Fast odour dynamics are encoded in the olfactory system and guide behaviour 2

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- 11 Abstract

12 Odours are transported in turbulent plumes resulting in rapid concentration fluctuations^{1,2} that contain rich information about the olfactory scenery, such as 13 14 odour source composition and location²⁻⁴. Yet whether the mammalian olfactory 15 system has access to the underlying temporal structure to extract information 16 about the environment remains unknown. Here, we show that 10 ms odour pulse 17 patterns result in distinct responses in olfactory receptor neurons. In operant 18 conditioning experiments mice discriminated temporal correlations of rapidly 19 fluctuating odours at frequencies of up to 40 Hz. In imaging and 20 electrophysiological recordings, such correlation information could be readily 21 extracted from the activity of mitral and tufted cells, the output of the olfactory bulb. Furthermore, we show that temporal correlation of odour concentrations⁵ 22 23 reliably predicts whether odorants emerge from the same or different sources in naturalistic environments with complex airflow. Training mice on such tasks and 24 25 probing with synthetic correlated stimuli at different frequencies suggests that 26 mice can indeed use the temporal structure of odours to extract information about 27 space. Our work thus demonstrates that the mammalian olfactory system has access to unexpectedly fast temporal features in odour stimuli. This in turn endows 28 29 animals with the capacity to overcome key behavioural challenges such as odour 30 source separation⁵, figure-ground segregation⁶ and odour localisation⁷, by 31 extracting information about space from temporal odour dynamics.

32 Main text

The turbulent nature of air^{1,2,4,8} as well as water^{9,10} flow results in complex temporal 33 fluctuations of odour concentrations that depend on the distance and direction of odour 34 35 sources^{1-4,8,10}. Insects are thought to use the temporal structure of odour plumes to infer e.g. odour source location^{4,7,11-13} or composition¹³⁻¹⁵. Mammalian olfaction on the other 36 hand has generally been considered a slow sense. Individual sniffs are thought to be the 37 unit of information¹⁶, implying that fast odour concentration changes (at sub-sniff 38 39 resolution) should be inaccessible to the mammalian olfactory system. However, the neural circuitry of e.g. the mouse olfactory bulb (OB) is in principle capable of 40

- 41 millisecond-precise action potential firing^{17,18}, and is endowed with rich computational
- 42 resources that could be employed to extract fine temporal information from dynamic
- 43 inputs¹⁹. Here, we show that the mouse olfactory system has access to fast, sub-sniff
- 44 temporal patterns in the odour scenery and that mice can use this information to detect
- 45 high-frequency odour correlations enabling source separation.

46 **Fast odour dynamics encoded in OB inputs**

47 Normal airflow is characterized by complex, often turbulent, flow patterns and imposes a rich temporal structure on odour concentration profiles with significant power in 48 frequencies well above typical sniff rates (Fig. 1a). To assess whether the mouse olfactory 49 system has access to this frequency regime, we designed an odour delivery system 50 51 capable of reliably presenting odours with a bandwidth beyond 50 Hz (Fig. 1b, **Supplementary Methods Fig. 1**). As prototypical, simplistic high frequency stimuli we 52 53 employed two 10 ms square pulses of odour separated by 10 or 25 ms (Fig. 1c). Olfactory 54 sensory neurons (OSN) are known to be slow in responding to odour stimuli²⁰. Both epithelial mucus and the biochemical transduction cascade act as low-pass filters^{16,20,21}, 55 suggesting that individual OSNs cannot directly follow rapidly fluctuating odour stimuli. 56 However, axons from up to 10s of thousands of OSNs that express the same olfactory 57 receptor converge onto one or a few glomeruli in the OB²². This organization resembles 58 the auditory system where, despite the relatively low temporal resolution of individual 59 cells, population responses faithfully report high-frequency signals²³. Thus, we built a 60 model of populations of noisy integrate-and-fire neurons with stimulus filtering and 61 62 neuronal dynamics matching experimental data to explore whether this large convergence could aid in detecting high frequency stimuli in OSNs (Extended Data Fig. 63 64 **1**). Our simulation results suggest that across the thousands of OSNs that express the 65 same OR - while still not directly following the odour profile - the population can 66 faithfully discriminate between such 10/25 ms stimulation (Extended Data Fig. 1d,f,h). Key high-frequency information in the odour profile might, therefore, be preserved in the 67 68 inputs to the OB.

69 To test this experimentally, we performed Ca²⁺ imaging experiments in anaesthetized and awake mice expressing GCaMP6f in OSNs (Fig. 1c-i, Extended Data Fig. 2) whilst 70 71 delivering odour pulses locked to inhalation (Fig. 1c,d). Overall, responses for all 72 glomeruli were highly correlated between the two stimuli (**Fig. 1f,g**). Glomerular activity did not directly follow the 10ms or 25 ms pulses (Fig. 1f). However, in 1/3 of glomeruli 73 74 (n = 33/100, p < 0.01), responses were consistently and significantly different for the two 75 stimuli (Fig. 1f-h, Extended Data Fig. 2) mirroring the simulation results (Extended 76 **Data Fig. 1**). Notably, just a few dozen randomly chosen glomeruli were sufficient to 77 discern between the stimuli at >80% success rate with a linear classifier (Fig. 1i, see also 78 Extended Data Fig. 1h). Expanding the stimulus set to different concentrations and 79 multiple pulses (Extended Data Fig. 2) confirmed that information about concentration and temporal patterns with features exceeding the 25 ms timescale is reliably and 80 independently preserved in the population of OSNs. 81

82 **Discrimination of correlation structure**

83 Can mice base behaviour on such high frequency stimuli? We trained mice in an automated go/no-go operant conditioning system ("AutonoMouse"²⁴, Fig. 2a, 84 **Supplementary Video 1**) to discriminate between high frequency stimuli. To ascertain 85 86 that the brief odour pulses were delivered during inhalation in freely moving mice we opted for 2 second pulse trains at different frequencies with constant airflow (Fig. 2b). 87 We found that mice can discriminate whether an odour is presented at e.g. 4 Hz or 20 Hz, 88 89 yet the apparent "critical flicker frequency" (Fig. 2c, Extended Data Fig. 3) was significantly lower than frequencies OSNs readily represent (Fig. 1, Extended Data Fig. 90 **1,2**). However, in both visual and auditory systems, conventional flicker fusion frequency 91 or gap detection thresholds substantially underestimate the temporal sensitivity, 92 particularly for tasks with multiple stimuli present^{23,25,26}: In vision, for example, flicker 93 fusion frequency is around 60 Hz, whereas thresholds for detecting synchrony between 94 stimuli has been reported to be 3 ms²⁶. Thus, we wanted to probe whether, similarly, 95 96 olfactory tasks involving multiple odours reveal behavioural access to higher frequencies. We presented stimuli composed of two odours fluctuating in a correlated or anti-97 98 correlated manner as the rewarded and unrewarded stimulus, respectively (and vice 99 versa, Fig. 2d-f). Mice readily learned to differentially respond to correlated or anti-100 correlated odours (Fig. 2h-k). Gradually increasing the correlation frequency showed 101 that animals were capable of reliably detecting the correlation structure of stimuli at 102 frequencies of up to 40 Hz (Fig. 2h,j,k). As a population, animal performance decreased 103 by approximately 5% per octave with performance significantly above chance at frequencies of up to 40 Hz (n = 33 mice in two cohorts of 14 and 19 mice, Fig. 2k). To 104 105 mitigate the risk of animals using non-intended cues for discrimination, odours were presented from changing valve combinations (Fig. 2g, Extended Data Fig. 4), odour flow 106 107 was carefully calibrated (Fig. 2e, Extended Data Fig. 4d-e) and additionally varied 108 randomly between trials such that neither flow nor valve clicking noises or average concentration provided any information about the nature of the stimulus (Extended 109 Data Fig. 4d-h). Consistent with this, when valve identities were scrambled, animals 110 performed at chance (grey, Fig. 2k). Finally, when odour presentation was changed to a 111 new set of valves, performance levels were maintained (Fig. 2g-i and Extended Data Fig. 112 113 **4i-k**), indicating that only intended cues (the temporal structure of odours) were used 114 for discrimination. Performance was independent of the odour pair used (Extended Data 115 Fig. 3g) and maintained for tasks discriminating correlated from uncorrelated (rather 116 than anti-correlated) odours (Extended Data Fig. 3e,f).

Mice tended to take more time to detect the correlation structure of stimuli with higher fluctuation frequencies (Extended Data Fig. 5j-l). This was most pronounced for animals with higher overall performance (Extended Data Fig. 5j). Accuracy strongly correlated with reaction time across all stimuli and animals (Extended Data Fig. 5k) despite the fact that total time of odour delivery was the same across all trials regardless of stimulus frequency. Consequently, when analysis was restricted to trials where mice sampled the stimuli long enough, e.g. for at least 750 ms, performance significantly increased across 124 frequencies (Extended Data Fig. 51). This indicates that the measured performance 125 might not be the psychophysical limit for discriminating fluctuating odour stimuli. Furthermore, this suggests that mice integrate information across large portions of the 126 127 presented stimuli, rather than e.g. detecting simultaneity of odour onset¹⁴ to determine whether odours were correlated or not. To directly test this possible strategy, we 128 interleaved training trials with probe trials where the onset characteristics were flipped 129 130 (Extended Data Fig. 5f-i). Notably, performance did not drop substantially (Extended **Data Fig. 5h,i**), consistent with a strategy that relies primarily on discerning the high 131 frequency correlation structure of the stimulus over several 100 ms rather than the onset 132

133 only (Extended Data Fig. 5f,g,i). Sniff rate in turn was independent of the correlation

134 frequency of stimuli presented (Extended Data Fig. 5a-e).

135 Odour correlation encoded in OB output

To assess how this high-frequency information is represented and reformatted in the 136 137 olfactory system, we imaged neural activity in response to high-frequency stimuli (Fig. 3). Ca²⁺ imaging of OSN responses to correlated and anti-correlated stimuli showed that 138 139 - unlike for two pulses with variable gaps (Fig. 1) - correlation structure of odour pulse trains was difficult to discern on the level of inputs to the OB using simple linear 140 141 classifiers (Extended Data Fig. 6). Directly imaging from the *output* of the OB, mitral and 142 tufted cells (M/TCs, Fig. 3a-g, Extended Data Fig. 7), showed that overall, M/TCs also 143 responded similarly to correlated and anti-correlated stimuli (Extended Data Fig. 7j-l). 17% of all M/TCs, however, showed significantly different integral responses (0-5 sec 144 145 after odour onset, p < 0.01) to the two stimuli (114/680 ROIs, Fig. 3d-f). As a result, correlated and anti-correlated odours were reliably discriminated by a linear classifier 146 using the M/TC population responses (somatic response Fig. 3g, dendritic response 147 **Extended Data Fig. 7d,i)** unlike for the OSN population response (Extended Data Fig. 148 **6k,**]). This is consistent with the idea that the OB circuitry implements a non-linear 149 150 transformation of OSN input where the representation of correlation becomes more 151 readily accessible in the OB output.

152 We employed odour stimuli rapidly fluctuating at frequencies that substantially exceeded the temporal resolution of Ca²⁺ imaging, which captures a low-pass filtered signal of 153 154 neural activity. Although the Ca²⁺ signal does not follow individual stimulus frequencies, the M/TC population response contained enough information to determine whether a 155 correlated or anti-correlated stimulus was presented. To probe whether additional 156 157 information about stimuli is present in the output of the OB at finer time scales, we turned to extracellular unit recordings (Fig. 3h-k, Extended Data Fig. 8) and whole-cell patch 158 recordings (Extended Data Fig. 9). Despite the kHz temporal resolution, single-units also 159 did not directly follow high-frequency stimuli. Average activity (summed spike count 160 161 during 500 ms after odour onset) was, however, significantly different between 162 correlated and anti-correlated stimuli in 24% of single-units (23/97, p < 0.01, Mann-163 Whitney U test, Fig. 3i, j, Extended Data Fig. 8b), consistent with the Ca²⁺ imaging results. 164 As few as 60 randomly selected units were sufficient to classify the odour stimuli with >80% accuracy (Fig. 3k). Additional information was contained at finer time scales as
 increasing the temporal resolution of analysis improved discriminability (Fig. 3k and
 Extended Data Fig. 8e-g). Together, these results demonstrate that information about
 high-frequency correlation structure in odours is accessible to the animal for behavioural
 decisions and readily available in the output of the OB.

170 **Correlations allow for source separation**

171 What could the detection of high-frequency correlations be useful for? Natural odours 172 consist of multiple different types of molecules, and a typical olfactory scene contains 173 several sources⁶. To make sense of the olfactory environment, the brain must be able to 174 separate odour sources, attributing the various chemicals present to the same or different 175 objects⁵. Motivated by the turbulent nature of odour transport, Hopfield suggested that the temporal structure of odour concentration fluctuations might contain location 176 177 information about odour sources⁵ - i.e. that chemicals belonging to the same source 178 would co-fluctuate in concentration. Detecting correlations in odour fluctuations would thus allow mice to discern which odours arise from the same object. To experimentally 179 180 probe the potential of odour correlation structure to facilitate odour source separation in air, we devised a dual-energy fast photoionisation detection method to simultaneously 181 measure the odour concentrations of two odours with high temporal bandwidth 182 183 (Methods, Fig. 4a,b, Extended Data Fig. 10a-e and Supplementary Methods Fig. 2). 184 When an odour was presented in a laboratory environment with artificially generated complex airflow patterns (Fig. 4a), to mimic the outdoor measurements (Fig. 1a), odour 185 186 concentration fluctuated with a spectrum extending beyond 40 Hz (Extended Data Fig. 187 **10a**). When two odours were presented from the same source, these fluctuations were 188 highly correlated (Fig. 4a,b and Extended Data Fig. 10b). When we separated odour 189 sources and presented the two odours 50 cm apart, odour dynamics were uncorrelated 190 (Fig. 4a,b) with intermediate correlations for closer distances (Fig. 4b). This pattern of 191 almost perfect correlation for the same source and virtually uncorrelated dynamics for 192 separated sources was maintained at closer and farther distances between odour source 193 and sensor (Extended Data Fig. 10d), independent of the odours used (Extended Data Fig. 10c) and was mirrored outdoors (Extended Data Fig. 10e). Thus, the correlation 194 195 structure of odorant concentration fluctuations indeed contains reliable information 196 about odour objects – e.g. whether odours emerge from the same or different sources.

197 Can mice make use of this information? We trained a new cohort of mice in a modified 198 AutonoMouse setting, presenting odours corresponding to the "same source" or "source separated" case as rewarded or unrewarded stimuli (Fig. 4c,d and Extended Data Fig. 199 200 **10**). Mice were able to learn to discriminate these stimuli (**Fig. 4d,e**). Once the task was 201 acquired, we probed their performance with artificially generated stimuli (Extended 202 Data Fig. 10f-k) that were derived from prior measurements with natural airflow but 203 perfectly correlated (Fig. 4e). Notably, they reliably responded to these probe trials with 204 correlated stimuli as they did to the "same source" stimuli they had been trained on (Fig. 205 **4e**, **Extended Data Fig. 10m**). To further ascertain that they were using the correlation structure to make these decisions, we probed with artificial square pulse stimuli (as in
Fig. 2, 3) at different frequencies. Mice performed significantly above chance in probe
trials at frequencies of up to 40 Hz (Fig. 4e, Extended Data Fig. 10s), implying that
learning about source separation directly translates to distinguishing temporal features
in correlated / uncorrelated stimuli.

211 Discussion

212 Here, we have shown that the mammalian olfactory system has access to temporal features of odour stimuli at frequencies of at least up to 40 Hz. We have demonstrated 213 214 access to information in rapid odour fluctuations using different behavioural 215 experiments (Fig. 2,4). We have shown reliable decoding from imaging and unit 216 recordings from different stages of the olfactory system using both correlated odour concentration fluctuations (Fig. 3, Extended Data Fig. 7, 8a-g) as well as simplistic 217 paired pulse stimuli with gaps as small as 25 ms (Fig. 1, Extended Data Fig. 2, 8h-l), 218 219 corroborated by computational modelling (Extended Data Fig. 1). Our results are 220 consistent with recent findings that the olfactory bulb circuitry not only enables highly 221 precise odour responses^{17,18} but enables detection of optogenetically evoked inputs with a precision of 10-30 ms²⁷⁻²⁹ with different projection neurons displaying distinct firing 222 patterns in response to optogenetic stimulation²⁹. While behavioural and physiological 223 224 responses to precisely timed odour stimuli have been observed in insects^{13,15,30}, in 225 mammals the complex shape of the nasal cavity was generally thought to low-pass filter 226 any temporal structure of the incoming odour plume. Our results show that while the 227 low-pass filtering in the nose and by OSNs might reduce the ability of neurons to directly 228 follow high-frequency stimuli, sufficient information about high-frequency content is 229 preserved and available such that mice can readily make use of this information.

230 What could such high bandwidth be useful for? We have shown that odour sources even 231 in close proximity differ in their temporal correlation structure. Thus, the ability to detect whether odorants are temporally correlated could allow mice to perform source 232 233 separation, solving the "olfactory cocktail party problem"^{5,6} without prior knowledge 234 about the odour scenery. We show that mice can indeed discriminate between "one source / separated source" stimuli. They readily translate this discrimination to artificial 235 236 correlated pulse trains demonstrating that they are using correlation structure to make 237 this distinction. Distinguishing between other environmental features, such as distance or direction of an odour source, could also be achieved by extracting temporal features 238 239 from odour fluctuations^{1-3,8} possibly in combination with strategies comparing information reaching the brain through the two nares^{31,32}. 240

How exactly is this temporal information extracted? While insects are able to detect the
simultaneity of onset of two odours^{14,33,34}, this strategy is unlikely to be the dominant
means that mice use to detect correlation (Extended Data Fig. 5). Similarly, mice do not
show adjustment of sniff strategies for discriminating high frequency odour correlations
(Extended Data Fig. 5, Supplementary Video 2). While individual mammalian OSNs are
thought to be quite slow and unreliable²⁰, the large convergence of OSN axons provides a

substrate to create the needed high temporal bandwidth³⁵ (Extended Data Fig. 1).
Biophysical heterogeneity of OSNs might improve how the population encodes
temporally structured stimuli^{36,37}. Intrinsic cellular biophysics (Extended Data Fig. 9),
local interneurons or long-range lateral inhibition^{5,38} might permit the extraction of
temporal correlation within the olfactory bulb circuitry and possibly result in individual
projection neurons tuned to specific temporal structures.

The turbulence of odour plumes has often been viewed as a source of noise for mammals. In contrast, we find that the mouse olfactory system has access to high-frequency temporal features in odour stimuli. This opens up a new perspective on how mice could make use of natural turbulence in order to obtain information about their spatial environment. This in turn provides new computational challenges for the mammalian olfactory system and an entry point into how information about space is extracted from sensory inputs.

