Collective behaviour enhances environmental sensing and decision-making in groups of animals\textsuperscript{1,2}. Experimental and theoretical investigations of schooling fish, flocking birds and human crowds have demonstrated that simple interactions between individuals can explain emergent group dynamics\textsuperscript{3-4}. These findings indicate the existence of neural circuits that support distributed behaviours, but the molecular and cellular identities of relevant sensory pathways are unknown. Here we show that \textit{Drosophila melanogaster} exhibits collective responses to an aversive odour: individual flies weakly avoid the stimulus, but groups show enhanced escape reactions. Using high-resolution behavioural tracking, computational simulations, genetic perturbations, neural silencing and optogenetic activation we demonstrate that this collective odour avoidance arises from cascades of appendage touch interactions between pairs of flies. Inter-fly touch sensing and collective behaviour require the activity of distal leg mechanosensory sensilla neurons and the mechanosensory channel NOMPC\textsuperscript{5-6}. Remarkably, through these inter-fly encounters, wild-type flies can elicit avoidance behaviour in mutant animals that cannot sense the odour—a basic form of communication. Our data highlight the unexpected importance of social context in the sensory responses of a solitary species and open the door to a neural-circuit-level understanding of collective behaviour in animal groups.

\textit{Drosophila melanogaster} is classified as a solitary species\textsuperscript{7} but flies aggregate at high densities (>1 fly per cm\textsuperscript{2}) to feed\textsuperscript{8} (Extended Data Fig. 1a, b and Supplementary Video 1), providing opportunities for collective interactions. Although groups affect circadian rhythms\textsuperscript{9} and dispersal\textsuperscript{10} in \textit{Drosophila}, how social context influences individual sensory behaviours is unknown. To study this question, we developed an automated behavioural assay to track responses of freely-walking flies to laminar flow of air or an aversive odorant, 5% carbon dioxide. Odour was presented to one half of a planar arena for 2 min (Fig. 1a and Extended Data Fig. 1c, d). Avoidance behaviour was quantified as the percentage of time a fly spent in the air zone during the second minute of a trial (Fig. 1b, c). Unexpectedly, isolated flies spent very little time avoiding this odour (Fig. 1d), despite the aversion to CO\textsubscript{2} observed in other assays\textsuperscript{11,12}. However, increasing the number of flies was associated with substantial increases in odour avoidance (Fig. 1d and Extended Data Fig. 1e). This effect peaked at 1.13 flies per cm\textsuperscript{2}, a density typical for fly aggregates (Extended Data Fig. 1b) and was only apparent for flies in the odour zone (Fig. 1e and Extended Data Fig. 1f). Time-course analysis revealed that, within only a few seconds after odour onset, a larger proportion of flies in high-density groups had left the odour zone compared to isolated individuals (Fig. 1f; comparing 0.06 against 1.13 flies per cm\textsuperscript{2}, P < 0.05 for a Mann–Whitney U-test from 0.6 s onwards). Additionally, the motion of flies after odour onset was coherent at higher densities, with flies moving in the same direction, out of the odour zone; this effect was not observed for flies in the air zone (Extended Data Fig. 1g, h).

To determine the basis of these global behavioural differences, we examined the locomotion of individual flies. Single animals are typically sedentary but walk more when exposed to CO\textsubscript{2} (Extended Data Fig. 2a, b). In groups, however, we discovered that 63% of the time, the first walking response of a fly after odour onset coincided with proximity to a neighbouring fly (an ‘Encounter’; distance to a neighbouring fly < 25% body length; Fig. 2a–c and Supplementary Video 2). These Encounters were more frequent with increasing group density (Fig. 2d). Moreover, walking bouts (velocity > 1 mm s\textsuperscript{-1}) initiated during an Encounter (‘Encounter Responses’) were significantly longer than those spontaneously initiated in isolation (Fig. 2e). These observations indicated that inter-fly interactions might contribute to the enhanced odour avoidance of groups of flies.

We examined this possibility initially by computational simulation of the olfactory assay. The dynamics of our simulation were driven by three phenomena observed in behavioural assays (Fig. 2f). First, flies initiate more spontaneous bouts of walking in odour than in air (Extended Data Fig. 2a, b). Second, flies are more likely to turn and retreat after entering the odour zone from the air zone (Extended Data Fig. 2c). Third, close proximity to another fly elicits Encounter Responses in stationary flies (Fig. 2e and Extended Data Fig. 2d). Importantly, these elements could reproduce collective behaviour: higher numbers of simulated flies exhibited greater avoidance (Fig. 2g). While changing the olfactory parameters preserved stronger responses in groups than isolated individuals (Extended Data Fig. 2e–h), diminishing the Encounter Response probability could abolish and even reverse collective behaviour (Fig. 2h). These results suggested that Encounter Responses are a crucial component of \textit{Drosophila} group dynamics.

To experimentally test the role of inter-fly interactions in collective behaviour, we sought to explain the mechanistic basis of Encounter Responses. Although our olfactory experiments were performed in the dark (Fig. 3a), the presence of light did not diminish Encounter Response frequency (Fig. 3a). Volatile chemicals are known modulators of many social behaviours\textsuperscript{13,14}, but putative anosmic flies (lacking known olfactory co-receptors) did not reduce Encounter Responses (Fig. 3a). By contrast, disruption of the mechanosensory channel NOMPC\textsuperscript{5-6} significantly diminished Encounter Response frequency (Fig. 3a). These data suggested that mechanosensing is required for Encounter Responses.

By observing groups of flies at high spatiotemporal resolution, we found that active flies elicited motion in stationary animals through gentle touch of peripheral appendages (legs and wings; Fig. 3b and Supplementary Video 3). Leg touches took place exclusively on distal segments (Fig. 3b, inset) and resulted in spatially stereotyped walking reactions (Fig. 3c). These reactions were kinematically indistinguishable from Encounter Responses (compare Extended Data Fig. 3c and e; two-sample Kolmogorov–Smirnov test, P = 0.07; see Methods). This analysis indicates that appendage touch is the stimulus that elicits Encounter Responses. The precise stereotypy of these locomotor responses, similar to cockroach escape reactions\textsuperscript{15}, implies their dependence upon somatotopic neural circuits linking touch with movement. As fly appendages also house taste receptors\textsuperscript{16}, we tested whether mechanical stimulation was sufficient to elicit Encounter Responses by tracking stationary flies following touch of appendages with a metallic...
spent in the non-odour zone during the second minute of the experiment. Two densities of flies are shown (0.06 and 1.50 flies per cm²). The scale bar is 2.5 mm. b, Schematic of the odour avoidance experiment. Flies in the odour zone at stimulus onset (t = 0) are measured for the time spent in the non-odour zone during the second minute of the experiment ('Time avoiding odour (%)'). c, Flies (white triangles) with a low (top) or high (bottom) percent time avoiding odour. d, The per cent time avoiding the odour (mean and s.d.) for five different densities of flies starting in the odour zone (black bars) and four densities of flies starting in the air zone (grey bars) (d) and their trajectories (dashed lines) during 2 s in a two-choice olfactory assay. 5% CO₂ ('Odour') flows through the right half while air flows through the left half. Two densities of flies are shown (0.06 and 1.50 flies per cm²). Notably, among the screened lines only R55B01-Gal4 drove expression in leg mechanosensory structures: the femoral and tibial innervating leg and wing neuropils of the thoracic ganglia (Extended Data Table 2), without reduced ability to produce sustained Encounter Responses compared to a gustatory neuron driver line (Extended Data Fig. 4a). 

