Differential Role for a Defined Lateral Horn Neuron Subset in Naïve Odor Valence in *Drosophila*

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Value coding of external stimuli in general, and odor valence in particular, is crucial for survival. In flies, odor valence is thought to be coded by two types of neurons: mushroom body output neurons (MBONs) and lateral horn (LH) neurons. MBONs are classified as neurons that promote either attraction or aversion, but not both, and they are dynamically activated by upstream neurons. This dynamic activation updates the valence values. In contrast, LH neurons receive scaled, but non-dynamic, input from their upstream neurons. It remains unclear how such a non-dynamic system generates differential valence values. Recently, PD2a1/b1 LH neurons were demonstrated to promote approach behavior at low odor concentration in starved flies. Here, we demonstrate that at high odor concentrations, these same neurons contribute to avoidance in satiated flies. The contribution of PD2a1/b1 LH neurons to aversion is context dependent. It is diminished in starved flies, although PD2a1/b1 neural activity remains unchanged, and at lower odor concentration. In addition, PD2a1/b1 aversive effect develops over time. Thus, our results indicate that, even though PD2a1/b1 LH neurons transmit hard-wired output, their effect on valence can change. Taken together, we suggest that the valence model described for MBONs does not hold for LH neurons.

In order to survive, animals must attach value to external stimuli, be it due to innate or learned behavior. In particular, the ability to accurately evaluate potential food resources is a critical trait for survival. To make such an assessment, animals use many senses, with olfaction often serving as a primary cue.

The *Drosophila* olfactory system resembles that of mammals, including our own, and uses similar principles to decode olfactory information1,2. Odors bind to olfactory receptor neurons (ORNs), which are located in the antennae and maxillary palps, where each ORN expresses a single type of odorant receptor (OR)3–5. All ORNs expressing the same OR converge onto the same region in the antennal lobe termed the glomerulus6–8. Second-order excitatory cholinergic projection neurons (ePNs) have dendrites that are restricted to a single glomerulus, whereas inhibitory GABAergic projection neurons (iPNs) are mostly multiglomerular9. Both PN types project to the lateral horn (LH), whereas only ePNs project to the calyx of the mushroom body (MB)9. Until recently, associative learning and memory processes were generally believed to occur in the MB, with innate behavior driven by the LH10,11. However, although the LH is still believed to contribute greatly to innate behavior, it has become apparent that the rigid functional distinction between the two neuropils cannot be upheld. There is now evidence that the MB also plays a role in some innate olfactory behaviors, mostly attractive12–14, while the LH is involved in some forms of associative memory15.

The LH compartment contains over 1300 cells that are categorized into over 150 types, each with individual morphology15. Cells that share morphological features are also more likely to share PN connectivity, although there is some variability17. Nine LH cell types could be distinguished by optogenetic activation to drive either attraction (3 cell types) or aversion (6 cell types)18. In the case of odor stimuli, effects on odor valence were demonstrated for only three types of LH neurons and under very specific conditions: I. AV1a1 LH neurons, which trigger aversion and are required for geosmin avoidance19 II. LH neurons, labeled by the R21G11- and R23C09-GAL4 driver lines, and which process CO2 avoidance20. III. PD2a1/b1 neurons (previously known as type I LH neurons21 or ML9 and ML817, respectively).

PD2a1/b1 neurons belong to the lateral horn output neurons (LHON)15. They have their somata in the lateral posterior dorsal protocerebrum, extend a short primary neurite towards the brain center and then bifurcate to...
connect their input regions in the LH (PD2a1/b1) and in the MB (PD2b1 only) with their presynaptic target areas in the superior intermediate protocerebrum (SIP) and superior medial protocerebrum (SMP) around the vertical MB stalk. About one third of input synapses in both LH and calyx derive from uniglomerular PNs, with another third provided by local LH neurons. In addition, reciprocal LHON input accounts for about 20%, and a varying amount of ipsi- and contralateral axoaxonic input in the SIP comes from the mushroom body output neuron (MBON)-\(\alpha 2sc\). PD2a1/b1 neurons were found to contribute to food odor approach at odor concentrations in the range of \(10^{-7}\) to \(10^{-5}\) dilution in starved flies. In addition, PD2a1/b1 neurons were also shown to be required for aversive conditioning and it was suggested that reduced activation of PD2a1/b1 neurons following aversive conditioning was responsible for the reduced odor approach. These observations are in agreement with current knowledge about learning and memory processes occurring at the MB and MBONs. Accordingly, MBONs are divided into neurons that drive either attraction or aversion, and plasticity between MB and MBONs shifts the balance between attraction and aversion for each odor. However, in contrast to the known plasticity of the synapse between MB neurons and MBONs, there is no information about any such similar plasticity between PNs and LH neurons. Furthermore, optogenetic activation of PD2a1/b1 neurons generated a mild aversion rather than attraction. In this context, odor valence can range from attractive to aversive, depending on the interplay of external factors (such as odor concentration) and internal states (such as satiation level), where a high odor concentration (even in the case of food odors) is usually associated with aversion. PN input to PD2a1/b1 neurons is positively correlated with odor concentration and as a result the activity of PD2a1/b1 neurons increases linearly with odor concentration. Thus, it is unclear how increased input to neurons which underlie attraction supports an overall aversive response to odors. This seemingly contradicts the notion that PD2a1/b1 neurons only drive attraction. We therefore examined whether PD2a1/b1 neurons always contribute to approach behavior or whether their role is context dependent. Thus, we examined the contribution of PD2a1/b1 neurons to behavior responses under broader conditions and in particular, higher odor concentrations and satiated flies. Here we show that PD2a1/b1 neurons, which were found to mediate an approach to odors in starved flies and at low odor concentrations, can also contribute to odor avoidance at higher odor concentrations in satiated flies. In accordance with these observations, the effects we saw on avoidance behavior are influenced not only by odor identity (there are differences even between odors with similar activation patterns) and odor concentration, but also by exposure time, and satiation. In particular, starvation abolished the contribution of PD2a1/b1 neurons to an aversive response to odors even though there was no effect on the PD2a1/b1 neuron odor responses.

Results
PD2a1/b1 neurons exhibit a spectrum of odor responses. Selecting suitable odors for behavioral experiments requires prior knowledge about the physiological responses of PD2a1/b1 neurons. We therefore used two GAL4 driver lines to characterize the response profile of PD2a1/b1 neurons. The first one, R37G11-GAL4, labels 6–7 PD2a1/b1 neurons and is a relatively clean line that is also suitable for behavioral experiments (Supplementary Fig. S1), while R48F03-GAL4 labels 18–22 PD2a1/b1 neurons and allows a larger coverage of the PD2a1/b1 population. Unfortunately, the R48F03-GAL4 driver line was also strongly expressed in additional neurons (e.g., in the subesophageal ganglion, the fan-shaped body, and the thoraco-abdominal ganglia). This line was therefore not suitable for all the characterizations and specifically could not be used for the behavioral experiments (Supplementary Fig. S1).

It was recently reported that the R37G11-GAL4 driver labels cholinergic lateral horn neurons. We confirmed these results and additionally examined whether the broader R48F03-GAL4 driver line also labels only cholinergic neurons. Indeed, ChAT-immunoreactivity exhibited an overlap with R37G11-GAL4 and R48F03-GAL4-labeled somata (Supplementary Fig. S1). Together with the complete lack of staining against dvlGlut and GABA in PD2a1/b1 neurons of both lines, it is assumed that all cells described here are homogeneously cholinergic (Supplementary Fig. S1).

Previous reports have shown that PD2a1/b1 neurons receive input from a small and stereotypic number of PNs. However, to what extent the population of all PD2a1/b1 neurons shows a correlated response to this stereotyped input was not fully examined. To investigate this issue, we examined the population response of PD2a1/b1 neurons in the same animal, the correlation between PD2a1/b1 neuron odor responses, and the percentage of neurons activated by any given odor in the two GAL4 lines. Odor responses were measured in PD2a1/b1 somata rather than at PD2a1/b1 axons in order to be able to record simultaneously from a number of neurons. Twenty odors (R37G11-GAL4) and a subset of 15 odors (R48F03-GAL4) were selected and delivered at a 5 x 10^-7 dilution (see Methods). Odors were chosen according to the Database of Odorant Responses (DoOR) to include odors predicted to activate PNs upstream to PD2a1/b1 neurons and, as controls, odors that are not predicted to activate the connected PNs.

PD2a1/b1 neurons exhibited varied response dynamics in their cell bodies. Some odors (e.g., geranyl acetate; Fig. 1a), elicited relatively rapid response kinetics, reaching peak response during the 5 second odor pulse. In contrast, other odors (e.g., 2-butanone; Fig. 1a), elicited relatively slow response kinetics, reaching peak response approximately 15 seconds after odor pulse onset, long after the end of the 5 second odor pulse. In addition, these prolonged odor responses often lasted throughout the subsequent 35 second recording. A similar variability was observed for neuron recruitment to an odor pulse, where certain odors activated almost all PD2a1/b1 neurons (e.g., 2-butanone, Fig. 1b), whereas others (e.g., ethyl benzoate, Fig. 1b) triggered a response in only a single neuron. Therefore, to better characterize the neuronal responses we analyzed five parameters: peak responses, percent of responding neurons, response persistence, time to peak, and the area under the curve (Fig. 1 and Supplementary Fig. S2, see methods).