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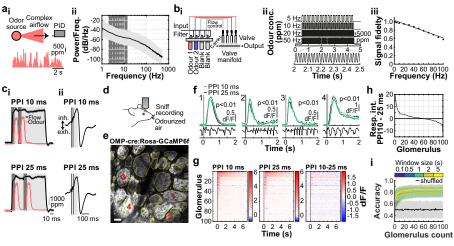
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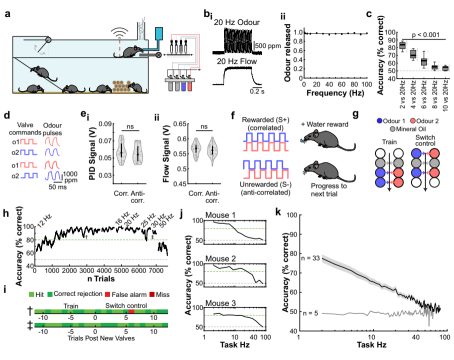
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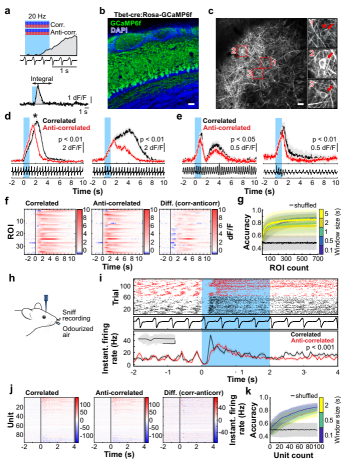
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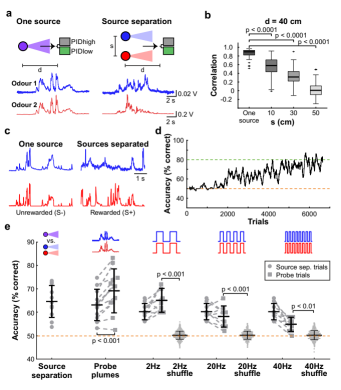


Fig. 1 | Sub-sniff detection of odour signals in olfactory bulb inputs. 348 349 **a**_i, Example odour plume recorded outdoors under natural, complex airflow conditions using a photoionisation detector (PID). aii, Averaged power spectrum of all recorded 350 351 odour plumes (n=37 plumes, mean±SD of log power), typical range of sniff frequencies observed in mice highlighted in dark grey. **b**_i, Schematic of multi-channel high bandwidth 352 odour delivery device. **b**_{ii}, Representative odour pulse recordings at command 353 354 frequencies between 5 and 50Hz. **b**_{iii}, Relationship of frequency and odour pulse signal fidelity (see Methods, n=5 repeats for each frequency, mean±SEM, see also 355 Supplementary Methods Fig. 1). c_i, Odour (red) and flow traces (black) of 10ms paired 356 pulse interval (PPI) stimuli for 10ms (top) and 25ms (bottom), valve commands are 357 358 shown in dark grey. c_{ii}, Stimuli are presented during the inhalation phase of the respiration cycle. **d**, Schematic of the two-photon imaging approach. **e**, GCaMP6f 359 360 fluorescence recorded in olfactory bulb glomeruli (maximum projection of 8200 frames, 361 marked glomeruli correspond to example traces shown in **f**). Scale bar: 50µm. **f**, Example calcium traces in response to 10 and 25ms PPI odour stimuli (mean of 10 trials±SEM, 362 unpaired two-sided t-test for 2s response-integral from odour onset). Bottom: Example 363 respiration trace. **g**, Calcium transients as colour maps for PPI 10ms (left), PPI 25ms 364 365 (middle), and the difference between both odour stimulations (right). Glomeruli are sorted by response magnitude to the PPI 10ms stimulus. h, Glomerular responses sorted 366 by magnitude of difference to PPI 10 vs. 25ms. i, Classifier accuracy over all glomeruli 367 when a linear classifier was trained on several response windows (colour-coded, black: 368 shuffle control) to PPI 10 vs. 25ms stimuli (n=up to 100 glomeruli from 5 individual 369 370 animals; mean±SD of 500 repetitions). Throughout, ethyl butyrate was used as the odour 371 stimulus.

372 Fig. 2 | Mice can discriminate odour correlation structure at frequencies up to **40Hz. a**, Schematic of the automated operant conditioning system ("AutonoMouse") 373 374 housing cohorts of up to 25 animals. **b**_i, Representative trace of a 20Hz odour pulse train 375 (top) and corresponding stable airflow (bottom). **b**_{ii}, Relationship of frequency and total 376 amount of odour released (n=5 repeats for each frequency, mean±SEM). c, Group 377 accuracy in frequency discrimination task (n=10 mice, p<0.001 for all stimuli compared 378 to chance accuracy (paired two-sided t-test); see also Extended Data Fig. 3). Boxes 379 indicate 25th-75th percentiles, thick line is median, whiskers are most extreme data points not considered outliers, see Methods. **d**, Left: Valve commands to release two odours 380 fluctuating at 20 Hz in a correlated (top) or anti-correlated (bottom) manner. Right: 381 382 Resultant odour concentration changes measured using dual-energy photoionisation detectors (Supplementary Methods Fig. 2). e, Odour (ei) and flow (eii) signal for 383 correlated and anti-correlated stimuli fluctuating at 20Hz (n=60 trials for each condition; 384 385 odour: p=0.19, flow: p=0.23, unpaired two-sided t-test). Median shown as black dot, first and third quartile are bounds of the black bar. **f**, Schematic of the discrimination stimuli; 386 387 mice were trained to discriminate between two odours presented simultaneously in 388 either a correlated (top) or anti-correlated (bottom) fashion in a standard go/no-go paradigm. g, Schematic of valve combinations for stimulus production. Train: 6 valves are 389

390 used to produce the stimulus through varying valve combinations. Switch control: two 391 extra valves are introduced and odour presentation switched over to the newly 392 introduced valves. **h**, Example animal performing the correlation discrimination task at 393 different frequencies. i, Trial response maps before and after switch to control valves (as described in **g**, n=12 trials pre-, n=12 trials post-new valve introduction). Symbols 394 indicate time point of valve introduction as marked in **h**, see also **Extended Data Fig. 4**. 395 396 j, Accuracy of 3 representative animals performing correlation discrimination where 397 stimulus pulse frequency is randomised from trial to trial. **k**, Group accuracy for the 398 experiment in **j** (black trace: standard training, band: SEM, grey trace: full scramble 399 control; n=33 training mice, n=5 control mice, n=9.3×10⁵ trials). Throughout, isoamyl 400 acetate and ethyl butyrate were used as odour stimuli.

401 Fig. 3 | Odour correlation structure is encoded by olfactory bulb output neurons. 402 a, Schematic of the two-photon imaging approach (see also Extended Data Fig. 7e). b, 403 Coronal olfactory bulb section showing GCaMP6f (green) expressed in projection neurons. Scale bar: 20µm. c, GCaMP6f fluorescence from mitral and tufted cells 404 405 (maximum projection of 8000 frames). Responses from ROI * in magnified inset 1 * is shown in **d**. Scale bar: 20µm. **d**, Example traces of ROIs that show differential response 406 407 kinetics to correlated (black) and anti-correlated (red) stimulation (mean of 24 408 trials±SEM, f=20Hz, unpaired two-sided t-tests on 5s response-integrals) in 409 anaesthetised and **e**, awake animals (mean of 16 trials±SEM, f=20Hz, unpaired two-sided 410 t-tests). Odour presentation indicated in light blue. **f**, Calcium transients as colour maps 411 for correlated (left) anti-correlated (middle) averaged trials and the difference between both odour stimulations (right) for the 5% of ROIs with the largest differential responses. 412 **g**, Accuracy of linear classifier trained on several response windows (colour-coded, black: 413 414 shuffle control) to correlated vs. anti-correlated stimuli at 20Hz (n=up to 680 ROIs from 415 6 individual animals; mean±SD of 500 repetitions). **h**, Schematic of the extracellular 416 recording approach. **i**, Example single unit of an odour response for correlated (black) 417 and anti-correlated (red) stimuli shown as raster plot (top) and PSTH (mean±SEM) of 418 spike times binned every 50ms (bottom); inset: average spike waveform (black) and 1000 individual spike events (grey), scale bar: 100µV and 1ms. Odour presentation 419 420 indicated in light blue. Two-sided Mann-Whitney U test comparing spike time distributions of correlated and anti-correlated trials during 4s after odour onset. j, Binned 421 422 spike discharge over time shown as colour maps for all units, correlated (left), anticorrelated (middle) and the difference between both odour stimulations (right). k, 423 424 Accuracy of linear classifier trained on the average 2s response to correlated vs. anticorrelated stimuli at 20Hz (yellow); green: 500ms window; blue 100ms window (n=up 425 426 to 97 units from 6 individual animals; mean±SD of 1000 classifier repetitions; see 427 Methods and Extended Data Fig. 8).

428 Fig. 4 | Source separation using correlations of odour concentration fluctuations.

a, Simultaneous measurement of two odours (Odour 1: α-Terpinene; AT, Odour 2: ethyl

430 butyrate; EB) using a dual-energy photoionisation detector (Extended Data Fig. 10a-e,

431 Supplementary Methods Fig. 2) at d=40cm, presented either from one source or 432 separated from each other by s=50 cm, with complex airflow in the laboratory. **b**, Correlation coefficients over all recordings for odours from the same source and for 433 434 odour sources separated by s=10-50cm (EB vs. AT; n=61 for Mix, n=71 for each individual distance; unpaired two-sided t-test). Boxes indicate 25th-75th percentiles, thick line is 435 median, whiskers are most extreme data points not considered outliers, see Methods. **c**, 436 437 Example plumes used for training animals on a virtual source separation task to discriminate between odour stimuli derived from the same one source (Unrewarded, S-) 438 and from separated sources recordings (Rewarded, S+). d, Example learning curve for a 439 mouse trained to perform the virtual source separation task. Isoamyl acetate and ethyl 440 butyrate were used as odour stimuli. **e**, Average accuracy over different variants of the 441 task, calculated over the last 2400 trials of virtual source separation training (n=11 mice, 442 443 p<0.0001, unpaired t-test, compared to chance performance), and subsequent stages 444 where probe trials containing novel plume types are interleaved with the training set. Responses are compared between probe and training plumes within each stage. Probe 445 plumes: odours fluctuate in a perfectly correlated manner, with a novel temporal 446 447 structure (120 probe trials, in a segment of 2400 trials, n=11 mice, paired t-test). Probe 448 2Hz, 20Hz, 40Hz: Correlated/anti-correlated square pulse trains (50 probe trials per frequency, in a segment of 1650 trials, n=9 mice). Responses to 2Hz, 20Hz and 40Hz 449 probe trials were shuffled 10000 times to calculate chance performance; data is 450 451 mean±SD; unpaired two-sided t-test.

452 Methods

453 Ethical compliance

454 All animal procedures performed in this study were approved by the UK government

455 (Home Office) and by the Crick Institutional Animal Welfare Ethical Review Panel.

456 **Mice**

All mice used for behavioural experiments were C57/Bl6 males (Fig. 2 and 4, Extended
Data Fig. 3-5 and 10). In vivo imaging experiments were performed in 12-20 week old
heterozygous OMP-cre (³⁹; JAX stock #006668; Fig. 1, Extended Data Fig. 2 and 6) or
Tbet-cre (⁴⁰; Jax stock #024507; Fig. 3, Extended Data Fig. 7) mice crossed with the
Ai95(RCL-GCaMP6f)-D line (⁴¹; JAX stock #028865) of either sex. Extracellular unit (Fig.
and Extended Data Fig. 8) and whole-cell patch-clamp recordings (Extended Data
Fig. 9) were performed in 5-8 week old C57/Bl6 males. Mice were housed up to 5 per

464 cage in a 12/12h light/dark cycle. Food and water were provided ad libitum.

465 **Reagents**

- All odours were obtained in their pure form from Sigma-Aldrich, St. Louis MO, USA. Unless
 otherwise specified, odours were diluted 1/5 with mineral oil in 15 ml glass vials (27160-
- 468 U, Sigma-Aldrich, St. Louis MO, USA).

469 Statistical analysis and data display

To test for statistical significance between groups where appropriate we used either 470 paired or non-paired student t-test or, for non-parametric data, the Mann-Whitney U test, 471 472 or the Kolmogorov-Smirnov test to test the equality of probability distributions. Statistical test details and p-values are provided in figures and / or their respective figure 473 474 legends. Unless specified otherwise, boxplots were plotted using the MATLAB boxplot function with the median depicted as a thick line and default maximal whisker length of 475 476 1.5 * (q3-q1) where q3 and q1 indicate the 75th and 25th percentile, respectively. If 477 points were located outside this whisker range they were displayed individually as 478 outliers. Violin plots show the median as a black dot and the first and third quartile by the 479 bounds of black adapted bars. Mouse cartoons from were 480 https://scidraw.io/drawing/123 and /49.

481 High-speed odour delivery device

The odour delivery device was based on a modular design of four separate odour 482 channels, and consisted of an odour manifold for odour storage, a valve manifold for 483 484 control of odour release and hardware for controlling and directing airflow through the 485 system (Fig. 1b). The odour manifold was a 12.2x3.2x1.5 cm³ stainless steel block with 4 milled circular indentations (1 cm radius). Within each of these indentations was a 486 threaded through-hole for installation of an input flow controller (AS1211F-M5-04, SMC, 487 488 Tokyo, Japan) and an output filter (INMX0350000A, The Lee Company, Westbrook CT, 489 USA). For each inset, the cap of a 15 ml glass vial (27160-U, Sigma-Aldrich, St. Louis MO, 490 USA) with the centre removed was pushed in and sealed with epoxy resin (Araldite Rapid, Huntsman Advanced Materials, Basel, Switzerland). This meant that glass vials could bescrewed in and out of the insets for rapid replacement.

- Solenoid valves typically limit high-fidelity odour stimulation resulting in odour rise 493 494 times of several 10s of milliseconds under optimal conditions⁴². We thus employed highspeed micro-dispense valves with custom electronics for pulse-width modulation to 495 maximize bandwidth: 4 VHS valves (INKX0514750A, The Lee Company, Westbrook CT, 496 497 USA) were installed in a 4-position manifold (INMA0601340B, The Lee Company, Westbrook CT, USA) with standard mounting ports (IKTX0322170A, The Lee Company, 498 499 Westbrook CT, USA). Each valve was connected to a corresponding odour position in the odour manifold with 10 cm Teflon tubing (TUTC3216905L, The Lee Company, Westbrook 500 501 CT, USA). Each valve was controlled by digital commands via a spike-and-hold driver. 502 Each digital pulse delivered to the spike-and-hold driver delivered a 0.5 ms, 24 V pulse to the valve (to open it), followed by a 3.3 V holding pulse lasting the rest of the duration of 503 504 the digital pulse. This spike-and-hold input allowed for fast cycling of the valve without switching between 0 and 24 V at high frequencies to prevent from overheating the valve. 505 Each valve was controlled by an individual spike-and-hold driver. Up to 4 drivers could 506 507 be controlled and powered with a custom-made PSU consisting of a 24 V power input and 508 a linear regulator to split the voltages into a 24 V and 3.3 V line, as well as control inputs 509 taking digital signal input and routing it to the appropriate valve. Pulse profiles for calibration and stimulus production were generated with custom Python software 510 (PyPulse, PulseBoy; <u>github.com/RoboDoig</u>) allowing to define pulse parameters across 511
- 512 multiple valves using a GUI.
- 513 To generate airflow through the olfactometer, a pressurised air source was connected to a filter (AME250C-F02, SMC, Tokyo, Japan) and demister (AMF250CF02, SMC, Tokyo, 514 Japan) and then split into two separate lines, the input line and carrier line. Both lines 515 516 were then connected to a pressure regulator (AR20-F01BG-8, SMC, Tokyo, Japan) and 517 flow controller (FR2A13BVBN, Brooks Instrument, Hatfield PA, USA). The main line was then connected to the input of the valve manifold. The input line was split into 4 separate 518 519 lines and connected to the input flow controllers (set to 0.25 L/min) on each odour 520 position of the odour manifold. The output of the valve manifold was fitted with MINSTAC 521 tubing (TUTC3216905L, The Lee Company, Westbrook CT, USA). Where the design was 522 scaled up (e.g. to include 8 odour positions) the valve manifold outputs were connected 523 and consolidated to a single output with 3-way connectors (QSMY-6-4, Festo, Esslingen 524 am Neckar, Germany). Shape and reliability of odour pulses depended strongly on low 525 volume headspace and low pressure levels (0.05 MPa). Flow change due to odour pulses 526 was always compensated by mineral oil presentation (e.g. light grey in Fig. 1ci).
- 527
- 528 Odour characterisation

529 Signal fidelities were calculated by first subtracting the average amplitude of troughs

530 from the average amplitude of peaks during a pulse train and then subsequently dividing

this peak-to-trough value by the difference of average peak amplitude subtracted by

- 532 baseline amplitude (*SignalFidelity* = (meanPeak meanTrough) / (meanPeak baseline)).
- This results in a value between 0 and 1, with 1 being perfectly modulated odour pulses.

534 Behaviour

535 Automatic operant conditioning of cohorts of mice (AutonoMouse)

In AutonoMouse, groups of mice (up to 25) implanted with an RFID chip are housed in a 536 537 common home cage (Fig. 2a, for detailed description see²⁴). Within the common home cage of AutonoMouse, mice have free access to food, social interaction and environmental 538 539 enrichment. Water is not freely available in the system, but can be gained at any time by completion of an operant conditioning go/no-go task. To access these behavioural tasks, 540 mice must leave the home cage and enter a behavioural area. This behavioural area 541 542 contains the odour port and a lick port through which water rewards can be released. The lick port is also connected to a lick sensor, which registers the animal's response (its lick 543 544 rate) in response to the task stimuli. As animals can only gain their daily water intake by completing behavioural tasks, mice are motivated to complete long sequences of trials 545 546 without manual water restriction.

547

548 Training on temporally structured odours

We aimed to probe whether mice could perceive a particular temporal feature of 549 550 naturally occurring odour signals: temporal correlations between odour signals. In 551 particular, we aimed to investigate this question with the simplest possible case: whether 552 mice could discriminate perfectly correlated from perfectly anti-correlated odour stimuli. 553 All tasks followed a standard go/no-go training paradigm. Animals were presented with 554 two odours presented in either a correlated pattern or an anti-correlated pattern (Fig. 2d and Extended Data Fig. 4a-c). For roughly half of all animals, the correlated pattern was 555 556 S+ (rewarded) and the anti-correlated pattern was S- (unrewarded); in the other half of 557 the group this reward valence was reversed. All stimuli were 2 s long. A water reward 558 could be gained by licking such that licking was detected for at least 10% of the stimulus 559 time during an S+ presentation (a "Hit"). Licking for the same amount of time during S-560 presentation resulted in a timeout interval of 7 seconds. In all other response cases, the 561 inter-trial interval was 3 seconds and no water reward was delivered.

