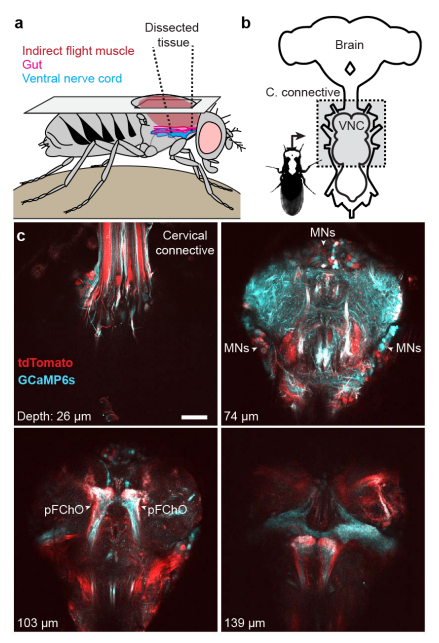
431 The authors declare no competing financial interests.

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## 433 **Corresponding author**

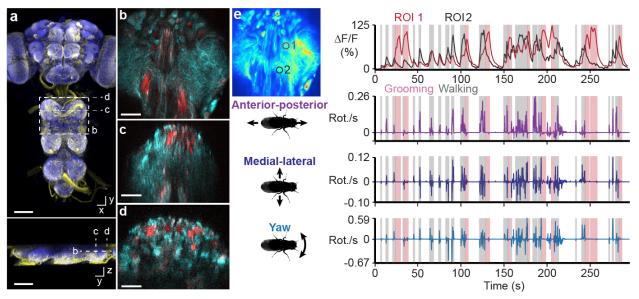
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## 435 **Figure Legends**



436

Figure 1 | Dissection for imaging the adult *Drosophila* ventral nerve cord (VNC).
(a) Schematic of the dorsal thoracic dissection. (b) Overview of newly accessible
nervous tissue following the thoracic dissection. (c) Horizontal sections of the VNC
imaged at different depths in an animal expressing GCaMP6s (cyan) and tdTomato
(red) throughout the nervous system (*GMR57C10>GCaMP6s; tdTomato*). Motor
neurons (MNs) and prothoracic (pFChO) femoral chordotonal organs are indicated by
white arrowheads. Scale bar is 30 µm.



445 Figure 2 | Recording populations of neurons in the VNC during behavior. (a) Confocal image of pan-neuronal driver line expression in the brain and VNC. Scale bars 446 are 90 µm. Neuronal GFP (yellow) and neuropil (nc82, blue) are labelled. Dashed lines 447 highlight the imaging modalities made possible by thoracic dissection: (b) horizontal 448 section imaging of the VNC (scale bar is 35 µm), (c) coronal section imaging of the VNC 449 (scale bar is 50 µm), and (d) coronal section imaging of the cervical connective (scale 450 bar is 35µm). All three modalities are illustrated by imaging flies expressing GCaMP6s 451 and tdTomato throughout the nervous system (GMR57C10>GCaMP6s; tdTomato). (e) 452 % AF/F image of a VNC horizontal section of the same animal in (b). ROI-associated 453 fluorescence signals during walking (gray) and grooming (pink) are shown on the top-454 right. Corresponding rotations of the spherical treadmill are shown on the bottom-right. 455

