Input Connectivity Reveals Additional Heterogeneity of Dopaminergic Reinforcement in *Drosophila*

**Highlights**
- Nanoscale anatomy reveals additional subtypes of rewarding dopaminergic neurons
- Connectomics reveals input specificity to subtypes of dopaminergic neurons
- Axon morphology implies dopaminergic neurons provide subcompartment-level function
- Unique dopaminergic subtypes serve aversive memory extinction and sugar learning

**Authors**
Nils Otto, Markus W. Pleijzier, Isabel C. Morgan, ..., Gerald M. Rubin, Gregory S.X.E. Jefferis, Scott Waddell

**Correspondence**
scott.waddell@cncb.ox.ac.uk

**In Brief**
Otto et al. use electron microscope resolution connectomics to describe the structure and organization of neurons providing synaptic input to functionally discrete subtypes of dopaminergic neurons. The nanoscale anatomy reveals further anatomical and functional specialization of dopaminergic neurons, which is confirmed with behavioral experiments.
Input Connectivity Reveals Additional Heterogeneity of Dopaminergic Reinforcement in *Drosophila*

Nils Otto,1,7 Markus W. Pleijzier,2,8 Isabel C. Morgan,1 Amelia J. Edmondson-Stait,2 Konrad J. Heinz,2 Ildiko Stark,2 Georgia Dempsey,2 Masayoshi Ito,3 Ishaan Kapoor,1 Joseph Hsu,3 Philipp M. Schlegel,2,4 Alexander S. Bates,2,4 Li Feng,3 Marta Costa,2 Kei Ito,1 Davi D. Bock,1,6,9 Gerald M. Rubin,3 Gregory S.X.E. Jefferis,2,4,10 and Scott Waddell1,11,12,*

1Centre for Neural Circuits and Behaviour, The University of Oxford, Tinsley Building, Mansfield Road, Oxford OX1 3SR, UK
2Drosophila Connectomics, Department of Zoology, University of Cambridge, Cambridge CB2 3EJ, UK
3Janelia Research Campus, Howard Hughes Medical Institute, Ashburn, VA 20147, USA
4Division of Neurobiology, MRC Laboratory of Molecular Biology, Cambridge CB2 0QH, UK
5Institute for Zoology, University of Cologne, Zu¨lpicher Straße 47b, 50674 Cologne, Germany
6Present address: Department of Neurological Sciences, University of Vermont Medical Center, Burlington, VT 05405, USA
7Twitter: @_Nils_Otto_
8Twitter: @mwpleijzier
9Twitter: @dddavi
10Twitter: @gsxej
11Twitter: @scottishwaddell
12Lead Contact
*Correspondence: scott.waddell@cncb.ox.ac.uk
https://doi.org/10.1016/j.cub.2020.05.077

**SUMMARY**

Different types of *Drosophila* dopaminergic neurons (DANs) reinforce memories of unique valence and provide state-dependent motivational control [1]. Prior studies suggest that the compartment architecture of the mushroom body (MB) is the relevant resolution for distinct DAN functions [2, 3]. Here we used a recent electron microscope volume of the fly brain [4] to reconstruct the fine anatomy of individual DANs within three MB compartments. We find the 20 DANs of the γ5 compartment, at least some of which provide reward teaching signals, can be clustered into 5 anatomical subtypes that innervate different regions within γ5. Reconstructing 821 upstream neurons reveals input selectivity, supporting the functional relevance of DAN sub-classification. Only one PAM-γ5 DAN subtype γ5(fb) receives direct recurrent feedback from γ5β2a mushroom body output neurons (MBONs) and behavioral experiments distinguish a role for these DANs in memory revaluation from those reinforcing sugar memory. Other DAN subtypes receive major, and potentially reinforcing, inputs from putative gustatory interneurons or lateral horn neurons, which can also relay indirect feedback from MBONs. We similarly reconstructed the single aversively reinforcing PPL1-γ1pedc DAN. The γ1pedc DAN inputs mostly differ from those of γ5 DANs and they cluster onto distinct dendritic branches, presumably separating its established roles in aversive reinforcement and appetitive motivation [5, 6]. Tracing also identified neurons that provide broad input to γ5, β2a, and γ1pedc DANs, suggesting that distributed DAN populations can be coordinately regulated. These connectomic and behavioral analyses therefore reveal further complexity of dopaminergic reinforcement circuits between and within MB compartments.

**RESULTS AND DISCUSSION**

In adult *Drosophila*, anatomically discrete dopaminergic neurons (DANs) innervate adjacent compartments of the mushroom body (MB) [23]. In some cases, different combinations of DANs serve discrete roles. However, there are instances where multiple functions have been assigned to DANs that innervate the same compartment. For example, DANs innervating the γ5 compartment reinforce short-term courtship memories and appetitive memories with sugar, and they also signal the absence of expected shock, to extinguish aversive memory [7–11]. Similarly, DANs innervating the β2a compartment have roles such as controlling thirst state-dependent water seeking and water memory expression, sugar reinforcement, and hunger-dependent modulation of carbon dioxide avoidance [9, 10, 12–14]. Moreover, individual PPL1-γ1pedc DANs, which innervate the γ1 compartment in both hemispheres, are required to reinforce aversive memories with electric shock, high heat, and bitter taste, and also provide hunger state-dependent motivational control of sugar memory expression [5, 6, 15, 16]. For an individual DAN to multi-task it must function in different modes. However, where a compartment is innervated by multiple DANs, different neurons
Determining the Nanoscale Structure of Reinforcing Dopaminergic Neurons

We used a recent EM dataset of a full adult fly brain (FAFB) [4] to identify, manually trace, and reconstruct the nanoscale anatomy of 20 DANs in the protocerebral anterior medial (PAM) cluster whose presynaptic arbors innervate the 5 compartment (PAM-5 DANs) [29] in the fly’s right brain hemisphere (we also reconstructed 9 PAM-γ5 DANs in the left MB). We identified 8 right hemisphere PAM-β2a DANs and reconstructed 6. We also reconstructed the 2 protocerebral posterior lateral (PPL)1-γ1pedc DANs that innervate the γ1 compartments of each MB (Figures 1A and S1A–S1D; Video S1). STAR Methods and the revision status table in Methods S1 detail metrics of quality control.

We noticed when reconstructing PAM-γ5 DANs that their dendrites occupied different areas of the superior medial protocerebrum (SMP) [17], that their somata were connected via 2 neurite tracts, and that each γ5 DAN had a contralateral projection crossing the midline of the brain in an upper, middle, or lower commissure (Figures 1A, 1C–1G, S1A, S1B, and S1E–S1H). We therefore used unbiased anatomical clustering to explore sub-organization of PAM-γ5 DANs (Figure 1B). This grouped PAM-γ5 DANs into 5 discrete clusters of 1–7 neurons (Figures 1C–1G). Importantly, a different clustering criterion produced an identical result (Figure S1I). Although we did not trace the finest axonal branches of PAM-γ5 DANs, it was evident that their major presynaptic arbors occupy different areas of the γ5 compartment. We therefore named the PAM-γ5 subtypes according to their defining morphological feature (Figures 1C–1G; Video S1).

The four reviewed PAM-β2a DANs could also be clustered into three groups with commissure crossed, overall morphology, and region of compartment innervation again being the distinguishing features (Figures 1B, 1H–1J, S1C, and S1E–S1H; Video S1). Tracing also identified a “non-canonical” PAM-β2a DAN, which mostly innervates β2a but also extends axons into γ5 (Figure 1J). The PAM-γ2a DAN dendrites were largely intermingled with those of PAM-γ5 DANs (Figures 1C–1J; Video S1), consistent with their roles in reinforcing appetitive memories.

Reconstructing the individual right hand PPL1-γ1pedc DAN revealed that its dendrites occupy locations distinct from those of PAM-5 and PAM-β2a DANs (Figure 1K). This suggests that PPL1-γ1pedc receives mostly different input, consistent with it signaling aversive rather than appetitive valence. The PPL1-γ1pedc DAN dendrite has four major arbors that extend into orthogonal locations in the brain. Postsynapses also clustered in each of these locations (Figures 1L, 1M, and S1H; Video S2).

Receiving branch-specific information may represent a solution for how a single PPL1-γ1pedc DAN isolates and prioritizes its discrete roles in reinforcement and state-dependent control [5, 6].

Mapping Neuronal Inputs onto Dopaminergic Neurons

We next traced 821 neurons providing input to postsynapses identified in the dendrites of the PAM-γ5, PAM-β2a, and PPL1-γ1pedc DANs (Figure 2A). Since connectivity is dense and manual tracing is labor-intensive, we traced between 45% and 97% of the inputs to the postsynapses annotated on all the reconstructed DANs. We prioritized upstream tracing to retrieve a comparable coverage of inputs to each of the DAN subclasses. Sampling criteria and metrics of quality control are detailed (Methods S1).