562

563 Stimulus structure

All anti-correlated and correlated stimuli on each trial followed a common pattern in their construction. Generally, wherever an odour position is inactivated a blank position should be activated to compensate for flow change. There should also be no consistent differences in the amount of odour or flow released during the stimulus between correlated and anti-correlated stimuli. The detailed algorithm for stimulus generation is as follows:

- 570
- 571 I. Correlated or anti-correlated/uncorrelated odour pulses (Fig. 2d-k and Extended
 572 Data Fig. 3 and 5)
- 573 1. The stimulus is chosen to be correlated or anti-correlated/uncorrelated.
- 5742. A set of 1-2 positions each for odour 1 and odour 2 and 2-3 positions for blank are
- 575 randomly chosen from a pre-defined subset of 6 of the 8 total positions. For

example, a valid combination could be odour 1 at position 1, 2; odour 2 at position 5; and blank at position 3 and 7. (see Fig. 2g and Extended Data Fig. 4b)

- 578 3. A guide pulse is created at the desired frequency (e.g. 2 Hz pulse with 50% duty, see
 579 Supplementary Methods Fig. 1c) for all positions that follows the chosen stimulus
 580 structure.
- 4. The relative contributions of each position to the total stimulus are randomly generated. At each time point in the stimulus, only two position types should be active (e.g. odour 1 and blank for an anti-correlated stimulus) so the maximum contribution for any position type is 50% of the total release amount. Where two positions have been chosen for a position type, their relative contributions should add to 50% (Extended Data Fig. 4b).
- 587 5. The guide pulses are pulse-width modulated according to the relative contributions
 588 of each position (Supplementary Methods Fig. 1c). Pulse-width modulation
 589 (PWM) is at 500 Hz with some added jitter in the duty to avoid strong tone
 590 generation.
- 591 6. For uncorrelated pulses, temporal offsets are added in one channel according to a
 592 distribution of time delays that follow the desired correlation structure between the
 593 two odour pulses (Extended Data Fig. 3e,f).
- 594
- 595 II. "One source" and "Source separated" naturalistic plumes (Fig. 4c-e and Extended
 596 Data Fig. 10)
- 597 1. The stimulus is chosen to be "One source" or "Source separated".
- 598 2. A plume bank of plume pairs obtained from indoor PID recordings is created. Each trial will contain 2 plumes, each representing one odour recording originating from 599 600 one source or from sources positioned 50 cm apart. To maintain consistency in trial 601 length between behaviour experiments, a 2 s time window from each plume was 602 selected from the middle of each 5 s recording, such that odour was always present in the first 500ms of the trial. Trials where the correlation of the 2 s window was 603 604 vastly different from the original 5 s were excluded from the plume bank. This procedure resulted in a plume bank containing 72 plume pairs for the separated 605 source condition and 48 plume pairs for the one source condition. 606
- 607 3. An odour plume pair is randomly selected from the plume bank, from the608 corresponding category.
- 4. The odour that will be used to replicate each plume in a pair and the positions in the odour delivery device that will be used for that purpose are randomly assigned, as described previously. For each odour valve active, a blank valve will also be activated to produce an "anti-plume" structure, to compensate for the changes in flow created by odour delivery.
- 5. Plumes are recreated from the chosen PID recordings. Each trace is normalised to
 between 0 and 1, and then converted into a series of binary opening and closing
 times. The length of the openings and closings relate directly to the value of the
 normalised signal, a value of one translates to a continuous opening, and a value of

- 618zero translates to continuously closed. This series of openings and closings are619relayed to the valves and the resulting output resembles the original plume.
- 620
- 621 III. Perfectly correlated plume trials (probe trials, **Fig. 4e**)
- A 2 s window is chosen from the source separated plume bank (plume structures previously associated with opposite reward valence 5 trials) or from independent plume recordings obtained in a different environment than the original recordings (completely novel plume structures 10 trials).
- 626 2. The chosen plume structure is replicated using both odour channels, resulting in a627 plume where both odour components fluctuate in a perfectly correlated manner.
- 628
- 629 IV. Frequency discrimination pulses (Fig. 2c and Extended Data Fig. 3a-d)
- 630 1. Two frequencies are chosen for discrimination (e.g. 2 Hz vs. 20 Hz).
- 631 2. For each trial one of the frequencies is chosen for presentation.
- 632 3. Valves are selected for presentation of both odours.
- 4. A guide pulse is created for each odour channel that pulses at the desired trial
 frequency with 50% duty, such that pulse alternates between channels at the given
 frequency.
- 5. Guide pulses are pulse-width-modulated as for correlated/anti-correlated stimuli.
- 637
- 638 *Task structure for the correlation experiment* (Fig. 2d-k, Extended Data Fig. 3-5)
- 639Task frequency was randomised from trial to trial in a range between 2-81 Hz. The choice640of frequency was with weighted probability divided into 3 frequency bands. E.g. this task641could be arranged such that 2-20 Hz would be chosen with P = 0.6, 21-40 Hz with P = 0.3642and 41-81 Hz with P = 0.1. Within each of these frequency bands, the choice of individual643task frequency was based on a uniform distribution. Thus, few trials were performed for
- 644 frequencies exceeding 40 Hz resulting in more "noisy" behavioural performance data in
- 645 **Fig. 2k**.
- 646
- 647 Onset detection
- For the onset detection experiments (**Extended Data Fig. 5f-i**) animals were trained to discriminate perfectly correlated (e.g. S+) from perfectly anti-correlated stimuli (e.g. S-) and probed with partially altered stimuli where the onset (first cycle) of the probe S+ stimuli was anti-correlated and probe S- stimuli where the onset (first cycle) was correlated. Performance during these probe trials is then compared to the average performance during training ($perf_{train}$).
- 654 We calculated the expected average animal performance on the probe trials based on two
- models (prediction data, **Extended Data Fig. 5f,g**): Model 1 assumed the animals were
- taking any part of the stimulus into account equally when making a decision. Model 2
- assumed that only the onset of the stimulus would contribute to discrimination. Thus for
- Model 1, a stimulus of frequency f (e.g. 10 Hz) that was sampled for t_{sample} consisted of a
- 659 "shifted" onset component of one cycle for S+ (1/f) and half a cycle for S- (0.5/f)

660 corresponding to a fraction of $frac_{onset} = 1/f/t_{sample}$ of the entire stimulus and a 661 "normal" residual ($frac_{res} = 1 - frac_{onset}$). Thus, the predicted probe trial performance 662 would be:

663 (1) predicted probe trial $perf_{entire} = perf_{train} \times frac_{res} + (1 - perf_{train}) \times frac_{onset}$

In Extended Data Fig. 5i this prediction was calculated for the following parameters:
Sniff frequency: 6 Hz, inhalation fraction: 0.2, stimulus sampling time: 0.7 s, In Extended
Data Fig. 5g sampling time was varied as indicated.

For Model 2, ignoring inhalation timing, the prediction would be that preference would 668 be reversed (as onset correlations during probe trials are reversed). However, this 669 670 ignores the fact that odour stimuli during the exhalation period might not be detected. 671 Thus, to more accurately predict animals' performance for Model 2, we assume that the part of the stimulus that is detected as the "onset" is the first odour pulse during an 672 673 inhalation phase. During the probe trial, this will be the "inverted" first cycle if the 674 stimulus begins either during the inhalation phase or at most 1/f before the inhalation (then inhalation would start during the inverted first cycle of the probe trial). The 675 probability of this occurring is $perf_{onset} = (dur_{inh} + 1/f) / dur_{sniff}$ with dur_{inh} and 676 dur_{sniff} being inhalation and sniff duration respectively (provided $dur_{inh} + 1/f < 1$ 677 *dur*_{sniff}). Predicted probe trial performance for an "onset only" model would thus be: 678

679 (2) predicted probe trial $perf_{onset} = perf_{train} \times (1 - perf_{onset}) + (1 - perf_{train}) \times perf_{onset}$

Extended Data Fig. 5f,g displays the predictions of these two models in comparison to
the experimental data for a broad range of respiration patterns. The "prediction data" in
Extended Data Fig. 5i shows model predictions assuming typical sniff and sampling
parameters as indicated above (Sniff frequency: 6 Hz, inhalation fraction: 0.2, stimulus
sampling time: 0.7 s).

685

667

686 Controls

687 Control valves could be automatically added to the random frequency task. These tasks
688 produced their stimuli based on a subset of 6 valves and control valves could be added
689 automatically after a set period of trials to force the algorithm to produce stimuli from all
690 8 valves (see Fig. 2g,i switch control and Extended Data Fig. 4i-k).

A subgroup of animals was created in which the valve map was scrambled, as an ongoing control against animals learning extraneous variables in the task (see **Fig. 2k**, scramble control). The valve map was scrambled in the following way: One blank to odour 1, one odour 2 to blank, one odour 1 to odour 2 and one odour 1 to blank. Every few days all odour bottles were cleaned and replaced, odour positions changed and valves reassigned⁴³.

697 *Airflow and sound recordings*

Airflow and sound were recorded in AutonoMouse during trials at different frequenciesto ensure that the temporal structure of the odour is the only parameter that varies over

trials and that no tactile or auditory cues were present in the stimulus. A flow sensor

- (AWM5101VN, Honeywell, USA) and a microphone (NTG1, RØDE, Australia) were placed
 in close proximity to the AutonoMouse odour port. In total, 286 trials were recorded (2
 Hz: n = 75 correlated, n = 70 anti-correlated; 40 Hz: n = 69 correlated, n = 72 anticorrelated) using Audacity for sound and Spike2 (Cambridge Electronic Design, UK) for
 flow signals. Airflow and sound signal underwent spectral analysis (Fourier transform),
 as well as linear classification analysis (Extended Data Fig. 4d-g).
- 707
- 708 Training on naturalistic plumes

One group of animals (n = 12) were trained to discriminate between plumes derived from those originating from one source (S-, unrewarded) or from separated sources (S+, rewarded), using 2 s long stimuli produced as described above from the recordings shown in Fig. 4a. An additional 12 animals trained simultaneously on the reverse reward valence did not pass the performance criterion within the given timeframe and were not carried forward to probe trials.

715 To test whether correlation structure was a feature used by mice to perform the virtual source separation task, probe trials were introduced randomly at a frequency of 716 717 approximately 1 in 11 trials, with every instance of a probe trial repeated every 330 trials. 718 Probe trials consisted of perfectly correlated plumes or correlated/uncorrelated square 719 pulses produced as described above, presented at 3 different frequencies: 2 Hz, 20 Hz, 40 720 Hz. The feedback for probe trials was the same as for a training trial, with a reward or 721 time-out given based on the response of the mouse. No change in performance across repeated presentation of the probe trials was observed indicating that performance was 722 723 not due to putative rapid re-learning. Of the 12 mice exposed to this protocol, a total of 9 724 mice reached all phases of the experiment.

- 725
- 726 Cohorts

727 The correlation discrimination experiment was performed in 3 separate experimental 728 cohorts (Fig. 2, Extended Data Fig. 3 -5: group 1, n = 14; group 2, n = 25 (one animal did 729 not successfully pass the pre-training); Fig. 4, Extended Data Fig. 10: group 3, n = 24, see above). Each cohort was organised into several subgroups, which performed slight 730 731 variations of the behavioural tasks in terms of reward valence and valves utilised, but 732 with the same underlying task aim. Half of the animals in each subgroup were trained on 733 correlated stimuli as the S+ rewarded condition, with the other half trained on anti-734 correlated as rewarded. Animals were further subdivided into groups, which were 735 trained on different subsets of valves as standard in the 8-channel olfactometer. For each 736 cohort, mice were once assigned to each of these subgroups based on performance in a simple pure odour discrimination at the beginning of the experiment - group 737 738 membership was randomised until no significant (ANOVA, Tukey-Kramer) differences in 739 performance could be extracted between these subgroups on this task.

740

741 Data analysis

AutonoMouse behavioural data was converted to MATLAB data format using the
Conversion module of the Python autonomouse-control package
(github.com/RoboDoig). All subsequent analysis was performed with custom-written
MATLAB scripts unless otherwise specified.

- 746 All behavioural performance within a specified trial bin was calculated as a weighted
- 747 average of S+ vs. S- performance:
- 748 $performance = \frac{(Hit / S+) + (CR / S-)}{2}$

Where S+ is the total number of rewarded trials, S- is the total number of unrewarded
trials, Hit is the total number of rewarded trials in which a lick response was detected, CR
(correct rejection) is the total number of unrewarded trials in which no lick response was
detected.

- For random stimulus pulse frequency experiments (e.g. **Fig. 2j,k**) trials were binned approximately by half-octave for performance analysis. The exact intervals were f(Hz) = [2, 3, 4, 5, 6:7, 8:10, 11:13, 14:17, 18:22, 23:29, 30:37, 38:48, 49:62, 63:81]. Reaction time **(Extended Data Fig. 5)** was calculated from S+ trials for each animal as the time to the first lick after stimulus onset. For presentation of learning curves (**Fig. 2h, 4d**) accuracy
- 758 was calculated over 100-trial sliding windows.
- Motion magnification of the respiration camera video recordings (Extended Data Fig. 5 759 and Supplementary Video 2) was performed with phase-based video motion 760 processing with correction for large body movements based on MATLAB scripts by ⁴⁴ 761 (phaseAmplifyLargeMotions). Parameters for phase amplification were: blurring $\sigma = 1$, 762 magnification a = 50, amplification in frequency band between 2-13 Hz. Following 763 magnification, static ROIs for each video were selected in Bonsai (http://www.kampff-764 765 lab.org/bonsai/, ⁴⁵) over the animal flank. An adaptive binary threshold was applied to 766 the ROI to segment the animal body from the video background. Respiration rate was 767 extracted from the total size of the ROI occupied by the body over time.

768 Olfactory sensory neuron population model

769 Overview

We modelled the olfactory sensor neuron (OSN) population as noisy integrate-and-fire
neurons integrating a filtered odour pulse and with independent (cell-specific) noise to
qualitatively match experimental data⁴⁶. The square of the resulting mean population
firing rate was convolved with a calcium imaging filter to produce a model of the observed
calcium imaging signal. All code and related data for the model can be found at
https://github.com/stootoon/crick-osn-model-release.

- 776
- 777 Odour Input Current
- 778 The olfactory input current I_t to each OSN was modelled as a filtered version of the odour 779 pulse input O_t :

$$\tau_C \ \frac{dI_t}{dt} = -I_t + O_t$$

781 This filtering models filtering of the nasal cavity, transport through the mucous, and 782 chemical transduction from odour concentration to receptor channel opening.

- 783
- 784 Olfactory Sensory Neurons
- Each OSN was modelled as a noisy integrate-and-fire neuron. Each OSN membrane performs a noisy integration of the olfactory input current I_t so that the membrane voltage V_t satisfies the following stochastic differential equation:
- 788 $\tau_V \, dV_t = (I_t V_t) dt + \sigma \, dB_t.$
- Here B_t is standard Brownian motion and σ is the standard deviation of the membrane voltage noise. The OSN generates a spike whenever its membrane voltage exceeds a spiking threshold θ :
 - $S_t = \begin{cases} 1 & If \ V_t \ge \theta; \\ 0 & otherwise. \end{cases}$

793

792

- 794Upon spiking the membrane voltage is clamped to a refractory voltage V_{ref} for a period795of t_{ref} seconds. The mean instantaneous firing rate of the population is computed as
- $\bar{S}_t = \frac{1}{N} \sum_{n=1}^N S_t^n$
- 797 where S_t^n is the spiking activity of OSN *n*.
- 798
- 799 Calcium imaging signal
- To model the calcium imaging signal the mean firing rate is squared and convolved with the imaging kernel h_t to form the calcium imaging signal C_t :

$$C_t = \left(\bar{S_t}^2\right) * h_t$$

803 The calcium imaging kernel is an alpha function:

805 804

802

A list of parameters is given in **Supplementary Table 1**. All parameters were fit manually: Parameters τ_C , τ_V and σ were set to produce a qualitative match in time courses between model membrane voltage traces and the suction current traces in Figure 2 of reference ⁴⁶. The remaining parameters were adjusted to produce a qualitative fit between model and the dynamics of the observed calcium imaging traces.

 $h_t = t \ e^{-t/\tau_h}$

811

812 Generating model glomeruli

813 Overview

814 We generated 100 model glomeruli by randomly varying a subset of the model OSN

parameters described in the previous section (τ_V , τ_C , σ , θ , *a*). Specifically, we picked the parameters of each glomerulus by selecting uniformly within ± 25% of the centre value

- of each parameter. All 5000 OSNs within each glomerulus had the same parameters, and
- 818 differed only due to the random noise applied to their membrane voltages. The range of
- 819 variation is shown in **Supplementary Table 2**.

820

- 821 The effect of concentration was modelled by linear scaling of the input waveforms. For
- 822 each setting of PPI and concentration the model was run to simulate 25 consecutive trials
- of length 2.5 seconds each, with the odour onset at 0.1 seconds into each trial. The first 5
- trials of data were discarded to allow the model to 'settle,' yielding 20 trials for each
- 825 condition that were used in subsequent analyses (Extended Data Fig. 1g,h, Extended
- 826 **Data Fig. 2j,k**).
- 827

828 Classifying glomerular outputs

829 Predictors

The predictors used for classification were the response integrals for each glomerulus, defined as the instantaneous mean firing rate of the OSNs in the glomerulus, filtered by the Ca2+ imaging filter (see section "Olfactory sensory neuron population model") and summed over the 2 seconds following odour onset in each trial. Since the scale of the responses is arbitrary, we scaled the response integrals by their overall standard deviation, computed over glomeruli and trials.

- 836
- 837 Labels

838 Trials were labelled by their paired-pulse interval (PPI), or a combination of PPI and839 concentration, depending on the task.

- 840
- 841 Classifiers

The classifiers used were support vector machines with linear kernels and l2 regularization as implemented by the 'LinearSVC' function of the Python scikit-learn library. The setting of the penalty parameter C and whether or not to learn an intercept were determined by cross-validation with scikit-learn's 'GridSearchCV'. The values of C considered ranged in powers of 10 from 10⁻⁴ to 10⁴.

847 Computing Decoding Accuracy

The decoding accuracy for a given subset of n glomeruli was computed as the average accuracy over 10 cross-validation trials for the results in **Extended Data Fig. 1h**, and 40 cross-validation trials for the results in **Extended Data Fig. 2j,k**. In each cross-validation trial, the classifier was trained on a random 90% of the trials, tested on the remaining 10%, and the accuracy recorded. The random subsets were stratified i.e. constrained to have the same fraction of trials from each class as the full dataset when possible. The mean accuracy across cross-validation trials was recorded as the accuracy for that subset.

- 855 To compute the shuffled performance, the labels of the training and test trials were
- 856 shuffled in each cross-validation trial before the classifier accuracy was computed.
- 857
- 858 Decoding PPI from the responses of model glomeruli
- To determine how decoding accuracy was affected by the size of the population used we
- selected a random subset of n glomeruli and computed the decoding accuracy as
- described above. This was repeated for 256 random subsets of n glomeruli generating

- 862 256 unshuffled and 256 shuffled accuracies. The subset size n was varied from 1 (using
- 863 only a single glomerulus) to 100 (using the full population; **Extended Data Fig. 1h**). Note
- that for $n \ge 99$, some subsets are likely to have been repeated because there are fewer
- than 256 possible subsets of size 99 and 100. The observed variability in accuracy in
 those cases is then due mainly to the random determination of training and testing trials.
- 867
- 868 Decoding PPI and concentration from the responses of model glomeruli
- To compute the decoding accuracy when decoding PPI and concentration, we followed a very similar procedure to the previous section, but fixed the population size at the maximum of 100 and varied the stimulus concentration from 0.5 to 5 in steps of 0.5 to cover a factor of 10 range in concentration as used in the experimental data (Extended Data Fig. 2j-m). For comparison, the results in Extended Data Fig. 1h were for a concentration of 1. Decoders were trained to extract just concentration, or PPI and concentration.