We next identified mechanosensory neurons required for touch-evoked Encounter Responses by driving tetanus toxin (Tnt) expression with a panel of candidate mechanosensory Gal4 lines (Extended Data Fig. 4a). R55B01-Gal4/UAS-Tnt flies exhibited significantly diminished Encounter Responses compared to a gustatory neuron driver line (Extended Data Table 2), without reduced ability to produce sustained high-velocity walking bouts (Extended Data Fig. 4b). R55B01-Gal4-driven expression of a UAS-Cδ46GFP reporter was detected in neurons innervating leg and wing neuropils of the thoracic ganglia (Extended Data Fig. 4a). Consistently, green fluorescent protein (GFP) labelled neurons in several leg mechanosensory structures: the femoral and tibial chordotonal organs, and distal leg mechanosensory sensilla neurons (Extended Data Fig. 4b). Notably, among the screened lines only R55B01-Gal4 drove expression in leg mechanosensory sensilla (Extended Data Table 2). A single asterisk (*) denotes P < 0.05 and a double asterisk (**) denotes P < 0.01. The per cent time avoiding the odour (mean and s.d.) for five densities of flies (n = 80 experiments for each condition). 

The sensitivity of simulated odour avoidance to Encounter Response probabilities ranging from 0 (never responding to Encounters, blue) to 1 (always responding, yellow). Each coloured line indicates the mean odour avoidance time (n = 10,902 experiments for each data point). The black line indicates Probability = 0.8, taken from real fly data in Fig. 1. Black circles indicate the mean fly avoidance times from Fig. 1d.

Fig. 4c, d), suggesting that these are the critical neurons for Encounter Responses. To ascertain the contribution to Encounter Responses of leg mechanosensory sensilla and/or chordotonal structures (which can also sense touch14–16), we identified additional Gal4 driver lines that drove expression in subsets of these neuron classes. By intersecting piezo-Gal4 with cha3-Gal80, a Gal4 suppression line, we could limit leg expression to mechanosensory sensilla neurons (termed 'Mechanosensory Sensilla driver' line) (Fig. 3f). Importantly, silencing neurons with this driver significantly diminished Encounter Response frequency (Fig. 3g). By contrast, silencing leg chordotonal organs alone had no effect on Encounter Response frequency (Extended Data Fig. 5a–c).

We tested the sufficiency of leg mechanosensory sensilla neuron activity to elicit Encounter Response-like walking by expressing channelrhodopsin-2 (ChR2) in each class of leg mechanosensory neurons.
Figure 4 | Encounter Responses are necessary and sufficient for collective odour avoidance. a, b. The per cent time avoiding the odour (mean and s.d.) for R55B01-Gal4 (a) or Mechanosensory Sensilla driver (b) flies expressing an inactive tetanus toxin control, or tetanus toxin. n = 22, 21, 22, and 19 experiments for R55B01-Gal4 and n = 23, 21, 21, and 21 experiments for the Mechanosensory Sensilla driver (genotypes: UAS-Tnt<sup>fl</sup>; R55B01-Gal4, UAS- Tnt<R55B01-Gal4, UAS-Tnt<sup>fl</sup>;piezo-Galcha3-Gal80<sup>+</sup>), UAS-Tnt<piezo-<br>Galcha3-Gal80<sup>+</sup>), c. The per cent time avoiding the odour (mean and s.d.) for heterozygous control, or homozygous <i>nompC<sup>1074</sup></i> mutants. n = 22, 22, 21, and 21, respectively. d, e. The per cent time avoiding the odour (mean and s.d.) for individual CO2 anosmic virtual and real flies (GR63a<sup>IR64a<sup>MB05268</sup></i>). Avoidance time is measured from a single ‘CO2 anosmic’ fly per experiment in a simulated model (d, n = 80 experiments each), or in <i>Drosophila</i> (e, n = 35, 37, 40 and 38 experiments) where single mutant flies were tested for CO2 avoidance in the context of wild-type flies.

and recording behavioural responses to blue light pulses. Optogenetic stimulation of flies expressing ChR2 in leg mechanosensory sensilla neurons, but not chordotonal organs, resulted in Encounter Response-like walking (Fig. 3h; Extended Data Fig. 5d, Supplementary Videos 5 and 6), consistent with natural elicitation of Encounter Responses by inter-fly touch of distal leg segments (Fig. 3b, inset).

Our identification of a neuronal basis for Encounter Responses allowed us to test our model’s prediction (Fig. 2h) that inter-fly interactions are required for collective odour avoidance. First, we silenced leg mechanosensory sensilla neurons by expressing Tnt with R55B01-Gal4 or the Mechanosensory Sensilla driver. Second, we studied <i>nompC<sup>1074</sup></i> mutants. Each of these perturbations abolished collective odour avoidance (Fig. 4a–c), supporting the link between mechosensation and group behavior.

Touch may enhance odour avoidance by increasing awareness of the stimulus. Alternatively, touch may produce an odour-independent Encounter Response reaction that initiates departure from the odour zone. To distinguish between these possibilities, we asked if odour-insensitive flies displayed increased avoidance in the presence of odour-sensitive animals. Indeed, both in simulations (Fig. 4d) and in real flies (Fig. 4e), increasing the number of odour-sensitive individuals led to greater avoidance behaviour of odour-insensitive individuals. Thus, in this context, touch-mediated modulation of odour awareness plays little, if any, role in collective avoidance.
Combining systems-level and neurogenetic approaches, we have uncovered a hierarchy of mechanisms that drive collective motion in *Drosophila*. Active flies elicit spatially stereotyped walking responses in stationary flies through appendage touch interactions, requiring the NOMPC mechanosensory channel and distal leg mechanosensory sensilla neurons. Through Encounter Responses, odour reactions of sensitive flies spark cascades of directed locomotion of less sensitive (or even insensitive) individuals, causing a coherent departure from the odor zone. This behavioural positive feedback and group motion are absent among flies in the non-odor zone since they are less likely to initiate walking and, consequently, have a reduced frequency of Encounters. Additionally, flies retreat when encountering the odor while transiting from the air zone. Together these behavioural phenomena cause flies to escape the odor zone and then remain in the air zone, resulting in higher odor avoidance for groups compared to isolated animals (Extended Data Fig. 6). When distal appendage mechanosensory touch detection is impaired, groups of flies cannot produce Encounter Responses, are less likely leave the odor zone, and instead behave like isolated flies. Encounters are likely to have widespread influence on sensory-evoked actions of individuals in groups. For example, movement of flies towards areas of high elevation is also increased in higher density groups (Extended Data Fig. 7).

Behaviour in animal groups arises from the detection and response to intentional and unintentional signals of conspecifics. While neural circuits controlling pairwise interactions, such as courtship, are increasing to intentional and unintentional signals of conspecifics. While neural density groups (Extended Data Fig. 7).

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Author Contributions P.R., R.B. and D.F. conceived, designed and supervised the project. P.R., S.C., P.L. and L.F. performed experiments; P.R., P.L., L.F. and W.T. analysed data. P.R. and R.B. wrote the paper with assistance from P.L. and D.F.

Author Information Reprints and permissions information is available at 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 R.B. (Richard.Benton@unil.ch) or P.R. (ramdy@gmail.com).
Methods

Drosophila lines. Actin88F-eGFP (ref. 22) backcrossed 5 generations to w1118 was used as the wild-type line enabling distinction from non-fluorescent mutant flies in Fluorescence Behavioural Imaging experiments (Fig. 3a, e) and Extended Data Fig. 5c).

Gr63a1;R66a1; AB3265/ mutants were used as the CO2-anomnic individuals (Fig. 4e).

IR86a;IR25a2;Gr63a1;ORCO4 quadruple mutant flies were used to measure the influence of olfaction on the frequency of Encounter Responses (Fig. 3a).

nanchung18 (ref. 23), piezo22 (ref. 24) and nompC20,21 (ref. 5) mutant flies were used to measure the impact of mechanosensory on the frequency of Encounter Responses (Fig. 3a).

