Plasticity-driven individualization of olfactory coding in mushroom body output neurons

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Although all sensory circuits ascend to higher brain areas where stimuli are represented in sparse, stimulus-specific activity patterns, relatively little is known about sensory coding on the descending side of neural circuits, as a network converges. In insects, mushroom bodies have been an important model system for studying sparse coding in the olfactory system3–5, where this format is important for accurate memory formation6–8. In Drosophila, it has recently been shown that the 2,000 Kenyon cells of the mushroom body converge onto a population of only 34 mushroom body output neurons (MBONs), which fall into 21 anatomically distinct cell types2–4. Here we provide the first, to our knowledge, comprehensive view of olfactory representations at the fourth layer of the circuit, where we find a clear transition in the principles of sensory coding. We show that MBON tuning curves are highly correlated with one another. This is in sharp contrast to the process of progressive decorrelation of tuning in the earlier layers of the circuit2,9,10. Instead, at the population level, odour representations are reformatted so that positive and negative correlations arise between representations of different odours. At the single-cell level, we show that uniquely identifiable MBONs display profoundly different tuning across different animals, but that tuning of the same neuron across the two hemispheres of an individual fly was nearly identical. Thus, individualized coordination of tuning arises at this level of the olfactory circuit. Furthermore, we find that this individualization is an active process that requires a learning-related gene, rutabaga. Ultimately, neural circuits have to flexibly map highly stimulus-specific information in sparse layers onto a limited number of different motor outputs. The reformatting of sensory representations we observe here may mark the beginning of this sensory-motor transition in the olfactory system.

The Kenyon cell (KC) axons are arranged in parallel bundles that form the output lobes of the mushroom body (MB). Mushroom body output neurons (MBONs) extend dendrites into those lobes, with different MBON types innervating distinct subregions3,4 (Fig. 1a). We expressed the calcium sensor GCaMP5 in MBONs using a series of split-GAL4 lines6 (Extended Data Table 1) and measured odour tuning using in vivo two-photon imaging, quantifying response magnitude as the area under the ΔF/F curves (F, fluorescence intensity; Fig. 1b). The high specificity of these drivers typically enabled us to track activity of individual MBONs. We thereby successfully collected data from 17 types/combination of types of MBONs, covering 18 out of 21 cell types, and one additional MBON from the calyx (Fig. 1c and Extended Data Figs 1 and 2; see Methods).

Consistent with high convergence at this stage of the circuit3–4, MBONs were generally broadly tuned to odours, as observed in other insects10–12, although there were a few exceptions (for example, α2p3p, β1 and MB-CP1 neurons; Extended Data Fig. 3). In the MBONs with axonal projections inside the MB lobes (β1, γ1 pedc, and γ4 neurons), we observed prolonged rise times (Extended Data Fig. 4).

One of the important factors governing the stimulus-specificity of population-level representations is how independent and decorrelated their sensory tuning is. Optimal coding theory dictates that a compact neuronal population most efficiently conveys stimulus-specific information if the tuning properties of different neurons are decorrelated, so the redundancy of their signalling is minimized13, which we refer to as tuning decorrelation. We confine our analysis here to a tuning-curve-based view of the system, and do not explore the role that temporal patterning of spikes might have in conveying information, as has been shown in other systems14,15. Overall, odour tuning of the MBON population was notable for its lack of diversity, showing high levels of correlation (Figs 1d and 2e). We found no obvious relationship between the degree of tuning correlation of different MBONs and their type of input KC, the neurotransmitter they release, or where they subsequently project (Fig. 1d and Extended Data Fig. 5a, b).

These highly correlated, dense response patterns were in sharp contrast to the KCs. The calcium responses of KCs to the same set of odours, measured at the cell body layer, were sparse and specific (Fig. 2a, b), with much lower levels of tuning correlation (Fig. 2e). To visualize how odour representations are transformed between the KCs and MBONs, we used principal component analysis (PCA) to represent population response patterns observed on each stimulus trial (Fig. 2c; see Methods and Extended Data Fig. 6). Although different odour clusters were well-separated in the KCs, in MBONs they were much closer to one another and often partially overlapping. Nevertheless, there was a coarse structure to the distribution of different odours, and some were well-separated. This basic structure was conserved when we analysed subpopulations of MBONs according to their axonal projection sites (Extended Data Fig. 5c). The close proximity of odour clusters visualized by PCA was reflected in a lower score of odour classification analysis in MBONs than KCs (Fig. 2d; see Methods). Importantly, this was not simply caused by the sharp reduction in the number of neurons, or their broad tuning compared to the KCs. When we held cell number and tuning breadth constant, but artificially decorrelated MBON tuning by assigning rearranged odour labels to each cell’s tuning curve, classification accuracy markedly increased (Fig. 2d, e; see Methods). Furthermore, when we examined the number of distinct odour clusters in MBON space, relatively few clusters were apparent, but artificial decorrelation of MBON tuning increased the number of clusters to match the number of odours, just like the KC representations (Fig. 2f; see Methods). These results clearly show that a neuronal population of this size and breadth of tuning is capable of representing odour identity accurately, however, the correlations in MBON tuning properties place an important limit on that capacity. We note that it is still possible that specific information about odour identity could be carried in the precise timing of MBON spike trains10,11,14.

