Locomotor and olfactory responses in dopamine neurons of the *Drosophila* superior-lateral brain

**Highlights**
- Specific dopamine neurons connect olfactory regions to locomotor networks
- Activity in these dopamine neurons is locked to spontaneous changes in walking speed
- Some of these cells respond to odors, but repeated stimuli suppress their responses
- The same cell can encode variable sensory-motor combinations in different individuals

**Authors**
Michael Marquis, Rachel I. Wilson

**Correspondence**
rachel_wilson@hms.harvard.edu

**In brief**
Marquis and Wilson describe the physiology of three types of dopamine neurons in the *Drosophila* brain. They show that all these neurons are correlated with the fly’s walking speed, and some also respond to odor stimuli. The same identifiable dopamine neuron can encode different combinations of locomotion and odor in different individuals.
Locomotor and olfactory responses in dopamine neurons of the *Drosophila* superior-lateral brain

Michael Marquis¹ and Rachel I. Wilson¹,²,*

¹Department of Neurobiology, Harvard Medical School, 220 Longwood Avenue, Boston, MA 02115, USA
²Lead contact
*Correspondence: rachel_wilson@hms.harvard.edu
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SUMMARY

The *Drosophila* brain contains about 50 distinct morphological types of dopamine neurons.¹⁻⁴ Physiological studies of *Drosophila* dopamine neurons have been largely limited to one brain region, the mushroom body,⁵⁻¹³ where they are implicated in learning.¹⁴⁻¹⁸ By comparison, we know little about the physiology of other *Drosophila* dopamine neurons. Interestingly, a recent whole-brain imaging study found that dopamine neuron activity in several fly brain regions is correlated with locomotion.¹⁹ This is notable because many dopamine neurons in the rodent brain are also correlated with locomotion or other movements²⁰⁻³⁰; however, most rodent studies have focused on learned and rewarded behaviors, and few have investigated dopamine neuron activity during spontaneous (self-timed) movements. In this study, we monitored dopamine neurons in the *Drosophila* brain during self-timed locomotor movements, focusing on several previously uncharacterized cell types that arborize in the superior-lateral brain, specifically the lateral horn and superior-lateral protocerebrum. We found that activity of all of these dopamine neurons correlated with spontaneous fluctuations in walking speed, with different cell types showing different speed correlations. Some dopamine neurons also responded to odors, but these responses were suppressed by repeated odor encounters. Finally, we found that the same identifiable dopamine neuron can encode different combinations of locomotion and odor in different individuals. If these dopamine neurons promote synaptic plasticity—like the dopamine neurons of the mushroom body—then, their tuning profiles would imply that plasticity depends on a flexible integration of sensory signals, motor signals, and recent experience.

RESULTS

Dopamine neuron morphology and connectivity

The mushroom body is a site of olfactory learning in *Drosophila*, whereas the lateral horn (LH) has long been assumed to mediate innate olfactory behaviors.²¹ However, like the mushroom body, the LH is innervated by multiple types of dopamine neurons,²² suggesting that dopamine may modulate olfactory innate olfactory behaviors.²¹ However, like the mushroom body, the LH is innervated by multiple types of dopamine neurons,²² suggesting that dopamine may modulate olfactory processing in this region, similar to its role in the mushroom body. To better understand dopamine's function in the LH, we investigated three dopamine neuron types (PPM1/2-1, PPL2-1, and PPL2-3). Collectively, these neurons tile the full volume of the LH, together with the superior-lateral protocerebrum (SLP), an adjacent olfactory region.

The hemibrain connectome³³ contains two PPM1/2-1 neurons, one PPL2-1 neuron, and one PPL2-3 neuron in the right hemisphere. Each of these neurons arborizes in several brain regions (Figures 1A–1F), without an obvious division of the cell into axon and dendrite (Table S1). Each of these neurons has a large number of synaptic partners (Figures 1J and 1K; Table S2), but their connectivity is nonetheless selective (Figure S1). For example, the connectome shows that PPM1/2-1 neurons (Figure 1G) receive input from an ascending neuron that carries locomotor signals from the ventral nerve cord,³⁴ and they provide output to a specific descending neuron that projects to the leg-control regions of the ventral nerve cord.³⁵,³⁶ They are also reciprocally connected with the mushroom body output neuron that has the strongest direct connections to descending neurons (MBON20).³⁷ Finally, they are reciprocally connected with specific olfactory local neurons in the antennal lobe. This connectivity pattern predicts that PPM1/2-1 activity is recruited by locomotion and odor.

