

1 Serotonergic Neurons Mediate 2 Operant Conditioning in *Drosophila* 3 Larvae

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12
13 **Abstract** Observed across species, operant conditioning facilitates learned associations
14 between behaviours and outcomes, biasing future action selection to maximise reward and avoid
15 punishment. To elucidate the underlying neural mechanisms, we built a high-throughput tracker
16 for *Drosophila melanogaster* larvae, combining real-time behaviour detection with closed-loop
17 optogenetic and thermogenetic stimulation capabilities. We demonstrate operant conditioning in
18 *Drosophila* larvae by inducing a bend direction preference through optogenetic activation of
19 reward-encoding serotonergic neurons. Specifically, we establish that the ventral nerve cord is
20 necessary for this memory formation. Our results extend the role of serotonergic neurons for
21 learning in insects as well as the existence of learning circuits outside the mushroom body. This
22 work supports future studies on the function of serotonin and the mechanisms underlying
23 operant conditioning at both circuit and cellular levels.

24 25 **Introduction**

26 Animals must rapidly alter their behaviour in response to environmental changes. An important
27 adaptation strategy is associative learning (*Dickinson, 1981; Rescorla, 1988*), in which an animal
28 learns to predict an unconditioned stimulus (US) by the occurrence of a conditioned stimulus (CS).
29 The US is often a punishing or rewarding event such as pain or the discovery of a new food source
30 (*Pavlov, 1927*). The nature of the CS distinguishes two major associative learning types: classical
31 conditioning (*Pavlov, 1927*) and operant conditioning (*Skinner, 1938; Thorndike, 1911*).

32 In classical conditioning, the CS is an inherently neutral environmental stimulus such as a sound,
33 odour, or visual cue. Pairing with an appetitive or aversive US leads to learned approach or avoid-
34 ance of the CS in the future. Many vertebrates (*Andreatta and Pauli, 2015; Brown et al., 1951;*
35 *Jones et al., 2005; Braubach et al., 2009*) and invertebrates (*Takeda, 1961; Vinauger et al., 2014;*
36 *Alexander et al., 1984; Wen et al., 1997; Scherer et al., 2003; Davis, 2005; Cognigni et al., 2018; Vogt*
37 *et al., 2014*) can make these associations. Across the animal kingdom, neural circuits have been
38 identified as convergence sites for the external CS and the rewarding or punishing US (*Heisenberg*
39 *et al., 1985; Hawkins and Byrne, 2015; Oswald and Waddell, 2015; Gründemann and Lüthi, 2015;*
40 *Caroni, 2015; Tonegawa et al., 2015*). In classical conditioning of both larval and adult *Drosophila*,

41 the mushroom body (MB) brain area serves this purpose (*Cognigni et al., 2018; Heisenberg et al.,*
42 *1985; Heisenberg, 2003; Rohwedder et al., 2016; Vogt et al., 2014; Saumweber et al., 2018; Oswald*
43 *and Waddell, 2015*). In each larval brain hemisphere, the CS is encoded by a subset of the ap-
44 proximately 110 Kenyon cells (KCs) (*Aso et al., 2014a; Honegger et al., 2011; Berck et al., 2016; Lin*
45 *et al., 2014; Oswald and Waddell, 2015; Campbell et al., 2013; Turner et al., 2008; Eichler et al.,*
46 *2017*), which synapse onto 24 MB output neurons (MBONs) driving approach or avoidance (*Aso*
47 *et al., 2014b; Oswald et al., 2015; Perisse et al., 2016; Séjourné et al., 2011; Saumweber et al., 2018;*
48 *Shyu et al., 2017; Plaçais et al., 2013; Eichler et al., 2017*). KC to MBON connection strength is
49 modulated by dopaminergic and octopaminergic neurons, which represent the rewarding or pun-
50 ishing US (*Schwaerzel et al., 2003; Schroll et al., 2006; Honjo and Furukubo-Tokunaga, 2009; Vogt*
51 *et al., 2014; Saumweber et al., 2018; Waddell, 2013*). Activation of the MB-innervating PAM cluster
52 dopaminergic neurons serves as both a necessary and sufficient reward signal in classical condi-
53 tioning (*Rohwedder et al., 2016; Liu et al., 2012; Vogt et al., 2014; Waddell, 2013; Cognigni et al.,*
54 *2018*).

55 In operant conditioning, the CS is an animal's own action (*Skinner, 1938; Thorndike, 1911*). Af-
56 ter memory formation, the animal can predict the outcome of its behaviour and bias future action
57 selection accordingly, usually to maximise reward and avoid punishment (*Skinner, 1938*). This be-
58 havioural adaptation can facilitate novel action sequences (*Topál et al., 2006; Nottebohm, 1991;*
59 *Fee and Goldberg, 2011*) and, in some cases, repetitive, high-frequency motor activity (*Olds and*
60 *Milner, 1954; Corbett and Wise, 1980; Jin and Costa, 2010; Lovell et al., 2015*). Such observations
61 have wider implications for understanding diseases including obsessive-compulsive disorder and
62 addiction (*Everitt et al., 2018; Balleine et al., 2015; Joel, 2006*). Invertebrates are also capable of
63 operant conditioning (*Brembs, 2003; Hoyle, 1979; Abramson et al., 2016; Nuwal et al., 2012; Booker*
64 *and Quinn, 1981*). Despite countless operant conditioning experiments across species using vari-
65 ous CS-US combinations, the underlying neural mechanisms remain poorly understood. For an
66 animal to associate an action with its outcome, behavioural information must converge with cir-
67 cuits encoding positive or negative valence. Although vertebrate basal ganglia-like structures ex-
68 emplify this (*Fee and Goldberg, 2011; Redgrave et al., 2011; Balleine et al., 2009*), some learned
69 action-outcome associations do not require the brain (*Booker and Quinn, 1981; Horridge, 1962;*
70 *Grau et al., 1998*). Operant conditioning may hence occur in more than one area of the central ner-
71 vous system (CNS). It is also unclear to what extent learning at these sites is mediated by synaptic
72 plasticity (*Lovinger, 2010; Surmeier et al., 2007; Reynolds and Wickens, 2002; Joynes et al., 2004;*
73 *Gómez-Pinilla et al., 2007*) versus changes in the intrinsic excitability of individual neurons (*Dong*
74 *et al., 2006; Shen et al., 2005; Nargeot et al., 1997; Brembs et al., 2002; Nargeot et al., 2009*). We
75 aim to establish the *Drosophila* larva as a tractable model system for studying the neural circuit
76 mechanisms underlying operant conditioning.