Peak odor responses varied greatly between different odors. While some odors did not elicit any, or only a very mild, response, others produced very robust responses (Fig. 1c,g and Supplementary Fig. S2a,c). We found marked differences between the ranked order of odor responses elicited by a dilution of 2.5 x 10^-4 as was...
Figure 1. PD2a1/b1 neurons display variable odor responses to high odor concentrations. (a) Representative single odor response traces obtained from the same neuron. Different odor response dynamics and amplitudes were observed. Odor pulse is highlighted in red. (b) Representative activity maps demonstrating variable population odor responses in the same fly to 2-butanone and ethyl benzoate. Pre-odor pulse and during odor pulse activity maps are depicted. (c,g) Peak ∆F/F responses during odor response for GCaMP6f-labelled cells driven by drivers R37G11-GAL4 (c) and R48F03-GAL4 (g), respectively. (55 ≥ n ≥ 20 and 46 ≥ n ≥ 14 cells, respectively. At least 5 different flies were imaged from each genotype.) (d,h) Percentage of responding neurons obtained from all neurons from all flies pooled for cells labeled by R37G11-GAL4 (d) and R48F03-GAL4 (h). (55 ≥ n ≥ 20 and 46 ≥ n ≥ 14 cells, respectively. At least 5 different flies were imaged from each genotype.) (e,i) Response persistence for R37G11-GAL4 (e) and R48F03-GAL4 (i). Response persistence was measured as the inverse of the ratio between the maximal peak of the odor response and the average response 5 seconds after the maximal peak time. (55 ≥ n ≥ 20 and 46 ≥ n ≥ 14 cells, respectively. At least 5 different flies from each genotype were imaged.) (f,j) Heatmaps of normalized selected parameters: peak response, number of responding cells, area under the curve, time to peak, and response persistence for R37G11-GAL4 and R48F03-GAL4, respectively. To compare different odors across several parameters, we normalized the results by calculating a Z-score for each odor value. The calculated Z-scores were further divided by the maximal Z-score for each parameter to yield a maximum value of 1. In general, a high correlation was observed between the different parameters. (k–m) Correlation between the parameters obtained for R37G11-GAL4 and R48F03-GAL4 for peak ∆F/F (k), the percentage of responding neurons (l) and the response persistence (m). (See Supplementary Table S1 for statistical analysis).
previously described and those obtained by us for $5 \times 10^{-5}$ (Fig. 1c). For example, whereas a $2.5 \times 10^{-4}$ dilution of benzaldehyde, 4-methylcyclohexanol (MCH), and linalool evoked a relatively robust odor response compared to other odors examined, a $5 \times 10^{-4}$ dilution of these odors hardly activated any PD2a1/b1 neurons. Similar results were obtained for both the R37G11–GAL4 and R48F03–GAL4 driver lines, with a high correlation between the response amplitude in neurons labeled by the two lines (Fig. 1g,k and Supplementary Fig. S2).

To examine the percentage of neurons activated by each odor, we pooled all the neurons that responded to any of the odors (Fig. 1d). As with the peak amplitude, the responses spanned the entire range, from odors that triggered only a handful of neurons (i.e., 6% in the case of linalool), to those that activated all neurons (i.e., 100% in the case of ACV). Amplitude of odor responses, and the percentage of neurons responding, was tightly correlated (Fig. 1f) and Supplementary Fig. S2g,h). Again, a high correlation was observed between the percentage of neurons responding in the two driver lines (Fig. 1b,j,l). Although the response dynamics to different odors varied enormously, it was noticeable that odors with a longer time to peak and a high response persistence (Fig. 1a,e,f and Supplementary Fig. S2) also generated the strongest responses with respect to both amplitude and the number of activated neurons (Fig. 1e,f and Supplementary Fig. S2). However, it is important to note that strong responses were also observed to certain odors, such as geranyl acetate and ACV that had a relatively short time to peak and low response persistence. These response dynamics were also highly correlated between the two driver lines (Fig. 1i,m and Supplementary Fig. S2f). Taken together, PD2a1/b1 neurons exhibit varied responses to different odors. This heterogeneity was seen across all the criteria used for analysis with a high correlation found between the different criteria for a given odor (Fig. 1f) and Supplementary Fig. S2).

PD2a1/b1 neurons respond in a coordinated fashion. In order to investigate the population response of PD2a1/b1 neurons further, we first examined whether the PD2a1/b1 neurons in an individual fly respond in a coordinated fashion. To this end, we measured the correlation between the odor response vectors of each neuron to a given odor. When PD2a1/b1 neurons were covered by the R37G11–GAL4 driver line, there was a very high correlation between the neurons for a given odor (Fig. 2a,c). It should be noted that this was due to similar response dynamics, even though the amplitude of the response was very variable between the different neurons (Fig. 2a). Repeating this analysis for all odors and all flies, revealed an average correlation of 0.71 (Fig. 2d). We consider it possible that this high correlation between neurons was a consequence of the relatively low temporal accuracy of our recording (i.e., two-photon functional imaging) and that this method is not sensitive enough to capture subtle differences in response dynamics. We therefore also examined the correlation between different odor responses in the same neuron. As expected from the results presented in Fig. 1, there was no correlation in odor responses whatsoever (0.17 Fig. 2b–d). The stereotyped connectivity predicts that odor responses should also be correlated between flies and indeed, for a given odor, there was a high correlation (an average of 0.52, Fig. 2e,f) between the odor responses of all neurons of all flies (Fig. 2e,f). We took two approaches to ensure that this high correlation was not a consequence of our calculation method (see Methods). As a first measure, we reevaluated the correlation between the flies’ odor responses after shuffling the vectors. Since the correlation between different odors was ~0.2, we expected that this procedure would yield the same result and indeed, the shuffled correlation was ~0.2 (Fig. 2e,f). As a second measure, in order to avoid the possibility that we generated a false high correlation by our selection of groups of correlated vectors, we generated an artificial data set with the same number of neurons, flies, and correlations between neurons of the same fly, but with no correlation between flies (the low odor correlation was used as the baseline for correlation). Again, this approach validated our calculations as only the low odor correlation was generated (Fig. 2f). Taken together, our results indicate that the small group of PD2a1/b1 neurons covered by the R37G11–GAL4 driver line exhibit correlated and stereotyped odor responses. Similar results were obtained for the PD2a1/b1 neurons covered by the R48F03–GAL4 driver line, although both the within-fly odor response correlations and consequently, the relationships in between-fly stereotyped odor responses, were much weaker.

Recently, different compartments in Drosophila neurons were discovered to exhibit different odor responses. For example, the odor responses of Kenyon cell (KC) somata exhibit longer odor responses than KC dendrites, with the suggestion that these prolonged responses may hold the memory of a recently sensed odor. To ensure that our analysis of PD2a1/b1 somata captures the essence of PD2a1/b1 neuronal output, we compared the odor responses obtained at PD2a1/b1 somata and axons. For this purpose, we selected an odor subset spanning the whole range of observed odor peak responses and response kinetics (Fig. 1). As expected from the pseudo-unipolar morphology of invertebrate interneurons, odor responses in PD2a1/b1 somata were low-pass filtered (Supplementary Fig. S3a). Nevertheless, we also seemed to capture relatively rapid response dynamics, albeit at a lower magnitude. For example, 2-butanol elicited a biphasic response in axons, with a small and fast response on odor onset, followed by a much stronger and slower response following odor offset. This biphasic response, although heavily filtered, was still observed in PD2a1/b1 somata (Supplementary Fig. S3a). As expected, different odors elicited statistically different responses for all three parameters examined. However, for a given odor, the responses in the axonal compartment were only slightly stronger than those in the somata (except for ACV, Supplementary Fig. S3b,c). In contrast, the response dynamics were markedly shorter in the axons than in the PD2a1/b1 somata (Supplementary Fig. S3d–g). In conclusion, the responses recorded in the somata tended to be slower, but still followed the same kinetics observed in the axon compartment.