876 In vivo two-photon imaging

877 Surgical and experimental procedures

- Prior to surgery all utilised surfaces and apparatus were sterilised with 1% trigene. Mice 878 879 were anaesthetised using a mixture of fentanyl/midazolam/medetomidine (0.05 mg/kg, 5 mg/kg, 0.5 mg/kg respectively). Depth of anaesthesia was monitored throughout the 880 procedure by testing the toe-pinch reflex. The fur over the skull and at the base of the 881 882 neck was shaved away and the skin cleaned with 1% chlorhexidine scrub. Mice were then placed on a thermoregulator (DC Temperature Controller, FHC, ME USA) heat pad 883 884 controlled by a temperature probe inserted rectally. While on the heat pad, the head of 885 the animal was held in place with a set of ear bars. The scalp was incised and pulled away 886 from the skull with four arterial clamps at each corner of the incision. A custom head-887 fixation implant was attached to the base of the skull with medical super glue (Vetbond, 888 3M, Maplewood MN, USA) such that its most anterior point rested approximately 0.5 mm 889 posterior to the bregma line. Dental cement (Paladur, Heraeus Kulzer GmbH, Hanau, Germany; Simplex Rapid Liquid, Associated Dental Products Ltd., Swindon, UK) was then 890 891 applied around the edges of the implant to ensure firm adhesion to the skull. A 892 craniotomy over the left olfactory bulb (approximately 2 x 2 mm) was made with a dental 893 drill (Success 40, Osada, Tokyo, Japan) and then immersed in ACSF (NaCl (125 mM), KCl 894 (5 mM), HEPES (10 mM), pH adjusted to 7.4 with NaOH, MgSO4.7H2O (2 mM), 895 CaCl2.2H2O (2 mM), glucose (10 mM)) before removing the skull with forceps. The dura was then peeled back using fine forceps. A layer of 2% low-melt agarose diluted in ACSF 896 897 was applied over the exposed brain surface before placing a glass window cut from a 898 cover slip (borosilicate glass 1.0 thickness) using a diamond knife (Sigma-Aldrich) over the craniotomy. The edges of the window were then glued with medical super glue 899 (Vetbond, 3M, Maplewood MN, USA) to the skull. 900
- 901 Following surgery, mice were placed in a custom head-fixation apparatus and transferred902 to a two-photon microscope rig along with the heat pad. The microscope (Scientifica
- 903 Multiphoton VivoScope) was coupled with a MaiTai DeepSee laser (Spectra Physics, Santa

904 Clara, CA) tuned to 940 nm (<50 mW average power on the sample) for imaging. Images
905 (512 x 512 pixels) were acquired in SciScan (Scientifica, UK) with a resonant scanner at
906 a frame rate of 30 Hz using a 16x 0.8 NA water-immersion objective (Nikon). The output
907 of a 4-channel version of the temporal olfactometer described above was adjusted to
908 approximately 1 cm away from the ipsilateral nostril to the imaging window, and a flow
909 sensor was placed next to the contralateral nostril for continuous respiration recording.

- 910
- 911 Awake recordings

For implantation of the head-plate, mice were anaesthetized with isoflurane in 95% oxygen (5% for induction, 1.5-3% for maintenance). Local (mepivacaine, 0.5% s.c.) and general analgesics (carprofen 5 mg/kg s.c.) were applied immediately at the onset of surgery. After surgery, animals were allowed to recover for 7 days with access to wet diet and, after recovery, habituated to the head-fixed situation for at least 15 min on three consecutive days preceding the imaging experiment

- 917 consecutive days preceding the imaging experiment.
- 918

919 *Odour stimulation*

For paired-pulse experiments, ethyl butyrate was diluted in mineral oil at the ratio of 1:5
and installed into a 4-channel version of the high-speed odour delivery device (15 ml per
vial) along with two blank positions (15 ml mineral oil). Odour concentration range was
adjusted over 10 steps on a logarithmic scale with a factor of 1.25 by modulating odour
pulse-width.

925 For correlated vs. anti-correlated stimulus experiments, stimuli were generated from 926 mixtures of physically mixed monomolecular odorants in order to ensure high 927 probability of finding odour responsive cells in the dorsal olfactory bulb using custom 928 Python Software (PulseBoy). Binary mixtures were diluted in mineral oil at the ratio of 929 1:5 and installed into a 4-channel version of the high-speed odour delivery device (15 ml per vial) along with two blank positions (15 ml mineral oil). Mix 1: ethyl butyrate + 2-930 hexanone, mix 2: isoamyl acetate + cineole. During glomerular imaging experiments 931 932 (Extended Data Fig. 6), six odours (A-F) where presented either individually or in pairs: 933 A (ethyl butyrate), B (2-hexanone), C (isoamyl acetate), D (cineol), E (ethyl tiglate) and F ((+)-fenchone). For all stimuli, odour valve offsets were compensated by opening a 934 935 corresponding blank position valve to ensure no global flow changes occurred over the 936 course of the stimulus. All stimuli were repeated between 16-50 times with at least 15 s 937 inter-stimulus interval.

- 938
- 939 Data analysis

940 For M/TC imaging, motion correction, segmentation and trace extraction were
941 performed using the Suite2p package (<u>https://github.com/MouseLand/suite2p</u>;⁴⁷).
942 Putative neuronal somata and dendritic segments were automatically identified by
943 segmentation and curated manually. Soma and neuropil fluorescence traces were
944 extracted and neuropil fluorescence was subtracted from the corresponding soma trace.
945 Further analysis was performed with custom written scripts in MATLAB.

946 M/TCs were recorded in 17 fields of view (FOV) from 6 individual Tbet-cre:Rosa-947 GCaMP6f animals, with 40 \pm 9.23 (mean \pm SD; range 27-48) cells per FOV and 30.25 \pm

- 948 12.97 (mean ± SD; range 7-53) M/TC dendrites,
- 949 For glomerular imaging experiments, ROIs corresponding to glomeruli were manually950 delineated based on the mean fluorescence image. Fluorescence signal from all pixels
- 951 within each ROI was averaged and extracted as time series. $\Delta F/F = (F-F0)/F0$, where F =
- raw fluorescence and F0 was the median of the fluorescence signal distribution.
- Glomerular signals from a total of 15 individual OMP-cre:Rosa-GCaMP6f animals were
 recorded with 28 ± 4.34 (mean ± SD; range 20-36) glomeruli per animal (Extended Data
 Fig. 6a).
- 956 Where the odour stimulus was not inhalation-triggered, traces were post hoc aligned to the first inhalation after odour onset. Calcium response integrals were calculated for a 957 958 range of window durations starting from odour onset (100-5000 ms). To analyse how 959 well odour responses predicted stimulus correlation on a trial-to-trial basis, we generated a linear discriminant classifier from the data set and analysed prediction 960 accuracy. For the classifier, we performed 50% holdout validation, splitting the data 961 randomly into a training set and test set with equal numbers of samples. We then 962 963 performed linear discriminant analysis on the training data set to determine the best linear boundary between 10 vs. 25 ms pulse interval stimulations or correlated vs. anti-964 965 correlated data. Classifier performance was then validated on the test data set. To determine the effect of number of ROIs used on classifier performance, we iteratively 966 trained multiple classifiers on random subsets of ROIs with increasing numbers of ROIs 967 968 within each set. For each ROI subset size, 100 classifiers were trained and the mean ± SD 969 of their performance accuracy was calculated. All classifier analysis was performed on 970 individual, unaveraged trials.
- 971

972 Glomerular imaging classifiers in Extended Data Fig. 6k,l

The classifiers used in **Extended Data Fig. 6k,l** were trained separately for each odour 973 pair, each frequency, and each time window. The inputs for classification were the 974 averaged responses of the 145 glomeruli in a given time window for 24 odour 975 976 presentation trials, where the odours were fluctuating in a correlated manner in half of the trials, and in an anti-correlated manner in the remaining half. Within the correlated 977 and anti-correlated subsets of trials, half had the first odour in the pair phase-shifted by 978 979 180 degrees, and the remaining half had no phase shift. The classification task was to determine whether the glomerular responses in a given trial were evoked by correlated 980 or anti-correlated odour fluctuations. 981

982

Because we had far fewer trials (24) than glomeruli (145) it was important to use regularized classifiers to avoid overfitting. To promote interpretability of the decision boundaries learned by the classifiers we opted for sparsity-promoting regularizers and settled on the Lasso, evaluated as a classifier by taking the sign of its output computed after the addition of a small amount of noise (to decide ambiguous classifications). The implementation of the Lasso we used was 'LassoLarsCV' provided by the Python scikitlearn library because it converged readily, gave very good classification performance, and
automatically tuned the weighting of the sparsity penalty. Inputs to the classifier were
standardized to have mean zero and unit variance across trials. We found that it was
important to learn the classification weights without intercept to avoid overfitting.

993

1006

994 The performance of a classifier was determined by cross-validation, where in each cross-995 validation iteration, the classifier was trained on a random $\sim 90\%$ of the trials (21 trials) and tested on the remaining $\sim 10\%$ (3 trials), and the test accuracy recorded. The random 996 997 subsets were selected in a stratified manner, meaning that the fraction of correlated and 998 anti-correlated trials in the subset were kept as close as possible to their fraction in the 999 full dataset (50/50). This meant that 10 of the 21 training trials were of one type and 11 of the other. This procedure was performed for 10 cross-validation iterations, and the 1000 1001 average performance over these repeats was recorded as the performance of the 1002 classifier. The shuffled performance was computed the same way but with training and test labels shuffled in each iteration. The entire procedure was then repeated for each of 1003 100 different random seeds to produce a distribution of classification accuracies, whose 1004 1005 means and standard deviations are plotted in Extended Data Fig. 6k,l.

1007 Because we used a sparsity-promoting classifier it was straightforward to determine which glomeruli were contributing to a particular classification decision. We found that 1008 if we used all 145 glomeruli available then frequently glomeruli would be selected for 1009 noisy fluctuations of their responses that were by chance 'informative' for the 1010 1011 classification. To avoid the inclusion of such noisy responses, we filtered glomeruli for responsivity. To determine the responsivity of a glomerulus, the mean $\bar{X}_{glom,baseline}$ and 1012 1013 standard deviation $\sigma_{glom, baseline}$ of its responses pooled across all baseline bins (defined as the 3 seconds before odour onset) and across all trials for the given odour pair and 1014 1015 frequency were first computed. A Z-score was then computed for its averaged response for the given time bin and for each trial by comparing this response to the baseline 1016 1017 activity according to

1018
$$Z_{glom,trial} = \frac{X_{glom,trial} - \bar{X}_{glom,baseline}}{\sigma_{glom,baseline}/\sqrt{n_{wnd}}}$$

where $X_{glom,trial}$ is the response of the glomerulus in the given time window and n_{wnd} is 1019 the number of time bins constituting the window. The scaling of the baseline standard 1020 1021 deviation is to account for the reduction in variance due to the averaging over time bins used to compute the response. A glomerulus was considered responsive in a given trial if 1022 1023 the absolute value of its Z-score as computed above was greater than 1 on three-quarters or more of the trials. Such a thresholding ensured that the number of responsive 1024 glomeruli was almost always zero before odour onset, but rose to a peak ~125 of the 145 1025 glomeruli available when 2 s windows were used. Reducing the window size reduced the 1026 peak number of glomeruli, but at least 25 glomeruli were used during the peak responsive 1027 1028 period in all cases, and frequently many more. This filtering also meant that some time

windows late in the response contained no responsive glomeruli for some window sizes,
which explains the `patchiness' observed in Extended Data Fig. 6k,l.

1031 Extracellular recordings

1032 Surgical and experimental procedures

5-8 week old C57BL/6Jax mice were anaesthetised using a mixture of ketamine/xyazline
(100mg/kg and 10mg/kg respectively) by intraperitoneal (IP) injection. An IP line was
inserted after the initial injection to allow for easier and more regular subsequent
injections of anaesthetics. Surgery was carried out as described above for two-photon
imaging, up until the application of agar and cranial window.

- 1038 Following surgery, mice and custom platform were transferred to the extracellular 1039 recording set up. A flow sensor (A3100, Honeywell, NC, USA) was placed in front of the 1040 contralateral nostril whilst an output from the temporal olfactometer was positioned in front of the ipsilateral nostril. A Ag/Ag⁺Cl⁻ reference coil was immersed in the well, over 1041 1042 the left hemisphere of the skull. The reference wire was connected to both the reference and ground of the amplifier board (RHD2132, intan, CA, USA), which was connected 1043 1044 (Omnetics, MN, USA) to a head-stage adapter (A32-OM32, NeuroNexus, MI, USA). A 32channel probe (A32-Poly3, NeuroNexus, MI, USA) was connected to the adapter, and the 1045 tip of the probe was manoeuvred to be positioned 1-2cm above the craniotomy. The 1046 1047 adapter and probe were held above the craniotomy using a micromanipulator (PatchStar, Scientifica, UK) set at 90 degrees to the surface of the brain. The probe was moved 1048 1049 towards the surface of the OB, whilst being observed through a surgical microscope. Once 1050 the probe was in contact with the surface, but had not entered the brain, the 1051 manipulator's Z position was set to zero. The signal from the probe was streamed through 1052 OpenEphys acquisition board and software (OpenEphys, RI, USA). The probe was 1053 inserted at $< 4 \mu m/s$ until the number and amplitudes of spikes began to decrease on deeper channels, indicating the tip of the probe was exiting the MC layer. This was found 1054 1055 to be between 400-600 µm from the surface of the OB. From here, the probe was left for 1056 10 minutes for neural activity to stabilise before recording began.
- 1057

1058 Odour stimulation

Odours were presented using an 8-channel version of the high-speed odour delivery
device, 4 of which contained odours (A: ethyl butyrate, B: 2-hexanone, C: isoamyl acetate,
D: eucalyptol) and 4 contained blank (mineral oil) which were used to compensate for
flow changes. Trials either paired A and B or C and D together. Stimuli were repeated 64
times and had an 8 s inter-trial interval. Onset of odour was recorded using TTL pulses
passed through additional channels in the OpenEphys acquisition board. Trial starts were
triggered on inhalation as detected by the flowmeter.

- 1066
- 1067 Data analysis

1068 Spikes were sorted using Kilosort2 (<u>github.com/MouseLand/Kilosort2</u>;⁴⁸) and classified 1069 as 'good' when they displayed a strong refractory period visible in their auto-1070 correlogram, a typical waveform and a stable firing rate, as 'MUA' (multi-unit activity) if 1071 they presented a typical waveform but a weak refractory period, or 'noise' if they were 1072 suspected of being electrical or mechanical interference. For a first-pass analysis units were classified as "differentially responding" to correlated and anti-correlated stimuli if 1073 1074 units were found to have significantly (p < 0.01, Mann Whitney u test) different spike time distributions during 4 s post odour onset. However, the cut-off for such distinction will 1075 1076 always be somewhat arbitrary. For the majority of the analysis, we therefore pooled *all* 1077 good units across experiments in a pseudo-population. All classifiers used for unit 1078 recording analysis were support vector machines (SVMs) with linear kernels with a low 1079 regularisation parameter, which translates to a greater freedom for a classifier to vary 1080 weights for any given component. Data was split into training and test sets prior to classification. Test sets either consisted of 26 trials (summed spike classifiers for 1081 correlated vs. anti-correlated) or 2 trials (PCA classifiers and short odour pulse 1082 1083 combinations). Data passed to the summed spike classifiers was pre-processed in one of 1084 two ways prior to classification.

Firstly, a rolling sum of detected spikes, within variable window sizes was used. The window sizes varied from 10 ms to 2000 ms. In addition to window size, window starts were also varied. Each window size was trialled with every possible window start from zero to four seconds minus the window size from odour onset, with 10 ms incremental changes. For example, a 500 ms window was tested with starts varying from 0 to 3500 ms from odour onset.

Secondly, the coefficients of PCs for units in each trial was used for classification. The PCs were found by applying PCA across all units and all training trials. Each PC represented a time series and hence the coefficients signify the strength at which that time series was followed by a given unit for a given trial. The two holdout trials were not used to find the PCs but were then projected onto them and their coefficients used as the test for the classifiers. All classifiers were repeated 1000 times with a random selection of holdout trials each time.

Finally, for the short odour pulse classification, classifiers were trained on summed spikes in windows of 500 ms post odour onset. Each classifier was trained on all but two hold out trials. To account for a varying number of trials between animals, training data was bootstrapped to 1000 trials of each type. Each trial was randomly selected from the initial pool of training trials, and each unit was bootstrapped independently. These classifiers were tested on the initial two hold out trials. This was repeated 1000 times with different bootstrapped datasets and different hold out trials.

Training data for all classifiers was scaled such that each feature (unit spike count / PC
coefficient) had a mean value of zero, and a standard deviation of one using the following
equation:

1109 where x is the initial value of the feature, u is the mean, s is the standard deviation, and z

is the scaled value. Means and standard deviations were calculated using the training dataso no information from the testing data could influence the scaling. The testing data was

- 1112 scaled using the same values as for the training. Scaling was applied during every repeat
- 1113 in this manner.