We then asked what features of sensory information become available at this layer. To address this, we calculated the correlations between neural representations of all pairs of odours in KCs and MBONs and compared the distributions (Fig. 2g). In KCs, this distribution showed a single sharp peak near zero, indicating that odour representations are largely decorrelated; in fact, artificially
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as well as the pattern of activity contribute to this distribution shape, because we observed a significant positive correlation between correlation coefficients in KCs and in MBONs (Fig. 2i). Notably, negative correlations between different odour representations, a rare feature in KCs, become relatively common in MBONs, especially in MBON subpopulations with axonal projections to particular downstream areas (Fig. 2h and Extended Data Fig. 5d). Thus, it seems that MBONs convey a sense of interrelationship between odours, be it positive or negative. In this context, it is interesting to note that two odour groups of opposite valence were reliably the most distantly located in MBON coding space (Fig. 2c and Extended Data Fig. 5c) and were never misclassified with each other (Extended Data Fig. 7). One of those groups was apple cider vinegar and yeast, which are attractive food odours for Drosophila\(^{15,16}\). The other was CO\(_2\) together with citronella, which are both reported to be natural repellents to Drosophila\(^{17,18}\).

We next focused on odour tuning properties at the single-cell level. Specifically, we asked whether cell types with uniquely identifiable anatomy also have consistent tuning properties in different animals. Such functional stereotypy is readily apparent in the projection neurons at the second layer of the circuit\(^{19,20}\). In MBONs, by contrast, we observed a range of similarity in tuning across animals depending on cell types (Fig. 3a). Some MBONs showed highly consistent responses; however, several had very diverse tuning. Among all the MBONs, the \(\alpha_{2c}\) neuron exhibited the greatest variability. This was not due to ambiguity in cell identification, since there is only one \(\alpha_{2c}\) neuron per hemisphere. Moreover, a similar level of inter-animal variability was also observed with whole-cell recordings (Fig. 3b–d).

Tuning patterns of individual KCs are not stereotyped across animals\(^{21}\), which is likely to arise from the probabilistic input connectivity of the projection neurons\(^{21,22}\). Is the variability in MBON tuning simply inherited from the probabilistic organization of the previous layer? If so, tuning properties of MBONs should be as variable across hemispheres in the same brain as they are across different animals. However, this was clearly not the case. Pairs of \(\alpha_{2c}\) neurons from opposite sides of the same brain exhibited strikingly similar tuning compared to those from different brains (Fig. 3e and g). We observed similar results in three other MBON types (Extended Data Fig. 8). Thus, tuning patterns of MBONs are individual-specific as a result of a process that is coordinated across hemispheres, rather than random wiring patterns.

We next set out to ask how this functional individuation of the circuit arises at the level of MBONs. The dense dendritic arbor of the MBONs implies that MBONs summate input from many KCs\(^{23}\). If so, does the variable tuning of MBONs derive from variable overall levels of population activity in KCs? However, while tuning profiles of individual KCs are not predictable\(^{20–22}\), the summed output from the overall KC population could be consistent across animals, since projection neurons from different glomeruli have relatively stereotyped numbers of output terminals in the MB, suggesting that the total excitatory drive to the KCs may be characteristic for each odour\(^{24,25}\). Furthermore, the number of responding KCs for a given odour is positively correlated with the total activity of olfactory receptor neurons responding to that odour. To directly examine the variability of bulk MB output, we imaged calcium responses in the KC axon bundle at the site where it contacts the \(\alpha_{2c}\) neuron. We found that the tuning of the bulk KC population was relatively consistent from fly to fly, in contrast to the \(\alpha_{2c}\) neuron (Fig. 3f, g). We thus found no sign of individuality in the summed activity of the KCs. But then, why do MBONs, thought to receive heavily convergent input from KCs, not end up with stereotyped tuning patterns? To better understand how KC activity is integrated by MBONs, we measured functional connectivity between \(\alpha/\beta\) KCs and \(\alpha_{2c}\) neurons by paired whole-cell recordings (Fig. 4a, b). Surprisingly, we found only 7 pairs out of 24 with an excitatory

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Figure 1 | Summary of olfactory tuning patterns in MBONs. a, Schematic of the KCs and MBONs. b, GCaMP5 imaging in the \(\alpha_{2}\) neuron. Left, example images of the odour responses from axonal arbor. Greyscale image, baseline fluorescence. Scale bar, 10 \(\mu\)m. Win, apple cider vinegar; Yst, yeast; Hep, 2-heptanone; Hex, 1-hexanol; Oct, 3-octanol; MCH, 4-methylcyclohexanol; Pep, peppermint; EtOH, ethanol; Cit, citronella. Middle, \(\Delta F/F\) time courses in the same cell. Grey, individual trials (\(n = 4\)). Red, mean. Shaded areas, 2-s odour stimulations. Right, odour tuning profiles. Bars, mean (± s.e.m.) normalized tuning (\(n = 5\) flies; see Methods). Heat map, tuning in each fly. c, Odour tuning in the remaining 16 MBON cell types. d, Correlation matrix of tuning patterns of MBONs (Pearson’s \(r\); middle). Cell types are arranged according to the dendrogram (left) based on the correlation distances (1 – \(r\)).  

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decorrelating KC tuning had little further effect. By contrast, in MBONs correlation coefficients ranged widely without an obvious peak around zero. Both the dense format of MBON representations, as well as the pattern of activity contribute to this distribution shape, since artificially decorrelating MBON tuning properties resulted in a wider distribution than in the KCs, but now with a clear peak around zero (Fig. 2g and Extended Data Fig. 5e). Some of the relationships between odours are partially inherited from previous layers of neurons, because we observed a significant positive correlation between correlation coefficients in KCs and in MBONs (Fig. 2i). Notably, negative correlations between different odour representations, a rare feature in KCs, become relatively common in MBONs, especially in MBON subpopulations with axonal projections to particular downstream areas (Fig. 2h and Extended Data Fig. 5d). Thus, it seems that MBONs convey a sense of interrelationship between odours, be it positive or negative. In this context, it is interesting to note that two odour groups of opposite valence were reliably the most distantly located in MBON coding space (Fig. 2c and Extended Data Fig. 5c) and were never misclassified with each other (Extended Data Fig. 7). One of those groups was apple cider vinegar and yeast, which are attractive food odours for Drosophila\(^{15,16}\). The other was CO\(_2\) together with citronella, which are both reported to be natural repellents to Drosophila\(^{17,18}\).