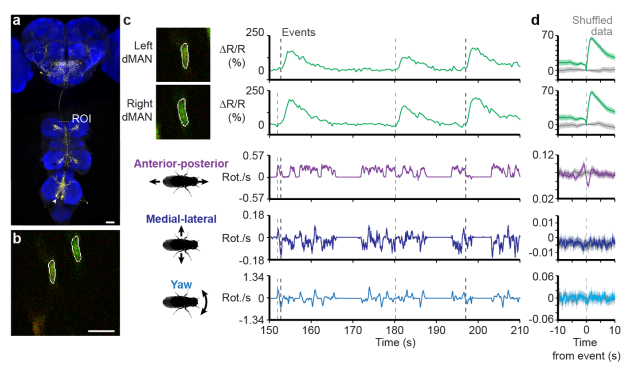


Figure 3 | Recording the activity of dorsal Moonwalker Ascending Neurons 457 (dMANs) during behavior. (a) Confocal image of MAN-Gal4 driver line expression in 458 the brain and VNC (scale bar is 40 µm). Neuronal GFP (yellow) and neuropil (nc82, 459 blue) are labelled. A dashed white line highlights the x-z plane imaged. (b) Coronal 460 section of the thoracic cervical connective in an animal expressing GCaMP6s and 461 tdTomato in MANs (MAN>GCaMP6s; tdTomato). Scale bar is 3.5 µm. (c) Separated 462 ROIs (top-left) and associated fluorescence signals from left and right dMANs (top-463 **right**). Corresponding rotations of the spherical treadmill are shown on the bottom right. 464 Events are indicated as dashed gray lines. (d) Summary of dMAN activity and spherical 465 treadmill rotations with respect to fluorescence events aligned to 0 s (dashed gray line). 466 Control data in which events are time-shuffled are overlaid in grey. Shown are the 467 means (solid line) and bootstrapped 95% confidence intervals (transparencies). 468

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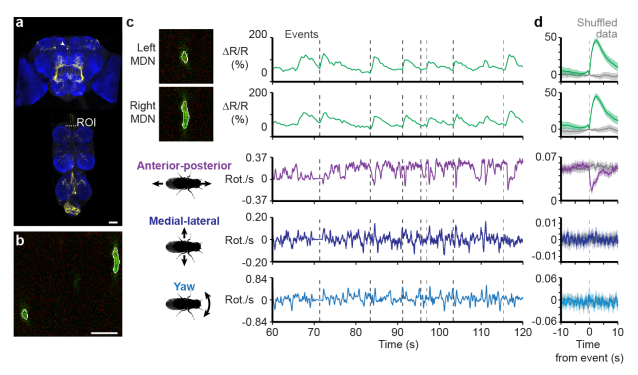
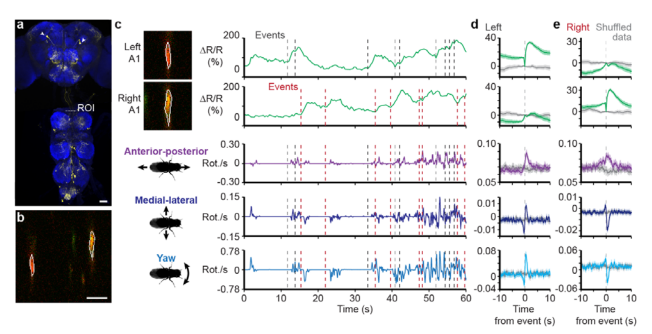


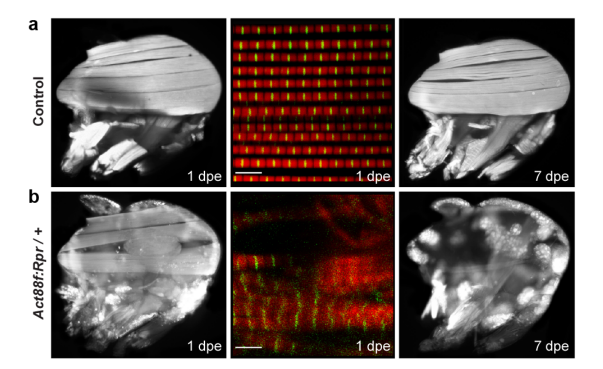
Figure 4 | Recording the activity of Moonwalker Descending Neurons (MDNs) 470 during behavior. (a) Confocal image of MDN-1-Gal4 driver line expression in the brain 471 and VNC (scale bar is 40 µm). Neuronal GFP (yellow) and neuropil (nc82, blue) are 472 labelled. A dashed white line highlights the x-z plane imaged. (b) Coronal section of the 473 thoracic cervical connective in an animal expressing GCaMP6s and tdTomato in 474 Moonwalker Descending Neurons (MDN-1>GCaMP6s: tdTomato). Scale bar is 6 µm. 475 (c) Separated ROIs (top-left) and associated fluorescence signals from left and right 476 MDNs (top-right). Corresponding rotations of the spherical treadmill are shown on the 477 bottom right. Events are indicated as dashed gray lines. (d) Summary of MDN activity 478 and spherical treadmill rotations with respect to fluorescence events aligned to 0 s 479 (dashed gray line). Control data in which events are time-shuffled are overlaid in grey. 480 481 Shown are the means (solid line) and bootstrapped 95% confidence intervals (transparencies). 482