Meanwhile, PPL2-1 neurons (Figure 1H) receive input from MBON20 as well as several other MBONs. They are reciprocally connected to several LH centrifugal neurons³⁶ as well as LH output neurons. Interestingly, they are also reciprocally connected to an octopaminergic neuron which is downstream of locomotor-related neurons in the central complex.³⁸,³⁹ Finally, they send output to several types of Kenyon cells in the mushroom body calyx. This connectivity pattern predicts that PPL2-1 is recruited by odor and locomotion.

Finally, PPL2-3 neurons (Figure 1I) are downstream of LH centrifugal neurons as well as LH output neurons. They also receive direct input from olfactory projection neurons of the antennal lobe. They send output to mushroom body Kenyon cells in the dorsal accessory calyx.³⁷ This connectivity pattern predicts that PPL2-3 is recruited mainly by sensory input.

Dopamine neuron locomotor responses

The distinct patterns of connectivity in these dopamine neurons motivated us to ask whether they also have distinct functional...
**Figure 1. Dopamine neuron morphology and connectivity**

(A–C) Morphology of each dopamine neuron type in the hemibrain:v1.2 dataset, with outlines of key brain regions. For brain region abbreviations, see Table S2 or https://neuprint.janelia.org/.

(D–F) Number of pre- and post-synaptic connections formed by each dopamine neuron type in every brain region that contains at least 100 synapses, sorted by input/output ratio. Synapse counts are averaged for the two PPM1/2-1 neurons.

(G–I) Selected synaptic partners of each dopamine neuron type.

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properties. Therefore, we expressed GCaMP in each cell type and monitored their activity using two-photon imaging while flies walked on a spherical treadmill (Figure 2A). We found that the activity of all three cell types increased just after the fly started to walk (Figures 2B–2D). These locomotor onset signals were significantly larger in PPM1/2-1 compared with the other two cell types (Figure 2C).

Moreover, these dopamine neurons continued to be active as flies walked, and we noticed that they tracked the spontaneous fluctuations in the fly’s walking speed (Figures 2E and 2F). Overall, the relationship between neural activity (ΔF/F) and walking speed was significantly stronger for PPM1/2-1 compared with the other two cell types (Figures 2G and S2). Changes in neural activity lagged changes in speed by about 300 ms (Figure S2), suggesting that these neurons are responding to speed changes rather than driving those changes. Note that LH/SLP dopamine neurons seem to be specifically locked to locomotion, as they are not active during non-locomotor leg movements (Figure S3). It is interesting that PPL2-1 and PPL2-3 have equally strong locomotor responses, given that the major inputs to PPL2-3 do not arise from brain regions generally associated with locomotion (Figure 1I).

Interestingly, we observed considerable individual variation in locomotor responses. In particular, PPM1/2-1 neurons had strong locomotor responses in some individuals, whereas in other individuals, they had much weaker responses (Figures 2D, 2G, and S2). Thus, locomotor input to these neurons appears to be flexible.

**Dopamine neuron odor responses**

Next, we investigated whether these dopamine neurons respond to odor stimuli (Figure 3A). We focused on appetitive stimuli (ethanol and vinegar) because some of the neurons in these dopamine cell clusters have been linked to appetitive behaviors.40–43 Unexpectedly, we found that the cell type with the strongest locomotor-related activity (PPM1/2-1) almost never responded to these odors, either in whole-cell recordings (Figure 3A) or in imaging experiments from locomoting flies (Figure 3B). Only in one experiment did this cell type respond to odor, and this response was relatively weak (Figure 3B). By contrast, the second cell type (PPL2-1) was consistently excited by odor. This was true in whole-cell recordings (Figure 3C) as well as imaging experiments in locomoting flies (Figure 3D). The last cell type, PPL2-3, also responded to odors: in whole-cell recordings, when flies were quiescent, it was typically inhibited at odor onset and excited at odor offset (Figure 3E). However, in calcium imaging experiments in locomoting flies, PPL2-3 odor responses had more diverse dynamics, with more individual variation and odor-dependent variation (Figures 3G and 3F).