PD2a1/b1 neurons contribute to aversive responses to odors. Based on our results, we selected 12 odors that matched three odor categories: odors that elicited strong and prolonged neuronal responses (methyl acetate, 2-butanol, ethyl lactate, ethyl acetate), odors that elicited strong but short neuronal responses (ACV, geranyl acetate, hexyl acetate, 2-heptanone), or odors that generated only weak neuronal responses (3-octanol, 1-hexanol, ethyl benzoate, MCH) in PD2a1/b1 neurons. We then examined whether silencing PD2a1/b1
neuronal output by expressing UAS-Shibirets1 using the R37G11-GAL4 driver line, affected the valence attached to these odors at 10^{-2} odor dilution in satiated flies. For the behavioral experiments, flies are placed in linear chambers where they can walk back and forth only along one axis and odors are delivered from both sides of the chamber with a clear decision area in the middle of the chamber (Fig. 3a). The flies in this apparatus constantly walk around the chambers and thus are making repeated decisions (see methods). All odors examined at a 10^{-2} dilution, elicited either avoidance or neutral behavioral responses. For 7 out of the 12 odors examined: methyl acetate

Figure 2. PD2a1/b1 neurons exhibit correlated population activity. (a) Odor responses from four PD2a1/b1 cell bodies in the same fly to three odors as indicated. A high correlation between the temporal pattern of the responses was observed. (b) Odor responses from one PD2a1/b1 neuron to seven odors. The temporal dynamics of the responses are not correlated. (c) Left, an example of the correlation values of six PD2a1/b1 neurons in the same fly obtained for the seven odors on the right. Right, an example of the correlation values of seven odors obtained for the six neurons on the left. (d) Within-fly mean correlation values for both driver lines (R37G11-GAL4 and R48F03-GAL4) for neurons and odors. High correlation values were obtained for the neurons and low correlation values for the odors (n = 26 flies and n = 9 flies, respectively). (e) Left, an example of the correlation values for a single odor obtained for all PD2a1/b1 neurons from all flies. Right, an example of the correlation values when the neuronal responses were shuffled (see Methods). (f) Left, between-flies mean correlation values for both driver lines (R37G11-GAL4, and R48F03-GAL4, n = 26 flies; the number of neurons ranged from 20 to 55 and n = 9 flies; the number of neurons ranged from 14 to 46, respectively). Right, Between-flies correlation for shuffled neurons and for artificial neurons (see Methods).
acetate, 2-butanone, ethyl acetate, ACV, geranyl acetate, hexyl acetate, and 2-heptanone, which elicited strong
peak responses (Fig. 1), silencing PD2a1/b1 neurons caused a significant decrease in odor avoidance behavior
(Fig. 3, Supplementary Table S1, which contains detailed statistical comparisons and tests, including all controls,
for all figures). Interestingly, despite eliciting strong neuronal responses in PD2a1/b1 neurons, one odor (ethyl
lactate, Fig. 3) exhibited no change in odor valence. For the odors that elicit weak peak responses (3-octanol,
1-hexanol, ethyl benzoate, and MCH, Fig. 1), we did not observe any significant change in odor valence (Fig. 3).
However, it is important to note that these odors became less aversive, owing to the temperature increase, which
might mask possible effects on valence. Although the R37G11-GAL4 driver line had almost no label in the
thoracico-abdominal ganglia, we still verified that fly movement was not impaired in R37G11-GAL4 flies express-
ing UAS-Shibirets1. Walking speed analysis revealed no defects in the flies’ locomotion (Supplementary Fig. S4a).
In addition, we obtained similar behavioral results with a sparser split-GAL4 line, LH98915,18 (see methods) which
is based on the R37G11-ZpGdbd and R29G05-p65ADZp promoter fragments (Supplementary Fig. S4b). These
results suggest that PD2a1/b1 neurons mediate avoidance behavior to odors in the case of satiated flies at the
relatively high odor concentration of 10−2 used in this experiment.

Odor concentration modulates the effect of PD2a1/b1 neurons on behavioral output. This
avoidance behavior raises the question of whether the PD2a1/b1 effect on valence is concentration dependent.
Since PD2a1/b1 neuronal odor responses increase linearly with odor concentration21, we first verified that lower
odor concentrations do elicit a response. Two-photon functional imaging was used to examine the neuronal
responses of PD2a1/b1 neurons to varying concentrations of six odors for which silencing PD2a1/b1 neurons
resulted in a phenotypic change. As previously described21, the results indicated a concentration-dependent
increase in the odor response of PD2a1/b1 neurons to all odors examined (Fig. 4a,b). In most cases we observed
saturation of the odor response (with no difference between odor responses at 1 × 10−2 and 5 × 10−2). Two
exceptions were 2-heptanone and apple cider vinegar (ACV) where there remained a significant increase in
odor response with increasing odor concentration (Fig. 4b) even at the highest concentrations. In addition to
the quantitative changes in neuronal response to different odor concentrations, certain odors (2-butanone and

Figure 3. PD2a1/b1 neurons contribute to aversive responses to high odor concentrations. (a) An example
of traces obtained for naïve flies for 2-butanone. Gray, parental control groups; red, the experimental group
at 32 °C; blue, the experimental control group at 23 °C. Naïve response to an odor (gray) was tested against
mineral oil (white). (b) Mean valence scores (see Methods) from experiments as in a, for the designated odors.
Odors are arranged according to their response profile as in Fig. 2 and Supplementary Fig. S2. A consistent
shift towards more positive valence values was observed during PD2a1/b1 inhibition for all odors that activate
PD2a1/b1 neurons, except for ethyl lactate. No effect during PD2a1/b1 inhibition was observed for odors that
only weakly activate PD2a1/b1 neurons. (137 ≥ n ≥ 21 flies for all conditions, * indicates a significant difference
of the indicated group from all other groups according to a multiple comparison test. The lowest value of all
multiple comparison tests is presented. # indicates a significant difference only between the two indicated
groups. * , p < 0.05, ** , **p < 0.01, ***p < 0.0005, ****p < 0.0001, see Supplementary Table S1 for statistical
analysis).
ethyl acetate, Fig. 4a) exhibited a dramatic change in the temporal dynamics. Based on this characterization, we selected four odors whose valence was affected by silencing PD2a1/b1 neurons (Fig. 3), and which produced neuronal odor responses at lower concentrations (Fig. 4a,b). Silencing PD2a1/b1 neurons had no effect on odor avoidance behavior at any of the odor concentrations tested, except for $10^{-3}$ ACV, which resulted in a significant...
increase in avoidance behavior (Fig. 4c). This apparent lack of contribution by PD2a1/b1 neurons to odor valence could not be attributed to a lack of ability to detect the odors, as the flies could be conditioned against the low odor concentration (Supplementary Fig. S5). The disappearance of the PD2a1/b1 effect on odor valence at lower odor concentration was surprising because of the high neural activity observed in PD2a1/b1 neurons even at low concentrations (ACV, Fig. 4a,b). Indeed, pooling the behavioral results presented in Figs. 3 and 4c, revealed no correlation between the magnitude of odor response observed in PD2a1/b1 neurons and the extent of the effect seen on behavior (Fig. 4d). In conclusion, the contribution of PD2a1/b1 neurons to odor avoidance behavior is concentration dependent.

**Starvation modulates the effect of PD2a1/b1 neurons on behavioral output but not their odor response.** In addition to odor concentration, there are a number of other factors that may affect the neural activity and the contribution to odor valence by PD2a1/b1 neurons. One such factor is the satiation state of the flies. Starvation has been shown to enhance attraction to food odors as well as modulate the neural activity in various glomeruli. In particular, the activity in PNs innervating glomeruli DM1 and DM4 (which are presynaptic to PD2a1/b1 neurons) has been reported to increase following starvation. In order to investigate the effect of context as manifested by starvation on the contribution of PD2a1/b1 neurons to odor valence we first examined whether starvation affects PD2a1/b1 neuron odor responses. With one exception (time to peak with acetic acid, Fig. 3a,b and Supplementary Fig. S6), there was no significant difference in any of the parameters analyzed. We then selected the seven odors that were affected by silencing PD2a1/b1 neurons in satiated flies (Fig. 3), and examined the effect of PD2a1/b1 neurons on behavioral responses of starved flies at 10−2 odor concentration. Surprisingly, although starvation did not affect PD2a1/b1 neuronal odor responses, it completely suppressed the previously observed behavioral effects in fed flies, so that the valence values for all tested odors remained unchanged after silencing PD2a1/b1 neurons (Fig. 5c) compared to controls. In the case of 2-heptanone we also observed a strong temperature effect in the starved flies. The reason for this observation remains unknown although it could be associated with the reports that heat reduces olfactory sensitivity or that temperature can generate increased odor attraction. Taken together, our results indicate that context, as manifested by starvation, diminishes the contribution of PD2a1/b1 neurons to odor avoidance behavior but without modulating the neuronal odor responses in the somata.