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Figure 5 | Recording the activity of A1 neurons during behavior. (a) Confocal image 484 of DN0a1-Gal4 driver line expression in the brain and VNC (scale bar is 40 µm). 485 Neuronal GFP (yellow) and neuropil (nc82, blue) are labelled. A dashed white line 486 highlights the x-z plane imaged. (b) Coronal section of the thoracic cervical connective 487 in an animal expressing GCaMP6s and tdTomato in A1 neurons (A1>GCaMP6s; 488 tdTomato). Scale bar is 5 µm. (c) Separated ROIs (top-left) and associated 489 fluorescence signals from left and right A1 neurons (top-right). Corresponding rotations 490 of the spherical treadmill are shown on the bottom-right. Events are indicated as dashed 491 gray and red lines for left and right A1 neuron events, respectively. (d) Summary of A1 492 neural activity and spherical treadmill rotations with respect to left A1 neuron 493 fluorescence events aligned to 0 s (dashed gray line). (e) Summary of A1 neural activity 494 and spherical treadmill rotations with respect to right A1 neuron fluorescence events 495 aligned to 0 s (dashed gray line). Control data in which events are time-shuffled are 496 497 overlaid in grey. Shown are the means (solid line) and bootstrapped 95% confidence intervals (transparencies). 498

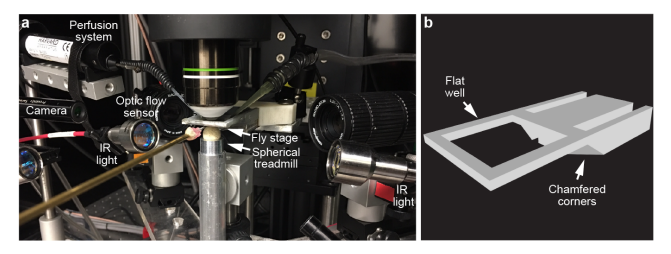


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Figure 6 | Indirect flight muscles in control and *Act88F:Rpr* animals. Confocal
images of dorsal longitudinal IFMs (DLMs) stained with TRITC-phalloidin at 1 dpe (left),
7 dpe (right), or wholemount confocal micrographs of myofibrillar structure (middle) for
(a) wild-type, or (b) *Act88F:Rpr* heterozygous flies. Scale bars are 5 µm.

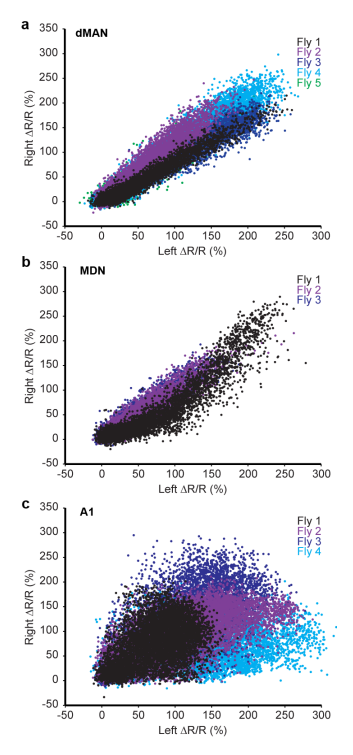
# 504 Supplementary Figures

505



506 Supplementary Figure 1 | System for VNC imaging. (a) Photograph of the spherical

507 treadmill system and **(b)** schematic of the custom fly holder used in this study.