Reassuringly, input selectivity of PAM-5, PAM-β2a, and PPL1-γ1pedc DANs largely reflected the relative overlap of their dendritic fields. All the DANs receive unique inputs (Figures 2B, 2C, and S2A–S2D). However, a greater share of the 553 identified inputs to PAM-γ5 DANs (181) also provided inputs to PAM-β2a DANs than to the PPL1-γ1pedc DAN (84). Likewise, more of the 281 PAM-β2a DANs inputs connect to PAM-γ5 DANs (181) than to the PPL1-γ1pedc DAN (67). In contrast more of the 275 traced PPL1-γ1pedc DAN inputs were unique (168) than also contacted PAM-γ5 DANs (84) or PAM-β2a DANs (67). Lastly, 5% (44) of the traced input neurons synapsed onto all three classes of traced DANs. These common input neurons suggest that activity in the valence-specific arms of the DANs...
system may be coordinated. It is, however, also possible that the different DANs respond in unique ways to the same input neurotransmitters.

Our sampling suggests that despite there being 20 PAM-γ5 DANs and one PPL1-γ1pdec DAN, there are only approximately twice as many inputs to all PAM-γ5 DANs compared to the PPL1-γ1pdec DAN (Figure 2B). However, the different DAN types have markedly different weighting of inputs, assuming that synapse number correlates with input strength (Figures 2D and S2E). Whereas the PPL1-γ1pdec DAN receives weakly connected inputs, each input to the PAM-γ5 DANs or PAM-β'2a DANs is more strongly connected and contributes a larger proportion of the individual neuron’s postsynaptic budget. Plotting a connectivity matrix for the most completely traced DANs reveals that certain groups of inputs preferentially synapse onto different PAM-γ5 and PAM-β'2a DANs, demonstrating that all individual PAM-γ5 and PAM-β'2a DANs have an element of input specificity (Figures 2E and S2F). Nevertheless, a matrix comparing input structure between DANs reveals significant similarities in input between particular groups of PAM-γ5 and PAM-β'2a DANs, which is more organized than random connectivity (Figure 2F; Methods S1). DAN clustering based on dendritic connectivity was identical using two different methods (Figure S2G). Moreover, clustering based on input connectivity correlated well with the prior clustering using full neuron morphology (Figure 2H).

The observed connectivity could be confounded by the incompleteness of reconstruction. However, plotting the synapses identified following standard and extensive review suggests that each iteration of review adds synapses that are evenly distributed across a DAN’s dendritic arbor (Figure S2H). Nevertheless, we tested whether connectivity clustering resulted from unintentional bias in input neuron tracing by repeated clustering following random downsampling of input connectivity from 5% to 50%. Cluster content remained largely robust in these analyses with DANs clustering within the same groups across 10,000 simulations where synapses were removed (Figures S2I–S2M). The stability of DAN clustering based on input structure and the similarity of connectivity and morphology clustering suggest that information conveyed by selective input is likely to be maintained in the activity of different DAN subtypes.

We next clustered the input neurons based on their morphology using a three-step approach (Methods S1). Using soma location and primary neurite layout revealed 20 coarse clusters (Figures 2I and S2N; Video S3). These could be further decomposed into 285 fine clusters distinguished by the anatomy of smaller neurites. Follow-up analyses were directed toward four classes of input neurons, for which we could postulate a functional role: MBONs; lateral horn (LH)-associated neurons that include lateral horn output neurons (LHONs); subsesophageal output neurons (SEZONs), potential gustatory projection neurons that ascend from the SEZ; and OTHERS, a variety of neurons conveying information from other brain areas (Figures 2I and 2J).

Annotating the DAN clustering with input neuron identity revealed that the γ5β'2a MBONs specifically provide feedback (fb) input from the MB to our previously defined PAM-γ5(d) subtype (Figure 2K). We therefore renamed PAM-γ5(dd) neurons as PAM-γ5(fb). Other MBONs also provide selective input to different DANs. In contrast, as a group the SEZONs and LHONs provide input to the PPL1-γ1pdec DAN and all PAM-γ5 and PAM-β'2a DANs, although the relative proportions vary considerably.

Toward assigning functional relevance to the identified input pathways, we used nBLAST [18] to screen traced neuronal skeletons against a collection of confocal volumes of GAL4 and split-GAL4 lines for those that potentially drive expression

Figure 2. Input Specificity to Dopaminergic Neurons Matches Anatomical Subtypes
(A) Representation of all 821 input neurons to PAM-γ5, PAM-β'2a, and PPL1-γ1pdec DANs identified in this study. Cell bodies (black spheres) and processes (gray). DANs and the MB outline shown for reference (see Figure 1A).
(B) Venn diagram of unique and common input neurons to the analyzed DANs. PPL1-γ1pdec receives largely different input to PAM-γ5 and PAM-β'2a DANs. PAM-γ5 and PAM-β'2a DANs have many common inputs and some are also shared by PPL1-γ1pdec.
(C) Pie charts showing percentage of postsynaptic budget occupied by shared and unique input neurons to PAM-γ5 and PAM-β'2a DANs. Percentage of shared inputs across all three groups is 16%, 15%, and 8% for PPL1-γ1pdec, PAM-β'2a, and PAM-γ5 DANs, respectively.
(D) Bar chart showing DANs have many inputs with very low edge weight and each representing a small fraction of their overall postsynaptic budget. Inputs contributing more of the postsynaptic budget (to the right of the graph) are more abundant for PAM-γ5 and PAM-β'2a DANs; PPL1-γ1pdec distribution is strongly left shifted (bars show mean ± SD).
(E) DANs can be clustered by input connectivity (rows correspond to F). Heatmap shows every DAN has a group of unique input neurons represented by unique left shifted (bars show mean ± SD).
(F) A matrix where DANs are grouped by the similarity of their input connectivity (comparison to null model, p < 0.0001; see Methods S1).
(G) Representation of traced input neurons labeled using the unique and common input anatomy determined in (B) (see also Figures S2A–S2D).
(H) Tanglegram comparing DAN clustering by morphology (from Figure 1B) and clustering by input connectivity (left of E). Connectivity and morphology are not significantly independent of each other (Pearson’s correlation between the corresponding distance matrices, r = 0.864; Mantel test, p < 10⁻¹⁵; pα < 10⁻⁷ within only γ5 or β'2a group).
(I) DAN input neurons clustered by morphology. Dendrogram below shows single neurons allocated to 20 major coarse clusters based on soma position and primary neurite tract. Approximate neuropil of origin is indicated: antennal lobe (AL), SMP, SEZ, LH/SIP, and SMP/SIP are marked. Many neurons originate from less explored neuropils (misc).
(J) Fine clusters of exemplary neurons for the MBON, LHON, SEZON, and OTHERS classes of DAN inputs.
(K) Bar plot showing respective number of MBON, LHON, SEZON, and OTHERS inputs to individual DANs, ordered according to cluster identity (Figure 1). In general, PAM-γ5 DANs receive about 35% of their input from SEZONs, and about 20% from LHONs. Only the 3 PAM-γ5(fb) DANs receive significant direct input from MBON-γ5(2a) (green segments), PAM-β'2a DANs receive about 15% from SEZONs and 35% from LHONs. One PAM-β'2a DAN also receives minor direct input from MBON-γ5(2a). PPL1-γ1pdec DANs receive roughly equal LHON and SEZON input.
(L) NBLAST compares CATMAID generated neuronal skeletons from FAFB to neurons labeled in confocal images of GAL4 expression patterns. See also Figure S2, Methods S1, and Video S3.
Figure 3. Functional Analyses of DAN Input Neurons

(A) Forty-nine GAL4 driver lines with identified DAN input neurons were used to drive UAS-CsChrimson and screened for memory implantation by pairing neuronal activation with odor exposure. Flies were starved 18–26 h prior to training and tested for immediate memory performance. Lines emphasized in this study (mean ± SEM; individual data points are displayed as dots, either P.I. > 0.1 or P.I. < −0.1, or connecting from a neuropil of prior interest) are labeled (Figure S3A, fully labeled version).

B, C, D

E, F

G

Connectivity to PPL1-γ1pedc dendritic compartments

Legend continued on next page.
in the relevant neurons [19–21] (Figure 2L). We identified >50 driver lines with putative expression in the SEZONs, LHN01, MBONS, and OTHERS groups of DAN inputs.

**Functional Analyses of Input Pathways to DANs**

Pairing odor exposures with optogenetic activation of PPL1-γ1pedc or PAM-γ5, PAM-β2a DANs can produce either aversive or appetitive odor memories, respectively [22]. We therefore assumed that neurons providing significant input to reinforcing neurons should generate similar phenotypes if artificially engaged, instead of the relevant DANs. We combined the GAL4 driver lines with the red-light activated UAS-CsChrimson [23] optogenetic trigger and screened them for their potential to reinforce olfactory memories (Figures 3A and S3A). Whereas activation of some GAL4 lines produced appetitive odor memories, others produced aversive memories, and some had no consequence. We next correlated the identity of neurons labeled in each GAL4 line with their implanted memory performance and their respective DAN connectivity (Figure 3B; Video S3; Data S1). These correlations revealed good concordance with the valence of the memories formed by MBONS and SEZONs (see below and Figures 3E, 3F, and S3B). Three different LHON types connect to γ5 DANs and not PPL1-γ1pedc DANs and form appetitive memories, LHON01, LHON02, and LHON-AD1b2 [24, 25] (Figure 3A), whereas the OTHERS15 line formed aversive memory and preferentially synapsed with PPL1-γ1pedc (Figures 3A and 3B).