**The effect of PD2a1/b1 neurons on odor valence develops with time.** The results obtained for starved flies and different odor concentrations indicate that the contribution of PD2a1/b1 neurons to odor avoidance behavior is varied and inconsistent. Another parameter that may affect the contribution of these neurons to odor avoidance is the temporal dynamics of the odor response. Since odor responses of ORNs and PNs are known to change over time, it was of interest to examine PD2a1/b1 neurons effect on valence with time. As our behavioral apparatus provides constant tracking of the flies, we were able to re-analyze the data at different time points. We first analyzed the velocity at the first exposure to the odor (positive values represent movement towards the odor source, see methods), and verified that this analysis indeed captures flies' preferences as previously reported. For attractive odors, such as ACV and geranyl acetate, we observed an initial movement towards the odor source, see methods), and verified that this analysis indeed captures flies' preferences as previously reported. For attractive odors, such as ACV and geranyl acetate, we observed an initial movement towards the odor source. The velocity of this movement was significantly reduced towards 2-heptanone and ethyl acetate, which generate aversion in our behavioral assay (Fig. 6a–c). When examined across all odors and genotypes, silencing PD2a1/b1 neurons had no effect on the initial preference to any odor that generated an effect after 120 seconds exposure (compare Figs. 3 and 6d). These results suggest that while PD2a1/b1 neurons do not affect the initial preference for an odor, an effect might develop over time. When we re-analyzed our data for the seven odors selected previously, at two intermediate time points (Fig. 6e), the results appeared to support this hypothesis. While 40 seconds exposure produced no significant effect on odor valence, after 80 seconds exposure, three of the seven odors examined did display a significant effect (Fig. 6e). Thus, the contribution of PD2a1/b1 neurons to odor valence is also dependent on the time of odor exposure. Odor conditioning has already been used to demonstrate the development of an effect of PD2a1/b1 neurons on odor valence with time, however, in that case, the time scale was measured in hours, as opposed to the current changes in naïve behavior, which occurred within seconds.

**PD2a1/b1 neuron output can interfere with aversive conditioning.** We have found that PD2a1/b1 neurons contribute to aversion even though they also receive input from glomeruli reported to contribute to attraction. We therefore sought to verify our results using another assay. To this end, we used classical olfactory conditioning (Supplementary Fig. S7a). In this assay, an odor is assigned a negative value by pairing it with an electric shock (conditioned odor, CS+) and is compared to another odor (unconditioned odor, CS−). Electric shock pairing adds a learned behavioral drive to the original naïve odor valence. If there is a strong naïve aversion to the CS−, it is possible that the learned aversion towards the CS+ will not be apparent above the naïve aversion. We reasoned that in such a case silencing PD2a1/b1 neurons might improve aversive conditioning by decreasing the naïve aversion of the CS−. Out of the odors we tested, only ethyl lactate elicited a profound odor response in PD2a1/b1 neurons (Fig. 1) without a change in valence after silencing (Fig. 3). We therefore selected ethyl lactate as the CS+ and 2-butanol as the CS− whose avoidance is decreased when PD2a1/b1 neurons are silenced (Fig. 3). Prior to conditioning flies robustly avoided the CS−, 2-butanol (Supplementary Fig. S7b) and, following conditioning, control flies still avoided the CS− (Supplementary Fig. S7b). Thus, the conflict between learned and naïve drives resulted in only a weak change in avoidance (Supplementary Fig. S7b). However, silencing PD2a1/b1 neurons, which reduces the CS− naïve aversion (Fig. 3), dramatically increased the change in avoidance, allowing them to better avoid the CS+ (Supplementary Fig. S7b). This enhanced conditioning capability was not due to a change in odor discrimination, since naïve flies showed no change in odor preference between 2-butanol and ethyl lactate even when PD2a1/b1 neurons were silenced (Supplementary Fig. S7b, pre-training avoidance).
Figure 5. Starvation abolishes the effect of PD2a1/b1 neurons on behavioral output without changing their odor responses. (a) Averaged traces of odor responses for fed (black) and starved (light blue) flies; no effect of starvation was observed. (b) Analysis of peak odor responses and response persistence for the designated odors in fed and starved flies. No statistical difference was observed (49 ≥ n ≥ 18 cells, from at least 5 different flies; see Supplementary Table S1 for statistical analysis). (c) Mean valence scores of starved flies for the designated odor. Gray, parental controls; light red, the experimental group at 32°C; blue, the experimental control group at 23°C. Starvation abolished the effect of silencing PD2a1/b1 neurons. (87 ≥ n ≥ 47 flies, *indicates a significant difference of the indicated group from all other groups according to a multiple comparison test. The lowest value of all multiple comparison tests is presented. # indicates a significant difference only between the two indicated groups. *p < 0.05, **p < 0.01, ***p < 0.0005, ****p < 0.0001, see Supplementary Table S1 for statistical analysis.).
Figure 6. The effect of PD2a1/b1 neurons on odor valence develops with time. (a) Heat maps of the initial walking velocity of single flies at the onset of the odor pulse, 3 seconds prior to the odor pulse, and 10 seconds following the odor pulse (black line). A positive velocity indicates movement towards the odor pulse. For the attractive odors (ACV and geranyl acetate), a clear positive velocity was observed for most flies. (b) Mean velocity of the heat maps presented in a. A clear movement towards the odor source was observed for the attractive odors. Odors are according to the color coding in panel a. (c) Analysis of the integral of the response for the flies presented in a for the first 3 seconds following odor onset. Odors are according to the color coding in panel a (**p < 0.01, see Supplementary Table S1 for statistical analysis). (d) Mean velocity scores during the first 3 seconds following odor onset for the designated odor. Gray, parental controls; red, the experimental group at 32 °C; blue, the experimental control group at 23 °C. No significant difference was observed (53 ≥ n ≥ 18 flies). (e) Mean valence scores (see Methods) obtained over time for the designated odors. Gray, parental controls; red, the experimental group at 32 °C; blue, the experimental control group at 23 °C. PD2a1/b1 neurons contribution to odor valence develops with time. Values at 120 seconds were similar to those presented in Fig. 2 (155 ≥ n ≥ 17 flies for all conditions, *p < 0.05, **p < 0.01, ****p < 0.0001, see Supplementary Table S1 for statistical analysis.).
As a control, we used ethyl lactate as the CS− since it does not change its valence value following the silencing of PD2a1/b1 neurons (Fig. 3), together with geranyl acetate as the CS+. As expected, prior to conditioning flies robustly avoided the CS−, ethyl lactate (Supplementary Fig. S7b) and after conditioning, control flies still avoided the CS− although to a slightly lesser extent than seen in the previous odor pair (Supplementary Fig. S7b). However, in contrast to the observations made with the 2-butanone/ethyl lactate odor pair, silencing of PD2a1/b1 neurons did not cause any change in avoidance for the ethyl lactate/geranyl acetate odor pair (Supplementary Fig. S7b). This is consistent with the observation that ethyl lactate exhibited no change in odor valence when PD2a1/b1 neurons were silenced. Taken together, we can conclude that silencing PD2a1/b1 neurons can contribute to aversive conditioning. This contribution is probably not a result of the direct participation of these neurons in the conditioning process, as was recently demonstrated15, but is more likely to be due to changes in the innate odor valence of the chemical compounds.

**PD2a1/b1 downstream neurons include MB KCs and neurons in the fan shaped body.** Our results suggest that context as manifested by a range of parameters (i.e. satiation, concentration, and odor exposure time) affects the contribution of PD2a1/b1 neurons to odor valence. This probably occurs without affecting the neural activity of PD2a1/b1 neurons themselves (satiation), or with no correlation between neural activity and the extent of their effect (concentration). These observations thus suggest that context is integrated downstream of the PD2a1/b1 neurons.

Recent EM data did not clearly locate potential downstream neurons15,18, although PD2a1/b1 axons were thought to be in close proximity to the axons of three MBONs (α2/1, α2, and β3/2 mp), which may suggest a common downstream integrator. Hence we attempted to identify downstream targets of PD2a1/b1 using trans-Tango55 with R37G11-GAL4. This labeled mostly PD2a1/b1 neurons (Fig. 7a–a’), in accordance with the report that 18.4% of the input synapses are reciprocal between PD2a1/b1 LH output neurons15. Additionally, strong trans-Tango signals appeared in varying subsets of all three main Kenyon cell types (Fig. 7b,c). The only site that allows for direct interaction of γ KCs with R37G11-GAL4-expressing neurons is the MB calyx, which is innervated by PD2b1 neurons35. We therefore examined the cellular polarity of these contacts. R37G11-GAL4 expression of the dendritic marker DenMark46, and the synaptic marker Dsyp-137 indeed suggests the presence of presynaptic sites of PD2b1 in the MB calyx (Supplementary Fig. S8). Furthermore, PD2b1 neurites exhibited colocalization with ChAT-immunoreactivity in the calyx (Supplementary Fig. S9), supporting the argument for active synapses and in line with the cholinergic identity of this cell type (Supplementary Fig. S1). Despite all these indications, we are very cautious about deducing functional calycal connectivity between PD2b1 neurons and KCs, given that a previous EM study15 did not report any presynapses between PD2b1 neurons and KCs in the calyx of their examined specimen. Functionally, a good candidate site for downstream integration of olfactory information independently processed by the MB and LH would be a premotor center like the fan-shaped body (FSB). Indeed, in some preparations we did see a trans-Tango signal driven by R37G11-GAL4 in layer 6 of the FSB (Fig. 7d). We should note, however, that the trans-Tango signal in the FSB was less prominent than the label in reciprocal LH cells and MB KCs. Recent data suggest that the FSB layer 6 is downstream of at least two MBON types (MBON-α2 and MBON-β3/2 mp), with yet another MBON apparently synapsing onto the FSB in layers 4 & 5 (MBON−β3/2 mp)58.