508

509 Supplementary Figure 2 | Covariance in fluorescence signals between bilateral 510 pairs of neurons. Scatter plots comparing  $\Delta R/R$  signals for left and right (a) dMAN,

511 (b) MDN, or (c) A1 neuron pairs. Data from each animal are color-coded.

## 513 Supplementary Videos

514 **Supplementary Video 1** | **Extent of VNC imaging volume.** 2-photon imaging of 515 horizontal sections across the dorsal-ventral extent of the VNC and cervical connective. 516 GCaMP6s (cyan) and tdTomato (red) are expressed throughout the nervous system 517 (*GMR57C10>GCaMP6s; tdTomato*). Imaging depth is indicated on the top-left.

518

519 Supplementary Video 2 | Horizontal VNC imaging. 2-photon imaging of a single horizontal section of the VNC in a walking and grooming fly. GCaMP6s (cyan) and 520 tdTomato (red) are expressed throughout the nervous system (GMR57C10>GCaMP6s; 521 tdTomato). Shown are synchronized raw fluorescence images (top-left),  $\Delta F/F$  images 522 523 (top-right), behavior images (bottom-left), and spherical treadmill rotations along the anterior-posterior ('AP'), medial-lateral ('ML'), and yaw axes (bottom-right). 524 Experimenter-administered air puffs are indicated by the appearance of red boxes 525 above behavior video images. Video is 4X faster than real-time. 526

527

Supplementary Video 3 | Coronal VNC imaging. 2-photon imaging of a single coronal section of the VNC in a walking fly. GCaMP6s (cyan) and tdTomato (red) are expressed throughout the nervous system (*GMR57C10>GCaMP6s; tdTomato*). Shown are synchronized raw fluorescence images (top-left),  $\%\Delta$ F/F images (top-right), behavior images (bottom-left), and spherical treadmill rotations along the anterior-posterior ('AP'), medial-lateral ('ML'), and yaw axes (bottom-right). Video is 4X faster than realtime.

535

Supplementary Video 4 | Coronal cervical connective imaging. 2-photon imaging of a single coronal section of the cervical connective in a walking fly. GCaMP6s (cyan) and tdTomato (red) are expressed throughout the nervous system (*GMR57C10>GCaMP6s; tdTomato*). Shown are synchronized raw fluorescence images (top-left),  $\%\Delta$ F/F images (top-right), behavior images (bottom-left), and spherical treadmill rotations along the

anterior-posterior ('AP'), medial-lateral ('ML'), and yaw axes (bottom-right). Video is 4X
faster than real-time.

543

Supplementary Video 5 | Coronal cervical connective imaging of dorsal 544 **Moonwalker Ascending Neurons.** 2-photon imaging of a single coronal section of the 545 cervical connective in a behaving fly. GCaMP6s (cyan) and tdTomato (red) are 546 expressed in MANs (MAN>GCaMP6s; tdTomato). Raw fluorescence images of the left 547 548 and right dMANs are presented and outlined by ROIs (top-left). These images are used to calculate % AR/R traces for each neuron (top-right). Corresponding behavior 549 550 videography (bottom-left) and spherical treadmill rotations along the anterior-posterior ('AP'), medial-lateral ('ML'), and vaw axes (bottom-right) are shown. 551

552

Supplementary Video 6 | Behavioral responses associated with dorsal Moonwalker Ascending Neuron activity events. Three example behaviors (rows) for each of three flies (columns) produced at the onset of dMAN fluorescence events. Red square indicates the time of each fluorescence event (t = 0 s). Video is 3X slower than real-time.