These results illustrate how difficult it can be to predict a cell’s tuning based on connectome data. On one hand, PPM1/2-1 is generally unresponsive to odors, although its top input is an olfactory mushroom body output neuron (MBON20).37 On the other hand, PPL2-3 has highly individualized odor responses, although it receives almost no direct MBON input, and MBONs are the cells within the olfactory system where individual variations have been most clearly documented.11,44 Thus, the anatomical inputs to these dopamine neurons might be regarded as flexible rather than fixed.

**Locomotor and olfactory contributions to dopamine neuron dynamics**

Thus far, we have seen that dopamine neuron activity can depend on both locomotion and odor. To quantify the relative contributions of these factors, we fit a linear regression to each cell’s activity pattern. These models allowed us to compactly summarize and compare the properties of different cells, both within and across cell types.

First, we found that the fly’s walking speed explained a remarkable amount of the variance in PPM1/2-1 activity (Figures 4A and 4B), with $R^2$ values approaching 0.6 in many individuals. Including the odor stimulus in the linear regression produced no improvement in the fit, except in the one individual that had obvious odor responses. This result supports our conclusion that PPM1/2-1 dopamine neurons are primarily locomotor-related.

Next, we turned to the two other dopamine neuron types (PPL2-1 and PPL2-3), whose properties are more complicated. Both these cell types respond to combinations of walking speed and odor, with the latter cell type often showing both excitation and inhibition in response to a single odor pulse. Interestingly, in some individuals, we noticed that the odor responses of these cells were suppressed by repeated odor presentations (Figures 4C–4E). We could accelerate this suppression by switching to a high odor pulse rate (Figures 4C and 4D).

To model the odor responses of these cells, we first computed each cell’s average response to an odor pulse. Then, we caused odor responses to decrement by a certain fraction with every odor pulse, recovering with an exponential time course; the fractional decrement per pulse was fit as a free parameter in each cell, with uniform recovery dynamics for all cells. Adding this type of odor response suppression to the model allowed us to capture much of the odor response variation in many cells (Figures 4F and 4G).

We also noticed that the onset of the high odor pulse frequency caused a downward shift in baseline fluorescence in some experiments (Figure 4D). We therefore augmented the model with subtractive inhibition that appears with each odor pulse and decays exponentially over time. Adding this component further improved the model fit in some cells (Figures 4F and 4G).

Interestingly, we found that dopamine neuron odor responses evolved with different dynamics in different individuals. In some individuals, we found clear evidence of odor response suppression and/or subtractive inhibition.

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(J) Number of upstream cell types versus total number of input synapses for each dopamine neuron type, along with a variety of other cell types for comparison. Each point represents a single neuron and is colored-coded by cell type.

(K) Equivalent plot for downstream connectivity. See also Figure S1 and Tables S1 and S2.
whereas there was no indication of suppression or inhibition in other individuals. Our results suggest that all these dopamine neurons have access to speed and odor information, but these variables are weighted differently in different cell types and individuals.

**DISCUSSION**

Here, we describe three types of dopamine neurons that collectively tile the superior-lateral brain. They are all part of the ‘TH-C’ group, whose activity promotes food-dwelling...
behavior\(^{40}\) and causes females to choose sucrose-rich locations for egg-laying.\(^{41}\) In particular, the PPL2 cluster within the TH-C group facilitates odor responses in mushroom body Kenyon cells\(^{45}\) and promotes male courtship behavior,\(^{42,43}\) which has a large olfactory component. Together, these findings suggest a general role in appetitive behaviors.