1114 Whole-cell patch recordings

- 1115 Whole-cell recordings were performed as described previously^{38,49}. Borosilicate pipettes
- 1116 (2 x 1.5 mm) were pulled and filled with (in mM) KMeSO3 (130), HEPES (10), KCl (7),
- 1117 ATP-Na2 (2), ATP-Mg (2), GTP-Nax (0.5), EGTA (0.05) (pH = 7.3, osmolarity ~290
- 1118 mOsm/kg). The OB surface was submerged with ACSF containing (in mM) NaCl (135),
- 1119 KCl (5.4), HEPES (5), MgCl2 (1), CaCl2 (1.8), (pH = 7.4 and ~300 mOsm/kg. Signals were 1120 amplified and low-pass filtered at 10 kHz using an Axoclamp 2B amplifier (Molecular
- 1121 Devices) and digitized at 40 kHz using a Micro 1401 analogue to digital converter
- 1122 (Cambridge Electronic Design).
- 1123 After zeroing the pipette tip position at the OB surface, the tip was advanced to reach a
- 1124 depth of \sim 200 μ m from the surface. Whole-cell patch-clamp recordings were obtained as
- 1125 described in⁵⁰. Series resistance was compensated and monitored continuously during
- 1126 recording. Neurons showing series resistance >25 M Ω were discarded from further
- analysis. To estimate the input resistance, a -50 pA current step was delivered at the start
- 1128 and end of each recording.
- 1129 The vertical depth of recorded neurons was calculated as the vertical distance from the
- 1130 brain surface. Respiration was recorded using a mass flow sensor (A3100, Honeywell, NC,
- 1131 USA) and digitized at 10 kHz. Odours were prepared and delivered as described above
- 1132 and triggered to the beginning of inhalation.
- 1133
- 1134 Data analysis
- 1135 Change in membrane potential
- 1136 Recordings were spike-clipped using a custom script written in Spike2 (Cambridge 1137 Electronic Design) and analysed in MATLAB (Mathworks, USA). All recordings were 1138 baseline subtracted as described previously⁵¹. The average change in membrane 1139 potential was defined as the difference between the average membrane potential over a 1140 2 s period before odour onset and the average membrane potential in the first 500 ms 1141 (~2 sniffs) after odour onset.
- 1142

1143 Change in spike frequency

- Action potentials were counted from raw traces, converted into spike frequency in 50 msbins and plotted as peri-stimulus time histograms (PSTH). The net change in spike
- 1146 frequency was defined as the difference of the average spike frequency during 2 s before
- 1147 onset and 500 ms after onset.
- 1148
- 1149 Arithmetic sum
- 1150 Baseline-subtracted traces obtained from independent component odour (A and B)
- 1151 presentations were either summed and averaged in an in-phase manner to generate the
- 1152 arithmetic sum equivalent of the correlated response or phase-shifted to generate the
- 1153 equivalent of the anti-correlated response.

1154	
1155	Arithmetic sum(correlated) = $(Vm_{odourA} + Vm_{odourB})/2$
1156	
1157	Arithmetic sum(anticorrelated)
1158	$= (Vm_{odourA} + shifted Vm_{odourB} + shifted Vm_{odourA} + Vm_{odourB})/4$
1159	

1160 Dual-energy fast photoionisation detection (defPID, Supplementary Methods Fig. 1161 2)

Two photoionisation detectors (200B miniPID, Aurora Scientific, Aurora ON, Canada) 1162 fitted with UV lamps of emission energy 10.6 eV (PID high) and 8.4 eV (PID low) were 1163 used to discriminate ethyl butyrate (EB, ionisation energy = 9.9 eV) from α -Terpinene 1164 1165 (AT, ionisation energy = 7.6 eV) or ethyl valerate (EV, ionisation energy = 10.0 eV) from tripropyl amine (TA, ionisation energy = 7.2 eV). To accommodate the lower voltage UV 1166 lamp, resonance circuitry in the PID headstage electronics was adjusted according to 1167 1168 manufacturer's recommendations. Specifically, potentiometer 'PT1' was adjusted up to the point where the 8.4 eV lamp began to glow. Further, we tested if the now converted 1169 1170 PID low was now sensitive to only AT and TA while not detecting EB and EV. The PID 1171 inlets were connected with a 3-way connector to detect incoming odours by both PIDs 1172 simultaneously from a common point. PID heads were held on lab stands with the PID 1173 inlet at approximately 4 cm above ground level.

- 1174
- 1175 Odour delivery

Odours were held in ceramic crucibles (5 cm diameter, 6 ml volume) covered in an airtight fashion using glass lids. Odours were released for 5 s with an inter-trial interval of 15 s by Arduino-based robots programmed to lift the lids from the crucibles using a servo motor (TowerPro SG-5010, Adafruit, UK). Lid lifting events were triggered by the Arduino board, recorded in Spike2 and defined as the onset of odour for analysis. Both the Arduino board and Spike2 were controlled by a portable computer and used the same clock for synchronisation. Experiments were carried out in a large open space, both indoors and

- 1183 outdoors (Supplementary Methods Fig. 2f,g).
- 1184 *Outdoors setup:* PIDs and odour delivery system as described above were used to record
- 1185 for multiple trials in different conditions on a day with low wind (\sim 8-12 mph \approx 3-5 m/s, 1186 recorded with a 2-axis ultrasonic wind sensor (Gill Instruments, Hampshire, UK)).
- 1187 Outdoor experiments were performed on a ~ 6 m x 10 m wooden patio structure
- 1188 surrounded by trees. Measurement of odour correlations in the outdoor setting were
- 1189 complicated by the presence of background odours: If background odours are detectable
- 1190 by both PIDs, measured correlation will be artificially inflated; if they can be ionized only
- 1191 by the "PIDhigh" they will artificially decrease the measured correlation.
- 1192 *Indoors setup:* A digitally controlled fan (2214F/2TDH0, ebm-papst, Chelmsford, UK) was
- 1193 placed at a distance of 325 cm facing the PID inlet. An exhaust line was situated behind
- the PID inlet to ensure the direction of air from the fan towards the PID inlet. During a
- recording, the fan was set to maximum speed such that it pushed approximately 550

1196cf/min (cubic feet per minute, ≈ 260 l/s) of air towards the PID inlet. A 25x25x25 cm1197Thermocool box was placed 200 cm downwind of the fan acting as an obstacle to air1198movement, promoting complex air movement patterns at the PID inlet. The pump at each1199PID was set to ≈ 0.02 l/s suction speed, unlikely to perturb overall airflow dynamics1200substantially.

- 1201
- 1202 *Recording conditions*

6 ml of the desired odour(s) were filled in two crucibles and placed in different locationsbased on the experimental conditions as described below:

1205

1206 1. Low energy only: The 'low-energy odour' (AT or TA) was placed 40 cm (radial distance 1207 d) away from the PID inlet, and displaced either 25 cm left or 25 cm right of the midline 1208 (the line between the PID inlet and the centre of the fan). The odour source was alternated 1209 between left and right positioning relative to the midline to remove any possible bias 1210 from positioning in the air stream. The purpose of this recording condition was to 1211 generate data to calculate the linear transformation from the low energy signal to the 1212 birth energy signal (Supplementary Methods Fig. 20 d)

- 1212 high energy signal (**Supplementary Methods Fig. 2c,d**).
- 1213 2. Mix: 3 ml EB + 3 ml AT (or 3 ml EV + 3 ml TA) was pipetted in one crucible and placed
 1214 either 25 cm left or 25 cm right of the midline at radial distances of 20 cm, 40 cm and 60
- 1215 cm. The purpose of this recording condition was to determine how the temporal structure
- 1216 of individual odours in a plume behaved when the odours were emitted from the same 1217 source.
- source.
 3. Separate: 3 ml EB and 3 ml of AT (or 3 ml EV and 3 ml TA) were individually pipetted
 in two different crucibles and placed at a radial distance of 40 cm from the PID inlet. For
- 1219 In two different cruciples and placed at a radial distance of 40 cm from the PID linet. For 1220 the s=50 cm apart condition, one odour source was placed 25 cm left of the midline while
- 1221 the other was 25 cm on the right of the midline and vice-versa (equal number of trials for
- 1222 both cases) separating the odour sources by 50 cm. This procedure was repeated for
- 1223 lateral distances of s=30 cm and s=10 cm. The '50 cm apart' case was repeated for radial
- 1224 distances of d=20 cm and d=60 cm. The purpose of this recording condition was to
- 1225 determine how the temporal structure of individual odours in a plume behaved when the
- 1226 odours were emitted from separated sources but were still free to mix in air.
- 1227
- 1228 Data analysis

1229Decomposition procedure: The low energy odour (AT) was recorded using both PIDs as1230described above. Assuming a linear relation between the recorded signals from the 21231PIDs, we plotted the recorded events with a linear regression fit (Supplementary1232Methods Fig. 2c) and calculated slope and R² value of the fit. The scaling factor (6.82 ±12330.356, mean ± SD) was calculated as the average slope of all linear fits for R² ≥ 0.9.

- 1234 The 'PID low' traces were multiplied by this scaling factor which was termed 'estimated
- 1235 low energy odour' (Supplementary Methods Fig. 2e). The 'estimated high energy odour'
- 1236 was calculated by subtracting the estimated low energy odour from the 'PID high' traces.

- 1237 Correlation calculation: Custom written scripts in MATLAB (Mathworks, USA) were used
- 1238 to calculate the correlation coefficient between the estimated low energy odour and the
- estimated high energy odour for all conditions. Box plots were obtained from these values
- 1240 using Igor Pro 6 (WaveMetrics, USA).

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- 1275

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1293 Author Contributions

ATS conceived the project, TA, AE, DD, ATS designed experiments with input from ACM,
TW, JH, and ST; TA (in vivo imaging, plume measurements), AE (plume measurements,
frequency and correlation behaviour), DD (plume measurements, whole-cell recordings),
ACM (source separation behaviour), TW (unit recordings), ST (analysis for OSN imaging),
JH (source separation behaviour) performed experiments and analysed data, IF
contributed tools and to experimental design, ST performed simulations, TA, AE, ATS
wrote the manuscript with input from all authors.

1301 Author Information

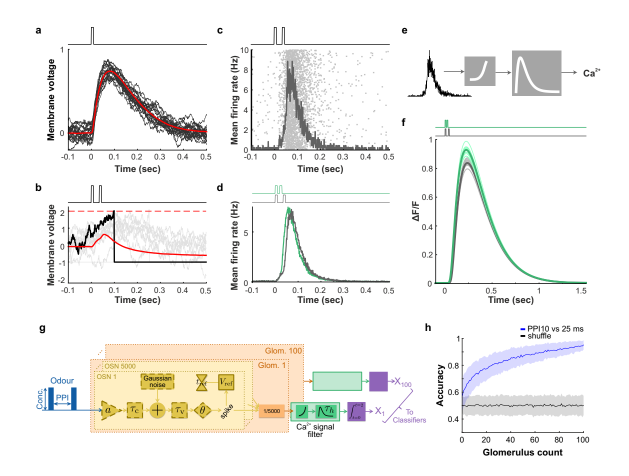
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www.nature.com/reprints. The authors declare no competing financial interests.
Readers are welcome to comment on the online version of the paper. Correspondence
and requests for materials should be addressed to ATS (andreas.schaefer@crick.ac.uk).

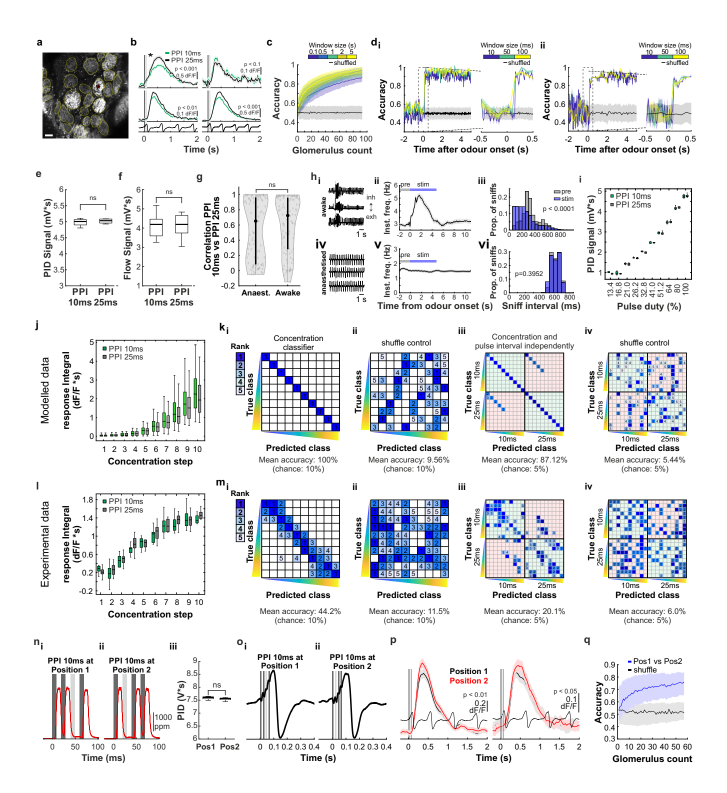
1306 Data availability

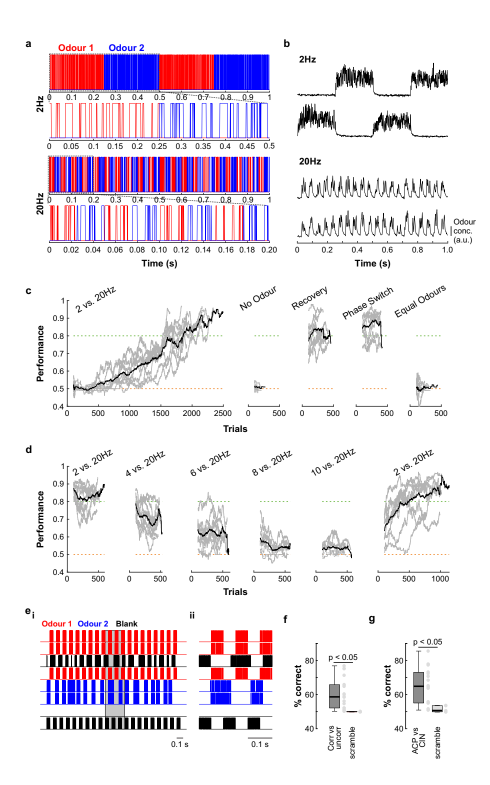
Data related to the OSN model (Extended Data Fig. 1) is available 1307 at 1308 https://github.com/stootoon/crick-osn-model-release. Data related to the glomerular 1309 classifier analysis (Extended Data Fig. **6**) is available at 1310 https://github.com/stootoon/crick-osn-decoding-release. The remaining data that 1311 support the findings of this study will be made available by the authors upon request.

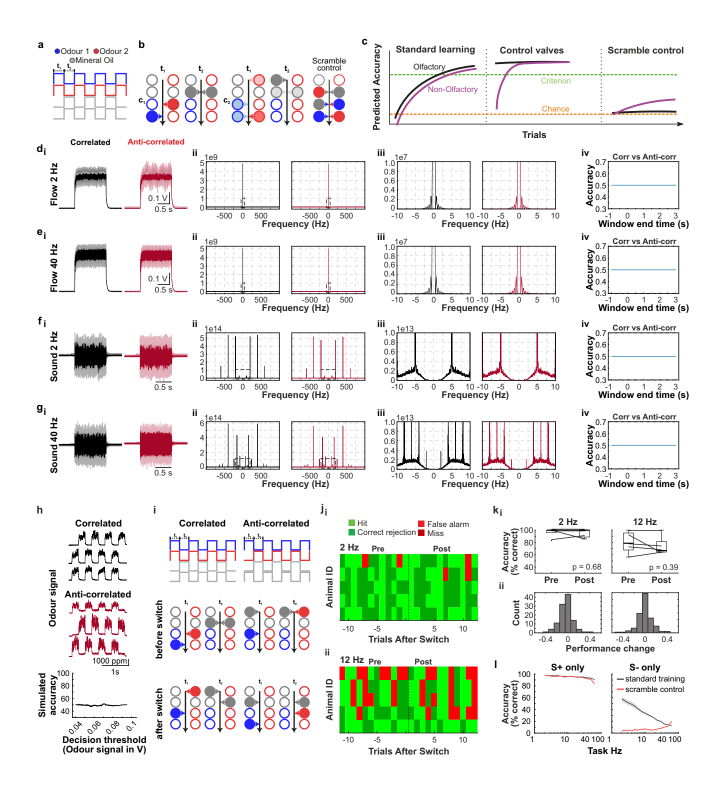
1312 *Code availability*

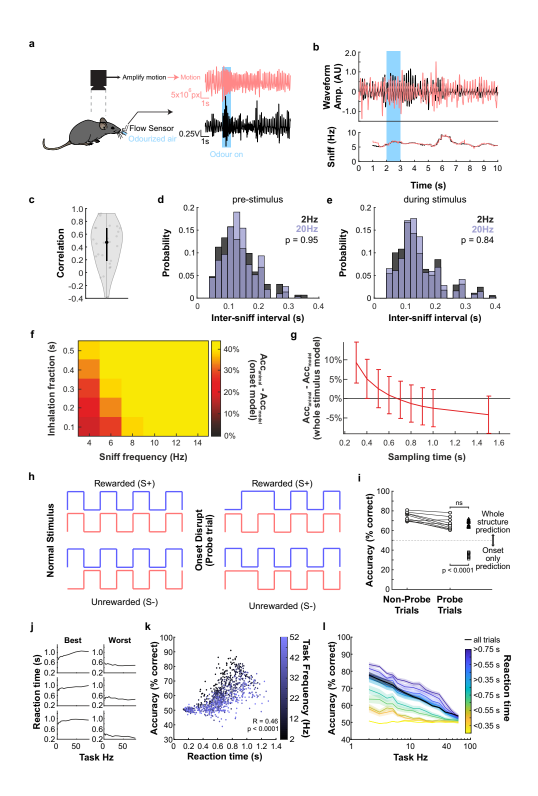
All custom Python scripts to generate pulses (PyPulse, PulseBoy) are available at 1313 http://github.com/warnerwarner. 1314 http://github.com/RoboDoig and Code for 1315 controlling AutonoMouse is available at https://figshare.com/articles/AutonoMouse Code/ 7616090. Code related to the OSN 1316 1317 model is available at https://github.com/stootoon/crick-osn-model-release. Code glomerular 1318 related the classifier analysis available to is at https://github.com/stootoon/crick-osn-decoding-release. 1319 1320

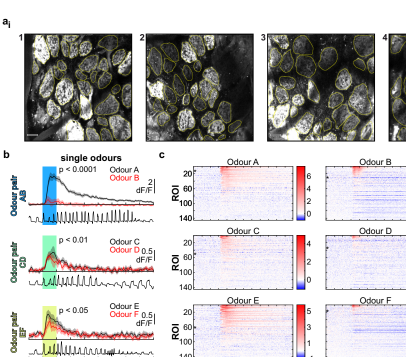










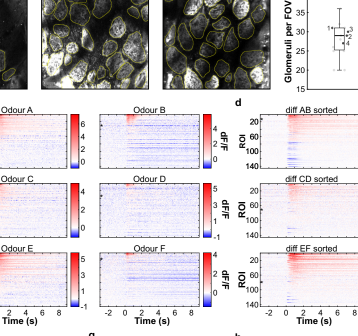


140

-2 0

1 2 3 4 5 6 7 8 Time (s)