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Figure 2 | Transformation of population representations from KCs to MBONs. a, Representative odour responses of KC somata in a single fly with pseudo-coloured ΔF/ΔF images overlaid on greyscale basal fluorescence. Scale bar, 20 μm. b, Summary of responses from the fly in a. Columns, KCs; Rows, trials sorted by odours. c, Odour representations in a single representative fly (KCs, same fly as a) or virtual fly (MBONs) projected onto the first three principal component axes (PC; 100 data points per odour; see Methods). Colours indicate odours as in b, d. Accuracy of odour classification (100 classification scores for both KCs and MBONs; see Methods). Red lines indicate medians, boxes interquartile ranges, notches 95% confidence intervals, whiskers data ranges, and crosses outliers. Grey line, chance level. e, Correlation matrices of tuning curves. Data are from a single representative fly (KCs) or virtual fly (MBONs). f, Clustering analysis of population activity patterns in KCs and MBONs (KCs, n = 10 flies; MBONs, n = 20 virtual flies; see Methods). Highest silhouette values indicate optimal clustering. Silhouette value ranges are KCs, 0.37–0.84; MBONs, 0.35–0.74; decorrelated MBONs, 0.27–0.65. g, Correlation coefficients (Pearson’s r) of neural representations of ten odours (45 odour pairs), calculated using the response patterns averaged across trials in KCs (n = 10 flies, orange) and MBONs (n = 1,000 virtual flies, blue). Mean distributions are shown. Distributions of artificially decorrelated data are in lighter colours. h, Same as g, but for MBON subpopulations with axonal projections to different areas. See also Extended Data Fig. 5. CRE, crepine; SIP, superior intermediate protocerebrum. i, Correlation coefficients in KCs and MBONs are positively correlated. Each dot corresponds to correlation coefficient of a specific odour pair (mean ± s.d.). Grey line, linear regression (R² = 0.42, P < 10⁻⁵).

Figure 3 | Individualization of tuning in MBONs. a, Correlations of MBON tuning patterns across different flies (n = 5 flies for each cell type). b, Sample traces from whole-cell recordings from 2sc2c neurons. Red dots, spikes. Shaded areas, 1-s odour presentations. c, Raster plots of 2sc2c odour responses. Iaa, isoamyl acetate. d, Tuning curves (mean ± s.e.m.) from c, e, Odour tuning of pairs of 2sc2c neurons on the left and right hemispheres of five representative flies, from GCaMP5 imaging (normalized to the strongest response, mean ± s.e.m.). BtOH, 1-butatol; Hxa, 1-hexanal. f, α/β KC population tuning across different hemispheres of the same fly, recorded at the α2 region of the MB lobes. g, Tuning patterns of 2sc2c neurons from different hemispheres of the same fly are more correlated than those from different flies (n = 8 flies, P < 10⁻⁴, Tukey’s post-hoc test following two-way analysis of variance (ANOVA)). Correlations of KC population tuning were indistinguishable within versus across flies (n = 7 flies, P > 0.95, Tukey’s post-hoc test). Interactions between cell types (2sc2c versus KCs) and comparison types (within versus across flies) were significant (P < 0.05, two-way ANOVA).
Figure 4 | Connection probability between KCs and a MBON. a, Schematic of paired whole-cell recording. b, Post-hoc immunohistochemistry of biocytin infused via whole-cell electrodes (green, maximum intensity projection). Axon of α/β KC, extending within α/β lobes (dotted line), and dendritic arbor of α2sc neuron (arrow) are visible. Magenta, anti-nC82 staining. Scale bar, 20 μm. c, A directly connected KC-α2sc neuron pair. Top, a train of KC spikes (Pre) and the evoked response in a α2sc neuron (Post) recorded simultaneously. Middle, discrete excitatory postsynaptic potential (EPSP) steps visualized by spike-trigger-averaging MBON membrane potential on the first KC spike of each trial. Bottom, enlarged spike-trigger-averaged EPSPs for the first, second and third spikes in the train reveal a short synaptic delay (<2 ms). d, The four categories of synaptic connectivity (see Methods). The first-spike-trigger-averaged responses for all pairs in each category are overlaid. Timing of the KC spikes is indicated by greyscale bar, whose intensity shows the mean relative spike rate.

connection, only 5 of which were likely to be monosynaptic (Fig. 4c, d and Extended Data Fig. 9; see Methods). This gives a probability of connection of <30%, far more selective than the all-to-one convergence suggested by the dendritic anatomy. Thus, our results suggest that α2sc neurons are capable of extracting very different information in different animals, even from presynaptic KCs that have similar overall population coverage of the population, combined with our back-to-back comparison of tuning at any layer of the olfactory system. This extensive one process that could plausibly underlie such flexible wiring is synaptic plasticity; indeed plasticity has been observed at this synapse in other insects36. To test this we examined whether Rutabaga, an adenylyl cyclase required for learning4, is involved in generating the variability of the α2sc post-hoc test). Interactions between the genotypes and comparison types (within versus across flies) were significant (P < 0.005, two-way ANOVA). Within-fly tuning correlations were not statistically different between the two genotypes (P > 0.65, Tukey’s post-hoc test).