Notably, we show that one of these cell types (PPM1/2-1) is highly correlated with rapid spontaneous fluctuations in walking speed. This is reminiscent of some dopamine neurons in the Drosophila mushroom body lobes, which are correlated with locomotion or leg movement.\(^{5,11,12,13}\) Similarly, in the rodent brain, some dopamine neurons correlate with forward acceleration of the body or rotational velocity of the head.\(^{20-30}\)

Figure 3. Dopamine neuron odor responses
(A) Spike rasters showing responses of one example PPM1/2-1 neuron to three different odors and a solvent control (paraffin oil) during whole-cell recording. Shading indicates odor delivery (cVA, cis-vaccenyl acetate).
(B) PPM1/2-1 GCaMP responses (\(\Delta F/F\)) to ethanol or vinegar. Each trace shows the trial-averaged response for one individual (6 flies total). Only trials when the fly was not walking were included.
(C and D) Same but for PPL2-1. Each trace in (D) is from a different fly \((n = 15\) total).
(E) Same but for PPL2-3. Each trace in (F) is from a different fly \((n = 29\) total).
(G) Trial-averaged responses of four PPL2-3 neurons to two different odors presented in interleaved trials. Each pair of traces shows data from an individual fly.
Figure 4. Locomotor and olfactory contributions to dopamine neuron dynamics

(A) One example PPM1/2-1 imaging session. Walking speed was used to predict ΔF/F.

(B) Variance explained by linear models fit to PPM1/2-1 for each experiment (adjusted R²). Black line is the mean (n = 8 flies). Open circle is the example from (A).

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The dopamine neurons we describe here track locomotor speed with a lag; this suggests that these neurons are not controlling locomotor speed. Rather, these neurons may promote learning about the consequences of speed changes. For example, dopamine could serve as a “when-to-learn” signal that modifies the rate of associative synaptic plasticity. Locking dopamine to locomotion could be a way to accelerate learning when the organism is moving through the environment, and thus, new information is likely to be available (unpublished data).

We also found odor responses in some of these dopamine neurons, akin to the odor responses in many mushroom body dopamine neurons. Notably, we found that odor responses in PPL2-1 and PPL2-3 neurons were suppressed by repeated odor encounters. The same type of suppression has been shown previously for a specific mushroom body dopamine neuron (PPL1-4, also known as PPL-α3). Because odor response suppression is quite variable across individuals, it is unlikely to be purely due to peripheral adaptation. Interestingly, some dopamine neurons in the mammalian brain also respond preferentially to unexpected or novel stimuli. If dopamine functions as a generalized when-to-learn signal, it is logical that dopamine should be linked to novelty. Experience-dependent suppression of dopamine neuron sensory responses may be due to feedback loops that allow these dopamine neurons to monitor the activity of their target cells.

A notable feature of these neurons is their inter-individual variability. This stands in contrast to many other neurons in the LH and/or SLP, which tend to have fairly stereotyped functional properties. Rather, it is more similar to physiology of mushroom body dopamine neurons, which have plastic odor responses.

In the future, it will be interesting to determine how these dopamine neurons influence appetitive olfactory behaviors. Our results predict that the learned aspects of these behaviors will depend on the fly’s locomotor state: faster locomotion should promote more dopamine release and thus faster learning. This might be a way to facilitate learning when a fly is exploring a new environment.

**STAR Methods**

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**SUPPLEMENTAL INFORMATION**

Supplemental information can be found online at https://doi.org/10.1016/j.cub.2022.11.008.

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**AUTHOR CONTRIBUTIONS**

M.M. performed all experiments and all analyses. M.M. and R.I.W. designed the study and wrote the manuscript.

**DECLARATION OF INTERESTS**

The authors declare no competing interests.