-1 0



ii

40 [n = 15 animals

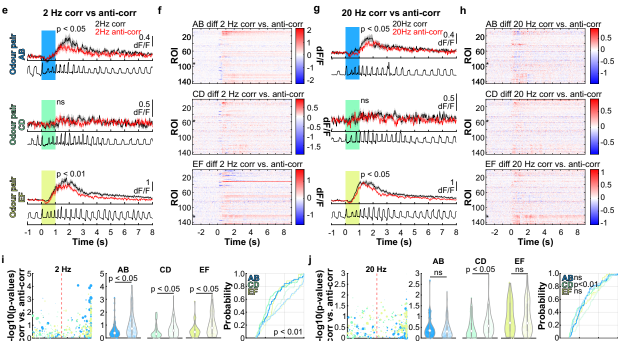
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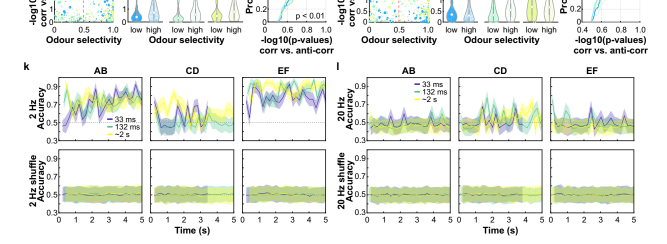
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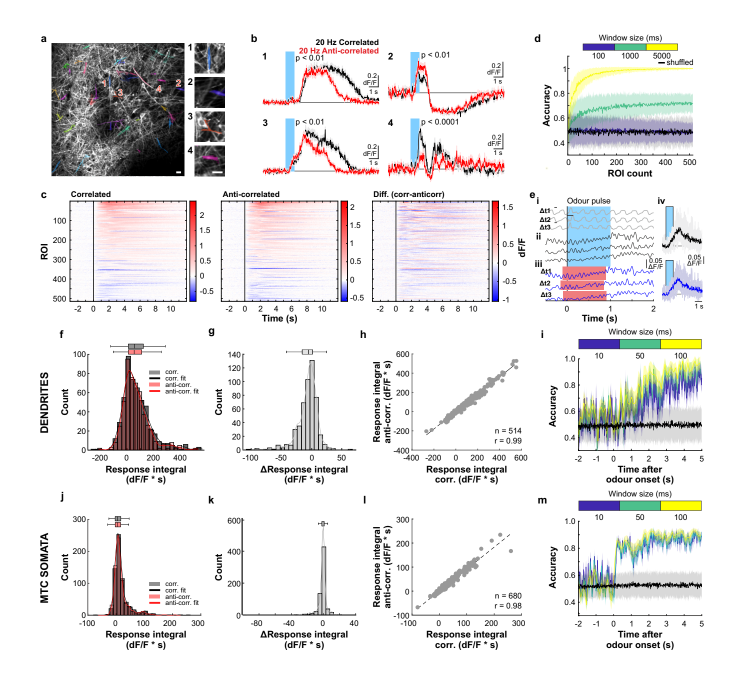
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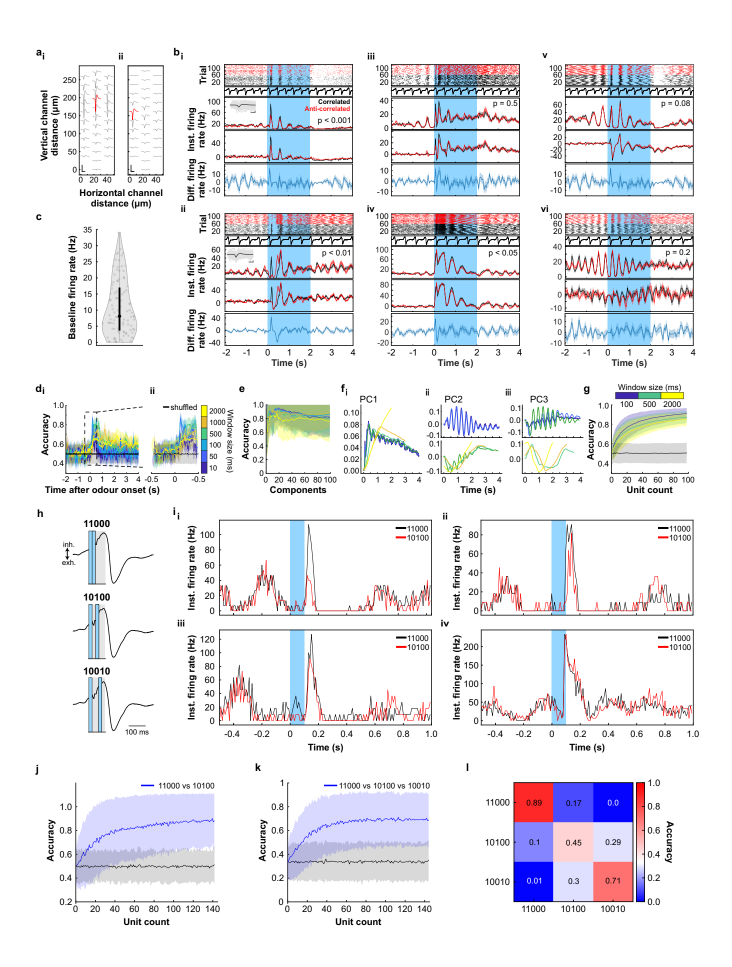
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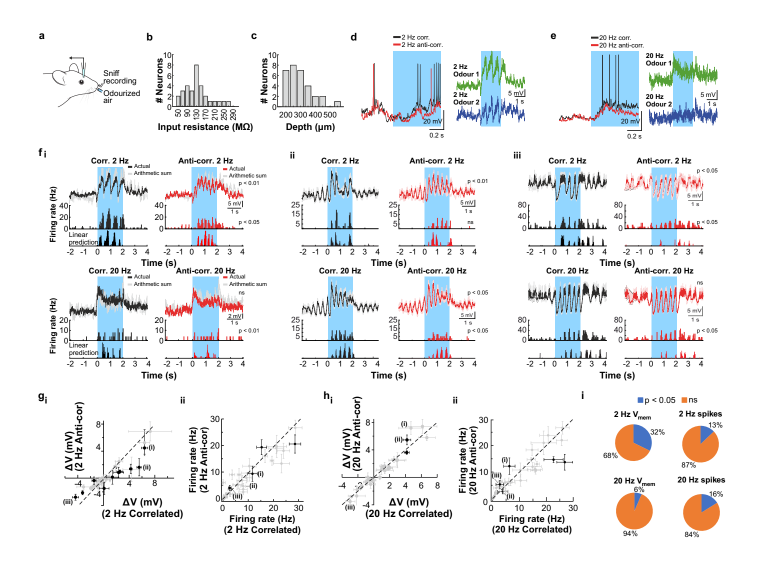
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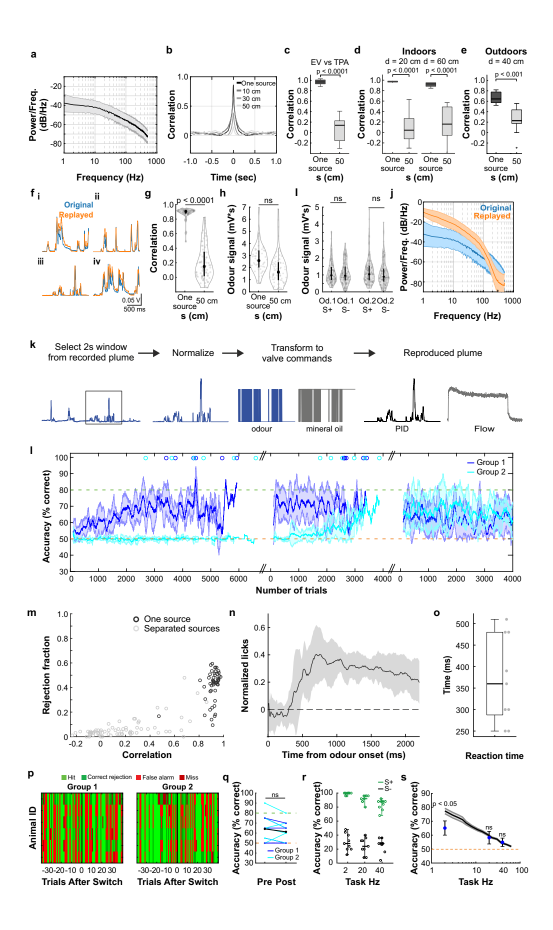












1321 Extended Data Fig. 1 | Distinguishing fast odour stimuli with slow OSNs. 1322 **a**, Membrane voltage relative to baseline of a single model OSN in response to a 10 ms odour pulse. Black traces are individual trials; red trace is average over 20 trials. OSN 1323 1324 spike threshold has been set high enough to prevent spiking to illustrate the subthreshold voltage time course. b, Membrane voltages (grey traces) of ten OSNs from a population of 1325 5000 in response to a paired odour pulse with pulse width 10 ms and PPI of 25 ms. The 1326 1327 voltage time course for one example OSN is in black. Several OSNs reach the OSN spike threshold (dashed red line) and are temporarily reset to the refractory voltage of -1. The 1328 population average membrane voltage (red) reveals membrane charging in response to 1329 1330 odour stimulation and the subsequent discharging and refractory period. c, Raster showing the spike times (dots) of the full population from **b** and the corresponding mean 1331 firing rate (trace) estimated in 1 ms bins. **d**, Mean firing rates computed over 20 trials in 1332 response to paired odour pulses of width 10 ms and PPIs of 10 ms (green) and 25 ms 1333 1334 (black). e, Model calcium signals are produced by squaring the instantaneous mean firing rate and filtering the result with a calcium imaging kernel. f, Model calcium responses to 1335 the paired odour stimulus with a PPI of 10 ms (green) and 25 ms (black). Thin traces are 1336 1337 single trials, thick traces are averages over 15 trials. g, Schematic of the OSN model. 1338 Variables in dashed bounding boxes are changed for each glomerulus (see Methods). h, 1339 Linear classifier analysis over an increasing subset size of glomeruli (1-100; plotted is mean ± SD, 256 repeats for random subsets of n glomeruli generating 256 unshuffled and 1340 256 shuffled accuracies). 1341

1342 Extended Data Fig. 2 | Sub-sniff odour information in the olfactory bulb input layer. a, GCaMP6f fluorescence recorded in olfactory bulb glomeruli in an anaesthetised OMP-1343 cre:Rosa-GCaMP6f mouse (maximum projection of 8200 frames, glomerulus marked with 1344 1345 red asterisk corresponds to first example trace shown in **b**). Scale bar: 50 µm. **b**, Example calcium traces in response to 10 and 25 ms PPI odour stimuli (mean of 50 trials ± SEM). 1346 Bottom: Example respiration traces. P-values derived from unpaired two-sided t-tests 1347 comparing responses of individual trials integrated over 2 s windows to paired odour 1348 pulse stimulation. c, Classifier accuracy over an increasing number of glomeruli when a 1349 linear classifier was trained on several response windows (colour-coded black: shuffle 1350 control) to PPI 10 vs. 25 ms stimuli (mean ± SD of up to 93 glomeruli from 4 individual 1351 1352 animals; 500 repetitions). d_i, Classifier accuracy when trained on all glomeruli in 1353 response to PPI 10 vs. 25 ms stimuli recorded in anaesthetised animals (n = 93 glomeruli, 1354 mean ± SD from 4 individual animals) with a sliding window of different durations (colour-coded; black: shuffle control; 100 repetitions) starting at 2 s before odour onset 1355 (left) and time period between -0.5 and 0.5 s from odour onset shown at higher 1356 1357 magnification (right). \mathbf{d}_{ii} , Same as \mathbf{d}_i for awake animals (n = 100 glomeruli, mean ± SD 1358 from 5 individual animals). **e**, Odour and **f**, flow signal integrated over 2 s for PPI 10 ms and PPI 25 ms stimuli (10 repeats each, odour: p = 0.1841, flow: p = 0.1786, unpaired 1359 1360 two-sided t-test). g, Correlation coefficients of glomerular calcium responses to PPI 10 vs. 25 ms in anaesthetised (n = 93 glomeruli from 4 individual animals) and awake (n = 1001361 1362 glomeruli from 5 individual animals) mice (p = 0.3187, unpaired two-sided t-test, measured as in Fig. 1 from OMP-Cre:Rosa-GCaMP6f mice). Violin plots show the median 1363

as a black dot and the first and third quartile by the bounds of the black bar. **h**_i, Example 1364 1365 respiration traces recorded using a flow sensor from awake mice. Inhalation goes in the 1366 upwards, exhalation in the downwards direction. \mathbf{h}_{ii} , Average instantaneous sniff 1367 frequency from one example animal plotted as a function of time (n = 24 trials, mean \pm SEM). The odour stimulus consisted of two 10 ms long odour pulses either 10 or 25 ms 1368 apart (see Fig. 1c). **h**_{iii}, Distribution of sniff intervals during a 2 s window before (grey) 1369 and a 5 s window after (blue) odour stimulus onset (p = 1.02e-189, two-sample 1370 1371 Kolmogorov-Smirnov test). h_{iv-vi} , Same but for the anaesthetised condition (p = 0.3952, two-sample Kolmogorov-Smirnov test). i, Mean odour signal for PPI 10 and 25 ms for 10 1372 increasing concentration steps defined by modulating valve pulse duty (see Methods and 1373 Supplementary Methods Fig. 1). There were no significant differences in odour 1374 concentration between both stimuli (unpaired two-sided t-tests). **j**, Modelled response 1375 1376 integrals to PPI 10 vs. 25 ms stimulations over a 10-fold concentration range pooled over all 20 trials and 100 glomeruli (see Methods). Box plots show median and extend from 1377 the 25th to 75th percentiles, whiskers extend to the 5th and 95th percentiles. \mathbf{k}_{i} , Confusion 1378 matrix of support vector machine (SVM)-based classification results of modelled 1379 glomerular signals in response to a range of 10 odour concentrations ranked and colour-1380 1381 coded (n = 100 glomeruli). \mathbf{k}_{ii} , Shuffle control with labels assigned randomly. \mathbf{k}_{iii} , Confusion matrix showing the ranked and colour-coded results of glomerular responses 1382 1383 independently classified for 10 ms vs. 25 ms PPI and across the range of 10 odour 1384 concentrations. \mathbf{k}_{iv} , Shuffle control for \mathbf{k}_{iii} with labels assigned randomly. **I**, Same as **j** but 2 s response integrals are derived from Ca^{2+} imaging data (10 repeats for each 1385 concentration). **m**, Same as **k** for Ca²⁺ imaging data (n = 57 glomeruli, from 2 individual 1386 animals, 10 repeats for each concentration). Note that 10 ms PPI could be reliably 1387 distinguished from 25 ms PPI with only few instances where a response to e.g. a 10 ms 1388 PPI stimulus was misclassified as 25 ms or vice versa (compare light red quadrants to 1389 light green quadrants). **n**, Shifting the position of 10 ms PPI within a single inhalation. **n**_i, 1390 PPI 10 ms at Position 1 or **n**_{ii}, at Position 2 of three 10 ms odour pulses. Odour pulses as 1391 1392 recorded with a PID shown in red, valve commands are shown in dark grey. Light grey 1393 area shows additional compensatory blank valve command to keep the flow profile indistinguishable between stimuli. niii, Total odour concentration was independent of the 1394 pulse profile (10 repeats, p = 0.57, unpaired two-sided t-test). **o**, Both the 10 ms PPI at 1395 Position 1 (o_i) and at Position 2 (o_{ii}) are presented during the inhalation phase 1396 (respiration shown in black, inhalation upwards, exhalation downwards). **p**, Example 1397 calcium traces in response to 10 ms PPI at Position 1 (black) and Position 2 (red), shown 1398 is the mean of 10 trials ± SEM. P-values derived from unpaired t-tests comparing 2 s 1399 1400 integrated responses of individual trials to odour pulses. q, Classifier accuracy over increasing number of glomeruli when a linear classifier was trained on the 2 s response 1401 1402 to PPI 10 ms at Position 1 vs. Position 2 (mean ± SD of up to 57 glomeruli, from 2 1403 individual animals, 500 repetitions; blue: PPI 10 ms at Position 1 vs. Position 2, black: 1404 shuffle control). Boxes in **e,f,n**_{iii} indicate 25th-75th percentiles, thick line is median, whiskers are most extreme data points not considered outliers (see Methods). 1405

Extended Data Fig. 3 | Frequency discrimination experiments. a, Frequency
discrimination stimuli are produced by alternating presentation of two odours to
generate a desired odour change frequency. During odour delivery, valves are not held

1409 open but rather randomly opened and closed over time to produce slight variation in 1410 odour amplitude for each pulse. This means that odour concentration cannot be used as a cue to learn the task and odour switching frequency is the primary stimulus signal. 1411 1412 Furthermore, valve clicking is randomised to minimize any acoustic cues. **b**, Replacing 1413 one odour channel with blank, un-odourised air and recording the frequency stimuli with a PID reveals that the desired odour pulse frequency is being produced. **c**, Mice readily 1414 1415 learn to discriminate 2 vs. 20 Hz pulse frequency stimuli in a go/no-go task. Replacing the odours with blank channels results in chance-level performance (No odour), which 1416 1417 recovers when odours are replaced (Recovery) showing that mice were likely discriminating the odour switching frequency rather than any extraneous cues such as 1418 1419 valve noise. The order of odour presentation in the stimuli had no effect on behaviour as 1420 when it was shifted (Phase switch) no decrease in performance was observed. 1421 Additionally, performance was dependent on the alternation between *different* odours as 1422 when the experiment was repeated with the same odours in each channel (Equal odours) performance was at chance level. **d**, To determine the perceptual limit of frequency 1423 1424 discrimination, the floor frequency used in the task over successive experiments was 1425 increased such that the difference in frequency between the stimuli progressively 1426 narrowed. Overall performance decreased as the difference in frequency grew smaller, 1427 reaching near-chance level with a frequency difference of 10 Hz (10 vs. 20 Hz). Switching back to the original discrimination (2 vs. 20 Hz) recovered performance quickly, showing 1428 1429 that the drop in discrimination ability was truly due to the frequency difference rather than general deterioration of performance over time. e, Example uncorrelated stimuli. 1430 1431 Combinations of odour 1 (red) and odour 2 (blue) valves are opened with temporal offsets and randomised pulse timing resulting in a correlation of 0 (see Methods). Blank 1432 (black) valves are used to keep total airflow constant throughout the stimulus. **e**_{ii}, Higher 1433 1434 magnification of the area in \mathbf{e}_{i} marked in grey. **f**, Animals show similar average accuracy 1435 as shown in Fig. 2k when probed to discriminate correlated from uncorrelated odour pulses at 10 Hz (n = 19 mice, mean \pm SEM of average accuracy = 0.6506 \pm 0.0016; after 1436 scrambling stimulus identity: 0.4997 ± 0.0032 ; p = 0.0175, unpaired two-sided t-test). g, 1437 1438 Animals show similar average accuracy when discriminating the correlation structure of 1439 a different odour pair (Acetophenone vs. Cineol) at 10 Hz (n = 19 mice, mean ± SEM of 1440 average accuracy = 0.6558 ± 0.0026 ; after scrambling stimulus identity: 0.5165 ± 0.0048 ; p = 0.0129, unpaired two-sided t-test). Grev dots mark average performance of individual 1441 1442 animals. Boxes in **f**,**g** indicate 25th – 75th percentiles, thick line is median, whiskers are most extreme data points not considered outliers (see Methods). 1443

Extended Data Fig. 4 | AutonoMouse stimulus and experimental design. a, Detailed
schematic of stimulus production; odour presentation (Odour 1: blue, Odour 2: red) is
always offset by clean air (Mineral Oil: grey) valves at the same flow levels, to ensure that
total flow during the stimulus is constant. b, Schematic of the use of valve subsets to
produce the desired stimulus. t1 and t2 represent valve openings at the corresponding
time points shown in a. c1 (b, left) and c2 (b, middle) represent two possible
configurations that could be used to produce the same resulting stimulus at the two time