To further investigate the FSB as a potential downstream integrator region, we performed trans-Tango experiments with MBON− α2/1 as these neurons arborize in close proximity to axons of PD2a1/b1 neurons19. Surprisingly, instead of FSB labeling, we identified MBON-β3/2 mp as a postsynaptic target of MBON− α2/1 (Fig. 7e). We also observed weak labeling in putative LHONs, which is in accordance with the recent report of axoaxonic contacts of MBON− α2/1 onto the LHON AD1b24. These results suggest that not only MBON-α2, but also MBON− α2/1 converges onto MBON-β3/2 mp, making it an integrator of highly processed MB output. A repetition of the trans-Tango experiment with MBON-β3/2 mp without antibody staining confirmed FSB labeling (Fig. 7f’). Additional trans-Tango labeling was observed in PD2 LHONs, but in contrast to the case with PD2a1/b1 neurons, the signal was mainly associated with arborizations in the ventral LH (Fig. 7f’). Taken together, our results suggest that LH neurons, KCs, and FSB neurons are possible downstream integrators of PD2a1/b1 neurons and that MB-processed information is returned to the LH via MBON-β3/2 mp.

**Discussion**

The results described here indicate that PD2a1/b1 neurons can mediate aversive behavioral responses of satiated flies to odor at the relatively high concentration of 10−2. This effect is context dependent and is influenced by odor identity (even between odors with a similar activation pattern), odor concentration, exposure time, and fly satiation. Interestingly, starvation blocks the PD2a1/b1 aversive effect but not by directly affecting the activity of PD2a1/b1 neurons.

The temporal dynamics of the PD2a1/b1 neuron response vary dramatically, with some odors eliciting a prolonged and robust response but only at high odor concentrations. The most likely explanation for this prolonged odor response is that this is not an intrinsic property of PD2a1/b1 neurons but can rather be attributed to properties of the ORNs59. The most likely candidate is OR59b, which, according to the DoOR database, is the OR shared by all odors that generate a prolonged odor response25. This notion is also supported by results indicating prolonged odor responses to ethyl acetate in DM4, the glomerulus cognate to OR59b46, which was shown to be connected to PD2a1/b1 cells15,17.

The results presented here concerning the role of PD2a1/b1 neurons in driving aversive behavioral responses to odors, seem to contradict a previous report that claimed that PD2a1/b1 neurons exclusively drive attraction15. It is important to note that this previous study used 3 day old flies for the behavioral experiments, rather than the 7–14 day old flies used here. However, this difference is unlikely to explain the difference in the effects on naïve odor valence observed in the two studies following silencing of PD2a1/b1 neurons. This apparent discrepancy
Figure 7. trans-Tango downstream targets of PD2a1/b1, MBON-γ2α’1 and MBON-β’2mp neurons. (a) R37G11-GAL4-driven myrGFP (green) labels 6-7 cells in the LH, where as the trans-Tango signal (magenta) appears to be strongest in overlapping neuropil regions and in random sparse KC subsets in the MB (boxed area enlarged in a’ and a”). (a’) R37G11-GAL4-driven myrGFP (green) labels 6-7 cells (arrowhead) in the LH (stippled). (a”) R37G11-GAL4-driven trans-Tango signal (magenta) marks cells (arrowhead) in the LH (stippled). (b) Confocal stack of MB (stippled), highlighting the PD2a1/b1 neurites in proximity to the vertical lobe and a trans-Tango signal in all sublobes, i.e., α\(^{\alpha}\), α\(^{\beta}\) and γ. (c) Confocal stack of MB calyx (stippled) showing innervation by R37G11-GAL4-driven GFP (green) and trans-Tango signal in both KC somata and the calyx neuropil. (d) Confocal stack of left hemibrain highlighting R37G11-GAL4-driven myrGFP (green) and postsynaptic trans-Tango signal in layer 6 of the FSB (stippled). (e) MB077B-GAL4-driven myrGFP (green) labels 3-4 MBON-γ2α’1 neurons, whereas a fairly restricted putative downstream trans-Tango signal (magenta) most prominently labels MBON-β’2mp (boxed area enlarged in e’). (e’) Confocal stack of left hemibrain highlighting MB077B-GAL4-driven mCD8::GFP (green) and postsynaptic trans-Tango signal (magenta) in β’2mp of the MB and in putative LHON neurites (arrowheads). (f) MB011B-GAL4-driven myrGFP (green) labels MBON-γ5β2α and MBON-β’2mp, whereas the putative downstream trans-Tango signal (magenta) labels several different cell types (boxed area enlarged in f’). (f’) Confocal stack of right hemisphere highlighting the postsynaptic signal of MBON-β’2mp in PD2 LHONs (arrowhead) in the LH (stippled) and in layer 4 & 5 of the FSB (stippled). (a–f’): Confocal stack views of 1-2 µm-thick optical section data. Scale bars: a,e,f: 50 µm; a”,a”,a”,d,e,e”,f’: 20 µm.

may be explained if PD2a1/b1 neurons reinforce a valence value that is generated by other neurons in the fly’s brain. Thus, for an odor with a positive valence value, indicating attraction, PD2a1/b1 neurons further strengthen the attractive response. Conversely, for an odor with a negative value, indicating aversion, PD2a1/b1 neurons further strengthen the aversive response. If this is indeed the case, silencing PD2a1/b1 neuronal output should push the flies towards a random choice (i.e., 0 preference index) rather than driving either attraction or aversion. However, the results obtained with ACV and geranyl acetate negate this option. Satiated flies exposed to high concentrations of ACV or geranyl acetate showed no preference to the odor and had a preference index around zero (Fig. 3). However, silencing PD2a1/b1 neurons resulted in a marked increase in attraction towards these odors, pushing the preference index to positive values.

Another possible explanation for the apparent discrepancy in results may be the different behavioral approaches (with some marked differences) used in the two studies. Specifically, flies in behavioral chambers (as used here) are in constant motion, so that they repeatedly enter and exit the odor plume, and make multiple decisions each time\(^{4,30,61}\). In contrast, flies in the T-maze (as used in\(^{15}\)) make fewer decisions and reach steady state more rapidly\(^{32}\). Another difference is that flies in a T-maze make their decision within 3–4 seconds\(^{63}\), whereas the standard analysis of the behavioral chambers takes into account a two minute window with repeated decisions\(^{4,30,61}\). Given the strong context dependent effect of PD2a1/b1 neurons it is very plausible that PD2a1/b1 neurons contribute to both attraction and aversion. Consistent with this notion, while we did not identify a role for PD2a1/b1 neurons in attraction, starvation and low odor concentration abolished the aversive effect of activating these neurons. In addition both studies demonstrated that PD2a1/b1 neurons contribute differentially to odor valence even if the neural activity elicited by the odors is similar (Figs. 1 and 3). In this context, it is important to
note that optogenetic activation of PD2a1/b1 neurons resulted in a mild yet significant aversive response\textsuperscript{18}. Thus, the combined results of our study and those of Dolan et al.\textsuperscript{15,18} suggest that PD2a1/b1 neurons may affect odor valence in a number of apparently contradictory ways. This conclusion is further supported by recent results demonstrating that PD2a1 neurons are essential for context-dependent long term memory, but not for "classical" MB-mediated, context-independent long term memory\textsuperscript{64}.

PN input to PD2a1/b1 neurons consists of neurons innervating predominantly six glomeruli: DM1, DM4, VA2, VM3, DP1m, and DP1\textsuperscript{15,17}. These glomeruli are considered to drive attraction\textsuperscript{15} as they are activated by appetitive and food odors\textsuperscript{65-67}. This raises the question of how an input, which presumably generates an attraction behavior, can generate an aversive response to odor. However, these glomeruli were also reported to drive aversion (VM3\textsuperscript{69}), to respond to aversive odors (DM4, VA2\textsuperscript{69}) or to have no effect on attraction (DM4\textsuperscript{47}). Furthermore, it was suggested that in the case of *Drosophila* larvae, hyperactivation of Or42b neurons, which are the cognate neurons to glomerulus DM1 can trigger repulsion from odors\textsuperscript{64}. Thus, it is possible that glomeruli that innervate PD2a1/b1 neurons also participate in aversive behavioral responses to odors.