Figure 5 | Cross-individual variability is lost in rutabaga mutants. a, Odour tuning of a pair of α2sc neurons on the left and right hemispheres in the same fly, recorded with GCaMPs imaging (mean ± s.e.m.). Data from five wild-type males. b, Same as a but rut2080 males. c, Control males show higher variability across flies than within flies (n = 9 flies, P < 10^-4, Tukey’s post-hoc test following two-way ANOVA). This difference is lost in rut2080 hemizygous males, which show similar levels of variability both within and across flies (n = 8 flies, P > 0.995, Tukey’s post-hoc test). Interactions between the genotypes and comparison types (within versus across flies) were significant (P < 0.005, two-way ANOVA). Within-fly tuning correlations were not statistically different between the two genotypes (P > 0.65, Tukey’s post-hoc test).

This work presents the most complete population-level characterization of tuning at any layer of the olfactory system. This extensive coverage of the population, combined with our back-to-back comparison to the previous layer, enabled us to demonstrate that the progressive decorrelation of neuronal tuning that marks the ascending layers of the sensory circuit29 comes to an end immediately downstream of the KCs, as the network converges. Fine temporal patterns of spiking, or subtle correlations in signalling across multiple MBONs within each animal, neither of which we could detect here, could contribute to the specificity of odour representations at this layer (but see Extended Data Fig. 6). Nevertheless, our results clearly show that positive and negative correlations arise in MBONs, clustering some odour representations together and pushing others apart. This grouping might be useful when it comes to making a behavioural choice, since the general categorization of stimuli, rather than detailed, stimulus-specific information would be more important at this stage. Thus, MBON representations may be well-suited to control motor outputs55. Interestingly, similarly prominent network convergence occurs with cortical projections to the basal ganglia56, whose main function is to select an appropriate action plan by interpreting the available sensory information.

The plasticity-driven individualized tuning of MBONs is a counterpoint to the highly stereotyped responses of the output neurons of the lateral horn (LH), an olfactory centre that lies in parallel with the MB29 and is implicated in innate responses to odour37. In contrast, the influence of plasticity on MBON tuning highlights the flexibility of this branch of the circuit, where odour representations could be reshaped either to fine tune, or perhaps to override the innate responses driven by the LH pathway, reflecting each fly’s individually unique olfactory experience.
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Author Contributions T.H. and G.C.T. designed the experiments with help from Y.A. and G.M.R; T.H. performed all imaging and electrophysiology experiments and data analyses. Y.A. and G.M.R created fly strains and collected anatomical data for MBONs. T.H. and G.C.T. wrote the manuscript.

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METHODS

Fly stocks. Flies were reared at room temperature on conventional cornmeal-based medium. All flies were reared from adult females, 2–5 days post-emergence, unless otherwise noted. Note that we used the same animal-rearing conditions for all flies when we examined the variability of tuning across different flies, and that we always compared flies of same gender. In several cases, we even recorded on the same day from progeny of the same cross, raised in the same food medium. No randomization or blinding methods were used. For calcium imaging, flies bearing UAS-GCaMP6f (ref. 31; KC cell body imaging) or GCaMP5f (ref. 32; MBON and KC axon imaging) were crossed with appropriate GAL4 or split-GAL4 driver lines (refs. 9, 33, 34) for KC imaging (both cell body and lobe imaging) and a series of split-GAL4 lines for MBON imaging (Extended Data Fig. 2; Extended Data Table 1; see also http://www.janelia.org/split-gal4 for the expression patterns as well as enhancer fragments used to construct these split-GAL4 lines). For experiments in rutabaga mutants, we used the fact that rutabaga is X-linked. rutabaga females were crossed with UAS-GCaMP5f; MB080C males, and their male progeny, hemizygous for rutabaga, were used for imaging. For electrophysiological recording from the 2sc neuron, we crossed UAS-2eGFP with MZ160-GAL4 (ref. 36). For dual whole-cell recording from a/b KCs and the 2sc neuron, we crossed UAS-2eGFP; MZ160(2sc) with c739, an a/b-specific driver14. UAS-GCaMP6f flies were provided by V. Jayaraman, rutabaga females were crossed with UAS-2eGFP with MZ160-GAL4 (Extended Data Fig. 2; Extended Data Table 1; see also http://www.janelia.org/split-gal4 for the expression patterns as well as enhancer fragments used to construct these split-GAL4 lines). For experiments in rutabaga mutants, we used the fact that rutabaga is X-linked. rutabaga females were crossed with UAS-GCaMP5f; MB080C males, and their male progeny, hemizygous for rutabaga, were used for imaging.

Nomenclature of MBONs. The set of 21 cell types of MBONs from the MB lobes and one from the calyx are defined by their morphology, having dendrites inside the MB and axonal projections elsewhere9. For simplicity, in this paper we call each MBON according to its dendritic region in the MB region, as well as X, Y, and Z (see Extended Data Table 1).

Odour delivery. The following monomolecular odors were purchased from Sigma and used as stimuli: 2-heptanone (Hept; CAS# 110-43-0), 1-hexanol (Hex; 11-27-3), 3-octanol (Oct; 588-98-0), 4-methylcyclohexanol (MCH; 589-91-3), ethanol (EtOH; 64-17-5), isomyl acetate (Iaa; 123-92-2), 1-hexanal (Hexa; 66-25-1), 1-butanal (B120; 71-36-3) and hexyl acetate (Hy; 142-92-7). We also used natural essential oils from peppermint (Peppermint) and citronella (Cit; Aura Cacia) as well as other natural odours, apple cider vinegar (Vine; Richfood) and yeast (Yst; Lessafre). In imaging experiments, odours were presented through a custom-built device as described previously13. 40-ml vials were loaded with 5-ml pure odours, except for essential oils, which were diluted with mineral oil at 1:100. Yeast was prepared by adding 5 ml distilled water to 1 g dry yeast. Saturated headspaces were vaped using two steps of air dilutions down to 10% (experiment in Figs 1–3a) or 5% (other experiments). CO2 was directly taken from the house-line and presented at 1%. Final flow rate of the air stream was set to 40 ml/min. In all cases, odorant delivery was independent across the two recordings. This ensured that the higher correlations we observed within animals could not be attributed to noise correlations, that is, coordinated changes in neuronal responses based on sensory transitions in the internal state of the animal, such as its level of arousal15.