77 *Drosophila melanogaster* larvae perform various different actions. Typically, when exploring an
78 environment, a larva alternates between crawling via forward peristalsis (*Heckscher et al., 2012*)
79 and bending its head once or more to the left or right (*Gomez-Marin et al., 2011; Luo et al., 2010;*
80 *Kane et al., 2013; Figure 1A*). In the presence of nociceptive stimuli, larvae exhibit escape behaviour.
81 While the most common response is an increase in bending away from undesirable conditions,
82 including extreme temperature (*Luo et al., 2010; Lahiri et al., 2011*), light (*Kane et al., 2013*), or
83 wind (*Jovanic et al., 2019*), larvae also retreat from aversive sources using backward peristalsis
84 (*Masson et al., 2020; Kernan et al., 1994; Heckscher et al., 2012; Vogelstein et al., 2014; Figure 1A*).
85 The fastest escape response is rolling, where the larva moves laterally by curling into a C-shape and
86 quickly turning around its own body axis (*Robertson et al., 2013; Hwang et al., 2007; Ohyama et al.,*
87 *2013; Figure 1A*). In nature, rolling is only observed after exposure to a strong noxious stimulus,
88 such as heat or a predator attack (*Ohyama et al., 2015; Robertson et al., 2013; Tracey et al., 2003*).

89 Powerful genetic toolkits have advanced the observation and manipulation of larval behaviour
90 at the cellular level, making *Drosophila* larvae particularly well-suited for studying the neural mech-
91 anisms underlying learning. In *Drosophila*, individual neurons are uniquely identifiable, with mor-

92 phology and function preserved across animals (*Skeath and Thor, 2003; Wong et al., 2002; Marin*
93 *et al., 2002; Jefferis et al., 2007*). Together with tissue-localised protein expression afforded by
94 the GAL4-UAS binary expression system (*Fischer et al., 1988; Brand and Perrimon, 1993*), this has
95 yielded neuron-specific GAL4 drivers (*Jenett et al., 2012; Luan et al., 2006; Pfeiffer et al., 2010*) that
96 reproducibly target the same group of cells in each individual. Adding fluorescent markers helps
97 pinpoint a neuron's location and reveal its anatomical features (*Lee and Luo, 1999*), while producing
98 light-sensitive channelrhodopsins and temperature-sensitive ion channels facilitates optogenetic
99 (*Zemelman et al., 2002; Lima and Miesenböck, 2005*) or thermogenetic (*Hamada et al., 2008; Ki-*
100 *tamoto, 2001*) modulation of neural activity. Furthermore, the larva's compact CNS has made it
101 feasible to manually reconstruct neurons and their synaptic partners from a larval electron mi-
102 croscopy (EM) volume (*Berck et al., 2016; Eichler et al., 2017; Fushiki et al., 2016; Ohyama et al.,*
103 *2015; Schlegel et al., 2016; Larderet et al., 2017; Jovanic et al., 2016, 2019*), giving rise to a full wiring
104 diagram of the MB (*Eichler et al., 2017; Eschbach et al., 2020a,b*).

105 There is overwhelming evidence that larvae are capable of classical conditioning. They can
106 be trained to approach an odour paired with a gustatory reward (*Schleyer et al., 2011; Hendel*
107 *et al., 2005; Kudow et al., 2017; Niewalda et al., 2008*), or avoid an odour paired with light (*von*
108 *Essen et al., 2011*), electric shock (*Aceves-Piña and Quinn, 1979; Tully et al., 1994*), heat (*Khurana*
109 *et al., 2012*), vibration (*Eschbach et al., 2011*), or the bitter compound quinine (*Gerber and Hendel,*
110 *2006; Apostolopoulou et al., 2014*). Light can also be a CS: innate avoidance of light and prefer-
111 ence for darkness (*Sawin-McCormack et al., 1995*) can be modulated when paired with reward or
112 punishment (*Gerber et al., 2004; von Essen et al., 2011*). It has remained an open question, how-
113 ever, whether *Drosophila* larvae can form action–outcome associations and where in the CNS these
114 memories are formed.

115 Conducting operant conditioning with larvae requires real-time behaviour detection such that
116 reward or punishment can be administered with minimal delay (*Figure 1B*). Single-animal closed-
117 loop trackers have recently been developed (*Schulze et al., 2015; Tadres and Louis, 2020*). However,
118 the efficiency of training paradigms would improve with automated US delivery and simultane-
119 ous conditioning of multiple animals. Therefore, we here introduce a high-throughput tracker for
120 *Drosophila* larvae with real-time behaviour detection and closed-loop stimulation. Efficiency of the
121 setup stems from the simultaneous, real-time, behaviour detection for up to 16 freely moving lar-
122 vae, and targeted closed-loop optogenetic and thermogenetic stimulus delivery with full intensity
123 control and minimal delay.

124 Results

125 High-throughput closed-loop tracker

126 Hardware design

127 Designing an automated operant conditioning protocol for the *Drosophila* larva was challenging
128 due to the larva's physical characteristics. We excluded partial immobilisation protocols similar
129 to the ones used to condition adult *Drosophila* navigation through virtual environments (*Nuwal*
130 *et al., 2012; Wolf and Heisenberg, 1991; Wolf et al., 1998; Brembs, 2011*). We instead built a high-
131 throughput multi-larva tracker combining live computer vision behaviour detection with closed-
132 loop control of US delivery in response to unrestricted larval behaviour.

133 All hardware resided within an optically opaque enclosure to ensure experiments were per-
134 formed without environmental light. Larvae moved freely on an agarose plate, backlit from be-
135 low by an infrared LED and observed from above through a high-resolution camera (*Figure 1C*).
136 A Camera Link communication protocol interfaced with a high-performance field-programmable
137 gate array (FPGA), which itself interacted with the host computer. The FPGA and the host computer
138 performed image processing, behaviour detection, and stimulus calculation (*Figure 1D*).

139 Our operant conditioning paradigm targeted individual larvae performing specific behaviours.
140 Optogenetic stimulation was achieved by directing red light through two digital micromirror de-