How can a single neuron class generate opposing behavioral responses? Without knowledge and genetic access to the downstream neurons, this is difficult to answer. However, one plausible explanation is that recruitment of downstream neuron types depends heavily on PD2a1/b1 neuronal activity. Thus, a low odor concentration that activates PD2a1/b1 neurons only weakly, may efficiently recruit one neuronal pathway leading to attraction. In contrast, a high odor concentration that produces a robust activation of PD2a1/b1 neurons may recruit a different neuronal pathway leading to aversion. This notion was demonstrated with salt, where low and high concentrations of salt, (leading to attraction and aversion, respectively), are encoded by different neuronal pathways\textsuperscript{66}. Similarly, while a low concentration of ACV triggers innate attraction by activating two glomeruli, DM1 and VA2, a higher concentration of ACV recruits an additional glomerulus, DM5, which leads to reduced attraction\textsuperscript{67}. For mechanosensory processing, the bandpass filtering mechanism of *Drosophila* antennal vibration sensation requires downstream neurons to have different levels of voltage-gated channels and different membrane resting potentials\textsuperscript{30}.

What may be the neurons downstream of PD2a1/b1 neurons? Unfortunately, the trans-Tango experiments did not provide a clear identification of the neurons downstream of PD2a1/b1. In accordance with a previous EM analysis\textsuperscript{15}, we saw strong labeling of LH neurons, but in contrast, our experiment also detected a signal in the MB, although the labeled subsets were not consistent. We did not detect dopaminergic neurons or MBONs, which were suggested to be innervated by PD2a1/b1 neurons. In some preparations we detected trans-Tango labeling in layer 6 of the FSB. Together with the recently described trans-Tango label in the same layer by MBONs\textsuperscript{28}, this would imply a downstream integrator of both MB and LH information in a premotor center. The FSB would represent an ideal candidate for a downstream integration site for olfactory information processed independently by both the MB and the LH. As part of the central complex, the FSB belongs to the main premotor center of the central brain, analogous to the basal ganglia in vertebrates\textsuperscript{61}. The central complex itself serves as the main hub, integrating innate and conditioned multisensory information to relay behavioral decisions to the respective downstream effectors\textsuperscript{12,22}.

A number of models have been proposed to explain odor identity and the associated valence in order to explain how a fly associates valence to a specific odor\textsuperscript{27}. One approach, termed "labeled lines", demonstrated that odors of particular relevance are often detected by ORNs expressing highly selective receptors that respond to only a single compound. In the case of *Drosophila*, this type of preferential relationship was demonstrated for sex pheromones, harmful substances, oviposition cues, and even food\textsuperscript{74-78}. According to this model, activation of the labeled line generates a hardwired behavioral response. However, since most ORNs are broadly tuned to odors\textsuperscript{77,88,79}, the labeled line approach is probably insufficient to explain behavioral responses in these cases. Indeed, behavioral responses to general odors are dependent on population neural activity\textsuperscript{80}. Nevertheless, even under the population neural activity approach, a semi-labeled line assumption exists, according to which, a neuron always drives either attractive or aversive behavior (but not both), and it is the sum of the attractive and aversive signals from all the responding neurons that eventually determines the odor valence\textsuperscript{66,67}. More recently, this notion was also demonstrated using optogenetic mapping of MBONs. Accordingly, an attractive or aversive behavior is observed following optogenetic activation of MBONs\textsuperscript{24}, and appetitive or aversive associative conditioning will cause an odor to activate MBONs that drive attractive or aversive behavior, respectively\textsuperscript{22,24}. This model is extremely suitable for the MB, where odor tuning of MBONs is modified by plasticity based on experience\textsuperscript{23-25}. Recently, similar optogenetic mapping of LH neurons classified them as driving either attractive or aversive behavior\textsuperscript{48}. However, it is well established that context affects innate behavioral output, which is mediated by the LH\textsuperscript{11}. Thus, while the dynamic nature of KC-to-MBON connectivity allows for context-dependent changes, it is less clear how the LH, which as far as we know receives mostly hardwired inputs from PNs, makes context-dependent changes in neuronal activity (but see\textsuperscript{15}). Our results suggest that at least for the LH, the notion that a neuron can be categorized as driving either attraction or aversion may need to be reconsidered.

**Methods**

**Fly strains.** Flies were grown on cornmeal agar under a 12 h light/12 h dark cycle at 25 °C. Experimental flies carrying transgenes were crossed with w\textsuperscript{118} flies. The following transgenic flies were used: UAS-GCaMP6\textsuperscript{90}, UAS-Shibire\textsuperscript{81}, R37G11-GAL4\textsuperscript{15} (Bloomington #49539), R48F03-GAL4\textsuperscript{21} (Bloomington #50373), UAS-myrGFP-tdTomato(3XHA);trans-Tango\textsuperscript{55}, MB011B, MB077B\textsuperscript{24} (Bloomington #68294, #68283, respectively). LH989\textsuperscript{15,18} was produced by combining R37G11-ZpGdbd and R29G05-p65ADZp promoter fragments. UAS-CD8::GFP, lexAop-rCD2::RFP\textsuperscript{82}, nSyb-lexA.DBD::QF 4D (Bloomington #51953), UAS-DenMark\textsuperscript{56}, UAS-GFP::Dsyl-1\textsuperscript{57}.

**Full methods.**
During training, an odor (usually the more attractive one) was paired with 12 equally spaced 1.25 s electric shocks, as (preference for the left side when it contains odor) – (preference for the left side when it contains air). The learning index was calculated as (preference for odor before training) – (preference for odor after training).

The air flow from both sides of the chamber converged at the center of the chamber. Chambers were stacked in two columns, and were backlit by LEDs (940 nm, Vishay TSAL6400). A MAKO CMOS camera (Allied Vision Technologies) equipped with a Computar M0814-MP2 lens was used to obtain the images. The entire apparatus was placed in an incubator (Panasonic MIR 154) to control the temperature.

Each fly position over time was extracted from the video images using custom written software (LabVIEW 7.1, National Instruments) which also controlled the delivery of odors and electric shocks. Data were analyzed using custom written software (MATLAB 2015b, The MathWorks) and Prism 6 (GraphPad).

Preference was calculated as the percentage of time spent in one half of the chamber. Odor avoidance and training protocols were as presented in Fig. 2 and Supplementary Fig. S7. The naïve avoidance index was calculated as (preference for the left side when it contains odor) – (preference for the left side when it contains air). During training, an odor (usually the more attractive one) was paired with 12 equally spaced 1.25 s electric shocks at 60 V. The learning index was calculated as (preference for odor before training) – (preference for odor after training).

A fly’s initial mean velocity was calculated from the first 3 seconds after the first odor pulse. Velocity was calculated only for flies that were on the odor side upon odor presentation. Fly position was recorded every 175 ms and data was smoothed across 5 data points. Vectors of the fly’s heading were calculated based on fly position and were divided by 175 ms to deduce velocity. To calculate differences between odors, the integral of the response and data was smoothed across 5 data points. Responsiveness was calculated as the percentage of cells responding, out of the total number of cells examined for the specific odor. Response persistence was calculated as the inverse of the ratio between peak odor and the inter-odor ∆F/F response 5 seconds after peak odor response was detected. Responsiveness was calculated as the percentage of cells responding, out of the total number of cells examined for the specific odor.

Strain dissection, fixation, and immunostaining were performed as previously described. In the case of native GFP and RFP fluorescence, brains were fixed at room temperature for 20 minutes in 4% (w/v) paraformaldehyde in PBS (1.86 mM NaH₂PO₄, 8.41 mM Na₂HPO₄, 175 mM NaCl). Samples were then washed three times for 20 minutes in PBS containing 0.3% (v/v) Triton-X-100 (PBT). The primary antisera used were mouse monoclonal anti-GFP IgG2a (1:500, Invitrogen), rabbit polyclonal anti-GFP (1:500, Invitrogen), mouse monoclonal anti-ChAT (1:100, 4B1, DSHB), rabbit polyclonal anti-dvGlut (1:10,000), and rabbit polyclonal anti-GABA (1:1,000, A2052, Sigma-Aldrich). Secondary antisera were Alexa488 coupled to goat anti-rabbit or goat anti-mouse IgG2a, Alexa647 coupled to goat anti-mouse (1:500, all Invitrogen), and STAR.
RED coupled to goat anti-rabbit (1:500, Abberior). Counterstaining was performed using nc82 (DSHB) and goat anti-mouse Alexa647 or Alexa564. Primary antisera were applied for 1–2 days and secondary antiserum for 1–2 days in PBT at 4 °C, followed by embedding in Vectashield. Images were taken using a confocal microscope (Leica TCS SP5, SP8, Zeiss LSM 800, or a Nikon A1). Images were processed using ImageJ or Amira (ThermoFisher).

For trans-Tango experiments, R37G11-GAL4, MB077B, or MB011B males were crossed with UAS-myrtGF PQUAS-mdTomato. trans-Tango virgin females and reared at 18 °C. Three-week-old flies were anesthetized using ice and then dissected and processed as described above.