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STAR METHODS

KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Experimental models: Organisms/strains** | | |
| P[w+mC]=ple-Gal4,F3 | Bloomington Drosophila Stock Center (BDSC) | RRID: BDSC_8848 |
| PBac[y+17.7[w+mC]=20XUAS-IVS-jGCaMP7f]VK00005 | BDSC | RRID: BDSC_79031 |
| 76F02-AD #B (attP40) | Mark Wu (Johns Hopkins University) | N/A |
| TH-D-AD #1 | Mark Wu (Johns Hopkins University) | N/A |
| 55C10-DBD #1 (attP2) | Mark Wu (Johns Hopkins University) | N/A |
| 61H03-DBD #1 (attP2) | Mark Wu (Johns Hopkins University) | N/A |
| TH-C-Gal4 | Mark Wu (Johns Hopkins University) | N/A |
| TH-G-KZ #1 | Mark Wu (Johns Hopkins University) | N/A |
| P(20XUAS-IVS-mCD8::GFP)attP2 | Gerry Rubin (Janelia) | RRID: BDSC_32194 |
| P(20XUAS-IVS-Syn21-opGCaMP6sp10)su(Hw)attP5 | Gerry Rubin (Janelia) | FlyBase: FBti0195669 |
| P(20XUAS-IVS-Syn21-opGCaMP6sp10)su(Hw)attP1 | Gerry Rubin (Janelia) | N/A |
| pJFRC7-20XUAS-cyRFP]VK000037 | Tom Clandinin (Stanford University) | N/A |
| **Deposited data** | | |
| Drosophila hemibrain v.1.2.1 | Scheffer et al. | https://neuprint.janelia.org |
| **Software and algorithms** | | |
| ScanImage 2018 | Vidrio Technologies | N/A |
| FicTrac | Moore et al. | http://rjdmoore.net/fictrac/ |
| Matlab R2016a, R2017b, and 2019b | MathWorks | N/A |
| NoRMCorre | N/A | https://github.com/flatironinstitute/NoRMCorre |
| R | R Foundation for Statistical Computing, Vienna, Austria | https://www.r-project.org/ |
| neuprint | Plaza et al. | https://neuprint.janelia.org |
| ImageJ 1.52 | Schneider et al. | https://imagej.nih.gov/ij/ |
| Data analysis code | This paper | https://github.com/wilson-lab |

RESOURCE AVAILABILITY

Lead contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Rachel Wilson (rachel_wilson@hms.harvard.edu).

Materials availability
This study did not generate new unique reagents.

Data and code availability

- Access to the imaging data, electrophysiology data, or behavioral data is available from the lead contact upon request. This study analyzes existing, publicly available connectome data; accession information for this dataset is detailed below (Cell type accession) and in the key resources table.
- Original code has been deposited in a Zenodo repository and is publicly available as of the date of publication, indexed in the key resources table.
Although this labeling was not noted in a previous report (where PPL2-3 is called "SLP"), we used this line to drive GCaMP expression in PPL2-3 neurons. This line consistently labels PPL2-3, PPL2c, and PPL1 clusters. We used this line to drive GCaMP expression in PPM1/2-1 and PPM1/2-3 neurons. (Note that PPM1/2-3 neuron targeted by this driver that arborizes in the central lateral horn. Therefore, by analyzing fluorescence specifically in the central lateral horn, we could use this line to measure activity in PPL2-1 neurons.

For imaging from PPM1/2-1 in Figures 2 C, 2D, 2F, 2G, 3D (responses to EtOH), 4C, 4E, S2, S3A, and S3B:

\[\text{w}^+ / \text{w}^-; \text{TH-C-Gal4} / \text{PBac}[^{17.7}w%^{+mC}=20XUAS-IVS-jGCaMP7f]VK00005\]

We found that the split-Gal4 combination TH-D-AD #1; 61H03-DBD #1 (attP2) targets several dopaminergic neurons in the PPM2, PPL2c, and PPL1 clusters. We used this line to drive GCaMP expression in PPM1/2-1 and PPM1/2-3 neurons. (Note that PPM1/2-3 neurons appear only in Figure S4 B.)

We found that the split-Gal4 combination TH-D-AD #1; 61H03-DBD #1 (attP2) targets several dopaminergic neurons in the PPL2ab and PPL1 clusters. We used this line to drive GCaMP expression in PPL2-3 neurons. This line consistently labels PPL2-3, although this labeling was not noted in a previous report (where PPL2-3 is called “SLP”).

TH-C-Gal4 targets a large subset of dopamine neurons. One of the cells targeted by this driver is PPL2-1, and this is the only neuron targeted by this driver that arborizes in the central lateral horn. Therefore, by analyzing fluorescence specifically in the central lateral horn, we could use this line to measure activity in PPL2-1 neurons.