P[GMB;Gal4]attP2 transgenic flies15,16 were used to identify neural populations with deficient touch responses (Fig. 3g, and Extended Data Figs 4, 5). Ga4 drivers were selected by pre-screening a large panel (http://bweb.janelia.org/) for those displaying sparse expression in neurons that projected from the legs to the thoracic ganglia and neurons innervating the antennal mechanosensory and motor centre in the brain26. To identify R55B01-Gal4, we compared the frequency of Encounter Responses in animals bearing these transgenes against that of a control driver, R27B07-Gal4, which drives a gustatory pattern of expression in the legs27 and Thoracic Ganglia (Extended Data Fig. 4a, green and data not shown). Brain expression in R55B01-Gal4 was limited to neurons projecting to the Antennal Mechanosensory and Motor Centre, and weaker expression in those innervating the subesophageal ganglion (which receives both gustatory and mechanosensory input from the labellum) and several visual areas (optic lobes and optic tubercle; Extended Data Fig. 5a). These weakly marked neural populations are likely to contribute only minimally to Encounter Responses as labellar touch was never observed and all experiments were performed in the dark. Finally, we also observed fan-shaped body expression in R55B01-Gal4. However, inhibiting the fan-shaped body cannot explain Encounter Response reductions in R55B01-Gal4 since silencing these neurons alone has no effect on Encounter Response frequency (Extended Data Fig. 5c, R65C03-Gal4).

piezo-Gal4,ch43-Gal80 flies were used to target mechanosensory sensilla neurors.

UAS-Tnt and UAS-TntMP flies were used to measure the effects of neuronal knockdown on Encounter Responses and collective behaviour (Figs 3g, 4a, b, and Extended Data Fig. 5c).

UAST-ChR2(T159C) flies were generated by cloning ChR2(T159C) (ref. 28) into pgUASTntgb8 (refs 29, 30) and inserting this transgene into attP22 (Genetic Services, Inc., Cambridge, MA, USA). UAST-ChR2 flies were then crossed with Ga4 driver lines for channelrhodopsin-2 stimulation experiments (Fig. 3h, and Extended Data Fig. 5d).

P{Gal4-1}III; UAST-ChR2 flies were imaged in the olfactory arena using the following illumination protocol: (1) infrared/blue light; air both sides (10 s); (2) infrared light, 5% CO2/air (2 min); (3) infrared/blue light; air both sides (5 s); (4) infrared/air; air both sides (10 s).

UAS-ChR2 flies were then crossed with Gal4 drivers for channelrhodopsin-2 stimulation experiments (Extended Data Fig. 5c).

pBac(y+;+mDint2) w;+ nc = UAS-Cd4tdGFP/VK00033 flies31 were used to visualise Ga4 driver expression in leg, brain, and thoracic ganglia neurors (Fig. 3f and Extended Data Figs 4d, e and 5a, b).

PromE(800-Gal4) [4M]; Tub-Gal80ts flies, UAS-StingerII, UAS-Hid/Cyo flies and UAS-StingerII flies were used for oenocyte ablation experiments (Fig. 3e) as described previously25.

Experimental and statistical conditions. Experiment sample sizes were chosen based on preliminary studies. If sample size constraints and proper experimental conditions were met, all experiments were included for subsequent analysis. Experiments for different conditions and genotypes were interleave to maximise the effects of time-of-day on behavioural results. Owing to the automated nature of almost all data acquisition and analysis, the experimenter was not blinded. For data meeting the criteria of normality, bar plots are presented and parametric and non-parametric statistics were used. Groups with similar variance are compared throughout the study.

Arena design and flow simulation. Arenas were designed using the 3D CAD software, SolidWorks (Dassault Systemes, Waltham, Massachusetts, USA) and CNC machined from polyoxymethylene and acrylic glass. Arena flow patterns were simulated using EasyCFD (http://www.easyCFD.net) incorporating measured physical and flow parameter values (Extended Data Fig. 1d).

Behavioural imaging and tracking. For low-resolution behavioural imaging, we used Fluorescence Behavioural Imaging (FBI)22 acquisition software and hardware. In all cases, we used Ctxr2 for tracking and data analysis was performed using custom Matlab scripts (The Mathworks, Natick, Massachusetts, USA).

Behavioural experiments. All experiments were performed on adult female Drosophila raised at 25 °C on a 12 h light:12 h dark cycle 2–4 days post-eclosion, with the exception of experiments in Extended Data Fig. 1b, which used male flies). Experiments were performed in a temperature-controlled room at 22 °C, except for those in Fig. 3d,h, which were performed at 22 °C. In all cases except for aggregation measurements, flies were starved in empty 50 mm Petri dishes for 3–6 h in humidiﬁed 25 °C incubators. Experiments were performed in either the morning or late afternoon, Zeitgeber time.

Aggregation measurements (Extended Data Fig. 1f). Flies were starved for 24 h at 25 °C in tubes humidiﬁed with moist Kimwipes. Ripe banana paste was prepared on the day of experiments and placed into a 12.78 cm2 dish. Experiments were performed in either the morning or late afternoon Zeitgeber time. Experiments for summary data (Extended Data Fig. 1b) were filmed with a webcam (Microsoft LifeCam Studio, Redmond, USA) for 90 min with images acquired every 10 min. Flies were placed into a clear transparent box (23.5 cm × 23 cm × 37.5 cm) with only red light illumination. To calculate densities, the number of flies on the food source was calculated for each image and averaged from the 30th to 90th minute.

Collective odour avoidance – wild-type, neural knockdown, nompC, and mixed wild-type/anomnic (Fig. 1 and Fig. 4a–c). For olfactory stimulation, pre-mixed 5% CO2, or air (Messer Schweiz AG, Lenzburg, Switzerland) was flowed through Mass Flow controllers (PKM SA, Lyss, Switzerland) at a regulated flow rate of 300 ml min−1 via computer controlled solenoid valves (The Lee Company, Westbrook, CT, USA). A custom-fabricated circuit board and software22 (sQuad, http://lis.epfl.ch/squid/) controlled valves, illumination LEDs (Super Bright LEDs Inc. St Louis Missouri, USA), and acquisition cameras (Allied Vision Technologies, Stadtroda, Germany). Flies were imaged in the olfactory arena using the following illumination/olfactory stimulation protocol: (1) infrared/blue light; air both sides (10 s); (2) infrared light, 5% CO2/air (2 min); (3) infrared/blue light; air both sides (10 s).

The arena half with CO2 was varied across experiments to eliminate the effects of other possible environmental asymmetries on behavioural results. Blue light was used in all cases to keep experiments consistent with mixed genotype FBI collective behaviour experiments (Fig. 4e).

Collective negative gravitaxis (Extended Data Fig. 7). For negative gravitaxis experiments, we tilted the behavioural arena at a 22.5° incline for 2 min. Flies were placed near the lower portion of the arena and were illuminated with red light. The Negative Gravitaxis Index was calculated by averaging their position along the long axis of the arena (with values ranging from 0 (bottom of the arena)) to 100 (highest point of the arena) during the second minute of the experiment.

Encounter Response modality screen (Fig. 3a). 12 wild-type flies (‘light’) or mixtures of 6 wild-type and 6 mutant flies (using a GFP reporter and Fluorescence Behavioral Imaging to distinguish genotypes22) were imaged in the olfactory arena using the following illumination/odorant protocol: (1) infrared/blue light; air both sides (10 s); (2) infrared light, 5% CO2/air (2 min); (3) infrared/blue light; air both sides (10 s).

Blue light was used in all cases to keep experiments consistent with mixed genotype Encounter Response experiments.

High-resolution inter-fly touch response (Fig. 3b, c and Extended Data Fig. 3d, e). Four flies were imaged in a small arena (1 cm × 5 cm) backlit with infrared light (Super Bright LEDs Inc. St Louis Missouri, USA). Images were continuously acquired at 125 frames per second (fps) using a high-speed video camera (Fastec Imaging, San Diego, CA, USA). The experimenter captured a video if a stationary fly exhibited touch-elicited walking.

High-resolution mechanical touch response (Fig. 3d and Extended Data Fig. 3f, g). Individual flies were imaged in a small arena (3 cm × 3 cm) illuminated by a red ring light (FALCON Illumination MV, Offenau, Germany). Images were continuously buffered at 20 fps using a high-resolution video camera (Gloor Instruments, Uster Switzerland). A small magnetic metallic disc (1 mm diameter) was directed to individual leg or wing appendages using a larger permanent magnet. The experimenter captured a video if a stationary fly exhibited touch-elicited walking.