Correlation analysis. Correlation analysis was performed using MATLAB R2015a. Odor response vectors (AF/F) were compared from the onset of odor response to the end of the recording (due to prolonged odor responses with some odors), 35 seconds later. To determine the within-fly correlation for a given odor vector, the odor responses of all neurons recorded (3–6 neurons) were compared. The averages of such correlation values for all odors and all flies were used. For odor correlation, all odor responses of a given neuron were compared. Similarly, the averages of such correlation values for all odors and all flies were used. To determine between-fly correlations, a matrix with all neurons from all flies for a given odor was constructed and all correlation values were calculated. An average correlation value was then calculated for each odor. The averages of such values for all odors were used. For control calculations, two approaches were used. First, a 3D matrix was composed as above but for all odors. Response vectors were then shuffled and the above calculation was repeated. We then repeated the calculation on an artificial data set with the same number of neurons, flies, and a correlation between the neurons of the same fly, but not between flies (the low odor correlation was used as the baseline for correlation).

Statistics. All statistical analyses were performed using GraphPad Prism as described in the figure legends and in Table S1. In general, no statistical methods were used to predetermine the sample sizes.

Data availability
The datasets generated and analyzed for the current study are available from the corresponding author on request.

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References
1. Touhara, K. & Vosshall, L. B. Sensing Odorants and Pheromones with Chemosensory Receptors. Annu. Rev. Physiol. 71, 307–332 (2009).
2. Li, Q. & Liberles, S. D. Aversion and Attraction through Olfaction. Curr. Biol. 25, R120–R129 (2015).
3. Buck, L. & Axel, R. A novel multigene family may encode odorant receptors: A molecular basis for odor recognition. Cell 65, 175–187 (1991).
4. Vosshall, L. B., Amrein, H., Morozov, P. S., Zheshkys, A. & Axel, R. A Spatial Map of Olfactory Receptor Expression in the Drosophila Antenna. Cell 96, 725–736 (1999).
5. Hallem, E. A. & Carlson, J. R. The odor coding system of Drosophila. Trends Genet. 20, 453–459 (2004).
6. Gao, Q., Yuan, B. & Chess, A. Convergent projections of Drosophila olfactory neurons to specific glomeruli in the antennal lobe. Nat. Neurosci. 3, 780–785 (2000).
7. Mombaerts, P. et al. Visualizing an olfactory sensory map. Cell 87, 675–686 (1996).
8. Vosshall, L. B., Wong, A. M. & Axel, R. An olfactory sensory map in the fly brain. Cell 102, 147–159 (2000).
9. Tanaka, N. K., Endo, K. & Ito, K. Organization of antennal lobe-associated neurons in adult Drosophila melanogaster brain. J Comp Neurol 520, 4067–4130 (2012).
10. Schulthhaus, J. N., Saleem, S., Carney, G. E. The role of the Drosophila lateral horn in olfactory information processing and behavioral response. Journal of Insect Physiology 98, 29–37 (2017).
11. Strutz, A. et al. Decoding odor quality and intensity in the Drosophila brain. Elife 3, e04147 (2014).
12. Bräcker, L. B. B. et al. Essential Role of the Mmushroom Body in Context-Dependent CO2 Avoidance in Drosophila. Curr. Biol. 23, 1228–1234 (2013).
13. Xia, S. & Tully, T. Segregation of odor identity and intensity during odor discrimination in Drosophila mushroom body. PLoS Biol. 5, 2398–2407 (2007).
14. Wang, X. et al. Blockade of Neurotransmission in Drosophila Mushroom Bodies Impairs Odor Attraction, but Not Repulsion. Curr. Biol. 13, 1900–1904 (2003).
15. Dolan, M.-J. et al. Communication from Learned to Innate Olfactory Processing Centers Is Required for Memory Retrieval in Drosophila. Neuron 100, 651–668.e8 (2018).
16. Prechter, S. et al. Functional and anatomical specificity in a higher olfactory centre. Elife https://doi.org/10.7554/eLife.44590 (2019).
17. Jeanne, J. M., Fişek, M. & Wilson, R. I. The Organization of Projections from Olfactory Glomeruli onto Higher-Order Neurons. Neuron 98, 1198–1213.e6 (2018).
18. Dolan, M.-J. et al. Neurogenetic dissection of the Drosophila lateral horn reveals major outputs, diverse behavioural functions, and interactions with the mushroom body. Elife https://doi.org/10.7554/eLife.43079 (2019).
19. Huoviala, P. et al. Neural circuit basis of aversive odor processing in Drosophila from sensory input to descending output. bioRxiv 394403. https://doi.org/10.1101/394403 (2018).
20. Varela, N., Gaspar, M., Dias, S. & Vasconcelos, M. L. Avoidance response to CO2 in the lateral horn. PLOS Biol. 17, e2006749 (2019).
21. Fişek, M. & Wilson, R. I. Stereotyped connectivity and computations in higher-order olfactory neurons. Nat. Neurosci. 17, 280–8 (2014).
22. Aso, Y. et al. The neuronal architecture of the mushroom body provides a logic for associative learning. Elife 3, e04147 (2014).
23. Higo, T., Ito, Y., Rubin, G. M. & Turner, G. C. Plasticity-driven individualization of olfactory coding in mushroom body output neurons. Nature 526, 258–262 (2015).
24. Aso, Y. et al. Mushroom body output neurons encode valence and guide memory-based action selection in Drosophila. Elife 3, e04580 (2014).
25. Pelsenberg, J. et al. Integration of Parallel Opposing Memories Underlies Memory Extinction. Cell 175, 709–722.e15 (2018).
26. Sjöström, J. R. et al. Mushroom body efferent neurons responsible for aversive olfactory memory retrieval in Drosophila. Nat. Neurosci., https://doi.org/10.1038/nn.2846 (2011).
27. Berry, J. A., Phan, A. & Davis, R. L. Dopamine Neurons Mediate Learning and Forgetting through Bitirectional Modulation of a Memory Trace. Cell Rep., https://doi.org/10.1016/j.celrep.2018.09.051 (2018).
28. Perisse, E. et al. Aversive Learning and Appetitive Motivation Toggle Feed-Forward Inhibition in the Drosophila Mushroom Body. *Neuron*, https://doi.org/10.1016/j.neuron.2016.04.034 (2016).
29. Owald, D. et al. Activity of defined mushroom body output neurons underlies learned olfactory behavior in Drosophila. *Neuron*, https://doi.org/10.1016/j.neuron.2015.03.025 (2015).
30. Lewis, I. P. C. et al. A Higher Brain Circuit for Immediate Integration of Conflicting Sensory Information in Drosophila. *Curr. Biol.*, https://doi.org/10.1016/j.cub.2015.07.015 (2015).
31. Root, C. M., Ko, K. L., Iafati, A. & Wang, J. W. Preynaptic facilitation by neuropeptide signaling mediates odor-driven food search. *Cell* 145, 133–144 (2011).
32. Heimbek, G., Bugnon, V., Gendre, N., Keller, A. & Stocker, R. F. A central neural circuit for experience-independent olfactory and courtship behavior in Drosophila melanogaster. *Proc. Natl. Acad. Sci.*, https://doi.org/10.1073/pnas.011314898 (2001).
33. Devaud, J. M., Keane, J. & Ferrús, A. Blocking sensory inputs to identified antennal glomeruli selectively modifies odorant perception in Drosophila. *J. Neurobiol.*, https://doi.org/10.1002/neu.10236 (2003).
34. Wang, Y. et al. Genetic manipulation of the odor-evoked distributed neural activity in the Drosophila mushroom body. *Neuron*, https://doi.org/10.1016/S0896-6273(01)00196-9 (2001).
35. Giang, T., He, J., Beladì, S. & Scholz, H. Key odorants regulate food attraction in Drosophila melanogaster. *Front. Behav. Neurosci.*, https://doi.org/10.3389/fnbeh.2017.00160 (2017).
36. Daniels, R. W., Gelfand, M. V., Collins, C. A. & DiAntonio, A. Visualizing glutamatergic cell bodies and synapses in Drosophila larval and adult CNS. *J. Comp. Neurol.* 508, 131–152 (2008).
37. Münch, D. & Galizia, C. G. DoOR 2.0 - Comprehensive Mapping of Drosophila melanogaster Olfactant Responses. *Sci. Rep.* 6, 21841 (2016).
38. Lüdke, A. et al. Calcium in Kenyon Cell Somata as a Substrate for an Olfactory Sensory Memory in Drosophila. *Front. Cell. Neurosci.* 12, 128 (2018).