Electrophysiology. Previously reported methods for in vivo whole-cell recordings in projection neurons18 were adapted for MBONs and KCs. The patch pipettes were pulled for a resistance of 4–5 MΩ (MBON) or 6–7 MΩ (KC) and filled with pipette solution containing (in mM): 1-potassium aspartate, 125; HEPES, 10; EGTA, 1; CaCl2, 0.1; Mg-ATP, 4; Na-GTP, 0.5; biocytin hydrazide, 13; with pH adjusted to 7.3 with KOH (265 mM). Bath solution was the same as in imaging experiments. Single or dual whole-cell current-clamp recordings were made using the Axon MultiClamp 700B amplifier (Molecular Devices). Cells were held at around ~60 mV by injecting hyperpolarizing current (20 ± 5 pA for MBONs, < 5 pA for KCs). Signals were low-pass filtered at 5 kHz and digitized at 10 kHz. Specific cell types were visually targeted by GFP signal with a 60× water-immersion objective (UMPlanFl/IR; Olympus) attached to an upright microscope (BX51WI; Olympus). Although MZ160-GAL4 labels multiple types of MBONs in the MB-V2 cluster, we were typically able to distinguish the 2sc neuron, which is a single unique neuron16, from the others based on the distinct size and location of its cell body. The morphology of all recorded cells, both 2sc neurons and a/b KCs, were visualized by post-hoc immunohistochemistry with biocytin19; any data from incorrectly targeted cells were discarded. In dual whole-cell recording from a/b KCs and the 2sc neuron, we took several steps to maximize the chance of detecting weak connections. Since KCs are immunonegative for choline acetyltransferase20 and unlikely to be cholinergic21–25, we applied the cholinergic-specific toxin to the MBON (100 μM) to the bath saline to minimize unrevealed circuit activity that could obscure weak connections. In addition, we tested for connections using current injection (25 ± 5 ± 5.5 pA, 175 ± 9.0 ms, mean ± s.e.m.) to drive high frequency spike trains in the KCs (10.6 ± 0.8 spikes, mean ± s.e.m.).

Data analysis. All data analyses were performed in MATLAB (R2008b, MathWorks). All sample sizes were enough for robust statistical tests, which are appropriately chosen based on the distribution of data. No statistical methods were used to predetermine sample size.

Analysis of imaging data. For the analysis of MBONs, a region of interest (ROI) was manually set for each trial based on the mean baseline image. Response magnitude was calculated as integrated fluorescence change (ΔF/F) in the time window between stimulus onset and 5 s after stimulus offset. Analysis of KC imaging required motion corrections and frame alignments in order to retain the identity of each ROI (that is, each KC cell body) throughout a whole imaging session, as described previously26. For the measurement of tuning similarity, we used Pearson’s correlation coefficient. When we used Euclidean distance instead, the results of the statistical tests remained the same in all cases (data not shown) except for one case (Extended Data Fig. 8e), where we did not detect a significant difference with the Euclidean distance measurement (data not shown).

Analysis of electrophysiological data. Spikes were automatically detected by examining spike counts in the frames. The frames were binned on amplitude, after removing slow membrane potential deflections with a high-pass filter, and verified by visual inspection. Response spike rates were calculated in a window of 0.5–3.5 s after odor valve opening, and baseline spike rates were subtracted. To calculate the statistical significance of a connection between an a/b KC and an 2sc neuron, we determined the difference in calcium channels at the soma and/or to the extremely long primary neurite connecting the soma to the axon and dendrites. On the other hand, calcium responses at dendrites and axons were consistently observed even when there was no somatic response (Extended Data Fig. 1a). Notably, the time course and the magnitude of the responses were largely similar between the dendrite and axon in the same neuron. We tested how general this was across different MBONs by comparing calcium responses in axons and dendrites in five different cell types. These five types were all the ones in which we could distinguish both axonal and dendritic signals for individual neurons (Extended Data Fig. 1a–e). In all cases, the responses were highly similar (Pearson’s r = 0.92 ± 0.013, mean ± s.e.m.; n = 10 cells). Furthermore, tuning breadths were also identical (Extended Data Fig. 1f, g). Therefore, to measure tuning curves in MBONs, we imaged at either dendrite or axon, whichever maximized the isolation of the signals from the MBON of interest (Extended Data Fig. 2; Extended Data Table 1), except for Extended Data Figs 1 and 8. KC imaging was performed at the cell body layer as described previously16. In both cases, imaging frames were typically 300 × 300 pixels, acquired with a pixel dwell time of 1.6 μs, yielding frame rates around 4 Hz, which slightly varied across experiments depending on the optical zoom factor. For each odor presentation trial, data were acquired for 20 s with an odor pulse (2 s in duration for experiments in Figs 1–3a, 1 s for other experiments) triggered 8 s after trial onset. Inter-stimulus interval was 25 s. When we compared tuning patterns of 2sc neurons across hemispheres, we imaged the two hemispheres sequentially rather than simultaneously, so stimulus presentations were independent across the two recordings. This ensured that the higher correlations we observed within animals could not be attributed to noise correlations, that is, coordinated changes in neuronal responses based on sensory transitions in the internal state of the animal, such as its level of arousal15.