These genetic driver lines allowed us to isolate GCaMP signals from individual cell types by analyzing regions of interest (ROIs) that were specific to the neuron in question. We found that different arborizations within the same neuron generally had highly correlated activity, and so we can use an ROI targeted to one arbor to infer the activity of the cell as a whole.

\[\text{w}^+ / \text{w}^-; \text{TH-D-AD #1} / \text{PBac}[^{17.7}w%^{+mC}=20XUAS-IVS-jGCaMP7f]VK00005\]

We used this line to drive GCaMP expression in PPM1/2-1 and PPM1/2-3 neurons. (Note that PPM1/2-3 neurons appear only in Figure S4 B.)

We found that the split-Gal4 combination TH-D-AD #1; 61H03-DBD #1 (attP2) targets several dopaminergic neurons in the PPL2ab and PPL1 clusters. We used this line to drive GCaMP expression in PPL2-3 neurons. This line consistently labels PPL2-3, although this labeling was not noted in a previous report (where PPL2-3 is called “SLP”).

\[\text{w}^+ / \text{w}^-; \text{TH-D-AD #1} / \text{PBac}[^{17.7}w%^{+mC}=20XUAS-IVS-jGCaMP7f]VK00005\]

For imaging from PPL2-3 (Figures 2C, 2D, 2F, 2G, 3C, 3E, and S3):

\[\text{w}^+ / \text{w}^-; \text{TH-C-Gal4} / \text{PBac}[^{17.7}w%^{+mC}=20XUAS-IVS-jGCaMP7f]VK00005\]

For imaging PPL2-1 in Figure 3 D (responses to vinegar) and Figure S3 B:

\[\text{w}^+ / \text{w}^-; \text{TH-C-Gal4} / \text{PBac}[^{17.7}w%^{+mC}=20XUAS-IVS-jGCaMP7f]VK00005\]

Fly preparation and dissection

All experiments were performed on female flies 12-24 hours (for electrophysiology) or 24-48 hours (for imaging) post-eclosion, except for a small number of imaging experiments which used slightly older flies (48-96 hours post-eclosion). In some imaging experiments (4 of 8 PPM1/2-1 experiments and 19 of 48 PPL2-3 experiments), flies were deprived of food (but not water) for 12-24 hours prior to the experiment to promote walking behavior. Because we had a large number of starved and also non-starved flies for PPL2-3 experiments, we were able to perform a quantitative comparison of locomotor responses and odor responses in starved versus non-starved flies for this cell type; this analysis showed no systematic effect of starvation. No circadian restriction was imposed for the experiments, we were able to perform a quantitative comparison of locomotor responses and odor responses in starved versus non-starved flies for PPL2-3 experiments (4 of 8 PPM1/2-1 experiments and 19 of 48 PPL2-3 experiments), flies were deprived of food (but not water) for 12-24 hours prior to the experiment to promote walking behavior. Because we had a large number of starved and also non-starved flies for PPL2-3 experiments, we were able to perform a quantitative comparison of locomotor responses and odor responses in starved versus non-starved flies for this cell type; this analysis showed no systematic effect of starvation. No circadian restriction was imposed for the experiment.
with 95% O₂, 5% CO₂. External solution was continuously perfused over the brain during electrophysiology and until just before the start of imaging acquisitions.

**METHOD DETAILS**

**Patch-clamp recordings**

Patch pipettes made from filamented borosilicate glass (OD 1.5, ID 0.86 mm, Sutter) using a P-97 Sutter puller and then fire-polished using a microforge (ALA Scientific Instruments) to achieve a final resistance of 8-12 MOhm. The internal solution contained (in mM): 140 potassium aspartate, 10 HEPES, 4 MgATP, 0.5 Na3GTP, 1 EGTA, 1 KCl and 15 neurobiotin citrate (Vector labs), filtered through a 0.22 μm filter. To visualize cells for recording we used a CMOS camera (GS3-U3-51S5M-C, FLIR) mounted on an upright compound microscope (Olympus BX51WI) with a 40x water immersion objective (UPlanFL, Olympus). We used a 100-W Hg arc lamp (U-LH100HG, Olympus) and an eGFP longpass filter (U-N41012, Chroma) to detect GFP fluorescence. The fly was illuminated from below using bright-field transmitted light through the microscope condenser. Recordings were obtained using an Axopatch 200B amplifier and a CV-203BU headstage (Molecular Devices). Voltage signals were low-pass filtered at 5 kHz before acquiring at 20 kHz on a National Instruments USB-6343 DAQ. To counteract leak currents caused by the recording electrode, at the start of each experiment we applied a constant hyperpolarizing current that lowered the membrane potential by ~5mV; this hyperpolarizing current was then applied constantly for the full duration of the recording. The liquid junction potential was corrected post hoc by subtracting 13 mV from recorded voltages. In electrophysiological experiments, we recorded one cell per fly.