558

Supplementary Video 7 | Coronal cervical connective imaging of Moonwalker 559 **Descending Neurons.** 2-photon imaging of a single coronal section of the cervical 560 connective in a behaving fly. GCaMP6s (cyan) and tdTomato (red) are expressed in 561 MDNs (*MDN-1*>*GCaMP6s; tdTomato*). Raw fluorescence images of the left and right 562 MDNs are presented and outlined by ROIs (top-left). These images are used to 563 calculate  $\%\Delta R/R$  traces for each neuron (top-right). Corresponding behavior 564 videography (bottom-left) and spherical treadmill rotations along the anterior-posterior 565 ('AP'), medial-lateral ('ML'), and yaw axes (bottom-right) are shown. 566

567

568 Supplementary Video 8 | Behavioral responses associated with Moonwalker 569 Descending Neuron activity events. Three example behaviors (rows) for each of

three flies (columns) produced at the onset of MDN fluorescence events. Red square indicates the time of each fluorescence event (t = 0 s). Video is 3X slower than realtime.

573

Supplementary Video 9 | Coronal cervical connective imaging of A1 Neurons. 2-574 photon imaging of a single coronal section of the cervical connective in a behaving fly. 575 GCaMP6s (cyan) and tdTomato (red) are expressed in A1 neurons (A1>GCaMP6s; 576 577 tdTomato). Raw fluorescence images of the left and right A1 neurons are presented and outlined by ROIs (top-left). These images are used to calculate  $\%\Delta R/R$  traces for each 578 579 neuron (top-right). Corresponding behavior videography (bottom-left) and spherical treadmill rotations along the anterior-posterior ('AP'), medial-lateral ('ML'), and yaw axes 580 581 (bottom-right) are shown. Video is 2X faster than real-time.

582

Supplementary Video 10 | Behavioral responses associated with left A1 neuron activity events. Three example behaviors (rows) for each of three flies (columns) produced at the onset of left A1 neuron fluorescence events. Red square indicates the time of each fluorescence event (t = 0 s). Video is 3X slower than real-time.

587

Supplementary Video 11 | Behavioral responses associated with right A1 neuron activity events. Three example behaviors (rows) for each of three flies (columns) produced at the onset of right A1 neuron fluorescence events. Red square indicates the time of each fluorescence event (t = 0s). Video is 3X slower than real-time.

## 592 **References**

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- Wosnitza, A., Bockemühl, T., Dübbert, M., Scholz, H. & Büschges, A. Inter-leg coordination in the control of walking speed in *Drosophila*. *The Journal of Experimental biology* **216**, 480–491 (2013).
- Mendes, C. S., Bartos, I., Akay, T., Márka, S. & Mann, R. S. Quantification of gait
   parameters in freely walking wild type and sensory deprived *Drosophila melanogaster. eLife* 2, (2013).
- Niven, J. E. Visuomotor control: *Drosophila* bridges the gap. *Curr Biol* 20, R309–
  11 (2010).
- 4. Card, G. M. & Dickinson, M. H. Visually mediated motor planning in the escape response of *Drosophila*. *Current Biology* **18**, 1300–1307 (2008).
- 5. Pavlou, H. J. & Goodwin, S. F. Courtship behavior in *Drosophila melanogaster*. towards a 'courtship connectome'. *Curr Opin Neurobiol* **23**, 76–83 (2013).
- 610 6. Zwarts, L., Versteven, M. & Callaerts, P. Genetics and neurobiology of aggression 611 in *Drosophila*. *fly* **6**, 35–48 (2012).
- 6137.Seeds, A. M. *et al.* A suppression hierarchy among competing motor programs614drives sequential grooming in *Drosophila*. *eLife* **3**, (2014).
- 6168.Mann, K. J., Gordon, M. D. & Scott, K. A pair of interneurons influences the<br/>choice between feeding and locomotion in *Drosophila*. **79**, 754–765 (2013).
- 6199.Trimarchi, J. R. & Murphey, R. K. The shaking-B-2 mutation disrupts electrical620synapses in a flight circuit in adult *Drosophila*. *J Neurosci* **17**, 4700–4710 (1997).
- 62210.Ikeda, K. & Kaplan, W. D. Neurophysiological Genetics in Drosophila623melanogaster. American Zoologist 14, 1055–1066 (1974).
- Tuthill, J. C. & Wilson, R. I. Parallel Transformation of Tactile Signals in Central
  Circuits of *Drosophila*. *Cell* 164, 1046–1059 (2016).
- 62812.Hedwig, B. & Burrows, M. Presynaptic inhibition of sensory neurons during kicking629movements in the locust. J Neurophysiol **75**, 1221–1232 (1996).
- 13. Bässler, U. & Büschges, A. Pattern generation for stick insect walking
   movements—multisensory control of a locomotor program. *Brain Research Reviews* 27, 65–88 (1998).
- 63514.Maimon, G., Straw, A. D. & Dickinson, M. H. Active flight increases the gain of636visual motion processing in *Drosophila*. Nat Neurosci **13**, 393–399 (2010).