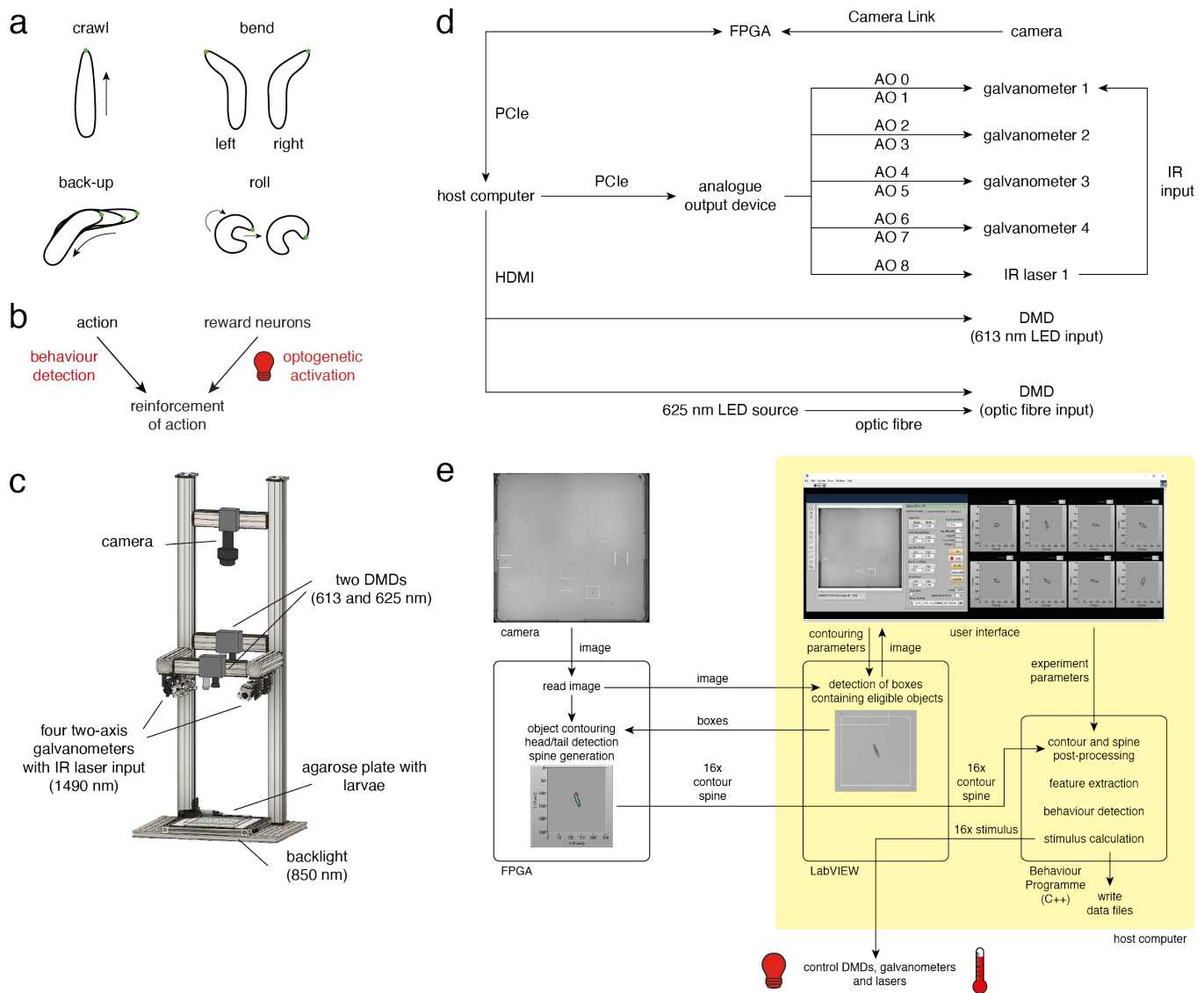


Figure 1. High-throughput operant conditioning in *Drosophila* larvae. **a.** Behavioural repertoire of *Drosophila* larvae. Schematics show the four most prominent actions displayed by *Drosophila* larvae (crawl, left and right bend, back-up and roll). The larval contour is displayed as a black outline with a green dot marking the head. **b.** In fully automated operant conditioning, an action of interest was reinforced by coupling real-time behaviour detection with optogenetic activation of reward circuits. **c.** High-throughput tracker schematic showing the relative positions of the agarose plate, backlight, camera, digital micromirror devices (DMDs), and galvanometers. IR: infrared. **d.** Block diagram of hardware components. AO: analogue output, FPGA: field-programmable gate array. **e.** Data flow between software elements.

Figure 1-Figure supplement 1. Contour calculation on field-programmable gate array (FPGA).

Figure 1-Figure supplement 2. Detecting head and tail.

Figure 1-Figure supplement 3. Calculating a smooth spine and landmark points.

Figure 1-Figure supplement 4. Calculating direction vectors.

Figure 1-Figure supplement 5. Features describing body shape.

Figure 1-Figure supplement 6. Velocity features.

Figure 1-Figure supplement 7. Temporal smoothing of features.

Figure 1-Figure supplement 8. Differentiation by convolution.

141 vices (DMDs) which were programmed to project small 1 cm² squares at the location of individual
142 larvae. Both DMDs, which were positioned to project over the entire plate area, were operated
143 simultaneously (**Figure 1C**).

144 Thermogenetic stimulation of individual larvae was achieved by directing a 1490 nm infrared
145 (IR) laser beam through a two-axis scanning galvanometer mirror positioning system (**Figure 1C**), a
146 technique previously used to stimulate single adult flies (**Bath et al., 2014; Wu et al., 2014**). Because
147 the 1490 nm wavelength is well-absorbed by water (**Curcio and Petty, 1951**), larvae exposed to the
148 IR beam were rapidly heated. We took advantage of the galvanometer's high scanning velocity to
149 rapidly cycle the beam between four larvae (**Figure 1D**).

150 Software architecture

151 Several computer vision algorithms exist for real-time tracking of freely behaving animals. **Stowers**
152 **et al. (2017)** and **Krynitsky et al. (2020)** developed software for tracking mice, and **Mischiati et al.**
153 **(2015)** developed high-speed tracking of single dragonflies in three-dimensional space. There are
154 numerous tracking frameworks for adult *Drosophila*, some requiring the flies to move within a two-
155 dimensional plane (**Straw and Dickinson, 2009; Donelson et al., 2012**) while others detect the three-
156 dimensional position of single (**Fry et al., 2008**) or multiple (**Grover et al., 2008; Straw et al., 2011**)
157 flies. The Multi-Worm Tracker (MWT) software developed by **Swierczek et al. (2011)** is suitable for
158 simultaneously tracking a large number of *Caenorhabditis elegans* and has been adapted to analyse
159 *Drosophila* larvae reactions in response to various stimuli (**Ohyama et al., 2013; Vogelstein et al.,**
160 **2014; Jovanic et al., 2019; Masson et al., 2020**).

161 Operant conditioning requires live behaviour detection to trigger delivery of reward or punish-
162 ment. Numerous algorithms have been developed to analyse offline behavioural recordings of an-
163 imals such as *C. elegans* (**Huang et al., 2006; Stephens et al., 2008; Gupta and Gomez-Marin, 2019**),
164 zebrafish larvae (**Mirat et al., 2013; Reddy et al., 2020**), adult *Drosophila* (**Katsov and Clandinin,**
165 **2008; Branson et al., 2009; Dankert et al., 2009; Robie et al., 2017; Berman et al., 2014; Klibaite**
166 **et al., 2017**), bees (**Veeraraghavan et al., 2008**), and mice (**Mathis et al., 2018; Luxem et al., 2020;**
167 **van Dam et al., 2020**). The *Drosophila* larva has also attracted attention due to analytical challenges
168 surrounding its deformable body and limited set of distinguishing features (**Luo et al., 2010; Gomez-**
169 **Marin et al., 2011; Gershow et al., 2012; Denisov et al., 2013; Vogelstein et al., 2014; Ohyama et al.,**
170 **2013, 2015; Masson et al., 2020**). Most of these approaches are not ideal to run in real time or re-
171 quire a mix of past and future information to provide reliable behaviour detection (**Gomez-Marin**
172 **et al., 2011; Masson et al., 2020**). More generally, machine learning based methods have gained
173 momentum in providing both supervised and unsupervised approaches to behaviour analysis. It
174 is worth noting a recent trend in developing unsupervised learning methods (e.g. **Graving and**
175 **Couzin, 2020; Luxem et al., 2020**).