**MBON-DAN Connectivity**

The γ5β2a and γ4γ5 MBONS both have a dendrite in the γ5 compartment [3, 11]. In the stimulus replacement screen, these MBONS appeared to convey opposite valence. Whereas MBON-γ5β2a activation formed appetitive memory, MBON-γ4γ5 reinforced aversive memory (Figure 3A; confirmatory 30 min memory experiment in Figure S3B). Connectivity supported these behavioral results. MBON-γ4γ5 is the strongest MBON input to PPL1-γ1pedc, but does not connect to PAM-γ5 or PAM-β2a DANs, while MBON-γ5β2a synapses onto PAM-γ5(bf) and a PAM-β2a DAN, but not PPL1-γ1pedc DANs (Figures 3B, 3C, 3S3, and S3J).

Tracing of DAN inputs also identified selective connectivity with a few other MBONS, some of which were unknown (Figure 3C). MBON-α2 was found to synapse onto PAM-β2a and PAM-γ5(ba) and γ5(uc) DANs, MBON-β2mp with PAM-γ5(uc), MBON-γ2α’1 with PAM-β2a, and MBON-γ3β’1 with PAM-β2a and PAM-γ5(uc). With few exceptions, the relatively sparse connectivity of MBONS to our traced DANs was largely maintained when other traced neurons were included as potential interneurons between them (Figure 3D). Most notably, the apparent bias of connection of MBON-γ4γ5 to PPL1-γ1pedc and selectivity of MBON-γ3β’1 remained and new selective clusters to PAM-γ5(bf), PAM-γ5(uc), and PAM-β2a groups became apparent. No indirect connections were detected between MBON-γ4γ5 and PPL1-γ1pedc in our dataset. Moreover, only one additional β2a DAN emerged downstream of MBON-γ4γ5 when putative indirect connectivity was considered. In contrast, although MBON-γ5β2a was directly connected to only the PAM-γ5(bf) DANs, it was indirectly connected to most of the traced DANs, and all of the inputs to the unique PAM-γ5(ba) neuron come from neurons downstream of MBON-γ5β2a (Figure 1G). Axo-axonic synapses are frequent (>11,000) in the DAN input network. For example, MBONS frequently make reciprocal synapses on the axons of SEZONs and LHN01s and the different classes of DAN input neurons are also highly interconnected within cluster (317).

**SEZON-DAN Connectivity**

Synthetic activation of the SEZON lines also produced different learning phenotypes. Activating SEZON01 neurons formed appetitive memory and these connect to PPL1-γ1pedc and some PAM-γ5, but not to γ5 DANs reinforcing sugar memory (see below). Stimulating SEZON03 neurons formed appetitive memory and these SEZONs synapse onto PAM-γ5, but not PPL1-γ1pedc. SEZON02 neuron activation did not implant significant memory of either valence and appears to connect weakly to all three classes of traced DANs (Figures 3A, 3B, and S3K).

Despite their specificity, we expect some of our identified GAL4 drivers to express in our traced neurons of interest, and additional similar neurons in a fascicle. For example, a SEZON line could label a collection of ascending neurons representing both tasteful and distasteful gustatory stimuli [26]. Labeling such a mixed population with contradictory value could explain the inability of SEZON2 to reinforce a memory with clear valence. We therefore used the dominant temperature-sensitive UAS-Shibirets1 transgene [27] to block neurotransmission from SEZONs during training with sugar or bitter taste reinforcement (Figures 3E, 3F, and S3E–S3J). We included R58E02-GAL4 as a positive control for sugar memory, which expresses in the majority of PAM DANs [28], and MB320C-GAL4 for bitter learning because it labels PPL1-γ1pedc [29]. Blocking R58E02 or

(B) Connectivity matrix between DANs ordered according to morphological cluster identity and neurons labeled in 10 GAL4 lines, corresponding to 11 input clusters (MBON-γ5β2a and MBON-γ4γ5, SEZON01-03, LHON01-02, LHON-AD1b2, and OTHERS15-16). Valence of memory formed is reflected by input connectivity.

(C) Direct MBON-DAN connectivity matrix. We identified several MBONS to provide input to specific DANs (note: we traced all inputs to 7 PAM-γ5, and 2 β2a DANs with extensive review; Figure S2; Methods S1). Numbers indicate total synapse counts between MBONS and DANs.

(D) Adding other traced DAN input neurons creates potential for indirect connectivity between some MBONS and specific subsets of DANs. Indirect connectivity matrix showing the number of DAN input neurons that are downstream of MBONS with at least 3 synapses between each. Columns are normalized by their sum.

(E) Olfactory learning with bitter (DEET) reinforcement. Schematic: experimental timeline and temperature shift protocol. Blocking neuron output during training abolished 30 min appetitive memory specifically in SEZON03-GAL4; UAS-Shiβ1 and R58E02-GAL4; UAS-Shiα1 files (mean ± SEM, p < 0.0241 and 0.0089, respectively; one-way ANOVA, with Dunnett’s post hoc test, n = 10).

(F) Olfactory learning with bitter (DEET) reinforcement. Schematic: experimental timeline and temperature shift protocol. Blocking neuron output during training impaired immediate appetitive memory in MB320C-, SEZON01-, and SEZON02-GAL4; UAS-Shiβ1, but not in SEZON03: UAS-Shiα1 files (mean ± SEM, p < 0.0009, 0.0344, and 0.0170, respectively; one-way ANOVA with Dunnett’s post hoc test, n = 12).

(G) Connectivity matrix to specific branches of the PPL1-γ1pedc dendrite reveals classes of input neurons have branch specificity.

See also Figure S3, Data S1, and Video S3.
Figure 4. Aversive Memory Extinction and Sugar Learning Require Different Subsets of PAM-γ5 DANs

(A) Brain from a VT006202-GAL4; UAS-GFP fly labels all 20 PAM-γ5 DANs and possibly some other PAM DANs (black). Three commissures are visible (asterisk). Brain co-stained with nc82 antibody (MB is outlined). Scale bar, 20 μm.

(B) MB315C-GAL4; UAS-GFP specifically labels 8 PAM-γ5 DANs per hemisphere that cross the midline in the lower commissure.

(C) 0804-GAL4 labels 5 PAM-γ5 DANs per hemisphere, previously “γ5 narrow,” that occupy the lower commissure. Scale bar, 20 μm.

(D) 0104-GAL4; UAS-GFP labels PAM-γ5 DANs, previously named “γ5 broad,” that cross the midline in the upper and middle commissures. 0104 also labels some other PAM DANs [9].

(E) Table summarizing DAN expression in GAL4 lines used for behavior, modified from [9]. R48B04GAL80 refines the 0104-GAL4 expression [12], shown in the Data S1.

(F) Aversive olfactory memory extinction. Schematic: experimental timeline and temperature shift protocol. Blocking neuron output during odor re-exposure impaired memory extinction in R58E02-, VT006202-, and 0804-GAL4; UAS-Shi[T1]. but not in MB315C- or 0104-GAL4 ± GAL80; UAS-Shi[T1] flies. Bars show mean ± SEM. Asterisks denote p < 0.035 (wild-type) and p < 0.0176 (0104); one-way ANOVA with Tukey’s post hoc test, n = 10–12.

(G) Olfactory learning with sucrose reinforcement. Schematic: experimental timeline and temperature shift protocol. Blocking neuron output during training impaired 30 min appetitive memory in R58E02-, VT006202-, and 0104-GAL4 ± GAL80; UAS-Shi[T1], but not MB315C- or 0804-GAL4; UAS-Shi[T1] flies. Bars show

(legend continued on next page)
SEZON03 neurons during training abolished 30 min memory re-inforced with sugar, but blocking SEZON01 and SEZON02 neurons had no impact. In contrast, blocking MB320C, SEZON01, or SEZON02, but not SEZON03, neurons impaired 30 min memory after bitter learning. These data support a role for SEZONs in relaying positive and negative gustatory valence to DAN subtypes.

Analyzing the location of identified inputs to PPL1-γ1pedc confirmed that it receives branch-specific information (Figures 3G, S3J, and S3K; Video S4). Whereas all aversively reinforcing input from SEZONs connects to the SMP arbor, OTHERS15 neurons, which can also produce aversive learning, connect to the PPL1-γ1pedc arbor in the ventral and dorsal crepine (CRE) [17]. MBONs in general and the strong MBON-γ5y input also mostly connect to the PPL1-γ1pedc CRE, and CREy branches. Since the CRE branches are closest to the primary axon, input from other MB compartments may be particularly salient to PPL1-γ1pedc (Figure S3J). Interestingly, the strongest input from MBON-γ5y2a to the PAM-γ5fb DANs is similarly placed on the DAN dendrite (Figure S4A).