Neuronal silencing Encounter Response screen (Extended Data Data Fig. 4a, b). 18 flies expressing inactive Tnt, or Tnt under the control of a specific Ga4 driver were imaged in the group arena using the following illumination protocol: (1) infrared/blue light (10 s); (2) infrared light (2 min); (3) infrared/blue light (30 s); (4) infrared light (2 min); (5) infrared/blue light (10 s).

Neuronal silencing Encounter Response frequency (Fig. 3g and Extended Data Fig. 5c, e). 6 flies expressing UAS-Tnt+/+[driver->Gal4+], UAS-Tnt+/+[driver->Gal4], or UAS-TntMP/[driver->Gal4] were imaged in the presence of 6 wild-type flies in the group arena using the following illumination protocol: (1) infrared/blue light; air both sides (10 s); (2) infrared light, air both sides (2 min); (3) infrared/blue light, air both sides (10 s).

Optogenetic stimulation (Fig. 3h and Extended Data Fig. 5d). Flies bearing UAS-ChR2(T159C) and the specified Ga4 driver were raised either in food mixed with 2 mM all trans-Retinal (ATR), Sigma-Aldrich, St Louis USA) or in the 95% ethanol solvent. Individual flies (2–4 days post-eclosion) were imaged in a small arena (3 cm × 3 cm) illuminated by a red ring light (FALCON Illumination MV, Offenau, Germany). Images were continuously buffered at 20 fps using a high-resolution video camera (Gloor Instruments, Uster Switzerland). An optically coupled red laser (Thorlabs, Newton, USA) was aligned to target the fly’s thoracic segment. Stimulation consisted of a short (1 s) pulse of blue laser light (Coherent,
Behavioral analysis. To determine threshold values for fly motion, Encounters, and Encounter Responses, we measured velocities, accelerations and distances that could conservatively account for a test data set of manually annotated events. To the best of our knowledge our results are qualitatively robust to small variations in these values.

Percent of time avoiding odour (Fig. 1d, e, Fig. 2g, h, Fig. 4, and Extended Data Fig. 2e–h). To calculate odour avoidance, we measured the per cent of time that flies spent within the odour (air) zone during the experiment’s second minute. This time period was chosen since we observed that flies tend to reduce exploration after one minute; see Fig. 1f. % of time avoiding odour = (time spent in the odour zone during the last minute/1 min) × 100. To report quantitatively equivalent values for experiments with different densities of flies, we resampled data using bootstrapping. This entailed randomly selecting a subgroup of experiments and, from these, one fly per experiment. We then averaged the odour avoidance for these flies to yield one result. We repeated this process a specified number of iterations to generate a distribution from which to report the mean and s.d. The number of iterations was closely linked to the average number of experiments. For example, in cases where $n = 40$, the number of bootstrapping iterations was 40. Walking bouts (Fig. 2e and Extended Data Fig. 2b). We measured activity bouts using a hysteresis threshold on forward velocity (Extended Data Fig. 2b) or velocity magnitude (Fig. 2e) to create a binary time-series. Bouts began when velocity exceeded a high threshold of 1 mm s$^{-1}$. Short bouts or pauses (<2 frames or 100 ms, see Extended Data Fig. 2b; <20 frames or 1 s, see Fig. 2e) were removed by merging the fly’s state with neighbouring measurements. Bouts were also terminated when moving flies encountered obstacles including other flies. This can explain the decreasing Encounter bout lengths observed at higher densities (Fig. 2e).

Coherent motion index (Extended Data Fig. 1g, h). To measure the coherence of group motion away or towards the odour zone, we calculated a coherent motion index (CMI). We did this by first identifying walking flies at every time-point. For these flies, we identified the orientation of walking in a binary fashion: within the half-circle pointing towards the odour half of the arena or within the half-circle pointing towards the air half of the arena. The CMI for each time-point is: \( \text{CMI} = \frac{\text{no. of flies moving towards the air} - \text{no. of flies moving towards the odour}}{\text{total no. of moving flies}}. \)

For a given experimental replicate, we average the CMI for the first ten seconds of odour presentation to capture the initial avoidance response. We report the distribution of this time-averaged CMI value across experimental replicates. For our analysis we examined the CMI for flies starting either in the air zone or the odour zone. Since the number of flies in a replicate can affect possible CMI values, comparisons should be limited to experiments with the same density of flies.

Encounter likelihood/frequency of Encounters (Fig. 2b–d). To calculate Encounter likelihood with respect to odour walking responses (Fig. 2b, c), we identified odour reactions as the time at which a stationary fly within the odour zone began moving (velocity magnitude $>1$ mm s$^{-1}$). As a control, a random time was selected from the entire experiment. We then determined the times at which each fly was undergoing any Encounter (distance to nearest neighbour $<25\%$ long-axis body length). Using these two data sets, we performed an event-triggered average of the Encounter time-series for all flies. Notably, the timing of the peak in Encounter likelihood is not of sufficient resolution to make inferences about causality. This is due to the inability to precisely define a touch encounter in low-resolution video for which the legs are not visible. With Encounters, we instead rely on an estimate based on the overlap between two circles defining the peripheral space of neighbouring flies. Therefore Encounters can continue past the onset of motion since neighbouring flies may not have become distant enough to terminate the Encounter. This is illustrated in Fig. 2b in which two flies (white circles) persisted past the times of ‘odour walking response’ (blue arrowheads) for both flies shown.

To calculate the frequency of Encounters for different group densities (Fig. 2d), we measured the proportion of time flies spent having Encounters during a given experiment. Notably, Encounters are a function of motion: flies that move are more likely to Encounter other flies.

Encounter Response frequency (Fig. 3a, e, Extended Data Fig. 4a, and Extended Data Fig. 3c). To calculate the Frequency of Encounter Responses, for each stationary fly (velocity magnitude $<1$ mm s$^{-1}$) undergoing an Encounter (distance to nearest neighbour $<25\%$ long-axis body length), we identified motion events (i.e., flight or angular velocity $>1$ mm s$^{-1}$). If there was continuous motion for the next half-second (mean velocity magnitude $>5$ mm s$^{-1}$) an Encounter Response occurred, otherwise not. The average frequency across all flies in a given experiment was used to calculate summary data. Notably, the Encounter Response frequency is normalized by the number of Encounters: Frequency of Encounter Responses = Encounters producing walking reaction/Encounters producing walking reaction + Encounters eliciting no reaction). Therefore this frequency is not a function of motion. For example, flies with high walking probabilities may generate more Encounters but reactions to these interactions—Encounter Responses—may be still more or less frequent. Similarly, flies that are predominantly stationary may have few Encounters but these too may result in a high or low frequency of Encounter Responses.

Encounter Response trajectories and kinematics (Extended Data Fig. 3b, c). To calculate Encounter Response trajectories (Extended Data Fig. 3b), for each stationary fly (velocity magnitude $<1$ mm s$^{-1}$ and angular velocity $<2$ rad s$^{-1}$) undergoing an Encounter (distance to nearest neighbour $<25\%$ long-axis body length) near the centre of the arena (distance to wall $>2$ mm), we identified motion events (angular velocity $\geq 2$ mm s$^{-1}$ or acceleration magnitude $\geq 15$ mm s$^{-2}$). The position of the fly was recorded for the remaining frames until it stopped (velocity $<1$ mm s$^{-1}$) or became close to a new fly (distance to nearest neighbour $<25\%$ long-axis body length) or to a wall (distance to wall $<2$ mm). Resulting response trajectories were pooled across experiments as a function of the octant of the Encounter (Extended Data Fig. 3a; the appropriate octant was identified as the region surrounding the fly that was bisected by a straight line between the fly’s centre of mass and that of the neighbouring fly). Encounter Response velocities were obtained for each of these trajectories and averaged to produce kinematic data. Boxplots were calculated by averaging over the first 500 ms of kinematic data (Extended Data Fig. 3c).