**Two-photon calcium imaging**

Imaging experiments were performed using a galvo-resonant two-photon microscope with a movable stage (Thorlabs Bergamo II), a 20x/1.0 NA objective (XLUMPFLN20XW, Olympus), and a fast piezoelectric objective scanner (Physik Instrumente P725) for volumetric imaging. We used a Chameleon Vision-S Ti-Sapphire femtosecond laser tuned to 940 nm for two-photon excitation of GCaMP. Emission fluorescence was collected on a GaAsP PMT (Hamamatsu) through a 525-nm bandpass filter (MDF-QUAD2, Thorlabs). We used ScanImage 2018 software (Vidrio Technologies) and custom Matlab scripts to control the microscope and collected the imaging data using National Instruments PXIe-6341 hardware.

In most experiments the imaging region was 256×128 pixels, with 12 slices in the z-axis for each volume (6-12 μm per slice) resulting in a 6-7 Hz volumetric scanning rate. In a few cases (namely the PPL2-1 imaging experiments in Figures 2C, 2D, 2F, 2G, 3D (EtOH only), 4C, 4E, and S2), we used a volume consisting of 4-6 512×256 imaging planes instead, with approximately the same volume rate.

**Measurement of fly locomotion**

The fly stood on a 9-mm ball made of white foam (FR-4615, General Plastics) painted with black shapes. The ball floated above a 3D printed plenum made of clear or white acrylic (Autotiv). Medical-grade breathing air flowed into the base of the plenum and out into a hemispherical depression to suspend the ball while allowing it to rotate freely. The ball was illuminated by two round boards of 36 IR LED lamps (SODIAL). The movement of the ball was tracked at either ~60 Hz (PPL2-1 data in Figures 2C, 2D, 2F, 2G, 3D (EtOH only), 4C, 4E, and S2) or 25 Hz (all other experiments) using a CMOS camera (CM-3-U3-1S2M-CS, FLIR) fitted with a Tamron 23FM08L 8-mm 1:1.4 macro zoom lens and an 875 nm shortpass filter (Edmund Optics #86-106) to block out the two-photon laser. Machine vision software (FicTrac) converted the image of the ball to an estimate of the ball’s position in all three axes of rotation as well as the fly’s fictive 2D position at each time point, and also recorded video of the fly for use in identifying grooming behavior.

**Delivery of odor stimuli**

In electrophysiology experiments, a stream of medical breathing air (2.5 L/min) was passed through an activated carbon filter and directed at the fly through a carrier tube (4 mm inner diameter) positioned ~1 cm from the front of the fly. A portion of this air stream (12 mL/min) was diverted away from the carrier and directed by 3-way solenoid valve into the headspace of a clean 1.5 mL vial containing 200 μL of a 1:100 solution of odor in paraffin oil, or else paraffin oil alone. The solenoid normally directed the odor stream to the empty vial and switched airflow into the odor vial after receiving a command. Valve commands were sent from Matlab via a digital line using a USB-6343 DAQ (National Instruments). After passing through either vial, the odor stream joined the carrier stream again. Paraffin oil was stripped of low–molecular weight volatiles by storing it under negative pressure for at least several days before use.