Functional Subdivision of PAM-γ5 DANs

Based on prior findings, we hypothesized that PAM-γ5(fb) DANs, which receive recurrent feedback from MBON-γ5y2a, would be required for memory revaluation [11] and other PAM-γ5 DANs receiving input from SEZONs labeled by SEZON03 would be required to reinforce sugar memory [9]. Testing this model required locating GAL4 drivers that label γ5 DAN subsets that at least partially correspond to functionally relevant subtypes. We therefore used commissure crossing to select GAL4 drivers expressing in subsets of PAM-γ5 DANs [9]. We reasoned that drivers labeling the lower commissure might express in γ5(fb)-DANs (Figure 1E) while others labeling the upper commissure could include γ5-DANs connected to sugar-selective SEZONs. We identified VT06202-GAL4 (Figure 4A), which expresses in γ5-DANs in all commissures; MB315C- (Figure 4B) and 0804-GAL4s (Figure 4C), which express in 8 and 3–5 γ5 DANs, respectively, in the lower commissure; and 0104-GAL4 (Figure 4D), which only expresses in upper commissure γ5 DANs [2, 9, 21, 30]. We also used genetic intersection with GAL80 to restrict 0104-GAL4 expression to γ5broad and β2m (Figure 4E; [9, 12]).

We next tested whether blocking the neurons labeled in these GAL4s with UAS-ShiR1 disrupted aversive memory extinction and/or sugar learning (Figures 4F and 4G). The PAM DAN expressing R58E02-GAL4 served as control. To assay extinction of aversive memory (Figures 4F and S4C), flies were differentially conditioned by pairing one of two odors with shock [31]. Then 30 min after training they received two un-reinforced exposures of the previously shock paired odor (CS+) or the other odor (CS−) at 15 min interval [11]. They were then tested 15 min later for olfactory memory. As previously established, only CS+ re-exposure diminished memory performance, demonstrating odor-specific memory extinction. We blocked subsets of DANs specifically during odor re-exposure by training flies at permissive 25°C, transferring them to restrictive 32°C immediately after training, then returning them to 25°C after the second odor re-exposure. Blocking R58E02, VT06202, and 0804 neurons abolished memory extinction, whereas memory was still extinguished in flies with blocked MB315C or 0104 neurons (+ and − GAL80). No extinction was observed in any line when flies were re-exposed to the CS− odor after training. These data support a role for the 0804-GAL4 group of lower commissure PAM-γ5(fb) DANs in memory extinction (Figure 4F).

In contrast, when these neurons were selectively blocked during sugar conditioning [32] (Figures 4G, S4D, and S4E), 30 min memory was impaired in R58E02, VT06202, and 0104 (+ and − GAL80) flies expressing UAS-ShiR1, but was unaffected in UAS-ShiR1-expressing MB315C and 0804 flies (Figure 4G). These data demonstrate that memory extinction and sucrose reinforcement are dissociable in the γ5 DANs and support the selective role for PAM-γ5(fb) DANs in memory extinction and SEZON-connected PAM-γ5 DANs in sucrose reinforcement (Figure 4H). We therefore propose that other PAM-γ5 DAN subtypes may serve different reward-related functions.

The morphologically distinct γ5 DAN subtypes innervate different regions of the γ5 compartment where they could depress or potentiate different parts of the KC-MBON network (Figures 1C–1G and S4B) [29, 33–35]. We do not currently understand the full relevance of the sub-compartment architecture. However, since the γlobe dorsal (γd) KCs carry visual information and the main γ KCs are olfactory [36], connections of these two streams of KCs to γ5 MBONs could be independently modified by PAM-γ5(da) (Figure 1G) and PAM-γ5(fb) DANs (Figure 1E), whose processes are confined to the respective subregions of the γ5 compartment (Figure S4F). DAN stratification may therefore maintain modality specificity of olfactory memory revaluation [11]. It is interesting to note that larvae only have one DAN per MB compartment [37] and that multiple DANs per compartment are an adult-specific specialization [2]. We expect this expansion reflects the additional behavioral demands of the adult fly [13, 38], and our work here suggests the larger number of DANs in each compartment provides additional functional capacity to the compact anatomy of the adult MB. We propose that the elaboration and specialization of γ5 DANs may permit the adult fly to individually represent the values of a broad range of rewarding events.

STAR METHODS

Detailed methods are provided in the online version of this paper and include the following:

- KEY RESOURCES TABLE
- RESOURCE AVAILABILITY
  - Lead Contact
  - Materials Availability
  - Data and Code Availability

mean ± SEM. Asterisks denote p < 0.0003 (R58E02), p < 0.0004 (0104), p < 0.0018 (VT06202), and p < 0.0010 (R48B04GAL80; 0104-GAL4); one-way ANOVA with Dunnett’s post hoc test, n = 10.

(H) Schematic of input pathways to PAM-γ5 and PAM-γ2a DANs and subtype innervation of the γ5 compartment. Colors correspond to clusters defined in Figure 1. Open circles represent γ5 DANs that were identified but not further analyzed.

See also Figure S4, Data S1, and Video S4.
Author Contributions

(789274). To S.W., G.S.X.E.J., D.D.B., and G.M.R. S.W. is also supported by a Wellcome Collaborative Award of the Waddell and Jefferis groups contributed to discussion throughout this project. We are grateful to Yaling Huang and Suewei Lin for confocal imaging during the COVID-19 shutdown in the UK. We also thank Marion Sillies, Tom Clandinin, Wes Grueber, and Wolf Huetteroth for access to confocal stacks. Members of the Waddell and Jefferis groups contributed to discussion throughout this project. This work was largely funded by a Wellcome Collaborative Award (203261/Z/16/Z) to S.W., G.S.X.E.J., D.D.B., and G.M.R. S.W. is also supported by a Wellcome Principal Research Fellowship (200846/Z/16/Z), the Gatsby Charitable Foundation (GAT3237), and an ERC Advanced Grant (789274).

Using BioCite, we can cite references in the text as follows:

1. Cognigni, P., Felsenberg, J., and Waddell, S. (2018). Do the right thing: neural network mechanisms of memory formation, expression and update in Drosophila. Curr. Opin. Neurobiol. 49, 51–58.

2. Aso, Y., Hattori, D., Yu, Y., Johnston, R.M., Iyer, N.A., Ngo, T.T., Dionne, H., Abbott, L.F., Axel, R., Tanimoto, H., and Rubin, G.M. (2014). The neuronal architecture of the mushroom body provides a logic for associative learning. eLife 3, e04577.

3. Aso, Y., Sitaraman, D., Ichinose, T., Kaun, K.R., Vogt, K., Belliart-Guerin, G., Placais, P.Y., Robie, A.A., Yamagata, N., Schnaitmann, C., et al. (2014). Mushroom body output neurons encode valence and guide memory-based action selection in Drosophila. eLife 3, e04580.

4. Zheng, Z., Lauterzen, J.S., Perlman, E., Robinson, C.G., Nichols, M., Milkie, D., Torrens, O., Price, J., Fisher, O.B., Shariff, N., et al. (2018). A complete electron microscopy volume of the brain of adult Drosophila melanogaster. Cell 174, 730–743.e22.

5. Krashes, M.J., DasGupta, S., Vrede, A., White, B., Armstrong, J.D., and Waddell, S. (2009). A neural circuit mechanism integrating motivational state with memory expression in Drosophila. Cell 139, 416–427.

6. Aso, Y., Siwanowicz, I., Bracker, L., Itt, K., Kitamoto, T., and Tanimoto, H. (2010). Specific dopaminergic neurons for the formation of labile aversive memory. Curr. Biol. 20, 1445–1451.

7. Kelemen, K., Vrontou, E., Krützner, S., Yu, J.Y., Kurtovic-Kozaric, A., and Dickson, B.J. (2012). Dopamine neurons modulate phenome responses in Drosophila courtship learning. Nature 489, 145–149.

8. Zhao, X., Lenek, D., Dag, U., Dickson, B.J., and Kelemen, K. (2018). Persistent activity in a recurrent circuit underlies courtship memory in Drosophila. eLife 7, e31425.

9. Huetteroth, W., Perisse, E., Lin, S., Klappenbach, M., Burke, C., and Waddell, S. (2015). Sweet taste and nutrient value subdivide rewarding dopaminergic neurons in Drosophila. Curr. Biol. 25, 751–758.

10. Yamagata, N., Ichinose, T., Aso, Y., Placais, P.Y., Friedrich, A.B., Sima, R.J., Preat, T., Rubin, G.M., and Tanimoto, H. (2015). Distinct dopamine neurons mediate reward signals for short- and long-term memories. Proc. Natl. Acad. Sci. USA 112, 578–583.