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between the mean n2scc membrane potential during the KC spike train and during the baseline period on each individual trial (51 ± 4 trials per pair), and used a t-test with a threshold significance value of P < 0.05. Out of 24 pairs recorded, we found only eight statistically significant postsynaptic responses. Five pairs were judged as monosynaptically connected because step-wise increments in membrane potential were obvious in spike-trigger-averaged traces and also because the delay between the KC spike and the onset of the EPSP was less than 2 ms (Extended Data Fig. 9a, d). In two other pairs, we observed smaller excitatory responses but could not detect unitary steps for each spike, even in spike-trigger-averaged traces (Extended Data Fig. 9b). Such connections could be either direct excitatory connections with unitary strength smaller than our detection limit or indirect feed-forward connections. The one remaining connection we found was, to our surprise, inhibitory (Extended Data Fig. 9c). The synaptic delay was too long for a monosynaptic connection via ionotropic transmission, suggesting that it represents a powerful feed-forward inhibitory connection. However, it remains possible that it is a monosynaptic connection with facilitatory synapses or slow metabotropic input. Although we cannot rule out the possibility that we are missing some extremely weak connections, we consider this unlikely given our experimental approach, namely quietening spontaneous network activity by blocking cholinergic transmission and then driving high frequency spike trains in the presynaptic KC (see the section above for details). Even if we assume that some connections went undetected, our results would nevertheless indicate that synaptic weights of KC–n2scc connections are highly heterogeneous, since some connections were strong enough for us to detect clear unitary synaptic events. Thus, in any case our results indicate that connectivity between n2scc KCs and n2scc neurons is highly selective.

**Population coding analyses.** To compare population coding in KCs and MBONs, we represented the odour response patterns from each stimulus trial as a point in an N-dimensional neuronal coding space, where each of the axes represents the response magnitude of each neuron. For KC representations, each axis corresponded to one of the KCs imaged in one fly. For MBONs, since different cell types were imaged in different flies, we combined data from 17 different flies to produce an aggregate MBON population reflecting activity in a single ’virtual fly’. Since we have recordings from 5 flies for each type of MBON, by combining different recordings we can construct many different virtual flies; one example is shown in Fig. 2c. This approach assumes that there is no specific correlation between the tuning patterns of different cell types in the same animal. However, if different MBON types tended to be positively or negatively correlated in individual flies, that could have an impact on the odour-specific information available in population-level activity patterns, which we would have overlooked by analysing virtual flies. To test whether this was the case, we imaged multiple types of MBONs in the same animal by combining different split-GAL4 drivers (Extended Data Fig. 6). We compared tuning patterns in four pairs of cell types. These pairs included MBONs receiving input from the same MB lobe (v1 vs n2scc), those with input from the same types of KCs (v1 vs n2scc), and those with input from different types of KCs (v1 vs b1), as well as MBONs with relatively higher variability of tuning across flies (v1 and n2scc; Fig. 3a). In no case did we find that the correlation of tuning across flies was significantly different from the correlation across different animals (Extended Data Fig. 6). This result justifies using virtual flies to analyse MBON population representations. Each recording consisted of 4–7 trials per odour, so for virtual flies, we generated random combinations of those trials by bootstrapping. For example we generated 100 bootstrapped trials in one virtual fly for the display in Fig. 2c. On the other hand, we could not combine KC data from multiple flies because, unlike MBONs, the identities of KCs cannot be matched up between different flies. To make our analysis of the KC population similar to the MBONs, we also applied trial bootstrapping to the KC data, so that noise correlations, absent from the individually recorded MBON data, are also present in the KCs.

**Odour classification.** For odour classification analysis, we adopted a linear classification algorithm based on Euclidean distances between odour representations in neuronal coding space. Odour-evoked activity patterns for each bootstrapped trial, generated as described above, were represented as points in this coding space, and the centroids of the points corresponding to each of the ten test odours were calculated. The data point for each trial was then classified as the odour with the minimum Euclidean distance from the trial. Generated as described above, we shuffled between trials, classification accuracy dropped to near chance level, indicating that patterns of activity are important for classification in both cases. They did not drop all the way since different odours evoke different overall levels of activity, which also contributes information about odour identity; this is eliminated by additionally shuffling odour labels (Fig. 2d).

**Cluster analysis.** For cluster analysis (Fig. 2f), we employed the k-means clustering algorithm. k-means clustering is an unsupervised clustering method that allows us to partition data points into arbitrary number of clusters such that the total distance from the individual points to their cluster centroids is minimized. The quality of such clusters can be evaluated by the mean silhouette value of all the data points(A). The silhouette value (ranging from −1 to 1) for the i-th data point can be calculated as $s(i) = \frac{(k(i) - a(i))/\sqrt{\sum_{j \neq i} d(i,j)^2}}{(a(i) + k(i))/\sqrt{\sum_{j \neq i} (k(i) + a(i))}}$, where $a(i)$ is the average distance to the other points in the same cluster, and $k(i)$ is the average distance to the points in the closest neighbouring cluster. Thus, if data points form compact clusters that are well separated from each other and are appropriately partitioned, the mean silhouette value approaches 1. On the other hand, if clusters are diffuse and/or overlapping each other, a lower value is obtained, even when the clusters are optimally partitioned. In addition, even when there are nicely separated compact clusters, it gives a lower value if the partitioning pattern is inappropriate (for example, dividing one cluster into halves or combining two clusters into one). Thus, by using k-means to explore a wide-ranging number of clusters (2 to 20), and comparing the silhouette width that gave the highest silhouette value, we were able to determine the optimal number of clusters in MBON and KC odour representations. We examined the clustering in the data with different dimensionalities by gradually increasing the number of principal components (PCs) for the projection. To make the analysis comparable between KCs and MBONs, which consist of profoundly different numbers of cells, we matched the total fraction of variance captured as we increased dimensionality, rather than directly increasing the number of PCs. In other words, we increased dimensionality by using a sufficient number of PCs to capture the same amount of variance in KC and MBON population activity as the first eight statistically significant postsynaptic responses. When too many PCs were added. This is presumably because these later PCs mainly contained noise rather than signals related to odours, which would simply make existing clusters more diffuse in higher-dimensional space. We used built-in functions of MATLAB to perform PCA and k-means clustering as well as to calculate silhouette values.