Touch response trajectories and kinematics (Fig. 3c, d, Extended Data Figs 3d–g). Trajectories were taken from raw tracking data of flies responding to touch. Trajectories ended when flies were near another fly or a wall. Each resulting response trajectory was pooled across experiments depending on the location of touch (for example, leg or wing). Touch response velocities were also obtained for each of these responses and averaged to recover kinematics. Boxplots were calculated by averaging over the first 160 ms (Extended Data Fig. 3e) or 500 ms (Extended Data Fig. 3g) of kinematic data. This discrepancy is due to differences in frame-rate between the two measurements.

Comparing response kinematics. Our aim was to compare the shape of kinematic data across Encounter responses, interfly touch responses, and mechanical touch responses. However, these data could be quite distinct with regards to spatial and temporal resolution. Therefore, we first concatenated the median value from each of the common seven octants (excluding the front octant in the Encounter response data set) across each of three velocity measures (forwards, sideways, and angular velocities) yielding a vector with 21 data-points. We then normalized these 21 element vectors to range from 0 to 1. These vectors were then compared using the 2-sample Kolmogorov–Smirnov test.

Simulations. Simulated flies. To verify our model of collective odour avoidance we used an agent-based simulation driven by probabilistic behaviours (Fig. 2f–h, Extended Data Fig. 2). The artificial flies had a circular body of 2.5 mm diameter and were placed in the arena of size 80 mm × 20 mm for 2,400 time-steps (corresponding to 120 s of ‘simulated’ time). The odour was presented on one half of the arena during the entire simulation. Simulated flies walked with a constant speed of 0.51 mm per time-step in straight bouts, which were separated by periods of inactivity. At the beginning of each bout, the fly was either entering an obstacle (a wall or another fly) or moving towards a random direction. Each fly randomly changed its walking direction. The bouts were initiated either spontaneously in isolation or during an Encounter.

Isolated bouts (Extended Data Fig. 2a, b). To estimate the propensities of flies to initiate walking in isolation, we performed 45 additional single fly experiments (density $= 0.06$) in which individual animals walked in the dark for 2 min. Flies were exposed to air throughout the entire arena in the first minute and odour during the second minute. For each fly ($i = 1, 2, \ldots, 45$), we integrated the differences between its consecutive positions during the first minute of the experiment (air) and separately during the second minute of the experiment (odour) at 20 Hz. The velocity magnitude of these data was treated as accumulated noise and subtracted from all 90 values. Consequently, we obtained the total distance travelled in air and in odour by each of the 45 flies (that is, $D_{\text{air}}^{\text{i}}$ and $D_{\text{odour}}^{\text{i}}$ for $i = 1, 2, \ldots, 45$). To estimate bout durations, we rescaled these 90 values such that their mean was equal to the mean duration of isolated bouts observed in the ‘six flies’ experiments (density $= 0.38$). Overall, we obtained 45 values of prototypical isolated bout lengths initiated spontaneously in air and 45 values of prototypical isolated bout lengths initiated in odour (that is, $L_{\text{air}}^{\text{i}} = 0.29D_{\text{air}}^{\text{i}}$ and $L_{\text{odour}}^{\text{i}} = 0.29D_{\text{odour}}^{\text{i}}$ for $i = 1, 2, \ldots, 45$). Of note, the estimated bout lengths varied between animals, and between air and odour for a single animal.

We used the 45 velocity values of $L_{\text{air}}^{\text{i}}$ and the 45 values of $L_{\text{odour}}^{\text{i}}$ to bootstrap the behaviour of simulated flies. A simulated fly performed a self-induced bout of length $L_{\text{air}}^{\text{i}}$ if initiated in air, and of length $L_{\text{odour}}^{\text{i}}$ if initiated in odour, where $s$ denoted which prototypical behaviour the simulated fly used. The value of $s$ was set at time-step 1 for each simulated fly independently and uniformly at random to an
integer value between 1 and 45. Thus, the values of s varied between the flies but it was possible for some flies to have the same value. The values of s were kept constant for each simulated fly during all 2,400 time-steps. Consequently, each simulated fly had a fixed propensity to move spontaneously. Moreover, within the same group, simulated flies usually differed in their propensity to move spontaneously.

Between bouts, a simulated fly following the prototypical behaviour s remained inactive for $(L_{\text{Max}} - L_{\text{Air}})/v$ time-steps when resting in air and for $(L_{\text{Max}} - L_{\text{Odour}})/v$ time-steps when resting in odour, where $L_{\text{Max}}$ is the maximal value of all 90 values of L (that is, $L_{\text{Max}} = 214$ mm) and v is the walking speed. We estimated the walking speed by measuring $D_{\text{L}}(t)$ over all 11 time-steps, from 1 to 120 s, which resulted in a value of $10.2 \text{ mm s}^{-1}$, or 0.51 mm per time-step.

Crossing the air–odour interface (Extended Data Fig. 2c). A simulated fly changed its direction of motion by 180 degrees when crossing from air to odour with probability $P(\text{turn away from odour}) = 0.4$, and when crossing from odour to air with probability $P(\text{turn away from air}) = 0.2$. The values of $P(\text{turn away from odour})$ and $P(\text{turn away from air})$ were estimated from 40 single fly (density = 0.06) experiments taken from Fig. 1d in which animals walked freely in the dark for 2 min with odour exposure on one half of the arena. We calculated the time flies spent in the odour after crossing from air, and vice versa. We classified a crossing from one half of the arena to another as a ‘turn around’ if the time spent in the new half was $\leq 3$ s. Overall, we observed 76 crossings from air to odour out of which 51 were classified as a ‘turn around’, and 72 crossings from odour to air out of which 16 were classified as a ‘turn around’.

Encounter-induced bouts (Extended Data Fig. 2d). In the simulation, at each time step t and for each walking fly we detected if the fly encountered an obstacle. If so, we checked whether in time-step $t + 1$ the fly’s body would overlap with a wall or with other flies’ bodies (assuming the fly would walk for 0.51 mm in the same direction it was heading). In these cases, the walking fly did not move in time-step $t$, but randomly changed its direction, and resumed the walk in time-step $t + 1$.

Moreover, if the walking fly encountered an inactive fly, it caused the encountered fly to initiate a bout with probability $P(\text{Encounter Response}) = 0.8$ (from Fig. 3a). The length of this Encounter Response bout was equal to $E(L_{\text{EncR}})$ and to $E(L_{\text{EncR}}+u)$ when initiated in air and in odour, respectively. The value $u$ is a random integer between 1 and 45, and $E$ is a mapping from the lengths of Isolated bouts to the lengths of Encounter Response bouts. Note that in contrast to the lengths of Isolated bouts (that is, $L_{\text{Air}}$ and $L_{\text{Odour}}$) where the value $s$ was fixed for each fly at the beginning of a simulation, here $u$ is a random variable redrawn independently for each Encounter Response. Consequently, simulated flies did not vary in their propensity to move due to Encounters.

We did not explicitly encode directionality in the Encounter Response angle. However, we observed that since virtual flies cannot occupy the same space, stationary flies would move on average away from the location of touch, an implicitly directional response.

We estimated $E$ using the data from the six fly experiments (density = 0.38) in which animals walked freely in green light illumination for 5 min without odour exposure. We observed 1,314 Isolated bouts and 618 Encounter-induced bouts. For all 1,932 bouts we calculated their lengths by integrating with temporal resolution of 20 Hz the differences between the consecutive positions of a given fly. Next, for both types of bouts, we calculated the 0th, 1st, 2nd, ..., 100th percentiles of their lengths, created a scatter plot and calculated a double linear mapping from the lengths of Isolated bouts to the lengths of Encounter Response bouts, which fit the data best (that is, $F(x) = 4.71x + 0.75$ when $x \leq 0.20$ and $F(x) = 1.04x + 70.69$ otherwise).