39. Tsao, C.-H., Chen, C.-C., Lin, C.-H., Yang, H.-Y. & Lin, S. Drosophila mushroom bodies integrate hunger and satiety signals to control innate food-seeking behavior. *Elife* 7, e35264 (2018).
40. Wang, S.-P. et al. Matting Experience and Food Deprivation Modulate Odor Preference and Dispersal in Drosophila melanogaster Males. *J. Insect Sci.*, https://doi.org/10.1673/031.014.131 (2016).
41. Ko, K. I. et al. Starvation promotes concerted modulation of appetitive olfactory behavior via parallel neuromodulatory circuits. *Elife*, https://doi.org/10.7554/eLife.08298.001 (2015).
42. Rivenor, J., Boto, T. & Alcorta, E. The effect of environmental temperature on olfactory perception in Drosophila. *J. Insect Physiol.*, https://doi.org/10.1016/j.jinsphys.2009.06.009 (2009).
43. Kreher, S. A., Kwon, J. Y. & Carlson, J. R. The molecular basis of odor coding in the Drosophila larva. *Neuron* 46, 445–456 (2005).
44. Amdor, O. & Laurent, G. Transient dynamics versus fixed points in odor representations by locust antennal lobe projection neurons. *Neuron* 48, 661–673 (2005).
45. Stopfer, M., Jayaraman, V. & Laurent, G. Intensity versus identity coding in an olfactory system. *Neuron* 39, 991–1004 (2003).
46. Seki, Y. et al. Olfactory coding from the periphery to higher brain centers in the Drosophila brain. *BMC Biol.* 15, 56 (2017).
47. Steck, K. et al. A high-throughput behavioral paradigm for Drosophila olfaction - The Flywalk. *Sci. Rep.*, https://doi.org/10.1038/srep00361 (2012).
48. Mohamed, A. A. M. et al. Odor mixtures of opposing valence unveil inter-glomerular crosstalk in the Drosophila antennal lobe. *Nat. Commun.* 10, 1201 (2019).
49. Tully, T. & Quinn, W. G. Classical conditioning and retention in normal and mutant Drosophila melanogaster. *J. Comp. Physiol. A.*, https://doi.org/10.1007/BF01350033 (1985).
50. Préat, T. Decreased odor avoidance after electric shock in Drosophila mutants biases learning and memory tests. *J. Neurosci.*, https://doi.org/10.1523/jneurosci.18-20-08534.1998 (1998).
51. Zars, T., Fischer, M., Schulz, R. & Heisenberg, M. Localization of a short-term memory in Drosophila. *Science* (80-.), https://doi.org/10.1126/science.288.5466.672 (2000).
52. Quinn, W. G., Harris, W. A. & Benzer, S. Conditioned behavior in Drosophila melanogaster. *Proc. Natl. Acad. Sci. USA*, https://doi.org/10.1073/pnas.71.3.708 (1974).
53. Pitman, J. L. et al. There are many ways to train a fly. *Fly*, https://doi.org/10.4161/fly.3.1.7772 (2009).
54. Clárđige-Chang, A. et al. Writing Memories with Light-Addressable Reinforcement Circuity. *Cell* 139, 405–419 (2009).
55. Talay, M. et al. Transsynaptic Mapping of Second-Order Taste Neurons in Flies by trans-Tango. *Neuron* 96, 783–795.e4 (2017).
56. Nicolai, L. J. et al. Genetically encoded dendritic marker sheds light on neuronal connectivity in Drosophila. *Proc Natl Acad Sci USA* 107, 20553–20558 (2010).
57. Owald, D. et al. A Syd-1 homologue regulates pre- and postsynaptic maturation in Drosophila. *J. Cell Biol.* 188, 565–79 (2010).
58. Scaplen, K. M. et al. Circuits that encode and predict alcohol associated preference. *bioRxiv* 1–15, https://doi.org/10.1101/578401 (2019).
59. Hallem, E. A. & Carlson, J. R. The spatial code for odors is changed by conditioning. *Neuron* 42, 359–361 (2004).
60. Parinas, M., Lin, A. C., Huetteloth, W. & Miesenbock, G. Odor Discrimination in Drosophila: From Neural Population Codes to Behavior. *Neuron* 79, 932–944 (2013).
61. Lin, A. C., Bygrave, A. M., De Calignon, A., Lee, T. & Miesenbock, G. Sparse, decorrelated odor coding in the mushroom body enhances learned odor discrimination. *Nat. Neurosci.* 17, 559–568 (2014).
62. Ichinose, T. & Tanimoto, H. Dynamics of memory-guided choice behavior in Drosophila. *Proc. Japan Acad. Ser. B.*, https://doi.org/10.2183/pjab.92.346 (2016).
63. Chabaud, M.-A., Preat, T. & Kaiser, L. Behavioral Characterization of Individual Olfactory Memory Retrieval in Drosophila Melanogaster. *Front. Behav. Neurosci.*, https://doi.org/10.3389/fnbeh.2010.00192 (2011).
64. Zhao, B. et al. Long-term memory is formed immediately without the need for protein synthesis-dependent consolidation in Drosophila. *Nat. Commun.*, https://doi.org/10.1038/ncomms14267-1 (2017).
65. Knaden, M., Strutz, A., Ahsan, J., Sachs, S. & Hansson, B. S. Spatial representation of odorant valence in an insect brain. *Cell Rep* 1, 392–399 (2012).
66. Badel, L., Ohta, K., Tsushimoto, Y. & Kazama, H. Decoding of Context-Dependent Olfactory Behavior in Drosophila. *Neuron* 91, 155–167 (2016).
67. Semmelhack, J. L. & Wang, J. W. Select Drosophila glomeruli mediate innate olfactory attraction and aversion. *Nature*, https://doi.org/10.1038/nature07983 (2009).
68. Kreher, S. A., Mathew, D., Kim, J. & Carlson, J. R. Translation of Sensory Input into Behavioral Output via an Olfactory System. *Neuron* 59, 110–124 (2008).
69. Zhang, Y. Y., Ni, J. & Montell, C. The Molecular Basis for Attractive Salt-Taste Coding in Drosophila. *Science* (80-.), 340, 1334–1338 (2013).
70. Azevedo, A. W. & Wilson, R. I. Active Mechanisms of Vibration Encoding and Frequency Filtering in Central Mechanosensory Neurons. *Neuron* 96, 446–460.e9 (2017).
71. Straussfeld, N. J. & Hirth, F. Homology versus convergence in resolving transsynaptic correspondences of brain organization. *Brain, Behavior and Evolution*, https://doi.org/10.1159/000356102 (2013).
72. Hu, W. et al. Fan-Shaped Body Neurons in the Drosophila Brain Regulate Both Innate and Conditioned Nociceptive Avoidance. Cell Rep., https://doi.org/10.1016/j.celrep.2018.07.028 (2018).
73. Haverkamp, A., Hasson, B. S. & Knaden, M. Combinatorial codes and labeled lines: How insects use olfactory cues to find and judge food, mates, and oviposition sites in complex environments. Front. Physiol. 9, 49 (2018).
74. Ruta, V. et al. A dimorphic pheromone circuit in Drosophila from sensory input to descending output. Nature 468, 686–690 (2010).
75. Stensmyr, M. C. et al. A conserved dedicated olfactory circuit for detecting harmful microbes in Drosophila. Cell 151, 1345–1357 (2012).
76. Kohl, J., Ostrovsky, A. D., Frechter, S. & Jefferis, G. S. X. E. A Bidirectional Circuit Switch Reroutes Pheromone Signals in Male and Female Brains. Cell 155, 1610–1623 (2013).
77. Dweck, H. K. M., Ebrahim, S. A. M., Farhan, A., Hansson, B. S. & Stensmyr, M. C. Olfactory Proxy Detection of Dietary Antioxidants in Drosophila. Curr. Biol. 25, 455–466 (2015).
78. Dweck, H. K. M. et al. Olfactory Preference for Egg Laying on Citrus Substrates in Drosophila. Curr. Biol. 23, 2472–2480 (2013).
79. Hallem, E. A. & Carlson, J. R. Coding of Odors by a Receptor Repertoire. Cell 125, 143–160 (2006).
80. Chen, T.-W. W. et al. Ultrasensitive fluorescent proteins for imaging neuronal activity. Nature 499, 295–300 (2013).
81. Kitamoto, T. Conditional modification of behavior in Drosophila by targeted expression of a temperature-sensitive shibire allele in defined neurons. J. Neurobiol. 47, 81–92 (2001).
82. Lee, T. & Luo, L. Mosaic analysis with a repressible cell marker for studies of gene function in neuronal morphogenesis. Neuron 22, 451–461 (1999).
83. Wang, J. W., Wong, A. M., Flores, J., Vosshall, L. B. & Axel, R. Two-photon calcium imaging reveals an odor-evoked map of activity in the fly brain. Cell 112, 271–282 (2003).
84. Ng, M. et al. Transmission of olfactory information between three populations of neurons in the antennal lobe of the fly. Neuron (2002).
85. Thevenaz, P., Ruttimann, U. E. & Unser, M. A pyramid approach to subpixel registration based on intensity. IEEE Trans. Image Process. 7, 27–41 (1998).
86. Olsen, S. R., Bhandawat, V. & Wilson, R. I. Divisive normalization in olfactory population codes. Neuron 66, 287–299 (2010).
87. Pitman, J. L., Huetteroth, W. et al. A pair of inhibitory neurons are required to sustain labile memory in the Drosophila mushroom body. Curr. Biol. 21, 855–863 (2011).
88. Wu, J. S. & Luo, L. A protocol for dissecting Drosophila melanogaster brains for live imaging or immunostaining.