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Extended Data Figure 1 | Comparison of odour-evoked calcium responses across multiple cell compartments. a, Odour responses were sequentially recorded at three different compartments (axon, blue; dendrite, orange and soma, black) in two different a2sc neurons with GCaMP5 imaging. Upper subpanel shows mean ΔF/ΔF traces from different compartments (4 or 5 trials for each). Shaded areas indicate 1-s odour stimulations. Lower subpanel shows tuning profiles in axon and dendrites (normalized to the strongest response, mean ± s.e.m.). Note that the time course, response magnitude and tuning profiles are largely consistent between the axon and dendrite in the same cell, while signals at the soma are small and slower (Cell2) or sometimes undetectable (Cell1). b–e, Data from 4 other types of MBONs. Scale of the traces is the same as in a. Correlation between axons and dendrites is 0.92 ± 0.013 (Pearson’s r, mean ± s.e.m.; n = 10 cells). Hya, hexyl acetate. f, Mean normalized tuning of the ten cells. Odours are sorted according to the rank order in each cell to visualize the tuning width. g, Kurtosis of tuning curves. There was no difference between axons and dendrites (P = 0.16, paired t-test).
Extended Data Figure 2 | Raw traces of calcium imaging in MBONs. For all 17 types/combination of types of MBONs, we show a projection image of confocal stacks from split-GAL4 lines (left), example ΔF/ΔF images of the calcium responses to ten odours and air (middle) and ΔF/ΔF time courses (right). Note that all the split-GAL4 lines used in this study label the target MBONs with extremely high specificity. The white squares indicate the approximate region scanned during calcium imaging. Greyscale images show baseline fluorescence (scale bar, 5 μm), while colour map images represent calcium responses (ΔF/ΔF, colour scale bottom right). Traces showing ΔF/ΔF time courses on individual trials (grey; n = 4–7) are overlaid with the mean (red), scale bar, 50% ΔF/ΔF.
Extended Data Figure 3 | Relationship between response intensity and tuning selectivity. Lifetime sparseness, an index of tuning selectivity\(^{1,44}\), was plotted against the magnitude of the largest odour response observed for each neuron (mean \(\Delta F/F\) during the response window). Each dot corresponds to one of the 17 MBON types (mean ± s.e.m.; \(n = 5\) flies). There was no correlation between the two variables (Spearman’s \(\rho = -0.16, P = 0.54\)). Thus, the relatively selective tuning patterns observed in four MBONs (blue) are likely not related to the intensity of the response. Rather, for three of these cells, we noticed that their narrow tuning was accompanied by unusual dendritic anatomy. \(\beta'1\) neuron is the only MBON with extensive dendritic processes outside the MB, and calyx neuron (MB-CP1) is the only MBON that samples from the MB calyx. \(\alpha2p3p\) neuron’s dendrites seem to contact solely the \(\alpha/\beta p\) KCs, whose dendrites arborize in the sub-region of the MB calyx known as the accessory calyx, where no olfactory input has been reported\(^7\). The narrow tuning patterns of the fourth neuron, \(\gamma5\beta'2a\), seem to be related to the transient inhibitory epochs uniquely observed in this neuron (Extended Data Fig. 4).
Extended Data Figure 4 | Diversity in response time courses across the MBON population. Normalized mean ΔF/F responses to yeast odour for all MBON types overlaid (shading indicates timing of odour presentation).

Note that three MBONs (β1, γ1pedc, and γ4) that have axonal terminals in the MB lobes show slower time courses (red) than the others (black). The cell with the characteristic inhibitory period is the γ5β'2a neuron (blue).
Extended Data Figure 5 | Population analysis of subpopulations of MBONs. 

**a,** Distribution of MBON terminals in the brain (dark colours), based on the localization of a presynaptic marker* (Syt:smGFP-HA). Anterior (top) and dorsal (bottom) views are shown. Axonal projections of MBONs converge heavily onto small areas inside four neighbouring neuropils (light colours), crepine (CRE; orange), superior medial protocerebrum (SMP; green), superior intermediate protocerebrum (SIP; purple) and superior lateral protocerebrum (SLP; blue). We have found numerous cells that send widely branching neurites into one or all of these areas (database associated with ref. 45), indicating that downstream neurons likely read out activity from populations of MBONs. Lateral horn (LH, red) also receives sparse input.

**b,** Pairwise correlations of MBON tuning patterns and the corresponding dendrogram are shown in the same way as Fig. 1d. The dots on the right show the axonal projection site of the different MBON types. None of the four major projection zones samples from particularly decorrelated set of MBONs.

**c,** Odour representations in MBONs from a single virtual fly visualized by PCA (different virtual fly from Fig. 2c). Representations from the full MBON population (All) and subpopulations with the same axonal projection zones (CRE, SMP, SIP and SLP) are shown. Representations by subpopulations tend to be noisier than those from the full population, but they retain grossly similar structures.

**d,** Correlation coefficients of neural representations of ten odours in KCs and MBONs (thick lines), as in Fig. 2g, h. Correlation values observed after artificially decorrelating KC and MBON tuning curves are shown for comparison (thin lines).