11. Felsenberg, J., Jacob, P.F., Walker, T., Barnstedt, O., Edmondson-Stait, A.J., Pleijzier, M.W., Otto, N., Schlegel, P., Shariff, N., Perisse, E., et al. (2018). Integration of parallel opposing memories underlies memory extinction. Cell 175, 709–722.e15.

12. Lin, S., Oswald, D., Chandra, V., Talbot, C., Huetteroth, W., and Waddell, S. (2014). Neural correlates of water reward in thirsty Drosophila. Nat. Neurosci. 17, 1536–1542.

13. Senapati, B., Tsao, C.H., Yuan, J.A., Chiu, T.H., Wu, C.L., Waddell, S., and Lin, S. (2019). A neural mechanism for deprivation state-specific expression of relevant memories in Drosophila. Nat. Neurosci. 22, 2029–2039.

14. Lewis, L.P., Sij, K.P., Aso, Y., Friedrich, A.B., Butteel, A.J., Rubin, G.M., and Grunwald Kadow, I.C. (2015). A higher brain circuit for immediate
integration of conflicting sensory information in Drosophila. Curr. Biol. 25, 2203–2214.

15. Galili, D.S., Dylla, K.V., Lüdke, A., Friedrich, A.B., Yamagata, N., Wong, J.Y., Ho, C.H., Szafrak, P., and Tanimoto, H. (2014). Converging circuits mediate temperature and shock aversive olfactory conditioning in Drosophila. Curr. Biol. 24, 1712–1722.

16. Das, G., Klappenbach, M., Vrontou, E., Perisse, E., Clark, C.M., Burke, C.J., and Waddell, S. (2014). Drosophila learn opposing components of a compound food stimulus. Curr. Biol. 24, 1723–1730.

17. Ito, K., Shinomiya, K., Ito, M., Armstrong, J.D., Boyan, G., Hartenstein, V., Harzsch, S., Heisenberg, M., Homberg, U., Jenett, A., et al.; Insect Brain Name Working Group (2014). A systematic nomenclature for the insect brain. Neuron 81, 755–765.

18. Costa, M., Manton, J.D., Ostrovsky, A.D., Prohaska, S., and Jefferis, G.S. (2016). NBLAST: rapid, sensitive comparison of neuronal structure and construction of neuron family databases. Neuon 91, 293–315.

19. Gohi, D.M., Silles, M.A., Gao, X.J., Bhalerao, S., Luongo, F.J., Lin, C.C., Poter, C.J., and Clandinin, T.R. (2011). A versatile in vivo system for directed dissection of gene expression patterns. Nat. Methods 8, 231–237.

20. Jenett, A., Rubin, G.M., Ngo, T.T., Shepherd, D., Murphy, C., Dionne, H., Pfeiffer, B.D., Cavallaro, A., Hall, D., Jeter, J., et al. (2012). A GAL4-driver line resource for Drosophila neurobiology. Cell Rep. 2, 991–1001.

21. Tirian, L., and Dickson, B.J. (2017). The VT GAL4, LexA, and split-GAL4 driver line collections for targeted expression in the Drosophila nervous system. bioRxiv. https://doi.org/10.1101/198648.

22. Aso, Y., and Rubin, G.M. (2016). Dopaminergic neurons write and update memories, diverse behavioural functions, and interactions with the mushroom body. J. Neurobiol. 88, 89–98.

23. Schlegel, C.M., Gerhard, S., Longair, M., Kazimiers, T., Li, F., Pietzsch, T., Preibisch, S., Rueden, C., Saalfeld, S., Schmid, B., et al. (2012). Neurogenetic dissection of the Drosophila lateral horn reveals major output pathways, diverse behavioural functions, and interactions with the mushroom body. eLife 5, e03079.

24. Harzsch, S., Heisenberg, M., Homberg, U., Jenett, A., et al.; Insect Brain Name Working Group (2014). A systematic nomenclature for the insect brain. Neuron 81, 755–765.

25. Owen, D., Felsenberg, J., Talbot, C.B., Das, G., Perisse, E., Huetteroth, W., and Waddell, S. (2015). Activity of defined mushroom body output neurons underlies learned olfactory behavior in Drosophila. Neuron 86, 417–427.

26. Cohn, R., Morantte, I., and Rut, V. (2015). Coordinated and compartmentalized neuromodulation shapes sensory processing in Drosophila. Cell 163, 1742–1755.

27. Potter, C.J., and Clandinin, T.R. (2011). A versatile in vivo system for spatial manipulation of neuronal function by combinatorial restriction of transgene expression. Neuron 72, 352–363.

28. Li, P.H., Lindsey, L.F., Januszewski, M., Zheng, Z., Bates, A.S., Taisz, I., Tyka, M., Nichols, M., Li, F., Perlman, E., et al. (2019). Automated reconstruction of a serial-section EM Drosophila brain with flood-filling networks and local realignment. bioRxiv. https://doi.org/10.1101/605634.
51. Prokop, A., and Meinertzhagen, I.A. (2006). Development and structure of synaptic contacts in Drosophila. Semin. Cell Dev. Biol. 17, 20–30.

52. Butcher, N.J., Friedrich, A.B., Lu, Z., Tanimoto, H., and Meinertzhagen, I.A. (2012). Different classes of input and output neurons reveal new features in microglomeruli of the adult Drosophila mushroom body calyx. J. Comp. Neurol. 520, 2185–2201.

53. Rousseeuw, P.J. (1987). Silhouettes: a graphical aid to the interpretation and validation of cluster analysis. Comput. Appl. Math. 20, 53–65.

54. Legendre, P., and Legendre, L. (2012). Numerical Ecology, Third Edition (Elsevier).

55. Fowlkes, E.B., and Mallows, C.L. (1983). A method for comparing two hierarchical clusterings. J. Am. Stat. Assoc. 78, 553–569.
## STAR METHODS

### KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Antibodies          |        |            |
| Anti-GFP            | Abcam  | RRID: AB_300798 |
| Alexa Fluor Goat anti-mouse | Invitrogen | RRID: AB_2633275 |
| nc82                | Developmental Studies Hybridoma Bank | RRID: AB_2314866 |
| Chemicals, Peptides, and Recombinant Proteins | | |
| Sucrose             | Sigma-Aldrich | Cat# S0389 |
| Mineral Oil         | Sigma-Aldrich | Cat#M5904 |
| 4-methylcyclohexanol (98%) | Sigma-Aldrich | Cat#218405 |
| 3-octanol (99%)     | Sigma-Aldrich | Cat#153095 |
| Retinal             | Sigma-Aldrich | Cat#R2500 |
| DEET                | Sigma-Aldrich | Cat#36542 |
| Xylose              | Sigma-Aldrich | Cat#X1500 |
| Normal goat serum   | Invitrogen | Cat#16210064 |
| VectaShield         | VectorLabs | SKU#H-1700-10 |
| Deposited Data      |        |            |
| Greyscale EM Data and raw neuronal traces | [4] | https://catmaid.virtualflybrain.org |
| Skeletonized traces – will be deposited on | This paper | https://v2.virtualflybrain.org; http://neuromorpho.org/ |
| Experimental Models: Organisms/Strains | | |
| Drosophila: Canton-S | Waddell Lab | N/A |
| Drosophila: R58E02-GAL4 | FlyLight, Bloomington Drosophila Stock Center (BDSC) | RRID: BDSC_41347 [20, 28] |
| Drosophila: MB504B-GAL4 | FlyLight, BDSC | RRID: BDSC_68329 [2, 20, 22] |
| Drosophila: MB315C-GAL4 | FlyLight, BDSC | RRID: BDSC_68316 [2, 20, 22] |
| Drosophila: MB320C-GAL4 | FlyLight, BDSC | RRID: BDSC_68253 [2, 20, 22] |
| Drosophila: 0273-GAL4 | InSITE collection | Waddell lab [19, 30] |
| Drosophila: MB077B-GAL4 | FlyLight, BDSC | RRID: BDSC_68283 [2, 20, 22] |
| Drosophila: VT006202-GAL4 | Vienna Tiles Project | RRID: BDSC_72445 [8, 21] |
| Drosophila: 0104-GAL4 | InSITE collection | RRID: BDSC_62639 [19, 30] |
| Drosophila: 0804-GAL4 | InSITE collection | RRID: BDSC_6325 [9, 19] |
| R48B04-GAL80; 0104-GAL4 | FlyLight/ InSITE | Waddell lab [12] |
| Drosophila: GMR88F08-GAL4 | FlyLight, BDSC | RRID: BDSC_47982 |
| Drosophila: GMR88E01-GAL4 | FlyLight, BDSC | RRID: BDSC_48397 |
| Drosophila: GMR66C08-GAL4 | FlyLight, BDSC | RRID: BDSC 49412 [11, 20] |
| Drosophila: SS46348-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: SS28305-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: SS27959-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: SS52050-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: SS45245-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: SS45251-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: SS52046-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: S49970-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: SS45221-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: SS47568-GAL4 | Ito/Rubin labs | unpublished data |