176 While real-time behaviour detection of casts and runs has been developed for a single animal
177 (**Schulze et al., 2015**), our study of operant conditioning in freely behaving *Drosophila* larvae re-
178 quired efficient, real-time behaviour detection of multiple animals. We built a system to simul-
179 taneously track up to 16 larvae in real time, using LabVIEW for the user interface and algorithm
180 implementation (**Figure 1E**). Instrumental to this software architecture was the fast image pro-
181 cessing speed afforded by FPGA-based parallelisation (**Soares dos Santos and Ferreira, 2014; Li**
182 **et al., 2011; Zhang et al., 2017**). Neuroscientists have adapted FPGA's real-time analysis capabili-
183 ties (**Shirvaikar and Bushnaq, 2009; Uzun et al., 2005; Chiuchisan, 2013; Yasukawa et al., 2016**) to
184 track rats (**Chen et al., 2005**), zebrafish larvae (**Cong et al., 2017**), and fluorescently labelled neu-
185 rons in freely behaving *Drosophila* larvae (**Karagoyozov et al., 2018**). In our system, the FPGA and
186 host computer worked together to read the raw camera images, detect eligible objects, and extract
187 and process object features (i. e. contour, head and tail position, and body axis) (**Figure 1E**). Larval
188 body shape, velocity, and direction of motion facilitated robust behaviour detection which, in turn,
189 drove closed-loop optogenetic and thermogenetic stimulation. All relevant experiment parame-
190 ters and time-series data were output for offline analysis through a custom MATLAB framework

191 (see Materials and methods).

192 Optogenetic and thermogenetic stimulation efficiency verified by behavioural readout
193 We conducted proof-of-principle experiments to ensure that our set-up could be successfully used
194 for optogenetic stimulation (**Figure 2A**). *Ohyama et al. (2015)* have identified two GAL4 lines ex-
195 pressed in neurons whose activation triggers strong rolling behaviour. *69F06-Gal4* drives expres-
196 sion in command neurons for rolling, whereas *72F11-Gal4* drives expression in the Basin neurons,
197 which integrate mechanosensory and nociceptive stimuli. *Klapoetke et al. (2014)* have developed
198 the red-shifted channelrhodopsin *CsChrimson*, which can be expressed under GAL4 control. We
199 tested whether *69F06-Gal4 x UAS-CsChrimson* and *72F11-Gal4 x UAS-CsChrimson* larvae rolled upon
200 exposure to red light (**Figure 2B**, see also Materials and methods). In each stimulation cycle, we
201 observed above-threshold rolls in over 40% of *69F06-Gal4 x UAS-CsChrimson* larvae and over 70%
202 of *72F11-Gal4 x UAS-CsChrimson* larvae. This behaviour significantly contrasted with that of *attP2 x*
203 *UAS-CsChrimson* control larvae (**Figure 2C**), suggesting that the DMDs could be used for optogenetic
204 stimulation without activating the animals' photoreceptors.

205 We also verified the efficacy of the galvanometer set-up for thermogenetic stimulation (**Fig-**
206 **ure 2D**). We tested whether *69F06-Gal4 x UAS-dTrpA1* and *72F11-Gal4 x UAS-dTrpA1* larvae rolled
207 upon exposure to the IR laser (**Figure 2E**, see also Materials and methods). In each stimulation
208 cycle, we observed above-threshold rolls in over 70% of *69F06-Gal4 x UAS-dTrpA1* larvae and over
209 35% of *72F11-Gal4 x UAS-dTrpA1* larvae; a significant contrast to the *attP2 x UAS-dTrpA1* control lar-
210 vae whose roll rate was close to zero. We concluded that these heating conditions were effective
211 for targeted *Trp* channel activation without larvae perceiving strong pain (**Figure 2F**).

212 **Operant conditioning of larval bend direction**

213 We chose optogenetic activation of reward circuits as a US for automated operant conditioning.
214 The main challenge was determining which neurons could convey a sufficient reinforcement sig-
215 nal, especially as the capacity for *Drosophila* larvae to exhibit operant learning was not yet demon-
216 strated. Across the animal kingdom, it has been observed that biogenic amine neurotransmitters
217 can provide such a signal (*Giurfa, 2006; Hawkins and Byrne, 2015; Meneses and Liy-Salmeron, 2012;*
218 *Fee and Goldberg, 2011*). It is also conceivable that the *Drosophila* PAM cluster dopaminergic neu-
219 rons that can signal reward in classical conditioning (*Rohwedder et al., 2016; Liu et al., 2012; Vogt*
220 *et al., 2014; Cognigni et al., 2018; Waddell, 2013*) may perform similarly in operant conditioning.
221 We therefore aimed to induce operant conditioning by stimulating a broad set of dopaminergic
222 and serotonergic neurons. If valence signalling relevant for operant conditioning is mediated by
223 one of these two neurotransmitters, activation of this large set of neurons paired with behaviour
224 should be sufficient to induce learning.

225 We expressed *UAS-CsChrimson* under the control of the *Ddc-Gal4* driver, which covers a large
226 set of dopaminergic and serotonergic neurons in the CNS (*Li et al., 2000; Sitaraman et al., 2008;*
227 *Lundell and Hirsh, 1994*), including the PAM cluster (*Liu et al., 2012; Aso et al., 2012*). Although the
228 function of most *Ddc* neurons is unknown, their collective activation can substitute for an olfactory
229 conditioning reward in adult flies (*Liu et al., 2012; Shyu et al., 2017; Aso et al., 2012*). The goal
230 of our paradigm was to establish a learned direction preference for bending, conditioning *Ddc-*
231 *Gal4 x UAS-CsChrimson* larvae to bend more often to one side than the other. Although stimulation
232 side was randomized across trials, we describe (for simplicity) the experiment procedure where
233 this predefined side was the left. Each experiment began with a one-minute test period where
234 no light was presented. What followed were four training sessions, each three-minutes long, in
235 which larvae received optogenetic stimulation when bending to the left. Between training sessions,
236 larvae experienced three minutes without stimulation. Larvae were periodically brushed back to
237 the centre of the agarose plate to mitigate the experimental side effects of reaching the plate's
238 edge (see Materials and methods for more details). Following the fourth training session was a
239 one-minute test period without stimulation (**Figure 3A**).