637

Chen, T.-W. et al. Ultrasensitive fluorescent proteins for imaging neuronal activity.

15.

638

Nature 499, 295–300 (2013). 639 640 16. Shaner, N. et al. Improved monomeric red, orange and yellow fluorescent proteins 641 derived from Discosoma sp. red fluorescent protein. Nature Biotechnology 22, 642 1567-1572 (2004). 643 644 17. Jenett, A. et al. A GAL4-Driver Line Resource for Drosophila Neurobiology. Cell 645 Reports 2, 991–1001 (2012). 646 647 18. Seelig, J. D. et al. Two-photon calcium imaging from head-fixed Drosophila during 648 optomotor walking behavior. Nature Methods 7, 535–540 (2010). 649 650 19. Hedwig, B. & Poulet, J. Complex auditory behaviour emerges from simple 651 reactive steering. Nature 430, 781-785 (2004). 652 653 20. Tsubouchi, A. et al. Topological and modality-specific representation of 654 somatosensory information in the fly brain. Science 358, 615-623 (2017). 655 656 21. Enriquez, J. et al. Specification of Individual Adult Motor Neuron Morphologies by 657 Combinatorial Transcription Factor Codes. Neuron 86, 955–970 (2015). 658 659 22. Namiki, S., Dickinson, M. H., Wong, A. M., Korff, W. & Card, G. M. The functional 660 organization of descending sensory-motor pathways in Drosophila. bioRxiv 1-67 661 (2017). 662 663 Bidaye, S. S., Machacek, C., Wu, Y. & Dickson, B. J. Neuronal control of 23. 664 Drosophila walking direction. Science 344, 97-101 (2014). 665 666 24. Sen, R. et al. Moonwalker Descending Neurons Mediate Visually Evoked Retreat 667 in Drosophila. Curr Biol 27, 766–771 (2017). 668 669 Luan, H., Peabody, N. C., Vinson, C. R. & White, B. H. Refined Spatial 670 25. Manipulation of Neuronal Function by Combinatorial Restriction of Transgene 671 Expression. 52, 425–436 (2006). 672 673 White, K. et al. Genetic Control of Programmed Cell Death in Drosophila. Science 674 26. **264,** 677–683 (1994). 675 676 Ramdya, P., Schaffter, T., Floreano, D. & Benton, R. Fluorescence Behavioral 27. 677 Imaging (FBI) Tracks Identity in Heterogeneous Groups of Drosophila. PLoS One 678 679 **7,** e48381 (2012). 680 28. Reedy, M. C., Bullard, B. & Vigoreaux, J. O. Flightin is essential for thick filament 681 682 assembly and sarcomere stability in Drosophila flight muscles. J. Cell Biol. 151, 1483-1499 (2000). 683