(Continued on next page)
| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Drosophila: SS49941-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: SS39616-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: SS27580-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: SS28308-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: SS21077-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: SS45222-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: SS42968-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: SS46613-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: SS35902-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: SS51316-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: SS31522-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: SS21206-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: SS31827-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: SS39070-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: SS25028-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: SS24588-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: SS27978-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: SS46337-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: SS32768-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: SS36651-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: SS46596-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: SS42983-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: SS24252-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: SS21208-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: SS32151-GAL4 | Ito/Rubin labs | unpublished data |
| Drosophila: LH991-GAL4 | Jefferis Lab | [24] |
| Drosophila: LH2241-GAL4 | Jefferis Lab | [24] |
| Drosophila: LH989-GAL4 | Jefferis Lab | [24] |
| Drosophila: LH1395-GAL4 | Jefferis Lab | [24] |
| Drosophila: LH2278-GAL4 | Jefferis Lab | [24] |
| Drosophila: LH1538-GAL4 | Jefferis Lab | [24] |
| Drosophila: LH1539-GAL4 | Jefferis Lab | [24] |
| Drosophila: LH1396-GAL4 | Jefferis Lab | [24] |
| Drosophila: UAS-Shibire™ | BDSC | RRID: BDSC_44222 [27] |
| Drosophila: UAS-CsChrimson.mVenus | BDSC | RRID: BDSC_55135 [23]; RRID: BDSC_55136 |

Software and Algorithms

- **Blender 3D v.2.79**: Blender Online Community [https://www.blender.org/](https://www.blender.org/)
- **Natverse** [39]: [https://github.com/natverse/nat/](https://github.com/natverse/nat/)
- **Dendoextras** [40]: [https://github.com/jefferis/dendoextras/](https://github.com/jefferis/dendoextras/)
- **R**: R Development Core Team [http://www.R-project.org/](http://www.R-project.org/)
- **Rcatmaid**: Jefferis and Cardona labs [https://github.com/natverse/rcatmaid](https://github.com/natverse/rcatmaid)
- **CATMAID** [41, 42]: [https://catmaid.readthedocs.io/en/stable/index.html#](https://catmaid.readthedocs.io/en/stable/index.html#)
- **Python**: Python Software Foundation [https://www.python.org/](https://www.python.org/)
- **PyMaid**: Philipp Schlegel [https://github.com/schlegelp/PyMaid](https://github.com/schlegelp/PyMaid)
- **Dendrogram code**: Markus Pleijzier [https://github.com/markuspleijzier/AdultEM/tree/master/Dendrogram_code](https://github.com/markuspleijzier/AdultEM/tree/master/Dendrogram_code)
- **Graphviz** [43]: [https://www.graphviz.org](https://www.graphviz.org)
RESOURCE AVAILABILITY

Lead Contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Scott Waddell (scott.waddell@cnbc.ox.ac.uk).

Materials Availability
The new split-GAL4 Drosophila lines described in this study were produced by Masayoshi Ito. They are available on request from the Lead Contact and will be sent from the Waddell lab or from the Janelia Research Campus, via K. Ito and G. M. Rubin.

Data and Code Availability
The datasets and code used for analyses in R and Python are mostly available through public repositories as indicated in this Methods section of the manuscript. Any other code is available on request and without restriction. Neuronal morphologies and connectivity data will be publicly available through VFB: https://v2.virtualflybrain.org/org.geppetto.frontend/geppetto and NeuroMorpho.org: http://neuromorpho.org/ and neuronal skeletons can be requested from the authors in .swc format.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Fly strains
All Drosophila strains were raised at 25°C on standard cornmeal-agar food at 50%–60% relative humidity in a 12:12 h light:dark cycle. For experiments, female flies bearing effector transgenes were crossed to male flies bearing GAL4 drivers. For driver line details see Key Resources Table and Data S1. Driver lines were obtained from the Janelia FlyLight collection [20] or split-GAL4 [48] collections, the Vienna Tiles Project [21], and the InSITE collection [19]. New split-GAL4 ‘SS’ lines were created in the Ito/Rubin labs (unpublished data). UAS-CsChrimson [23] and UAS-Shibirets [27] effectors were used to stimulate and block specific neurons.

METHOD DETAILS

Neuron reconstruction - ‘tracing’ in FAFB
Neurons were reconstructed from a serial section transmission electron microscopy (ssTEM) volume of a full adult fly (D. melanogaster) brain (FAFB) [4] using CATMAID, a web-based software for collaborative neural circuit reconstruction from large image datasets (https://catmaid.readthedocs.io/en/stable/) [41, 42]. Consistent with previous studies [3, 37, 49], tracing followed the centerline of a neuron’s profiles through the dataset to reconstruct neurite morphology and annotate synaptic sites. We used an established and tested iterative approach [42] where initial reconstruction is followed by a systematic proofreading by at least two expert reviewers (> 500 h of tracing experience). We also took advantage of recent automatic segmentation efforts of the FAFB dataset [50], where flood-filling algorithms create volumetric segmentations of the EM data. These segmentations are then skeletonised to produce neuron fragments that can be joined together to expedite reconstruction. Human proof-reading is still required to remove incorrect merges of skeletons. In this study auto-segmentation was only used to aid the tracing of DAN input neurons to identification (see below).

Synaptic sites were identified based on three, previously described criteria [51] and reviewed as above: an active zone with (1) T-bar(s) and (2) surrounding vesicle cloud, and (3) a synaptic cleft to which all postsynaptic neurons must have access. In Drosophila, presynapses have been found on fine axonal processes [42], boutons [52], and other neurites that are neither in the dendritic nor the axonal field. Postsynapses have been found on large or fine dendritic processes and spine-like twigs that are shorter than 3 µm [42].

REAGENT or RESOURCE SOURCE IDENTIFIER

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| NetworkX           | [44]   | https://networkx.github.io/ |
| Dendextend         | [40]   | https://github.com/jefferis/dendroextras |
| Fiji                | NIH    | http://fiji.sc/ |
| GraphPad Prism 6   | GraphPad Software, La Jolla, CA | https://www.graphpad.com/scientific-software/prism/ |
| Adobe Illustrator CC | Adobe Systems, San Jose, CA | https://www.adobe.com/uk/products/illustrator.html |
| Vegan              | Vegandevs - Jari Oksanen | https://github.com/vegandevs/vegan |
| SciPy              | [46]   | https://www.scipy.org/ |
| Nbclust            | [47]   | https://rdrr.io/cran/NbClust/man/NbClust.html |

OPEN ACCESS

e3 Current Biology 30, 3200–3211.e1–e8, August 17, 2020
has been estimated that the tracing approach employed finds 99.8% of presynapse and 91.7% of postsynapses [42]. The probability of identifying false-positive postsynapses is 2.2% and negligible for presynapses.

**DAN identifying, tracing and quality control**

DANs were first identified by selecting potential profiles in the midline commissure between left and right hemisphere MBs. These profiles were traced until axonal branches could be identified in the MB compartment of interest. We exhausted all possible profiles between the two MB compartments and in doing identified, both PPL1-γ1pedc DANs, and right-hand side (RHS) β'2a DANs, and γ5 DANs. Although the number of γ5 DANs has been estimated to be between 8 and 21 [2] we nevertheless considered the possibility of unilateral γ5 DANs, which would not have a process extending across the midline. To do this we also sampled neural profiles in the descending tract where the processes of γ5 DANs enter the MB lobe. However, we did not identify additional γ5 DANs. Following identification DANs were traced and reviewed (as described above). Full details of tracing and review are provided in the Revision Status Table, Methods S1. In brief, the RHS PPL1-γ1pedc DAN was completed and subjected to standard expert review. The γ1pedc dendritic field was further extensively reviewed. 20 PAM-γ5 DANs on the RHS and 9 on the left were reconstructed and the RHS neurons received standard expert review strategy as described above. 7 PAM-γ5 DANs received further extensive reviewed of their dendritic field. Of the 4 PAM-β'2a DANs, 2 underwent both standard and additional extensive review, one only received standard review and a fourth was only partially reviewed. Any neurons that were not reviewed to this standard were excluded from the analyses. We note that it was more challenging to reconstruct DANs than many other neurons in the Drosophila brain. DAN dendrites are very thin and have a dark/granular texture, which increases the likelihood of missing branches and synapses. We therefore scrutinized completion and postsynapse annotation for 7 PAM-γ5 DANs (representing all morphological clusters), 2 PAM-β'2a DANs and the PPL1-γ1pedc DAN. Following this extended reconstruction and revision effort, we are confident that we have annotated all identifiable postsynapses on these selected DANs. Comparing data obtained from the regular review protocol to that from our extended review effort showed that regular review captured ~30% of the postsynapses on more than 60% of all cable. We also analyzed the placement of old (regular review) and new (added following extensive review) synapses, by measuring their geodesic (along-the-arbor) distance to the dendritic root (Figure S2H). This analysis showed that each round of additional review adds new synapses that are distributed along the arbor. Lastly, we assessed whether uneven tracing of input connectivity altered the clustering of DANs by randomly downsampling (see below) the 9 extensively reviewed neurons to a level of inputs traced for the other regularly reviewed neurons. DANs could be similarly clustered following the downsampling, demonstrating that our DAN clustering results are unlikely to vary greatly with additional tracing of more input neurons.