Experiments and sensitivity analyses (Fig. 2h and Extended Data Fig. 2e–h). Overall, there were six experiments. We performed one main experiment to test the collective behaviour of flies in two conditions: (1) all flies in the group were odour-sensitive (Fig. 2h) and (2) the first fly from the group was odour-insensitive (Fig. 4d). To simulate an odour-insensitive fly s we used $L_{\text{odour}} = L_{\text{Air}}$ in place of $L_{\text{odour}}$ values. Additionally, we performed five experiments, each corresponding to a different sensitivity analysis. For each of these experiments there were eleven conditions, each corresponding to a different value of the investigated parameter.

In the first experiment we varied the propensity to move due to Encounters by setting $P(\text{Encounter Response})$ between 0 and 1 with a step size of 0.1 (104, 6, 3, 1, 0.34, 0.17, 0.09, 0.05, 0.03, 0.01). For each of these experiments there were eleven conditions, each corresponding to a different value of the investigated parameter.

In the second experiment we varied the propensity to move in air (Extended Data Fig. 2e). To this end, we used $L_{\text{air}} = L_{\text{Air}} + L_{\text{odour}}$ in place of $L_{\text{air}}$ value, and we set the damping coefficient $a_{\text{odour}}$ from 0 to 1 with a step size of 0.1. In the third experiment we varied the propensity to move in odour (Extended Data Fig. 2f). To this end, we used $L_{\text{odour}} = L_{\text{Air}} + L_{\text{odour}}$ in place of $L_{\text{odour}}$ value, and we set the damping coefficient $a_{\text{odour}}$ from 0 to 1 with a step size of 0.1. In the fourth experiment we varied the probability to turn back when crossing the interface from odour to air by setting $P(\text{turn away from odour})$ between 0 and 1 with a step size of 0.1 (Extended Data Fig. 2h). Odour avoidance. In both conditions of the main experiment, and for each pair of 5 sensitivity experiments and 11 conditions, we ran simulations with 9 different group sizes. We used groups composed of $n = 1, 3, 6, 9, 12, 15, 18, 21$, and 24 simulated flies (not all data reported). Overall, there were $1 \times 2 \times 9 \times (\text{main})$ and $5 \times 11 \times 9$ (sensitivity) lines of experiments. Each experimental line was replicated 22,000 times using a Mersenne Twister pseudo-random numbers generator31 with a seed set to 1, 2, ..., 22,000, respectively. The initial positions, initial directions and the prototypical behaviours of simulated flies were identical between corresponding replicates across different experimental lines.

The odour avoidance of a simulated fly was calculated as the proportion of time-steps the fly spent in air during time-steps 1,200 to 2,400 corresponding to the second minute of the experiment. To compare simulations’ outcomes between treatments, conditions and group sizes, we averaged in each experimental line the odour avoidance of the first simulated fly across all replicates in which the fly was initially placed in odour (there were 10,902 such replicates out of all 22,000 replicates). Note that we chose to compare experimental lines based on the first simulated fly because it was the only fly used in all experimental lines. For example, the second and the third simulated flies were present in all experimental lines with groups composed of 3 or more flies, but were not present when the group was composed of just one fly.

For more details see the simulation’s implementation in Java on-line at: http://documents.epfl.ch/users/rit/radamya/www/ramadya_collective_sim.html.

Anatomical imaging

Brain/thoracic ganglia staining and imaging (Extended Data Fig. 5a). Immunofluorescence on whole-mount brains and thoracic ganglia was performed as described previously34. The primary antibodies were mouse monoclonal nE82 (1:8 dilution; Developmental Studies Hybridoma Bank), rabbit anti-GFP (1:200, Invitrogen A-6455). The secondary antibodies were Alexa Fluor 488- and Cy3-conjugated goat anti-rabbit or anti-mouse IgG, respectively (Molecular Probes and Jackson ImmunoResearch) diluted 1:250. Microscopy was performed using an LSM 510 laser scanning confocal microscope (Zeiss).

Leg neuron imaging (Fig. 3f and Extended Data Figs 4c, d and 5b). Legs were removed from female adults 2 days post-emergence and mounted in VectaShield under a coverslip. Cuticle was imaged with a 543 nm laser while CD4G:GFP was imaged using a 488 nm laser. Microscopy was performed using an LSM 510 laser scanning confocal microscope (Zeiss). We reoriented leg images using a custom script to identify and crop the femur, tibia, and tarsal segments. Using these sub-images, we then quantified fluorescence values (excluding autofluorescence from the cuticle and surface debris) orthogonal to the long axis of each leg segment to produce a profile of leg mechanosensory structures. Chordotonal organs and mechanosensory sensilla neurons were distinguished by morphology: sensilla neurons had small somata with dendrites projecting to the base of leg sensilla (Extended Data Fig. 4d).
Extended Data Figure 1 | *Drosophila* aggregate and move coherently at high densities. **a**, Images at the start (left) and end (right) of a $3\text{ h}$ video recording with 100 flies (50 male and 50 female) moving within a large container containing a banana paste dish (left) and an agarose dish (right). **b**, Fly densities on the banana paste dish for each gender or mixture of genders averaged from the 30th through 60th minute of a 90 min experiment ($n = 4$ experiments for each genotype). **c**, The arena for simultaneous odour stimulation and behaviour tracking of *Drosophila* groups. **d**, Laminar flow and odour localization validation using simulated fluid dynamics. High velocity vectors (yellow/red) are present at the odour entry and exit ports while lower, uniform velocity vectors (green/blue) are located within the arena. **e**, A histogram showing the per cent of time avoiding the odour for all flies in all experiments and for each density (colour-coded). Data are from Fig. 1d. **f**, The per cent time avoiding the odour (mean and s.d.) for five different densities of the subset of flies starting in the odour zone that have at some point entered the air zone ($n = 37, 38, 36, 35,$ and $38$ experiments for $0.06, 0.38, 0.75, 1.13,$ and $1.5$ flies per cm$^2$ respectively). In contrast to Fig. 1d, the lack of density dependence suggests that flies that leave the odour zone tend not to return. **g**, The formula for a Coherent Motion Index that captures the degree of motion in the same direction (top) and an example of coherent motion away from the odour zone by 9 out of 11 flies total (bottom, cyan). **h**, The Coherent Motion Index for flies in the air (white boxes) or odour (grey boxes) zones during the ten seconds following odour onset. Data are from Fig. 1d. Shown are the results across all tested densities ($0.06–1.5$ flies per cm$^2$) for flies that began the experiment in the odour (grey boxes) or the air zone (white boxes). $n = 31–38$ experiments. A single asterisk (*) denotes $P < 0.05$ and a double asterisk (**) denotes $P < 0.01$ for a Bonferroni sign test comparing medians to 0.
Extended Data Figure 2 | Model parameter determination and the sensitivity of simulated collective behaviour to parameter variation. 

a, b, Individual freely walking flies were presented with 5% CO$_2$ (‘odour’) or air across the entire arena for 1 min. Mean (solid line) and s.e.m. (translucent shading) walking velocity magnitude (a) and forward bout probability (b) before, during, and after an odour impulse (black, $n=45$ flies) or an air impulse control (blue, $n=43$ flies). Bouts began when velocity exceeded a high threshold of 1 mm s$^{-1}$. Bouts ended when velocity dipped below a low threshold of 0.5 mm s$^{-1}$. Short bouts or pauses (<2 frames or 100 ms) were removed by merging the fly’s current behavioural state with neighbouring measurements. Grey indicates the period of odour presentation. 

c, Probability for *Drosophila* to turn back when crossing the interface from odour to air and vice versa after a given period of time. Data are from Fig. 1d (density = 0.06). 

d, Scatter plots of *Drosophila* bout lengths during isolation versus Encounter Response bout lengths (red dots) and the double-linear function fitting the data (blue line). $n=16$ experiments at density = 0.38 flies per cm$^2$. The graph on the right is a zoom-in of that on the left (dashed box). 