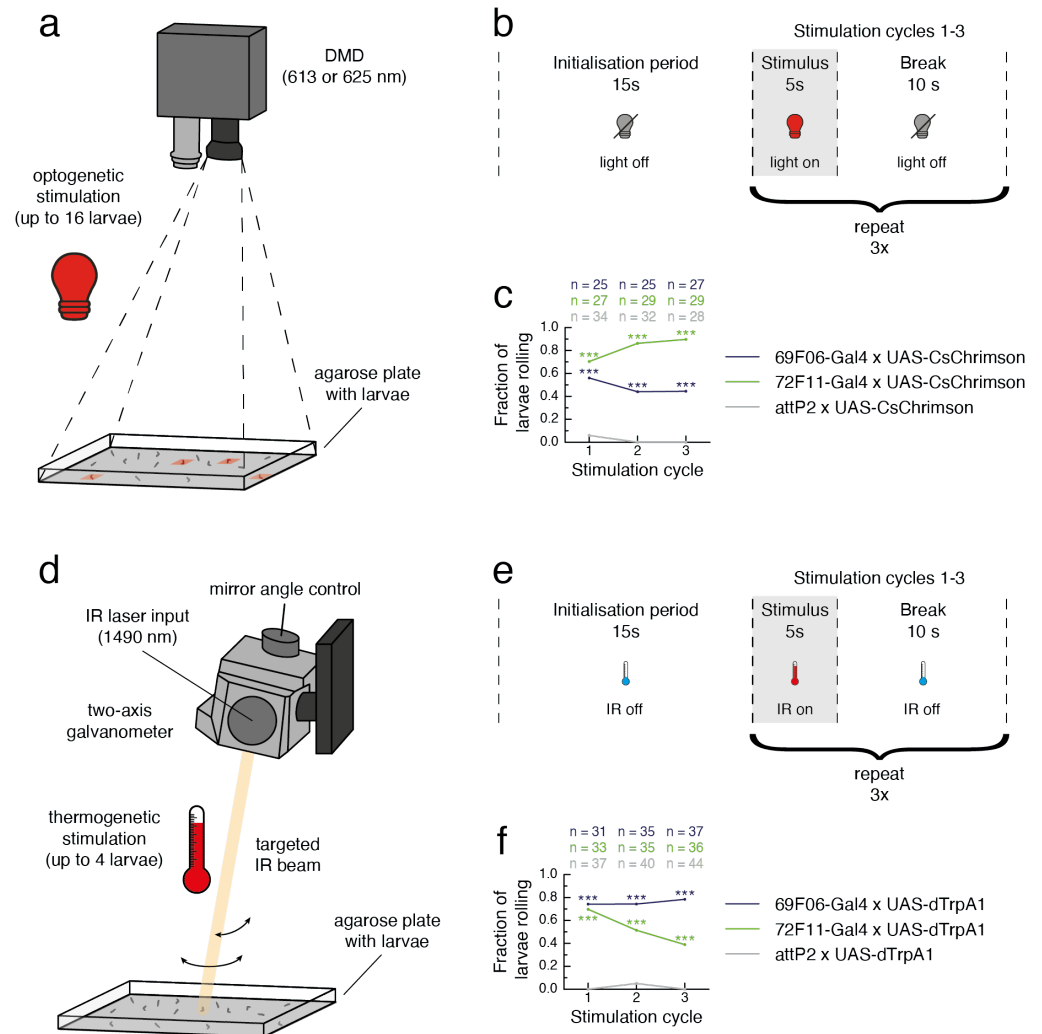


Figure 2. Optogenetic and thermogenetic stimulation with the high-throughput tracker. **a.** Hardware design schematic for optogenetic stimulation. Although the high-throughput tracker included two digital micromirror devices (DMDs), only one is shown for simplicity. **b.** Proof-of-principal experiment protocol for optogenetic stimulation. **c.** The fraction of larvae for which a roll was detected in each stimulation cycle. 69F06-Gal4 x UAS-CsChrimson and 72F11-Gal4 x UAS-CsChrimson larvae (CsChrimson expressed in neurons triggering roll behaviour; experiment groups) were compared to attP2 x UAS-CsChrimson larvae (no CsChrimson expression; control group). Fisher's exact test was used to calculate statistical differences between the experiment and control groups (***) $p < 0.001$. **d.** Hardware design schematic for thermogenetic stimulation. Although the high-throughput tracker included four two-axis galvanometers, only one is shown for simplicity. IR: infrared. **e.** Proof-of-principal experiment protocol for thermogenetic stimulation. **f.** The fraction of larvae for which a roll was detected in each stimulation cycle. 69F06-Gal4 x UAS-dTrpA1 and 72F11-Gal4 x UAS-dTrpA1 larvae (dTrpA1 expressed in neurons triggering roll behaviour; experiment groups) were compared to attP2 x UAS-dTrpA1 larvae (no dTrpA1 expression; control group). Fisher's exact test was used to calculate statistical differences between the experiment and control groups (***) $p < 0.001$.