**Tracing neurons providing inputs to DANs**

When a postsynapse was annotated on a DAN, a single-(seminal) node profile was placed in the center of the presynaptic cell, unless a neuron or fragment was already present. To reconstruct upstream neurons from these seminal nodes we randomized the sampling order from each postsynapse within the total population on a neuron-by-neuron basis. For the reviewed PAM DANs (18 γ5 and 4 β'2a) we typically traced over 85% of the input neurons to identification from annotated postsynapses on a DAN arbor (see Revision Status Table, Methods S1). For the PPL1-γ1pedc DAN, we traced from 50% of annotated DAN postsynapses to identify the input neurons. Tracing inputs to this collection of γ5, β'2a and γ1pedc DAN postsynapses recovered 821 upstream neurons, some of which connect to multiple DANs in the traced groups. The tracing of the upstream neurons also varies in level of completeness but all neurons were traced to identify their microtubule containing backbone and were followed to a soma to retrieve their gross morphology.

**Data visualization**

Reconstructed neurons were visualized using Blender 3D, an open-source 3D software (https://www.blender.org/) or natverse [39] and RCatmaid packages in R (http://jefferis.github.io/rcatmaid/). Neuron data from CATMAID were imported using an existing CATMAID plugin for Blender (https://github.com/schlegelp/CATMAID-to-Blender [49]).

**3D Representations and Videos**

3D representations of traced skeletons that were obtained as swc files from CATMAID after reconstruction were either plotted using the natverse R tool box, or 3D representations were created and rendered with blender v2.79. Video footage of skeletons was rendered to obtain 3D representations with blender v2.79 and cut with Adobe Premiere Pro2020.

**Matching genetic driver lines to EM skeletons**

EM skeletons were matched to published library GAL4 lines using registered Janelia FlyLight micrograph data from VirtualFlyBrain (https://v2.virtualflybrain.org). NBLAST similarity matrices were calculated comparing both data types and top hits were visualized and manually cross compared. In case of new split-GAL4 SS lines created by Masayoshi Ito (unpublished data), imaging stacks were newly registered with bridging and mirror registrations from the natverse package [39].

**Generating flies for behavioral experiments**

UAS-CsChrimson [23] and UAS-Shibire [27] female flies were crossed to males from a collection of GAL4 and split-GAL4 lines (Data S1). For wildtype controls Canton-S flies were used. For heterozygous controls, GAL4 lines were crossed to Canton-S flies. All behavioral experiments used 4 to 9-day old mixed-sex flies.
Immunohistochemistry and imaging
Following [9], Brains were dissected in cold Phosphate Buffered Saline (PBS) and fixed in PBS with 4% paraformaldehyde at 25°C for 40 min. They were washed 3X by quick PBS exchange and 3X for 20 min each, in PBS containing 0.5% Triton X-100 (PBT), followed by 30 min incubation in PBT containing 5% normal goat serum. Brains were either then imaged for endogenous expression of GFP or the buffer was exchanged with anti-GFP (1:200; Abcam) and anti-nc82 (1:50; DSHB) antibodies in PBS containing 0.5% Triton X-100 (PBT). Brains were incubated for 24–72 h at 4°C, washed 3X 20 min in PBT at 25°C, followed by incubation in PBT containing the appropriate Alexa Fluor secondary antibodies (Invitrogen) overnight at 4°C. Brains were then washed 3X 20 min in PBT at 25°C, before being mounted on slides with VectaShield (Vector Labs). Imaging was performed using a Leica TCS SP5X confocal microscope. Maximum intensity projections of Confocal stacks were generated using FIJI [45].

QUANTIFICATION AND STATISTICAL ANALYSIS

Analysis of neuroanatomical features and DAN connectivity
All analyses were performed in R and Python using open-source software. PyMaid (https://github.com/schlegelp/PyMaid) and RCatmaid (https://github.com/natverse/rcatmaid; https://github.com/natverse/elmr) were used to interface with CATMAID servers and perform morphological analyses. Neuron analyses were performed with natverse functionality (https://github.com/natverse/nat) or custom-written code, which is available on request. Hierarchical clustering was performed using base R functions and dendrogram visualizations with dendroextras (https://github.com/jefferis/dendroextras). Tanglegrams were generated using dendextend (https://github.com/talgalili/dendextend) [40]. Mantel test analyses were computed with the vegan R package (https://github.com/vegandevs/vegan).

Clustering

Euclidean and Manhattan distance metrics for clustering
To analyze and draw conclusions from differences and similarities in large amounts of connectivity or morphology data, the information is represented in the form of distance matrices between each data point in space. Euclidean distance is the direct (bee-line) distance between two points in a Cartesian coordinate system. Manhattan distance between two data points in a Cartesian system is the sum of distances between the coordinates.

Ward’s clustering criterion
Ward’s method was used for agglomerative hierarchical clustering (part of R base package). Each datapoint starts in its own cluster and pairs of clusters are merged, moving up the hierarchy. At each step the pair of clusters with minimum within-cluster variance are merged. Connectivity data, as well as morphology data was clustered using Ward’s criterion to compare to clusters formed using average linkage criterion.

Average linkage clustering criterion
Average linkage (also known as unweighted pair group method with arithmetic mean, UPGMA) is another criterion for agglomerative hierarchical clustering. With average linkage clustering pairwise dissimilarities between each element in cluster 1 and 2 are computed and the average of these dissimilarities are considered as the distance between the two clusters. Clusters separated by the smallest distance are merged during clustering, moving up the hierarchy. Both morphology and connectivity data were clustered using average linkage to compare to data clustered with Ward’s criterion.

Clustering DANs by morphology
To compensate for different levels of completeness of tracing, DANs were simplified to their longest tree with 200 branch points (the minimum number of branch points throughout the PAM DAN population). Morphological similarity matrices were calculated using NBLAST [18]. Hierarchical clustering was primarily performed using base R functions, taking Euclidean distance matrices of similarity scoring, with average linkage clustering criterion. Morphology clustering was performed with Ward’s and average linkage criteria for comparison.

Clustering DANs by input connectivity
Connectivity information was retrieved from CATMAID after synapse annotation and upstream tracing of input neurons. Only neurons upstream of the dendritic region of DANs with > 50 sampled profiles were included in the analyses. Before clustering the number of synapses annotated on each DAN was normalized to reduce bias in clustering that could arise from the varying levels of tracing completeness and/or natural differences in the number of inputs to the different DANs. Hierarchical clustering was primarily performed using the Manhattan distance between upstream connectivity profiles of DANs with Ward’s clustering criterion. Connectivity data was also clustered using the average linkage criterion for comparison.

DAN PPL-γ1pedc postsynapse clustering
For PPL1-γ1pedc the x, y and z coordinates of postsynapses on the dendrite were clustered using Ward’s hierarchical clustering in the SciPy package (https://www.scipy.org/ [46]).

Silhouette method to determine accuracy of the number of clusters
Silhouette is a graphical representation of the quality of clustering across a range of potential values for k (the number of clusters). Where possible Silhouette was used to select an appropriate number of clusters (it was less useful for morphology clustering). This
method measures how similar observations are to their own cluster and how dissimilar to other clusters. The average silhouette width ranges from 0 and 1, with 1 indicating observations are well clustered. To validate DAN clustering the average silhouette width was calculated using the nbcust R package [47, 53].

**Input neuron morphology clustering**

Morphology clustering of upstream neurons was performed using hierarchical clustering with average linkage criterion. This involved a multi-step approach to account for varying levels of tracing and for the morphological diversity of 821 neurons (Methods S1). Coarse clustering was performed taking the soma tract as the primary feature of neuron identity. Subsequently the larger primary clusters were subclustered by splitting neurons into the primary neurite and its complement/remainder. Similarity matrices were calculated using NBLAST and an element-wise mean (80:20) was used for clustering. For fine clusters, weighting methods were selected iteratively depending on overall sub-cluster morphology.

**Tanglegrams to compare clustering of 2 feature spaces**

Tanglegrams were generated to visually compare clustering dendrograms produced by different criteria (e.g., average linkage versus Ward’s) or clustering based on morphology versus those produced using connectivity. Dendrogram layouts were determined to minimize edge crossing (i.e., minimize Manhattan distance between corresponding DANs) using dendextend [40].