In most imaging experiments, clean air was passed through an activated carbon filter and directed at the fly through a 19g steel tube placed ~5 mm in front of the fly at 20 mL/min. A solenoid valve switched between this and a parallel air stream that had passed through the headspace of a 1.5 mL vial containing a 1:100 dilution of odor in paraffin oil. CO₂ was presented using a similar system, but diluting the CO₂ in a stream of clean air instead of passing it through an odor vial. In a subset of imaging experiments (PPL2-1 imaging experiments in Figures 2C, 2D, 2F, 2G, 3D (EtOH only), 4C, 4E, and S2), a wider 4.5 mm odor delivery tube and undiluted ethanol odor stimulus at 12 mL/min were used. Photo-ionization detector (PID) measurements (Aurora Scientific) from the position of the fly confirmed that the intensity and temporal dynamics of the two odor delivery methods were roughly equivalent.
Data analysis was performed using Matlab R2016a, R2017b, and 2019b, (MathWorks) and R 3.6.0; the details are described below. Some initial processing of calcium imaging and behavior video data was parallelized on a high-performance computing cluster (Harvard Medical School Research Computing O2 cluster).

**Calcium imaging alignment and processing**
Rigid motion correction in the x- and y-axes was performed for each acquisition using the NoRMCorre algorithm. Volumetric ROIs were defined by combining 2D ROIs drawn in multiple imaging planes. Fluorescence values were determined by averaging all pixels in the volumetric ROI. Baseline fluorescence (the denominator F in ΔF/F) was calculated using either the average fluorescence during the pre-stimulus or pre-movement baseline period (Figures 2B–2D, 3B, 3D, 3F, and S3), the 5th percentile of fluorescence values for an ROI throughout the entire experiment (Figures 2E–2G and S2), or the average the 5th percentile of fluorescence values within a sliding 60-second window centered on each time point (Figure 4). For display purposes, ΔF/F traces in all figures were smoothed with a Gaussian kernel 3 samples wide.

**Electrophysiology spike detection**
In whole-cell recording data, spikes were identified by setting a threshold on the second derivative of the recorded voltage signal.

**Morphology and connectomics analysis**
All connectomic data were obtained from the hemibrain:v1.2 dataset. Synapse counts in each brain region (Figures 1D–1F) were obtained by downloading the data from neuprint using a custom Neo4j Cypher query. Example neuron morphologies (Figure 1A) were created by capturing a 2D projection of each neuron from the hemibrain:v1.2 dataset and converting it to a silhouette using ImageJ. Segregation indices (Table S1) were calculated in R using the flow_centrality() function from the network toolbox) in each of two ROIs around the fly and the spherical treadmill, respectively, in the video of the fly’s behavior. All grooming epochs (Figures 2C, 2D, and S3) were de-
experiment. The kernel was then convolved with the onset of each odor stimulus to generate an estimated odor response term. To model odor-dependent subtractive inhibition, the time-integrated history of odor delivery for each time point was also calculated by integrating the total time in which the odor valve was open in the past 240 sec multiplied by a difference of exponentials function with an onset time constant of 0.5 sec and a decay time constant of 175 sec.

\[
f(x) = e^{\frac{x}{C_0}} - e^{\frac{x}{C_1}}
\]

To model suppression of the odor response, we used an approach that has been used to model short-term synaptic depression.\textsuperscript{75,76} Specifically, the odor response term \(f_{\text{odor}}\) was multiplied by a constant scaling factor \(F\) (\(0 < F < 1\)) at the onset of each odor stimulus, recovering exponentially towards baseline with a time constant of 150 sec. The parameters fit for each model consisted of the adaptation scaling factor \(F\), an intercept term \(C\), and the coefficients for the linear terms of walking speed, modeled odor response (with suppression), and subtractive inhibition calculated from the integrated odor history.

\[
\hat{y} = \beta_1(f_{\text{odor}}) + \beta_2(\text{speed}) + \beta_3(\text{subtractive inhibition}) + C
\]

Model performance was evaluated and \(R^2\) values and 95\% confidence intervals for each model were calculated using a bootstrapping approach with an 80\%/20\% train/test split and \(n=1000\) iterations. Alternative models used for the comparisons in Figures 4E and 4F were obtained using the same protocol, but without either the subtractive inhibition term, the suppression of the odor response term, or both.

To quantify the suppressive effect of repeated odor pulses for each PPL2-1 and PPL2-3 experiment, we first quantified the predicted \(\Delta F/F\) due to locomotor speed, and we subtracted this from the measured \(\Delta F/F\) to account for the contribution of locomotion. Then, we measured the peak \(\Delta F/F\) during a 2-sec window after each of the first 5 odor pulses. Finally, we normalized these values by the magnitude of the first peak.