e–h, Sensitivity of simulated collective behaviour to $P$(bout$_{air}$) ranging from $P$(bout$_{air}$) = 0 (blue, never initiating spontaneous walking in air) to $P$(bout$_{air}$) = 1 (yellow, always initiating spontaneous walking in air) (e), $P$(bout$_{odour}$) ranging from $P$(bout$_{odour}$) = 0 (blue, never initiating spontaneous walking in odour) to $P$(bout$_{odour}$) = 1 (yellow, always initiating spontaneous walking in odour) (f), $P$(turn around from air) ranging from $P$(turn around from air) = 0 (never turning around from the air zone, blue) to $P$(turn around from air) = 1 (always turning around from the air zone, yellow) (g), $P$(turn away from odour) ranging from $P$(turn away from odour) = 0 (never turning around from the odour zone, blue) to $P$(turn away from odour) = 1 (always turning around from the odour zone, yellow) (h). In all panels, each coloured line indicates the mean per cent time avoiding the odour across densities, the black line indicates the simulation result for parameter values taken from real fly data, $n=10,902$ for all data-points, and superimposed are the mean values for real flies (black circles).
Extended Data Figure 3 | Encounter Response kinematics for inter-fly or metallic disc touches. **a**, Schematic of octant colour-coding. Each Encounter Response trajectory is assigned to the perimeter octant bisected by a line drawn to the nearest neighbouring fly during an Encounter. A head octant (red) is included here but these responses likely represent front leg touches. **b**, The mean (solid lines) and standard error (translucent areas) for Encounter Response trajectories (right) colour-coded by the relative location of the neighbouring fly as in panel a. The scale bar is 1 mm. **c**, Boxplot of mean forward (top), sideways (middle), and angular (bottom) velocities for the first 0.5 s of Encounter Responses (n = 112–244 Encounters with duration >0.5 s) in the olfactory avoidance experiment from Fig. 1d (density = 0.75 flies per cm²). Velocities are colour-coded by octant. **d**, Schematic of touch-point colour-coding for high-resolution inter-fly touch response experiments. Each walking trajectory is colour-coded by the appendage touched by a neighbouring fly. Data are from Fig. 3c. **e**, Boxplot of mean forward (top), sideways (middle), and angular (bottom) velocities for the first 0.16 s of touch responses. Velocities are colour-coded by touch-point. **f**, Schematic of touch-point colour-coding for mechanical touch response experiments. Each touch response trajectory is assigned to the appendage touched by a metallic disc. Data are from Fig. 3d. **g**, Boxplot of mean forward (top), sideways (middle), and angular (bottom) velocities for the first 0.5 s of touch responses. Velocities are colour-coded by touch-point.
Extended Data Figure 4 | A behavioural screen for neurons mediating Encounter Responses and their leg expression patterns. a, Frequency of Encounter Responses for each Gal4 driver expressing UAS-Tnt. Driver lines are sorted by median frequency of Encounter Responses. A single asterisk (*) indicates $P < 0.05$ for a Bonferroni-corrected Mann–Whitney U-test comparing a given line against a gustatory neuron expression line, R27B07-Gal4 (green). Density = 1.13 flies per cm$^2$ and $n = 10$ experiments for each line. The selected line, R55B01-Gal4, drives expression in distal leg mechanosensory neurons (cyan). b, The fraction of flies in each experiment exhibiting walking velocities that meet the criteria for Encounter Responses (mean velocity magnitude greater than 5 mm s$^{-1}$ for more than 0.5 s) at any time during the experiment. Lines are sorted and colour-coded as in panel a. c, The identity and leg expression patterns of Gal4 drivers tested in the screen. Black boxes denote the presence of a given cell class. A cyan outline indicates distal leg mechanosensory neuron expression. A red outline indicates thoracic ganglion expression in lines with significant reductions in Encounter Response frequency. The expression pattern is also shown for piezo-Gal4, which was used in subsequent experiments to refine identification of the leg mechanosensory neuron class required for Encounter Responses. d, Tarsal segments for w;UAS-CD4:tdGFP,R55B01-Gal4 (left) and w;UAS-CD4:tdGFP,piezo-Gal4 (right) flies. Each tarsal segment is labelled from proximal to distal (T1-T5). Endogenous GFP fluorescence (green) is superimposed upon a transmitted light image (magenta). The scale bars are 30 μm. Below is a high-resolution image of a mechanosensory sensilla neuron on the tarsus of a w;UAS-CD4:tdGFP,R55B01-Gal4 fly. Endogenous GFP fluorescence (green) is superimposed on cuticular autofluorescence (magenta). The axon, cell body, and dendrite of this neuron are labelled. The scale bar is 10 μm.
Extended Data Figure 5 | Leg mechanosensory sensilla neurons, but not chordotonal organs, are necessary and sufficient for Encounter Responses. We identified five lines expressing Gal4 in different subsets of mechanosensory neurons (R55B01-Gal4, piezo-Gal4, piezo-Gal4;cha3-Gal80, R86D09-Gal4, and R46H11-Gal4) and one line expressing Gal4 in the fan-shaped body (R65C03-Gal4) as a control for fan-shaped body expression in R55B01-Gal4.

a, Brain and thoracic ganglion expression for Gal4 lines driving UAS-CD4:tdGFP. Immunostaining is shown for the neuropil marker nc82 (magenta) and CD4:tdGFP (green). Sensory neuron projections from the wings ('W') and legs (R1–R3 and L1–L3) are labelled for R55B01-Gal4. Importantly, neurons expressing GFP in the brains of R55B01-Gal4 and piezo-Gal4;cha3-Gal80 flies are different, implying that they are not responsible for the production of Encounter Responses. The scale bars are 40 μm. 

b, Transmitted light images, inverted GFP fluorescence images (GFP indicated in black), and summed fluorescence of Gal4 driver legs expressing CD4:tdGFP. Autofluorescent cuticle and pretarsus debris are indicated in black. GFP expression is shown in green. When present, the femoral chordotonal organ (‘fChO’), tibial chordotonal organ (‘tChO’) and mechanosensory sensilla neurons (‘MS’) are labelled. The scale bar is 100 μm.

c, The frequency of Encounter Responses for a parental control (‘Gal4’), Gal4 line neurons expressing an inactive tetanus toxin control (‘Gal4’ and ‘Inactive Tnt’), or Gal4 line neurons expressing tetanus toxin (‘Gal4’ and ‘Tnt’). n = 10–15 experiments for each condition.

d, Blue laser pulse stimulation responses of Gal4 line flies expressing UAS-ChR2 in the absence (left) or presence (right) of the essential cofactor all-trans-Retinal (n = 6–12 flies for each condition). Each box indicates the response for a single fly (‘walk’, ‘groom’, ‘leg shift’, ‘none’, or ‘jump’).
Extended Data Figure 6 | Schematic of collective odour avoidance in Drosophila. 