**Mantel test to determine dependence of 2 feature spaces**

The Mantel test was used to compare 2 sample spaces - here neuron morphology distance matrices obtained from all-by-all NBLAST and distances based on connectivity were used. To create distance matrices for connectivity, connectivity matrices were normalized by the postsynaptic budget of DANs. The implementation of the Mantel test was based on [54]. Pearson’s correlation between the two observed datasets was calculated, then one of the matrices was shuffled $10^7$ times and each event tested for correlation with the observed data. The number of events where the correlation is higher than between the two original datasets was divided by the amount of comparisons ($10^7$) to create a p value. When p values were lower than the significance level, it was concluded that the null model of independence between the two feature spaces could be rejected (see Methods S1).

**Downsampling connectivity to verify reproducibility of clustering**

To verify that clustering into the observed DAN groups does not result from bias in the relative completeness of tracing of the input network to each DAN, down-sampled the datasets so that all DANs were randomly stripped of 5%-50% of their input connections. Clustering resulting from 10,000 repetitions of this down-sampling were then compared to the clustering obtained from the full dataset. In addition, to exclude that more exhaustive reviewing of a few exemplary DANs might skew the clustering, we created a dataset of 10,000 repetitions where the connectivity of only the exhaustively reviewed DANs was reduced to the average of all the remaining DANs. Clustering obtained with this normalized dataset was also compared to that retrieved using the full dataset.

**Cluster similarity analyses**

In 10,000 iterations of resampling a dataset with reduced connectivity, each neuron has a different likelihood to cluster with the same original group that it did in the 100% connectivity dataset or, with any of the other original groups. We therefore also calculated the average likelihood (over the 10,000 trials) that a downsampled DAN clustered with the same group that it clustered with in the full 100% connectivity dataset. These values were then plotted as a stacked bar plot.

**Fowlkes-Mallows Index (FMI)**

The Fowlkes-Mallows Index measures the similarity of the content between two different clusterings. The performance of the first clustering is compared to that of the second clustering (which is assumed to be perfect). Exact matches/good performance result in an $FMI = 1$ [55].

**Neuropil of origin**

To identify SEZONs and LH-associated neurons, the cable length within the respective neuropil mesh (3D bounding box) was calculated. A cut off of > 60 nodes within the defined neuropil region was required for classification.

**Dendrogram representations**

Dendrogram representations of neurons were created as in [11]. Dendrograms are 2D representations of 3D neuronal reconstructions which preserve the topology of neuron and visualize specific synapses on specific branches. The neato layout (Graphviz, https://graphviz.gitlab.io/) [43] attempts to minimize a global energy function, equivalent to statistical multi-dimensional scaling to represent the neuron morphology as a graph. Code available (https://github.com/markuspleijzier/AdultEM/tree/master/Dendrogram_code) using the Graphviz library with Python bindings provided by NetworkX, (https://networkx.github.io/) [44].

**Marking MBON postsynapses on dendrograms by closest DAN cluster**

Euclidean distances between MBON-γ5b/2a postsynapses and the closest DAN presynapse were measured and marked with the identity of the morphological DAN clusters (Figure 1). The Euclidean distances were then thresholded to within 2 μm and the synapses
identified to be under that threshold were plotted on a neato dendrogram. The plot in Figure S4F therefore shows all postsynapses within a 2 μm diffusion distance from a dopaminergic presynapse.

**Edge weight distribution**

Edge weight distributions describe how many upstream neurons contribute a given number of presynapses to a connection with a postsynaptic neuron (frequency versus number of synapses). Normalizing by the total number of postsynapses details the percentage of the total postsynaptic budget a given number of synapses represents. For example, if a neuron makes 10 presynapses onto a postsynaptic neuron, which has a total of 100 postsynapses, then that upstream neuron contributes 10% of the postsynaptic budget.

**DAN-MBON direct connectivity**

Identified MBONs were collapsed by type. The number of synapses between DANs and MBONs was normalized by the number of all DAN-MBON connections of the given DAN. Connectivity matrices can be calculated for single branches of a neuron after defining the relevant branchpoints in CATMAID. For the PPL1-γ1pedc DAN, we manually split the dendrite into 4 postsynaptic clusters, as defined from cluster analyses, and recorded the specific connectivity to each of these clusters/branches.

**DAN connectivity similarity matrices**

A connectivity similarity score between 2 DANs was defined as one minus half of the Manhattan distance between their normalized connectivity patterns (normalized connectivity patterns of DANs shown in Figure 2E).

**Statistical analysis of DAN connectivity - comparison to a null model of random connectivity**

A DAN input connectivity matrix was first randomized 10⁴ times, respecting both DAN postsynaptic budget and input neuron presynaptic budget, so that after randomization each row sum and each column sum remained the same as in the observed data (i.e., each DAN gets the same number of inputs and each input neuron has the same number of outputs). Then the Manhattan distance between upstream connectivity profiles of DANs in the observed data and those in simulated random matrices, both normalized by DAN postsynaptic budget were calculated and means of these distances were compared to obtain a p value describing the similarity of these means. A p value lower than the significance level concluded that the null model of randomized connectivity could be rejected (see Methods S1).

**Behavioral experiments**

The odors used for US substitution, sucrose learning and DEET learning experiments were 10⁻³ dilutions of 3-octanol (OCT) and 4-methylcyclohexanol (MCH) in mineral oil. For extinction experiments odor concentrations of 10⁻⁶ were used to avoid pre-exposure effects [11]. Experiments were performed at 23°C and 55%–65% relative humidity, except for electric shock learning which occurred at 70% relative humidity.

**US-substitution experiments using CsChrimson**

In both the behavioral screen and follow-up experiments, neurons were artificially activated to substitute for an unconditioned stimulus in the training chamber of a T-maze. Prior to the experiments, 80-120 1-5 day old mixed sex flies were housed on standard food supplemented with 1% all-trans-Retinal for 3 days before a 20–28 h starvation period in vials containing 2 mL 1% agar as a water source and a 2x4 cm strip of Whatman filter paper. During training, groups of flies were exposed to the CS- for 2 min followed by 30 s rest with fresh air, then 2 min of CS+ odor with optogenetic activation of the genetically encoded Channel Rhodopsin with red light exposure. Three red (620-630nm) LEDs (Multicom, p/n OSW-4338) with 3 W maximum power were mounted on the training arm of a T-maze and 1ms pulses were driven at 1.2V with a stimulation frequency of 500Hz, which is flicker free red-light that flies cannot see. For screening, immediate memory testing followed. Flies were transferred back into their starvation vials after training before testing 30 min memory.

**Appetitive olfactory learning with sucrose reward**

Prior to the experiments, 80-120 3-8 day old mixed sex flies were starved for 20-28 h in vials containing 2 mL 1% agar as a water source and a 2x4 cm strip of Whatman filter. Flies were transferred to 32°C 30 min before training. During training, groups of flies were exposed to the CS- odor with dry paper for 2 min followed by 30 s of fresh air, then 2 min of CS+ odor exposure with dry sugar paper. Flies were either tested immediately after training or were transferred back into 25°C starvation vials after training prior to testing 30 min memory.

**Aversive olfactory learning with bitter reinforcement**

Flies were aversively trained with DEET as previously described [16]. In brief, prior 80-120 3-7 day old mixed sex flies were starved for 20-24 h in vials containing 2 mL 1% agar and a 2x4 cm strip of Whatman filter paper. Training and immediate testing were performed at 32°C. During training groups of flies were exposed to the CS- odor with 1% agar on filter paper for 2 min followed by 30 s fresh air, then 2 min of CS+ odor with 0.4% DEET, 3 M xylose and 100 mM sucrose in 1% agar on filter paper. Flies were tested for their odor preference immediately after training.
Aversive memory extinction
Extinction memory was tested as described [11]. In brief, mixed sex groups of 80-120 flies were transferred into vials with 2 mL corn-meal medium and a 2x4 cm strip of Whatman paper for 18-26 h before training. Aversive olfactory conditioning in the T-maze was conducted as previously described [31]. Flies were exposed to the CS+ odor for 1 min paired with twelve 90 V electric shocks at 5 s intervals. Following 45 s of clean air, the CS- odor was presented for 1 min without shock. Immediately after training flies were transferred to 32°C. 30 min later flies were re-exposed twice to either the CS- or CS+ odor with a 15 min interval. Flies were then returned to permissive 23°C and tested 15 min later for memory performance.

Memory testing and statistical analyses of behavioral data
To test memory performance flies were loaded into the T-maze and transported to the choice point where they were given two min to choose between the CS+ and CS- odors in the dark. A Performance Index was calculated as the number of flies in the CS+ arm minus the number in the CS- arm, divided by the total number of flies [31]. MCH and OCT, were alternately used as CS+ or CS- and a single sample, or n, represents the average performance score from two reciprocally trained groups. Statistical analysis was carried out with GraphPad, Prism (v8.1). All experiments were analyzed with a one-way ANOVA. For extinction CS- re-exposure flies were compared with CS+ re-exposure flies and Tukey’s post-doc analyses for multiple comparisons applied. For all other experiments statistical comparisons were performed between a wildtype control and Dunnett’s post hoc analysis carried out for multiple comparisons.