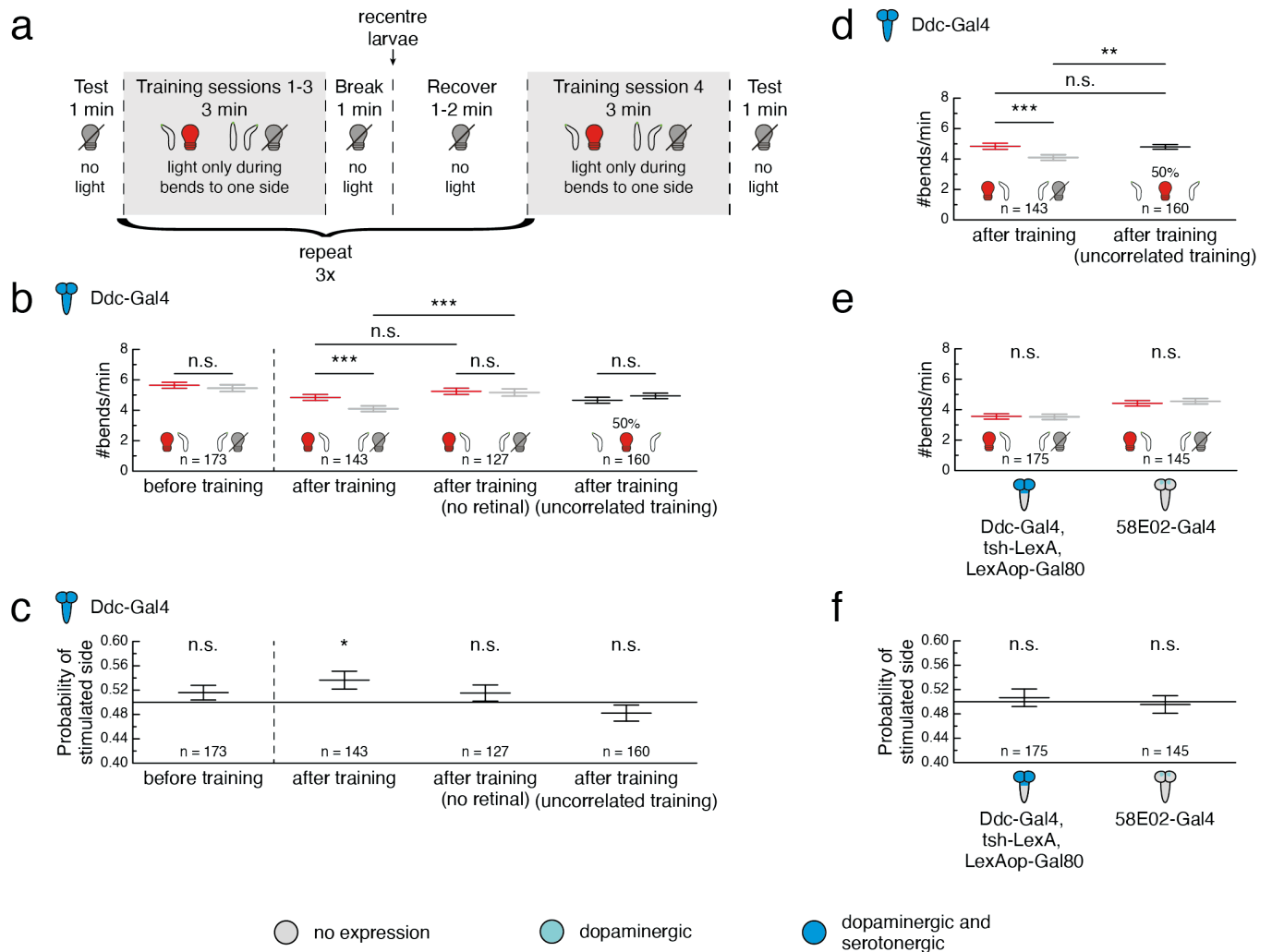


Figure 3. Operant conditioning of bend direction in *Drosophila* larvae requires the ventral nerve cord. **a.** Experiment protocol using the high-throughput closed-loop tracker. Behaviours are depicted as larval contours (black) with head (green). During training, the larva received an optogenetic stimulus (red light bulb) whenever it bent to one predefined side (here depicted as the left for simplicity), and light was switched off during all other behaviours (grey light bulb). **b,d,e.** Larval bend rate shown as the number of bends per minute, grouped by bend direction. The bend rate to the stimulated side (depicted as a left bend with a red light bulb for simplicity) is shown in red and the bend rate to the unstimulated side (depicted as a right bend with a grey light bulb for simplicity) is shown in grey. For larvae that received random, uncorrelated stimulation during 50% of bends, the bend rates to the left and right are shown in black. Statistical differences within groups were tested with a two-sided Wilcoxon signed-rank test; statistical differences between two groups were tested with a two-sided Mann-Whitney *U* test. **c,f.** Probability that a given bend was directed towards the stimulated side or, in the case of the uncorrelated training group, towards the left. Grey line indicates equal probability of 0.5 for bends to either side. Statistics calculated from a two-sided Wilcoxon signed-rank test. **b-f.** Gal4 expression depicted as color-coded CNS. All data is shown as (mean \pm s. e. m.). n. s. $p \geq 0.05$ (not significant), * $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$. **b.** Bend rate for *Ddc-Gal4* \times *UAS-CsChrimson* larvae. Data is shown from the test period before the first training session and the test period after the fourth training session. **c.** Data from same experiments as in **b.** **d.** Same data as in **b**, but bend rate for uncorrelated training group was calculated without stratification by bend direction. **e.** Bend rate for *Ddc-Gal4* \times *UAS-CsChrimson*; *tsh-LexA*, *LexAop-Gal80* and *58E02-Gal4* \times *UAS-CsChrimson* larvae. The effector, *UAS-CsChrimson*, is omitted from the figure for visual clarity. Data is shown from the test period immediately following the fourth training session. **f.** Data from same experiments as in **e.**

Figure 3-Figure supplement 1. *Ddc-Gal4* expression pattern without and with *tsh-Gal80* restriction.

240 For each larva, two measures served as a read-out for bend direction preference: i) the bend
241 rate, measured as the number of bends per minute performed towards a given side, and ii) the
242 probability that a given bend was directed towards the stimulated side, obtained by normalising
243 the bend rate with the total number of bends performed by the larva in that minute. Individual
244 larva variation in bend rate yielded different results for these measures at the population level. In
245 the one-minute test prior to the first training session, we observed no significant difference in larval
246 bend rate to either side and the likelihood of these naïve animals choosing one side over the other
247 was not significantly different from chance. In the one-minute test following the fourth training
248 session, larvae showed a preference for bends towards the side paired with red light stimulation
249 during training, and the probability of these larvae bending towards this previously stimulated side
250 was significantly greater than 50% (**Figure 3B**).

251 The light-dependent activation of neurons using *CsChrimson* requires a cofactor, retinal, which
252 we supplemented in the food during development (*Klapoetke et al., 2014*; see Materials and meth-
253 ods). A control group of larvae raised on food without retinal showed no significant difference
254 in absolute bend rate (**Figure 3B**) or bend direction probability (**Figure 3C**) throughout the exper-
255 iment. This suggested that the US, which triggered a learned direction preference for bends in
256 larvae raised on retinal, was indeed the collective activation of all *Ddc* neurons and not the red
257 light itself. Notably, when directly comparing larvae raised with retinal to this control group raised
258 without, the two groups showed no significant difference in the bend rate towards the stimulated
259 side. Instead, the bend rate towards the unstimulated side was significantly reduced in larvae that
260 received paired training compared to this control (**Figure 3B**). This raised the question whether
261 larvae were learning to prefer the side paired with the rewarding US, or rather to avoid the side
262 without the stimulus.

263 To confirm that the bend preference we observed after training was attributable to pairing
264 light with bends solely in one direction, we conducted another control experiment in which lar-
265 vae received random, uncorrelated stimulation during 50% of bends regardless of direction. After
266 training, larvae showed neither a difference in absolute left and right bend rates, nor a significant
267 probability of choosing one side over the other (**Figure 3B, Figure 3C**). These bend rates aver-
268 aged together were indistinguishable from those of pair-trained larvae as they bent to the previ-
269 ously stimulated side. However, larvae which received uncorrelated training showed a significantly
270 higher bend rate overall compared to pair-trained larvae bending to the previously unstimulated
271 side (**Figure 3D**).

272 **The mushroom body is not sufficient to mediate operant conditioning in larvae**

273 Our experiments showed that activation of *Ddc* neurons is a sufficient US for operant conditioning.
274 While we did not identify which individual neurons mediate the observed effect, we hypothesised
275 that not all *Ddc* neurons are involved. Some prior work in adult flies suggests that the MB is involved
276 in operant conditioning (*Sun et al., 2020*), while other studies in the adult suggest that operant con-
277 ditioning does not require the MB (*Booker and Quinn, 1981; Wolf et al., 1998; Colomb and Brembs,*
278 **2010, 2016**) and may instead involve motor neuron plasticity (*Colomb and Brembs, 2016*). The ex-
279 tent to which the MB is dispensable in larval operant conditioning is unknown. We investigated
280 whether smaller subsets of *Ddc* neurons in the brain and subesophageal zone (SEZ) could support
281 memory formation in our bend direction paradigm.