**e,** Variance of the correlation coefficients shown in **d** (dark bars; mean ± s.d.; n = 1,000 virtual flies), showing that values are more widely distributed in MBONs compared to KCs. Artificially decorrelating MBON tuning curves reduces the variance considerably (light bars), indicating that the pattern of activity in the MBONs contributes to the high variance. However, variance remains greater than that in the KCs, indicating that the breadth of tuning in the MBONs also contributes to wide range of correlation coefficients. The ratio of the contribution of those two factors seems to be different across subpopulations projecting different zones.
Extended Data Figure 6 | Correlation of tuning between multiple MBON types measured in the same animal. a, Odour tuning of α1 and α2sc neurons recorded sequentially in the same fly with GCaMP5 imaging (normalized to the strongest response, mean ± s.e.m.). Data from five representative flies are shown. Two split-GAL4 drivers, MB319C and MB080C, were combined to label the two cell types.

b, Similar to a but recordings are from α1, β1 and γ4 neurons. MB319C was combined with MB434B that labels both β1 and γ4 neurons.

c, Correlation of tuning between different cell types, calculated within and across flies. Four different pairwise combinations of cell types were examined. Although within-fly correlations tended to be slightly higher than across-fly correlations, none of the four cases was significant (Mann–Whitney U-test). This is in sharp contrast with the comparisons of the same cell type, where we observed much higher correlations within the same fly than across flies (Fig. 3).
Extended Data Figure 7 | Confusion matrices from odour classification analysis. Confusion matrices generated from the classification analysis in Fig. 2d. a, KCs show nearly perfect classification performance. b, MBONs fail to discriminate some odours. Although the confusion matrix was generated using 100 different virtual flies (see Methods), a high rate of misclassifications is observed only between certain odour pairs that form overlapping clouds in MBON space (Fig. 2c). Odour pairs that were never misclassified with each other are indicated with white asterisks. Note that the group of vinegar and yeast were never misclassified with the group of CO₂ and citronella.
Extended Data Figure 8 | Individualized tuning in multiple different MBON types. a–c, Odour tuning of pairs of MBONs in the same fly, from GCaMP5 imaging at axons (normalized to the strongest response, mean ± s.e.m.). Data from five representative flies are shown. For γ1pedc (a) and α’2 neurons (b), cells on the left and right hemispheres are compared. For α1 neuron (c), recordings were from ipsilateral cell pairs. d–f, For all three cell types, tuning patterns of the neurons from the same fly are more correlated than those from different flies (n = 8 flies per cell type; Mann–Whitney U-test).
Extended Data Figure 9 | Diverse functional connections between KCs and a MBON. a, Another example of a monosynaptically connected pair of α/β KC and the α2sc neuron from experiments shown in Fig. 4. a1, Sample traces of simultaneously recorded KC (Pre) and the α2sc neuron (Post). a2, The α2sc neuron’s membrane potential spike-trigger-averaged on the first KC spike of each trial. Raster plot of the KC spikes is shown below. a3, Enlarged spike-trigger-averaged EPSPs for the first, second and third spikes in the train. b, A pair with small excitatory connection that we could not confirm as monosynaptic. In total, two of such pairs were found. c, A pair with an inhibitory connection, which is likely to be polysynaptic. Only one such pair was found. d, Summary of all five monosynaptically connected pairs. The spike-trigger-averaged EPSPs from the first spike in the train from five different recordings are shown overlaid in grey. The mean across cells is shown in red.
Extended Data Table 1 | MBON types and imaging conditions.

| Type* (Dendritic site in MB) | Axonal terminals in MB* | No. of cells per hemisphere* | Imaged strain* | Signal isolation | Imaged locus |
|----------------------------|-------------------------|-----------------------------|----------------|-----------------|--------------|
| α1                         | -                       | 2                           | MB310C         | b               | dendrite     |
| α2p3p                      | -                       | 2                           | MB062B         | b               | dendrite     |
| α2Sc                      | -                       | 1                           | MB080C         | a               | dendrite     |
| α3                         | -                       | 2                           | MB093C         | b               | dendrite     |
| β1                         | α                       | 1                           | MB434B         | a               | dendrite     |
| β2β2a                      | -                       | 1                           | MB399B         | a               | dendrite shaft |
| α’1                      | -                       | 2                           | MB543B         | b               | axon terminals |
| α’2                         | -                       | 1                           | MB018B         | a               | axon arbor   |
| α3ap                      | -                       | 1                           | MB027B         | c               | dendrite     |
| α3m                         | -                       | 2                           | MB057B         | b               | axon arbor   |
| β’1                         | -                       | 8                           | MB002B         | a               | dendrite     |
| β’2mp                    | -                       | 1                           | N.A.           | N.A.            | N.A.         |
| γ1 pedc                   | α/β                     | 1                           | MB085C         | a               | dendrite shaft |
| γ1γ2                      | -                       | 1                           | N.A.           | N.A.            | N.A.         |
| γ2α’1                   | -                       | 2                           | MB051B         | b               | axon arbor   |
| γ3                      | -                       | 1                           | MB027B         | c               | axon terminals |
| γ3β’1                    | -                       | 1                           | MB298B         | a               | axon shaft   |
| γ4                      | γ1γ2                    | 1                           | MB242A         | a               | dendrite shaft |
| γ4γ5                      | -                       | 1                           | MB011B         | a               | γ5 dendrite  |
| γ5β’2a                    | -                       | 1                           | MB027B         | a               | dendrite     |

*Imaging signal is from single neuron. †Signal is from multiple cells of single cell type. ‡Signal is from multiple cell types. *Data from ref. 8.