1451 points. Opacity in the colours represents total concentration contribution to the resulting 1452 stimulus at the time point. For example, to produce the dual odour pulse at t1, configuration c1 can be used where odour 1 (blue) is delivered from one valve and odour 1453 1454 2 (red) from another valve. During t2 two valves contribute clean air. Alternatively, configuration c2 can be used in which during t1 odour 1 (blue) is generated by 50% 1455 opening of two valves, with odour 2 (red) produced by 70% / 30% opening of two other 1456 valves respectively. (b, right) Scramble control: valve maps (represented by arrow 1457 colour) are maintained compared to the training condition but odour vial positions are 1458 1459 scrambled resulting in odour stimuli uninformative about reward association whilst maintaining any non-odour cue such as putative sound or flow contributions. c, Predicted 1460 accuracy for animals in the case that they use solely olfactory temporal correlations 1461 (black) and in the case that they use extraneous non-olfactory cues or non-intended 1462 1463 olfactory cues (e.g. contaminations, clicking noises) (violet). Note that when switching stimulus preparations to a new set of valves (as in Fig. 2i and below in i-k), such non-1464 intended cues would not provide any information about stimulus-reward association, 1465 thus animals' accuracies would transiently drop back to chance. d_i, Average flow 1466 recordings (mean \pm SD) of 2 Hz correlated (black, n = 75) and anti-correlated (red, n = 1467 1468 70) trials taken from the AutonoMouse odour port. dii, Fourier transform of the flow plots from **d**_i, showing the power of the signal over a range of 1 kHz. **d**_{iii} A zoom in over the 1469 1470 range of 10 Hz indicated by the dotted box in **d**_{ii}. **d**_{iv}, Mean accuracy of a series of linear 1471 classifiers trained on an increasing window of the integrated signal starting from 1 s before trial shown in d_i. Classifiers were tested on two withheld trials, one correlated and 1472 one anti-correlated, and repeated 100 times. e, Same as d but for 40 Hz trials (n = 69 1473 correlated and n = 72 anti-correlated). f_{i} Average audio recording trace (mean ± SD) of 2 1474 Hz stimuli using a microphone placed in close proximity to the AutonoMouse odour port. 1475 1476 \mathbf{f}_{ii} , \mathbf{f}_{iii} , Fourier transforms of the audio signal from \mathbf{f}_i . Note, whilst there are notable peaks at specific frequencies, these are present in both correlated and anti-correlated trials. f_{iv} , 1477 Accuracy of a series of linear classifiers as shown in **d** but using the modulus of the audio 1478 1479 signal. g, Same as f but for 40 Hz trials. Note, whilst the sound profile and the Fourier 1480 transforms are different between 2 and 40 Hz, there is no difference detectable between correlated and anti-correlated trials. h, Example traces of odour signal (ethyl butyrate, 1481 isoamyl acetate, PID recorded) during correlated (top) and anti-correlated trials 1482 (middle). Simulated maximum accuracy based on differences in mean odour signal 1483 (bottom). Simulated accuracy was calculated as the fraction of trials correctly identified 1484 as correlated / anti-correlated based on a decision threshold set at some level between 1485 the minimum and maximum mean signal. Simulated accuracy was calculated for multiple 1486 1487 decision thresholds, increasing the decision threshold from minimum odour signal to maximum odour signal in steps of 1/5000th of the range between minimum and 1488 1489 maximum. **i**, Detailed schematic of correlated (top left) and anti-correlated (top right) 1490 stimulus production before (middle) and after (bottom) switching valves. For the switch 1491 control, a set of previously unused odour valves is introduced to rule out potential bias towards a specific valve combination when performing the odour correlation 1492 1493 discrimination task. **j**, Trial map of 5 representative animals during 2 Hz (**j**_i) and 12 Hz (**j**_{ii}) correlation discrimination tasks before and after introduction of control valves (n = 1494 1495 12 trials pre-, n = 12 trials post-new valve introduction, new valve introduction indicated 1496 by black vertical dotted line. Each row corresponds to an animal, each column within the

1497 row represents a trial. Light green: hit, dark green: correct rejection, light red: false alarm, 1498 dark red: miss. \mathbf{k}_{i} , Boxplots of mean accuracy for animals (n = 5 mice) pre- and postcontrol for 2 Hz (left) and 12 Hz (right). Box indicates 25th – 75th percentiles, thick line is 1499 1500 median, whiskers are most extreme data points not considered outliers, see Methods. Pvalues derived from unpaired t-tests. **k**_{ii}, Summary histograms of performance change for 1501 all animals during all "valve switch" control tests (see Methods) indicating that 1502 discrimination accuracy was based on intended olfactory cues. The five animals showing 1503 highest performance before the valve switch/bottle change (and thus the largest 1504 potential to drop in performance) were analysed. **I**, Discrimination accuracy (n = 33) 1505 animals, mean ± SEM) for rewarded S+ (left) and unrewarded S- (right) trials when 1506 odours were presented using standard training valve configurations (black) and 1507 scrambled valve identity (red), data from Fig. 2k. Note that frequencies >40 Hz were 1508 1509 presented predominantly in the last block of the training schedule and reduced licking in the control group (decreased S+ performance and increased S- performance) might be 1510 1511 due to decreased motivation at that point.

Extended Data Fig. 5 | Respiration recordings, stimulus onset model and reaction 1512 1513 time for correlation discrimination experiments. a, An overhead camera was used to image a head-fixed mouse during a sequence of odour presentations. Simultaneously, a 1514 1515 flow sensor was placed close to one nostril to monitor respiration to establish the validity of motion imaging-based respiration recording. Phase-based motion amplification was 1516 used to magnify motion on the animal's flank to capture body movements associated with 1517 1518 respiration. Right: example for simultaneous respiration measurement with motion imaging (red) and flow sensor (black; see Methods and Supplementary Video 2). **b**, Three 1519 1520 further example trials with respiration rate extracted from motion imaging (red) and simultaneous flow sensor recording (black). Below: instantaneous sniff frequencies 1521 1522 calculated from either sensor were tightly correlated. **c**, Correlation between respiration 1523 traces extracted from motion imaging and respiration captured by flow sensor (n = 26trials, 10 s duration each). Violin plot shows the median as a black dot and the first and 1524 third quartile by the bounds of the black bar. d, Probability distributions of inter-sniff 1525 1526 intervals for odour presentations (isoamyl acetate vs. ethyl butyrate, 2 Hz and 20 Hz) for freely moving animals in AutonoMouse before stimulus onset and **e**, during 2 s odour 1527 1528 stimulation (n = 605 sniffs for 2 Hz and n = 668 for 20 Hz, two-sample Kolmogorov-1529 Smirnov test). **f**, Heat map of accuracy difference between a model where animals rely on 1530 onset information only (see Methods) and actual animal accuracies across a range of sniff frequencies and inhalation fractions (n = 10 mice). No matter what assumed sniff 1531 1532 frequency and inhalation frequency, the "onset model" deviates substantially from the accuracy measured in the behavioural experiments (panels h,i). g, Difference between a 1533 1534 model where animals use the entire stimulus structure (see Methods) and actual behavioural accuracies across different stimulus sampling times (n = 10 repeats, mean ± 1535 SD). The "whole stimulus" model accurately describes animal behaviour indicating that 1536 1537 mice base a decision about the correlation structure of a stimulus not predominantly on 1538 the onset. Note the different scales in **f** and **g**. **h**, Schematic of experimental stimulus in which the first stimulus pulse was disrupted when presented on "probe trials". Top: 1539

1540 normal stimulus design, bottom: "onset disrupt" stimuli in which the first pulse in a correlated stimulus is disrupted to be anti-correlated; and vice versa for an anti-1541 correlated stimulus. i, Animals were trained on standard (non-probe) correlation 1542 1543 discrimination stimuli (f = 10 Hz) but onset disrupt (probe) stimuli were presented 1544 randomly on probe trials with a 1/10 probability. Accuracy was only slightly degraded on probe trials (mean ± SD of accuracy for non-probe trials 75.8 ± 4.4%; for probe trials 1545 1546 $67.8 \pm 6.1\%$; p = 0.001, paired two-sided t-test, n = 9 mice) but did not drop below chance (p = 7.3e-06, paired t-test). Importantly, accuracy on probe trials was consistent with 1547 whole-structure prediction (70.3 \pm 3.5%, p = 0.13, paired t-test of comparison to probe 1548 trials) and differed significantly from the accuracy of onset-only prediction $(41.6 \pm 1.5\%)$; 1549 p = 1.02e-6, paired t-test of comparison to probe trials). **j**, Mean reaction time (time from 1550 stimulus onset to first lick in S+ trials) plotted as a function of stimulus pulse frequency 1551 for the three animals with the best (left) and the worst (right) global accuracy (mean 1552 1553 accuracy across all trials). Better performing animals tend to increase their reaction time as stimulus pulse frequency increases. **k**, Scatter plot of mean accuracy vs. mean reaction 1554 time for each animal and stimulus pulse frequency condition (averaged over blocks of 1555 1556 100 trials). Points are colour-coded according to stimulus pulse frequency. Accuracy was 1557 significantly positively correlated to reaction time, suggesting that mice that sampled a greater portion of the stimulus made more accurate decisions about its correlation 1558 structure (Pearson correlation coefficient R = 0.49, p<1.1e-112). I, Accuracy (mean \pm 1559 SEM) is plotted as in Fig. 2k, but only trial blocks with reaction times above or below a 1560 certain threshold (colour code) are included in the analysis. Where only longer reaction 1561 1562 times are considered, global performance is higher than the case where only shorter reaction times are included, again suggesting that longer stimulus sampling improves 1563 1564 discrimination of odour correlation structure across all stimulus pulse frequencies.

Extended Data Fig. 6 | OSN imaging in response to correlated vs. anti-correlated 1565 odour stimulation. a, Four example fields of view (FOV) recorded from the dorsal 1566 olfactory bulb of individual mice. **a**_{ii}, Number of individual glomeruli per FOV in all 1567 experimental mice (n = 15). The number of individually delineated glomeruli ranges from 1568 20-36 with an average of 28 glomeruli per FOV. Labelled data points (1-4) correspond to 1569 FOVs shown in **a**_i. Scale bars: 50 µm. Edges of the box are the 25th and 75th percentiles, 1570 the whiskers extend to the most extreme data points not considered as outliers, see 1571 Methods. **b**, Example glomerulus response from OMP-Cre:Rosa-GCaMP6f mice to 1572 1573 presentation of individual odours plotted pairwise (AB, CD, EF; mean of 6 trials ± SEM). Stimulation period (1 s) is indicated by vertical bar (blue, green and yellow). Bottom: 1574 Typical example respiration trace. P-values derived from unpaired two-sided t-tests 1575 1576 comparing 2 s integrated responses between paired odours. c, Averaged calcium transients from all glomeruli (n = 145 from 5 individual animals) in response to 1577 1578 individual odours, plotted as colour maps sorted by response magnitude. **d**, Difference between glomerulus responses to individual odours plotted pairwise as colour maps. 1579 1580 Glomeruli are sorted by average magnitude of response difference. e, Example 1581 glomerulus response to presentation of correlated vs. anti-correlated odour pairs 1582 fluctuating at 2 Hz (mean of 12 trials ± SEM). Bottom: typical example respiration trace.

P-values derived from unpaired two-sided t-tests comparing 2 s integrated responses of 1583 individual trials to correlated and anti-correlated odour stimulation. f, Difference 1584 between glomerulus responses to 2 Hz correlated and anti-correlated odours as colour 1585 1586 maps sorted as shown in **d**. **g-h**, Same as in **e-f** but for 20 Hz correlated vs. anti-correlated. 1587 Example glomerulus from **b**,**e**,**g** indicated with an asterisk in colour maps in **c**,**d**,**f**,**h**. **i**, Left: P-values derived from comparing trials of the summed 2 s response to correlated vs. anti-1588 correlated odour stimulation at 2 Hz (unpaired two-sided t-tests) for three odour pairs 1589 1590 (colour-coded) as a function of glomerulus selectivity to individual odours (n = 145glomeruli). Selectivity is calculated as the difference between the absolute response to 1591 single odours scaled by the summed absolute response. A threshold is set at 0.5 defining 1592 1593 glomeruli as low or high selective. Dot size represents magnitude of the summed response. Middle: Comparison of p-values between low and high selective glomeruli (p < 1594 1595 0.05, unpaired two-sided t-test). Violin plots show the median as a white dot and the first and third quartile by the bounds of the grey bar. Right: Cumulative distribution function 1596 of p-values for low and high selective glomeruli (p < 0.01 for all pairwise comparisons, 1597 1598 two-sample Kolmogorov-Smirnov test). j, Same as i but for 20 Hz, (n = 145 glomeruli). k, Top row: Mean ± SD over 100 repetitions of classifier accuracy when trained on all 1599 1600 responsive glomeruli (n = 145 available, from 5 individual animals, see Methods) to discriminate 2 Hz correlated vs. anti-correlated stimuli, trained separately for each of the 1601 1602 three odour pairs and within sliding windows of different widths (colours); x-coordinates 1603 indicate latest extent of each window. Bottom row: same as top row but with labels 1604 shuffled as control. I, Same as k for 20 Hz correlated vs. anti-correlated odours. Some data 1605 points in **k**, **l** are absent because not all time points had responsive ROIs for every window size (see Methods). 1606

Extended Data Fig. 7 | Odour correlation structure is encoded in dendrites of 1607 olfactory bulb output neurons. a, GCaMP6f fluorescence from mitral and tufted cells 1608 and their dendrites recorded in the dorsal portion of the olfactory bulb of a Tbet-1609 1610 cre:Rosa-GCaMP6f mouse (maximum projection of 8000 frames). Dendritic ROIs are superimposed in colour. Four dendritic segments (1-4) are shown in higher 1611 magnification, scale bars: 20 µm. **b**, Four example calcium traces extracted from dendritic 1612 1613 segments shown in a that show differential response kinetics to correlated (black) and 1614 anti-correlated (red) stimulation (mean of 24 trials ± SEM, f = 20 Hz). In total, 24% of dendritic segments showed significantly different integral responses (0-5 s after odour 1615 1616 onset, p < 0.01, unpaired two-sided t-test; 121/514) to the two stimuli. **c**, Average calcium 1617 transients as colour maps for correlated (left) anti-correlated (middle) and the difference 1618 between both odour stimulations (right) of all analysed dendritic segments (n = 514, from 6 individual animals). d, Classifier accuracy over an increasing number of dendritic ROIs 1619 trained on several response windows (colour-coded) to discriminate correlated vs. anti-1620 1621 correlated stimuli at 20 Hz (n = up to 514, mean ± SD from 6 individual animals, black: shuffle control). e, Method of aligning calcium traces to first inhalation after odour 1622 stimulus onset. e_i, Representative respiration traces recorded using a flow sensor placed 1623 1624 in front of the nostril contralateral to the imaging window. The first inhalation peaks were 1625 detected and the time (Δt) to the first inhalation after odour onset was calculated for each 1626 trial individually. eii, Representative calcium transients in response to a single odour 1627 presentation (here: 20 Hz correlated). \mathbf{e}_{iii} , Transients are shifted according to Δt . \mathbf{e}_{iv} ,

1628 Individual calcium transients (faint colours, 24 trials) in response to 20 Hz correlated 1629 odour presentations with the average calcium signal (thick traces) superimposed. Top: before aligning to first inhalation after odour onset, bottom: after alignment. Blue bar 1630 1631 represents the odour presentation phase (approximate for the aligned data). f, Distribution of odour response integrals from all recorded ROIs (n = 514) for correlated 1632 (grey) and anti-correlated (red) stimulation. Box indicates 25th – 75th percentiles, thick 1633 line is median, whiskers are most extreme data points not considered outliers, see 1634 Methods. g, Histogram of the difference between correlated and anti-correlated odour 1635 responses. Box plots as in f. h, Comparison of correlated and anti-correlated odour 1636 responses of all dendritic ROIs (f = 20 Hz, n = 514 dendrites). **i**, Classifier accuracy when 1637 trained on all dendritic ROIs recorded with a sliding window of different durations 1638 1639 starting 2 seconds before odour onset (colour-coded, black: shuffle control, n = 514 from 6 individual animals; mean ± SD, 100 repetitions). **j-m**, same as **f-i** for projection neuron 1640 somata (f = 20 Hz, n = 680 cells; see Fig. 3). 1641

1642 Extended Data Fig. 8 | Projection neurons unit recordings in response to correlated 1643 vs. anti-correlated stimulation and short odour pulse combinations. 1644 a, Data from unit recordings as described in Fig. 3h-k. Average waveforms across all 1645 channels of two isolated units shown in $\mathbf{b}_{i,ii}$. Each waveform represents the average waveform for the unit on a specific channel. Red waveform indicates the channel with the 1646 1647 largest average waveform for the unit. Scale bar in the bottom left represents 100 μ V (vertically) and 1 ms (horizontally). **b**, Additional example single unit odour responses to 1648 correlated (black) and anti-correlated (red) stimuli shown as raster plot (top) and PSTH 1649 (mean of 64 trials for each condition ± SEM) of spike times before (second from top) and 1650 1651 after baseline subtraction (second to bottom), and the differential PSTH of correlated and anti-correlated (bottom, blue). Average spike waveform shown as insets in **b**_{i,ii}. Duration 1652 of odour presentation (2 s) is indicated in light blue. P-values are derived from a two-1653 1654 sided Mann-Whitney U test comparing the spike time distributions of correlated and anti-1655 correlated trials during 4 s after odour onset. **c**, Average baseline firing rate for all units (n = 97 from 6 individual animals). Baseline firing rates were calculated from 4 s to 0 s1656 before odour onset for each of the 1312 trials presented during all recordings. Violin plot 1657 1658 shows the median as a black dot and the first and third quartile by the bounds of the black bar. **d**_i, Classifier accuracy when trained on all baseline-subtracted units in response to 1659 20 Hz correlated vs. anti-corelated stimulation (n = 97 units, mean ± SD from 6 individual 1660 animals) with a sliding window of different durations (colour-coded; black: shuffle 1661 1662 control; 100 repetitions) starting at 2 s before odour onset. Time along the x-axis represents the end time of the window. d_{ii} , Time period between -0.5 and 0.5 s from 1663 1664 odour onset shown at higher magnification (n = 97 units, mean \pm SD from 6 individual animals). e, To take the entire temporal structure of responses into account we 1665 performed a principal component analysis (PCA) on the temporal evolution of the firing 1666 rate responses (see Methods). Shown here is the accuracy for linear SVM classifiers 1667 (mean ± SD) trained on increasing numbers of principal components (PCs). Classifiers 1668 were trained on all but two trials (one correlated, one anti-correlated). Training and 1669 testing were repeated 1000 times. The colour code represents the same window sizes as 1670 1671 defined in **d**. **f**, The first (**f**_i), second (**f**_{ii}), and third (**f**_{iii}) PCs found from PCA for different 1672 rolling window sizes (colour code as defined in **d**). In the second and third PCs, the

1673 windows have been split as to better compare the similarities in PCs for different window 1674 sizes. **g**, Average classifier accuracy of a set of classifiers trained on the PC weights of increasing number of units. Classifiers were trained on all but two trials (one correlated, 1675 1676 one anti-correlated). The number of PCs used for each window was selected by the peak accuracies in **e** (colour-coded; n = up to 97 units from 6 individual animals; mean ± SD of 1677 1000 classifier repetitions). h, Schematic of odour pulse stimuli timings in relation to the 1678 respiration cycle. Three combinations were presented, each trial 120 ms in length. For 1679 example, 11000 (top) consisted of a 40 ms odour pulse (light blue) followed by 80 ms of 1680 blank odourless air (grey); All trials were triggered at the onset of inhalation. i, PSTH from 1681 four example units (i_{i-iv}) showing their average firing rate prior, during, and after odour 1682 presentation (light blue vertical bar). Responses are either to 11000 trial (black) or 1683 10100 odour presentation (red). The instantaneous firing rate was calculated by 1684 summing the number of detected spikes in 10 ms windows and multiplying the value by 1685 100 to get Hz. j, Accuracy of linear classifiers as a function of the number of units available 1686 for training/testing (mean \pm SD of n = up to 145 units from 8 individual anaesthetised 1687 animals). Each classifier is trained on the summed spike count of the available units in a 1688 window of 500 ms starting at odour onset. The classifiers were trained on all but two 1689 trials, one 11000 and one 10100 trial and the number of repeats between animals varied 1690 1691 between 11 and 30. To account for this and to minimise the variability of the training set, 1692 trial number was bootstrapped to 1000 repeats. This was achieved by randomly selecting 1693 a repetition for each unit independently. The test set was isolated from the responses prior to bootstrapping and thus was not seen by the classifier until it was tested on it. 1694 Each classification was repeated 500 times with a different selection of units, and a 1695 different test set. The shuffled control (black) was accomplished by shuffling the training 1696 labels during each iteration of the classifier without shuffling test labels. **k**, Same as in **j** 1697 1698 but classifying all three odour pulse combinations shown in **h**. **l**, Confusion matrix showing the fractions that each trial type was classified as (n = 145 units from 8)1699 1700 individual animals). True labels are shown on the x-axis and labels predicted by the 1701 classifier on the y-axis. Accuracies correspond to maximum unit count shown in **c** and **d**. The classifiers can readily separate between trials containing a single 40 ms odour pulse. 1702 Accuracy is lower when distinguishing between an intermission of 20 or 40 ms but 1703 1704 remains above chance (chance = 0.33).