282 GAL80 under control of the *tsh* promoter suppresses GAL4 expression in the ventral nerve cord
283 (VNC), but not in the brain or SEZ (*Clyne and Miesenböck, 2008; Figure 3–Figure Supplement 1*).
284 When trained under our operant conditioning protocol (**Figure 3A**), *Ddc-Gal4 x UAS-CsChrimson; tsh-*
285 *LexA, LexAop-Gal80* larvae were equally likely to bend towards the side where they had previously
286 received the optogenetic stimulus as they were to bend towards the unstimulated side (**Figure 3E,**
287 **Figure 3F**). Activating these neurons was thus an insufficient rewarding US in this paradigm. The
288 loss of the operant conditioning effect we observed with *Ddc-Gal4 x UAS-CsChrimson* larvae high-
289 lighted the necessity of dopaminergic or serotonergic neurons in the VNC for the formation of a

290 bend direction preference. Their sufficiency was inconclusive, however, since perhaps two or more
291 distinct groups of *Ddc* neurons needed collective activation in order to form a memory.

292 We then assessed whether exclusively activating the PAM cluster dopaminergic neurons inner-
293 vating the MB could induce operant conditioning, as is the case for classical conditioning. *58E02-*
294 *Gal4* drives expression in the majority of these neurons (*Rohwedder et al., 2016*). *58E02-Gal4 x UAS-*
295 *CsChrimson* larvae did not develop any direction preference for bends following training (*Figure 3E,*
296 *Figure 3F*). It is unsurprising that activation of these neurons alone could not act as a rewarding
297 US in this paradigm, given our finding that *Ddc* neurons in the brain and SEZ are insufficient. It is
298 remarkable, however, because it suggests that the neural circuits signalling reward in operant con-
299 ditioning differ from those of classical conditioning. Although it remains to be seen whether these
300 PAM cluster neurons contribute to memory formation by interacting with other *Ddc* neurons, these
301 results further supported the idea that operant conditioning in *Drosophila* may not be mediated
302 by the MB.

303 **Serotonergic neurons in brain and SEZ are a sufficient reward signal in classical** 304 **conditioning**

305 Pairing an action with activation of numerous dopaminergic and serotonergic neurons across the
306 CNS was sufficient to induce operant conditioning of bend direction preference. Furthermore, our
307 results indicated that the VNC subset of these neurons was essential for memory formation in the
308 paradigm. It was an open question, however, whether this learning was mediated by dopamine,
309 serotonin, or both. Dopamine and serotonin receptors are necessary for different classical con-
310 ditioning tasks in honeybees, suggesting that the two neurotransmitters may carry out separate
311 functions (*Wright et al., 2010*). We conducted a high-throughput classical conditioning screen of
312 sparser dopaminergic and serotonergic driver lines to identify US candidates for comparison with
313 our operant conditioning paradigm.

314 We expressed *CsChrimson* under the control of different GAL4 driver lines and tested whether
315 pairing optogenetic activation of these neurons (US) with odour presentation (CS) could induce
316 olfactory memory. Conditioning was performed using a similar procedure to those described in
317 *Gerber and Hendel (2006)*, *Saumweber et al. (2011)* and *Eschbach et al. (2020b)*. In the paired
318 group, larvae were exposed to alternating three-minute presentations of ethyl acetate with red
319 light and air with no light. To ensure that any observed effects were a result of learning rather
320 than innate odour preference or avoidance, an unpaired group was trained simultaneously with
321 reciprocal stimulus presentation (odour/dark, air/light). Following training, larvae in both groups
322 were tested on their preference for the odour in the absence of light (*Figure 4A*). All learning scores
323 were compared to a negative control containing no GAL4 driver, *w¹¹¹⁸ x UAS-CsChrimson*, which
324 did not exhibit a learning phenotype (*Figure 4B*). Consistent with prior study results (*Rohwedder*
325 *et al., 2016*; *Eichler et al., 2017*; *Almeida-Carvalho et al., 2017*), *58E02-Gal4 x UAS-CsChrimson* larvae
326 showed appetitive olfactory learning with a significantly higher performance index than *w¹¹¹⁸ x UAS-*
327 *CsChrimson* larvae and so were used as a positive control (*Figure 4B*). *Ddc-Gal4 x UAS-CsChrimson*
328 larvae exhibited appetitive memory comparable to *58E02-Gal4* ($p = 0.1304$, two-sided Mann-Whitney
329 *U* test); an unsurprising result since the *Ddc-Gal4* expression pattern includes the PAM cluster neu-
330 rons. Consistent with previous studies in the larva (*Schroll et al., 2006*) and adult (*Aso et al., 2012*;
331 *Claridge-Chang et al., 2009*; *Liu et al., 2012*), *TH-Gal4 x UAS-CsChrimson* larvae exhibited significant
332 aversive olfactory learning. *TH-Gal4* covers most dopaminergic neurons, excluding the PAM clus-
333 ter (*Rohwedder et al., 2016*). The effect we observed may be mediated by punishment-signalling
334 dopaminergic neurons that project to the MB vertical lobes (*Eschbach et al., 2020b*; *Selcho et al.,*
335 *2009*). Isolating the locus of this effect may prove challenging, given the dearth of larval driver lines
336 targeting dopaminergic neurons without MB innervation.

337 Serotonergic signalling is required for associative learning in both larval (*Huser et al., 2017*) and
338 adult (*Johnson et al., 2011*; *Sitaraman et al., 2012*) *Drosophila*. We tested *Trh-Gal4* and *Tph-Gal4*,
339 two driver lines that target the majority of serotonergic neurons and no dopaminergic neurons