a, A group of flies experiences odour flow on the right half of the arena. The direction of odour or air flow is indicated by red and black arrows, respectively. Odour increases the probability of spontaneous walking (black fly). b, Walking increases the probability of encountering a stationary fly, producing an Encounter Response. c, Walking flies cause additional Encounters and a cascade of Encounter Responses in the odour zone. d, Walking flies pass into the non-odour zone through interactions with the arena walls and possibly by sensing the direction of odour flow. e, The influx of walking flies to the air zone results in additional Encounter Responses. f, The propensity to turn around at the air–odour interface (perhaps compounded by the effects of unknown aggregation pheromones) causes flies to remain in the air zone, resulting in odour avoidance.
Extended Data Figure 7 | Collective negative gravitaxis in *Drosophila*. a, A schematic of the negative gravitaxis experiment. Flies are placed at the lowest point of a behavioural arena tilted at 22.5°. The flies' positions are normalized to the long-axis of the arena ranging from 0 (arena bottom, lowest elevation) to 100 (arena top, highest elevation). b, Image of flies (black triangles) and their trajectories during 1 s (black dotted lines) in the negative gravitaxis experiment. Shown are representative images of an experiment with one fly (density = 0.06 flies per cm²) and an experiment with 18 flies (density = 1.13 flies per cm²). Negative Gravitaxis Index value positions of 0 (lowest elevation in the arena) and 100 (highest elevation in the arena) are shown (white-dashed lines). c, To obtain a Negative Gravitaxis Index for a given fly, its position was averaged during the second minute of the experiment. Shown are the mean and s.d. of Negative Gravitaxis Indices for wild-type animals at densities of either 0.06 or 1.13 flies per cm² (n = 28 and 30 experiments, respectively). d, Images of two flies (left, black triangles in black dashed box) undergoing an Encounter (middle, red dashed box) that results in an Encounter Response (right, blue dashed box) during a negative gravitaxis experiment.
Extended Data Table 1 | P values for data in main figures

| Figure | Comparison | P value (uncorrected) | Number of Comparisons |
|--------|------------|-----------------------|-----------------------|
| 1d     | 6 vs. 1 fly | 3.37 x 10^{-8}        | 4                     |
|        | 12 vs. 1 fly| 4.20 x 10^{-13}       |                       |
|        | 18 vs. 1 fly| 4.96 x 10^{-21}       |                       |
|        | 24 vs. 1 fly| 6.77 x 10^{-18}       |                       |
| 2d     | 12 vs. 6 flies | 1.67 x 10^{-11}    | 1                     |
|        | 18 vs. 12 flies | 3.91 x 10^{-11}     | 1                     |
|        | 24 vs. 18 flies | 4.50 x 10^{-11}     | 1                     |
| 2e     | 6 flies, Enc. vs. Iso. | 3.18 x 10^{-4}    | 1                     |
|        | 12 flies, Enc. vs. Iso. | 3.78 x 10^{-4}    | 1                     |
|        | 18 flies, Enc. vs. Iso. | 1.30 x 10^{-2}    | 1                     |
| 2g     | 6 vs. 1 fly | 3.69 x 10^{-8}        | 4                     |
|        | 12 vs. 1 fly | 1.87 x 10^{-17}    |                       |
|        | 18 vs. 1 fly | 4.99 x 10^{-10}    |                       |
|        | 24 vs. 1 fly | 2.27 x 10^{-8}     |                       |
| 3a     | Light vs. Dark | 1.73 x 10^{-2}    | 5                     |
|        | Anosmic vs. Dark | 4.27 x 10^{-1}    |                       |
|        | nanchung vs. Dark | 1.73 x 10^{-2}   |                       |
|        | piezo vs. Dark | 1.13 x 10^{-2}    |                       |
|        | npmpC vs. Dark | 1.00 x 10^{-3}    |                       |
| 3e     | oex vs. oe- | 5.55 x 10^{-1}      | 1                     |

| Figure | Comparison | P value (uncorrected) | Number of Comparisons |
|--------|------------|-----------------------|-----------------------|
| 3g     | Tnt vs. Gal4>Tnt | 1.28 x 10^{4}      | 3                     |
|        | Gal4 vs. Gal4>Tnt | 2.74 x 10^{4}      |                       |
|        | Inactive Tnt vs. Gal4>Tnt | 1.10 x 10^{3} |                       |
| 4a     | Inactive Tnt, 18 vs. 1 fly | 2.99 x 10^{-2}   | 1                     |
|        | Tnt, 18 vs. 1 fly | 5.54 x 10^{1}       |                       |
| 4b     | Inactive Tnt, 18 vs. 1 fly | 3.90 x 10^{2}      | 1                     |
|        | Tnt, 18 vs. 1 fly | 3.75 x 10^{2}       |                       |
| 4c     | npmpC^{--}, 18 vs. 1 fly | 1.12 x 10^{5}      | 1                     |
|        | npmpC^{--}, 18 vs. 1 fly | 1.62 x 10^{1}      |                       |
| 4d     | 3 vs. 1 fly | 3.98 x 10^{2}        | 3                     |
|        | 6 vs. 1 fly | 2.98 x 10^{2}        |                       |
|        | 12 vs. 1 fly | 3.12 x 10^{3}        |                       |
| 4e     | 3 vs. 1 fly | 8.87 x 10^{-18}      | 3                     |
|        | 6 vs. 1 fly | 1.76 x 10^{18}       |                       |
|        | 12 vs. 1 fly | 3.60 x 10^{18}      |                       |

The uncorrected P values for each main figure panel and its associated comparison are indicated. The number of comparisons used for post-hoc Bonferroni correction for multiple comparisons is also shown.
Extended Data Table 2 | P values for data in Extended Data figures

| Figure | Comparison                  | P value (uncorrected) | Number of Comparisons |
|--------|-----------------------------|-----------------------|-----------------------|
| 1h     | Air vs. Odour, 6 flies      | 8.90 x 10^-1         | 1                     |
|        | Air vs. Odour, 12 flies     | 2.90 x 10^-3         | 1                     |
|        | Air vs. Odour, 18 flies     | 2.16 x 10^-4         | 1                     |
|        | Air vs. Odour, 24 flies     | 1.40 x 10^-5         | 1                     |
| 4a     | R65A11 vs. R27B07           | 5.01 x 10^-4         | 19                    |
|        | R14F12 vs. R27B07           | 2.18 x 10^-4         |                       |
|        | R20C06 vs. R27B07           | 8.15 x 10^-6         |                       |
|        | R55B01 vs. R27B07           | 2.50 x 10^-3         |                       |
|        | R13E04 vs. R27B07           | 5.10 x 10^-3         |                       |
|        | R93A02 vs. R27B07           | 5.69 x 10^-2         |                       |
|        | R39D08 vs. R27B07           | 2.50 x 10^-3         |                       |
|        | R95A11 vs. R27B07           | 1.26 x 10^-2         |                       |
|        | R41A08 vs. R27B07           | 7.94 x 10^-2         |                       |
|        | R46H11 vs. R27B07           | 1.78 x 10^-2         |                       |
|        | R59C09 vs. R27B07           | 1.81 x 10^-2         |                       |
|        | R46D02 vs. R27B07           | 4.02 x 10^-2         |                       |
|        | R86D09 vs. R27B07           | 3.25 x 10^-1         |                       |
|        | R22A04 vs. R27B07           | 2.37 x 10^-1         |                       |
|        | R39A11 vs. R27B07           | 4.18 x 10^-2         |                       |
|        | R27E02 vs. R27B07           | 2.37 x 10^-1         |                       |
|        | R93D11 vs. R27B07           | 2.18 x 10^-1         |                       |
|        | R86G01 vs. R27B07           | 3.72 x 10^-1         |                       |
|        | R74F10 vs. R27B07           | 8.44 x 10^-1         |                       |
| 6c     | R55B01-Gal4                 | 1.10 x 10^-3         | 2                     |
|        | Gal4 vs. Gal4>Tnt           | 1.70 x 10^-3         |                       |
|        | piezo-Gal4                  |                        |                       |
|        | Gal4 vs. Gal4>Tnt           | 1.24 x 10^-3         |                       |
|        | Gal4>inactive vs. Gal4>Tnt  | 1.50 x 10^-3         |                       |
|        | piezo-Gal4, cha3-Gal80      |                        |                       |
|        | Gal4 vs. Gal4>Tnt           | 2.74 x 10^-4         |                       |
|        | Gal4>inactive vs. Gal4>Tnt  | 1.10 x 10^-3         |                       |
|        | R86D09-Gal4                 |                        |                       |
|        | Gal4 vs. Gal4>Tnt           | 2.07 x 10^-1         |                       |
|        | Gal4>inactive vs. Gal4>Tnt  | 8.94 x 10^-1         |                       |
| 7c     | 18 vs. 1 fly                | 3.63 x 10^-1         | 1                     |

The uncorrected P values for each Extended Data figure panel and its associated comparison are indicated. The number of comparisons used for post-hoc Bonferroni correction for multiple comparisons is also shown.