1705 Extended Data Fig. 9 | Whole cell recordings of projection neurons in response to correlated vs. anti-correlated odour stimulation. a, Schematic of the whole-cell patch 1706 1707 clamp recording approach. **b**, Distribution of input resistance and **c**, recording depth as measured from all recorded projection neurons (n = 31). **d**, Left: Example recording from 1708 1709 single cells with consecutive presentations of correlated (black) and anti-correlated (red) 1710 odour stimulus at 2 Hz. Duration of odour presentation (2 s) is indicated in light blue. 1711 Right: Baseline-subtracted and spike-clipped subthreshold voltage response from a single cell to odour 1 (green) and odour 2 (blue) for 2 Hz. e, the same as d but for 20 Hz 1712 1713 odour stimulation. **f**, Voltage response from three example cells for correlated (black) and anti-correlated (red) odour stimulus for 2 Hz (top) and 20 Hz (bottom). The cell shown 1714 1715 in \mathbf{f}_{i} corresponds to the cell shown in \mathbf{d} and \mathbf{e} . The grey overlaid traces correspond to the 1716 arithmetic sum estimated from the response to individual odours. Bottom: Linear 1717 prediction histogram calculated by thresholding the arithmetic sum of the subthreshold 1718 responses to the individual odours. Differences here suggest that correlation can be 1719 calculated based on a single cell level if the two individual odours engage overlapping 1720 OSN populations. P-values are derived from a paired two-sided t-test of the membrane 1721 potential and the firing rate in the first 500 ms after odour onset. **g**, Average change in voltage (g_i) and in instantaneous spike frequency (g_{ii}) in the first 500 ms after odour 1722 onset from baseline membrane potential for 2 Hz correlated vs. anti-correlated odour 1723 presentation and **h**, for 20 Hz. Each marker corresponds to a single cell, error bars 1724 1725 represent SEM. Data points in black represent cells where p < 0.05 between correlated 1726 and anti-correlated conditions. P-values are derived from a paired t-test of the membrane potential and the firing rate in the first 500 ms after odour onset. Indicators (i), (ii) & (iii) 1727 1728 represent cells shown in **f**. **i**, Pie charts depicting the proportion of cells showing significant difference as described above (blue) in subthreshold membrane potential 1729 1730 (left) and spike frequency (right) for all 2Hz (top) and 20Hz (bottom) cells. P-values are derived from a paired t-test of the membrane potential and the firing rate in the first 500 1731 1732 ms after odour onset.

1733 Extended Data Fig. 10 | Odour plume generation and additional analysis of source separation experiments. a, Power spectrum of all recorded odour plumes (mean ± SD 1734 of log power, n = 132 plumes). **b**, Cross correlation of all recordings at different lateral 1735 1736 separation distances. c, Correlation coefficients over all recordings for odours from the same source and for odour sources separated by 50 cm in a controlled laboratory 1737 environment with complex airflow (indoors; ethyl valerate (EV) vs. tripropylamine 1738 (TPA); n = 25 for same source, n = 27 for sources separated by 50 cm; p < 0.0001, unpaired 1739 two-sided t-test). Box indicates 25th – 75th percentiles, thick line is median, whiskers are 1740 most extreme data points not considered outliers; see Methods. d, Same as Figure 4b (for 1741 1742 odours α -Terpinene and ethyl butyrate) but for radial distances to the PID of 20 cm and 1743 60 cm (p < 0.0001, unpaired two-sided t-test). **e**, Same as **d** but measured outdoors (n = 7 for same source, 10 for sources separated by 50 cm; p < 0.001, unpaired t-test; Indoors 1744 versus outdoors, one source: p = 0.0060, s = 50 cm: p = 0.0632, unpaired two-sided t-1745 test). f, Example plume structures originating from the same one source or separated 1746 sources as recorded with a PID (blue) and replayed with the multi-channel high 1747 bandwidth odour delivery device (orange). g, Correlation coefficients over all recordings 1748 1749 of replayed plumes for one source (n = 53 plumes) and for sources separated by 50 cm 1750 from each other (n = 74 plumes; p = 2.27e-41, unpaired two-sided t-test). **h**, Odour signals 1751 integrated over 2 s for all recordings of replayed plumes for one source (n = 53 plumes) 1752 and for sources separated by 50 cm (n = 74 plumes; p = 0.75, unpaired two-sided t-test). 1753 i, Odour plume signals integrated over 2 s for rewarded and unrewarded trials (n = 150 trials each; Odour 1: p = 0.4739, Odour 2: p = 0.0923, unpaired two-sided t-test). j, 1754 1755 Overlaid power spectra (mean ± SD of log power) of all plumes (n = 127 plumes) recorded in complex, natural airflow conditions (blue) and replayed plumes (orange). k, Schematic 1756 of plume reproduction: First, a 2s long window is selected from the PID recording, 1757 starting around the middle of the trace and such that odour is present during the first 500 1758 1759 ms. Secondly, the trace is normalised between 0 and 1. Thirdly, the trace is converted into 1760 a series of binary opening and closing commands directly related to the value of the

1761 normalised signal. A value of 1 translates to a continuous opening, and a value of 0 1762 translates to continuously closed. This series of commands is relayed to an odour valve 1763 and an inverted version of the commands is relayed to a mineral oil valve to generate a 1764 compensatory airflow. The resulting output resembles the original plume, as measured with a PID, and there is constant airflow throughout the trial, as measured with a flow 1765 1766 meter. The same procedure is then applied to the accompanying odour, to create both 1767 plumes needed for each trial. I, Group learning curves (mean ± SD) for the two groups of 1768 animals trained on the virtual source separation task, but on different set of valves. Group 1769 1 (n = 6 mice, blue) were trained on the task from the start, while Group 2 (n = 6 mice, 1770 cyan) were first exposed to a scrambled version of the task and were later transferred to 1771 the same plumes as Group 1. This served as a control that the cue required for learning is indeed olfactory information contained in the odour plumes. For the 3rd stage of learning, 1772 1773 the plumes were refined to ensure odour was always present in the first 500 ms of the 1774 trial and performance stabilised for the two groups. Mice progressed through these learning stages as a group, based on time elapsed from the beginning of training. 1775 Therefore, some mice performed more trials than others. The last trial performed by a 1776 1777 mouse in each phase is represented by a colour-coded circle above the plot. Accuracy is 1778 calculated over a 100-trial sliding window. **m**, Rejection fraction (fraction of trials the mouse abstained from licking) calculated for each plume pair plotted in relation to the 1779 correlation between the two odour traces in that plume pair. Animals are trained to lick 1780 1781 (expected low rejection fraction) for source separated trials (low correlation) and abstain from licking (high rejection fraction) for one source trials (high correlation). **n**, Difference 1782 1783 in lick rates in response to source separation training trials (n = 9 mice, mean \pm SD), calculated for each mouse as lick rate (licks / 100 ms) in response to S+ trials minus the 1784 lick rate in response to S- trials, normalized to averaged lick rate for all trials across the 1785 1786 corresponding time period. **o**, Reaction times for each mouse, calculated as the time point 1787 when the difference in lick rate for each mouse crossed a threshold (mean + 3 SDs over the baseline, defined as the first 200 ms of the trace, when odour was not present). Box 1788 indicates 25th – 75th percentiles, thick line is median, see Methods. **p**, Trial map of all 1789 1790 animals during virtual source separation tasks before and after introduction of control 1791 valves similar to Extended Data Fig. 4 (n = 40 trials pre-, n = 40 trials post-new valve 1792 introduction, new valve introduction indicated by black vertical line). Each row corresponds to an animal, each column within the row represents a trial. Light green: hit, 1793 1794 dark green: correct rejection, light red: false alarm, dark red: miss. **q**, Mean performance of animals (n = 11 mice) that reached performance criterion during training during pre-1795 and post-control. r, Discrimination accuracy split by stimulus valence (green, S+; black, 1796 S-) for odour correlation fluctuation frequencies 2, 20 and 40 Hz (Fig. 4e; n = 9 mice, data 1797 is mean ± SD, unpaired two-sided t-test). **s**, Group performance for the square pulse probe 1798 1799 trials at different frequencies, in animals trained on the source separation task (blue dots, 1800 n = 9 mice, data is mean \pm SD), compared to group performance where animals were 1801 trained on correlated and anti-correlated square pulse trains (from Fig. 2k, black line and 1802 SEM band, n = 33 mice; 2 Hz: p = 0.0018, 20 Hz: p = 0.19, 40Hz: p = 0.94, unpaired twosided t-test). Violin plots in g-i show the median as a black dot and the first and third
quartile by the bounds of the black bar.

1 Supplementary Information Guide

Supplementary Video 1: Automated operant conditioning system ("AutonoMouse")
equipped with high speed odour delivery device.

Supplementary Video 2: Comparison between original video-based respiration
recording and phase-based motion amplification (red trace) in head-fixed condition on
the animal's flank to capture body movements associated with respiration.
Simultaneously, respiration was recorded with a flow sensor placed in front of one nostril
(black trace). Odour stimulus highlighted with blue bar.

Supplementary Methods Fig. 1 | Characterization of odorants presented with a 9 high-speed odour delivery device. a, Calculated signal fidelities for seven different 10 odours (colours, see legend in **b**) pulsed for 2 s over a frequency range of 2 to 100 Hz at 11 50% pulse duty (n = 5 repeats for each condition, mean ± SEM). **b**, Amount of released 12 odour (n = 5 repeats for each condition, mean \pm SEM). Odours are: AA (isoamyl acetate), 13 ACP (acetophenone), AT (α -Terpinene), CN (cineol), EB (ethyl butyrate), Hex (2-14 hexanone), PEA (phenylethyl alcohol). c, Left: Schematic of the pulse-width modulation 15 (PWM) method. For any period of odour release, maximum final concentration is 16 achieved by keeping the valve open for the entire time (top). The amount of odour 17 released can be reduced by cycling the valve at a high frequency (here 500 Hz) with a 18 different level of PWM (middle and bottom panel). Right: Odours were released over a 2 19 s period with different PWM duties at 500 Hz (n = 5 repeats for each condition, mean ± 20 SEM). The resulting amount of released odour is normalised to the maximum release 21 (PWM = 1). **d**, Average PID signal of single 100 ms pulses (pulse indicated in blue) for 22 seven different odours (n = 60 pulses for each odour, mean ± SEM). **e**, Summary table: 23 delay (time from start of the odour pulse to 5% of maximum signal amplitude), rise (time 24 from 5% to 95% of maximum signal amplitude), decay (time for the signal to decay back 25 to 5% of maximum amplitude after the end of the odour pulse). **f**, Effect of tubing length 26 attached to the valve manifold on signal fidelity at different pulse frequencies pulsed for 27 2 s at 50% pulse duty (ethyl butyrate). 28

Supplementary Methods Fig. 2 | Dual-energy fast photoionisation detection (defPID).

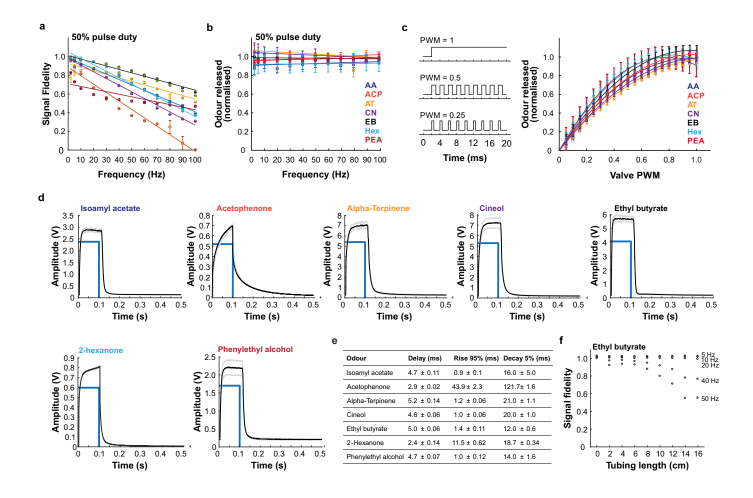
a, Schematic of the dual-energy fast photoionisation detection method. Two odours are 31 recorded simultaneously by two PIDs with different ionizing energies (different 32 wavelength UV light sources). The odours are chosen such that one odour (odour 2 (ethyl 33 butyrate), 9.5 eV) has an ionization energy greater than the low energy PID bulb, but less 34 than the high energy PID bulb, thus only being detectable by the high energy PID. The 35 other odour (odour 1 (α -Terpinene), 7.9 eV) is chosen such that its ionization energy is 36 lower than both PID bulbs (detectable by both PIDs, see also e). b, Method of 37 decomposing odour signals. Top panel: high energy PID signal (grey: recorded signal, 38 blue: calculated signal due to odour 1, red: calculated signal due to odour 2). Bottom 39 panel: low energy PID signal (green: recorded signal, red: calculated signal due to odour 40 2; the entire signal is due to odour 1). **c**, Single data points of the PID signal evoked by α -41 Terpinene in the two PIDs. The slope of the linear fit serves as a scaling factor to map the 42 low energy PID to the high energy PID signal. **d**, Histogram of R-squared values of all dual-43

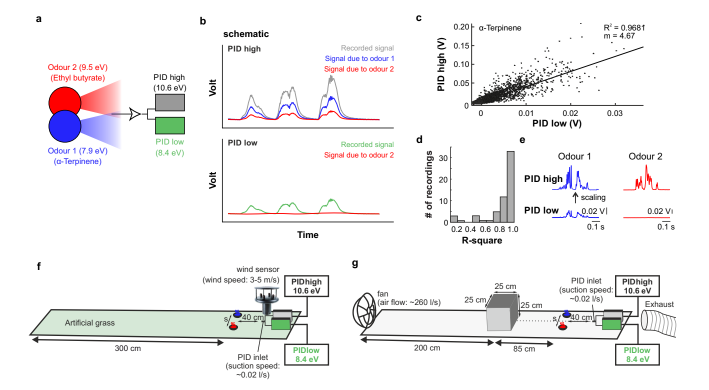
- PID α -Terpinene recordings to define the scaling factor (n = 59 recordings). **e**, Summary 44 of signal combinations for defPID recordings. The scaling factor for the PIDlow signal is 45 determined by the slope in **c**. **f**, Schematic of outdoors odour plume recording setup. PIDs 46 and odour delivery system were used to record for multiple trials at different lateral 47 distances (s) between odours held in ceramic crucibles. Data was collected on a day with 48 low wind (\sim 8-12 mph, equivalent to \sim 3-5 m/s, recorded with a 2-axis ultrasonic wind 49 sensor at the height of the PID inlet. Outdoor experiments were performed on a ~ 6 m x 50 10 m wooden patio structure surrounded by trees. There was >300 cm of unobstructed 51 space on an artificial grass mat in front of the PIDs to capture air movements. g, Indoor 52 setup: A digitally controlled fan was placed at a distance of 325 cm facing the PID inlet. 53 An exhaust line was situated behind the PID inlet to ensure the direction of air from the 54 fan towards the PID inlet. During a recording, the fan was set to maximum speed such 55 that it pushed approximately 552 cf/min (cubic feet per minute, \sim 260 l/s) of air towards 56 the PID inlet. A 25x25x25 cm Thermocool box was placed 200 cm downwind of the fan 57 acting as an obstacle to air movement, promoting complex air movement patterns at the 58
- 59 PID location. The pump at the PID was set to ~ 0.02 l/s suction speed, unlikely to perturb
- 60 overall airflow dynamics substantially.

61 **Supplementary Table 1:** Parameters of the olfactory sensory neuron population model.

62 **Supplementary Table 2:** Parameters of the olfactory sensory neuron population model

63 that were varied.





Supplementary Table 1: Parameters of the olfactory sensory neuron population model

Parameter	Description	Value
$ au_{C}$	Olfactory transduction time constant	75 ms
$ au_V$	OSN membrane time constant	75 ms
σ	Standard deviation of OSN membrane	0.25 V
	voltage additive noise	
а	Stimulus amplitude	15 V
θ	OSN spike threshold	2 V
t _{ref}	OSN refractory period	1 sec
V _{ref}	OSN refractory voltage	-1 V
$ au_h$	Ca2+ imaging filter time constant	150 ms
Δt	Euler integration step size	1 ms

Parameter	Centre	Minimum	Maximum
$ au_V$	75 ms	56.25 ms	93.75 ms
$ au_{C}$	75 ms	56.26 ms	93.75 ms
σ	0.25 V	0.1875 V	0.3125 V
θ	2 V	1.5 V	2.5 V
а	5	3.75 V	6.25 V

Supplementary Table 2: Parameters of the olfactory sensory neuron population model that were varied