684		
685	29.	Robie, A. A. et al. Mapping the Neural Substrates of Behavior. Cell 170, 393–406
686		(2017).
687		
688	30.	Harris, R. M., Pfeiffer, B. D., Rubin, G. M. & Truman, J. W. Neuron hemilineages
689		provide the functional ground plan for the <i>Drosophila</i> ventral nervous system.
690		<i>eLife</i> <b>4</b> , (2015).
691		
692	31.	Markstein, M., Pitsouli, C., Villalta, C., Celniker, S. E. & Perrimon, N. Exploiting
693	01.	position effects and the gypsy retrovirus insulator to engineer precisely expressed
694		transgenes. <i>Nat Genet</i> <b>40</b> , 476–483 (2008).
695		(101030100.140.140.100.100.12000).
696	32.	Nongthomba, U. & Ramachandra, N. B. A direct screen identifies new flight
697	52.	muscle mutants on the <i>Drosophila</i> second chromosome. <i>Genetics</i> <b>153</b> , 261–274
698		(1999).
699		(1999).
700	33.	Viswanathan, M. C., Blice-Baum, A. C., Schmidt, W., Foster, D. B. & Cammarato,
700	55.	A. Pseudo-acetylation of K326 and K328 of actin disrupts <i>Drosophila</i>
701		<i>melanogaster</i> indirect flight muscle structure and performance. Front Physiol 6,
		116 (2015).
703 704		110 (2013).
704	34.	Schindelin, J. et al. Fiji: An open-source platform for biological-image analysis.
703 706	54.	Nature Methods <b>9</b> , 676–682 (2012).
708		Nature Methods 9, 070–002 (2012).
707	35.	Weir, P. T. et al. Anatomical Reconstruction and Functional Imaging Reveal an
708	55.	Ordered Array of Skylight Polarization Detectors in <i>Drosophila</i> . <i>Journal of</i>
709		Neuroscience <b>36</b> , 5397–5404 (2016).
711		Neuroscience <b>30</b> , 5597–5404 (2010).
712	36.	Fayyazuddin, A. & Dickinson, M. H. Haltere afferents provide direct, electrotonic
	50.	input to a steering motor neuron in the blowfly, <i>Calliphora. J Neurosci</i> <b>16,</b> 5225–
713		5232 (1996).
714		5252 (1990).
715 716	37.	Horn, B. & Schunck, B. G. Determining Optical-Flow. Artificial intelligence 17,
	57.	185–203 (1981).
717 718		105–205 (1981).
	38.	Brox, T., Bruhn, A., Papenberg, N. & Weickert, J. High accuracy optical flow
719 720	50.	estimation based on a theory for warping. <i>Computer Vision - Eccv 2004, Pt 4</i>
720		<b>2034,</b> 25–36 (2004).
722 723	39.	Revaud, J., Weinzaepfel, P., Harchaoui, Z. & Schmid, C. DeepMatching:
723 724	53.	Hierarchical Deformable Dense Matching. Int J Comput Vis <b>120</b> , 300–323 (2016).
724		$\frac{1}{2010}$
725 726	40.	Boyd, S., Parikh, N., Chu, E., Peleato, B. & Eckstein, J. Distributed optimization
726	40.	and statistical learning via the alternating direction method of multipliers.
727		Foundations and Trends in Machine Learning <b>3</b> , 1–122 (2010).
728		$r$ oundations and mends in machine Learning $\mathbf{J}_{\mathbf{r}}$ $r$ $r$ $z$ (2010).
147		

730 731 732	41.	Unser, M., Soubies, E., Soulez, F., McCann, M. & Donati, L. GlobalBiolm: A unifying computational framework for solving inverse problems. in CTu1B.1 (OSA, 2017).
733		
734	42.	Sun, D., Roth, S. & Black, M. J. A quantitative analysis of current practices in
735		optical flow estimation and the principles behind them. Int J Comput Vis 106, 115–
736		137 (2014).
737		
738	43.	Guizar-Sicairos, M., Thurman, S. T. & Fienup, J. R. Efficient subpixel image
739		registration algorithms. Opt Lett 33, 156–158 (2008).
740		
741	44.	Weir, P. T. & Dickinson, M. H. Functional divisions for visual processing in the
742		central brain of flying Drosophila. Proc Natl Acad Sci USA 112, E5523-32 (2015).
743		
744	45.	Ikegaya, Y. et al. Synfire chains and cortical songs: temporal modules of cortical
745		activity. Science <b>304,</b> 559–564 (2004).
746		