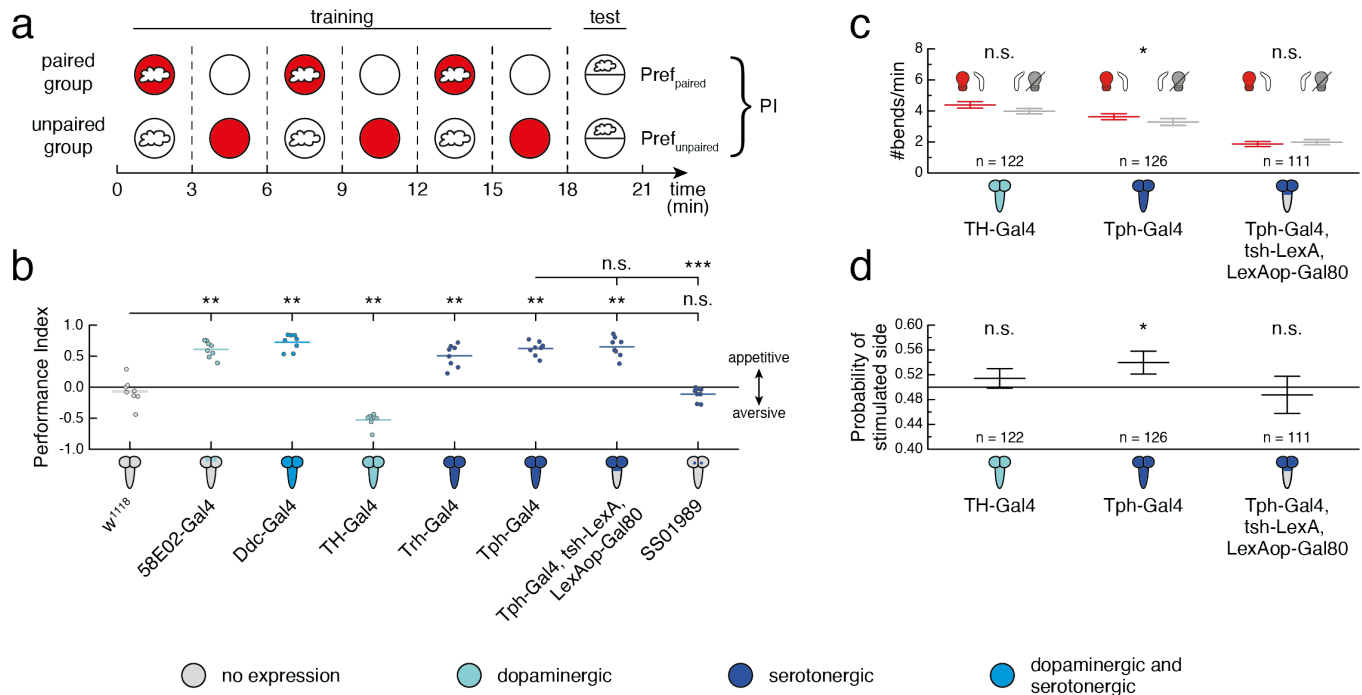


Figure 4. Different serotonergic neurons mediate classical and operant conditioning. All fly lines contained the *UAS-CsChrimson* effector, which is omitted from the figure for visual clarity. Gal4 expression depicted as color-coded CNS. **a.** Olfactory conditioning experiment protocol. During training, larvae in the paired group received three minutes of optogenetic red light stimulation (solid red circles) paired with the odour (white cloud) followed by three minutes of darkness (solid white circles) paired with air (no cloud). The unpaired group received reciprocal stimulus presentation (dark paired with odour, light paired with air). This procedure was repeated three times. In half of the experiments, the order of training trials was reversed, starting with air presentation instead of odour presentation. Both groups were then tested for learned odour preference in the dark with odour presented on one side of the plate and no odour on the other (PI = performance index). **b.** Performance indices following olfactory conditioning, plotted as raw data points and mean. *w¹¹¹⁸* *x UAS-CsChrimson* was the negative control (grey, $n = 8$), *58E02-Gal4 x UAS-CsChrimson* was the positive control (blue, $n = 8$). Statistical comparisons to *w¹¹¹⁸* *x UAS-CsChrimson* were calculated using a two-sided Mann-Whitney *U* test with Bonferroni correction; n.s. $p \geq 0.05/7$ (not significant), ** $p < 0.01/7$. Statistical comparisons to *Tph-Gal4 x UAS-CsChrimson* were calculated using a two-sided Mann-Whitney *U* test with Bonferroni correction; n.s. $p \geq 0.05/2$ (not significant), *** $p < 0.001/2$. **c,d.** All data is shown as (mean \pm s. e. m.), n.s. $p \geq 0.05$ (not significant), * $p < 0.05$. **c.** Experiments followed the protocol depicted in **Figure 3A**. Data is shown from the test period immediately following the fourth training session. Larval bend rate shown as the number of bends per minute, grouped by bend direction. The bend rate to the stimulated side (depicted as a left bend with a red light bulb for simplicity) is shown in red and the bend rate to the unstimulated side (depicted as a right bend with a grey light bulb for simplicity) is shown in grey. Statistical differences within a group were tested with a two-sided Wilcoxon signed-rank test. **d.** Probability that a given bend is directed towards the stimulated side. Grey line indicates equal probability of 0.5 for bends to either side. Statistics were based on a two-sided Wilcoxon signed-rank test.

Figure 4-Figure supplement 1. *Tph-Gal4* expression pattern without and with *tsh-Gal80* restriction.

Figure 4-Figure supplement 2. SS01989 exclusively drives expression in the CSD neuron.

Figure 4-Figure supplement 3. Paired and unpaired group data for olfactory conditioning experiments.

340 across the CNS of third-instar larvae (*Huser et al., 2012*). Consistent with previous reports (*Gan-*
341 *guly et al., 2020*), larvae expressing *CsChrimson* under either driver line formed strong appetitive
342 olfactory memory, highlighting the sufficiency of serotonin as a US in associative learning. *Tph-Gal4*
343 targets fewer serotonergic neurons than *Trh-Gal4*, making it valuable for narrowing down which
344 serotonergic neurons serve as a relevant reward signal. We eliminated all *Tph-Gal4* expression in
345 the VNC using *tsh-Gal80* (**Figure 4–Figure Supplement 1**). Activating the remaining *Tph* neurons in
346 the brain and SEZ was sufficient to induce strong appetitive memory (**Figure 4B**). This result was
347 notable and raised further questions: are serotonergic neurons in the brain and SEZ indirectly
348 connected to MB-innervating dopaminergic neurons or do alternative learning circuits exist that
349 altogether bypass the MB?

350 The contralaterally projecting serotonin-immunoreactive deutocerebral (CSD) neuron (*Roy et al.,*
351 *2007*) is one previously described serotonergic brain neuron within the *Tph-Gal4* expression pattern
352 (*Huser et al., 2012*) that innervates the antennal lobe and only has a few indirect connections to
353 the MB (*Berck et al., 2016*). Combining anatomical features from existing EM reconstruction (*Berck*
354 *et al., 2016*) with available lineage information facilitated identification of a split-GAL4 line (*SS01989*)
355 that drives expression exclusively in the CSD neuron (**Figure 4–Figure Supplement 2**). Pairing acti-
356 vation of *SS01989* with ethyl acetate was insufficient for inducing olfactory memory (**Figure 4B**), sug-
357 gesting that the classical conditioning phenotype we observed under *Tph-Gal4 x UAS-CsChrimson;*
358 *tsh-LexA, LexAop-Gal80* was mediated by at least one other group of serotonergic neurons in the
359 brain or SEZ.

360 **Serotonergic neurons in VNC are necessary for operant conditioning of bend direc-** 361 **tion**

362 Given their strong associative learning phenotypes, we used the *TH-Gal4* and *Tph-Gal4* drivers
363 to investigate whether operant conditioning of bend direction could be induced exclusively by
364 dopaminergic or serotonergic neurons, respectively. Under our high-throughput training proto-
365 col (**Figure 3A**), *TH-Gal4 x UAS-CsChrimson* larvae showed no difference in bend rate between the
366 previously stimulated and unstimulated sides in the one-minute test period (**Figure 4C**). Further-
367 more, the probability that any given bend was directed towards the previously stimulated side was
368 not significantly different from chance (**Figure 4D**). Activating these dopaminergic neurons was an
369 insufficient substitute for reward or punishment in operant conditioning.

370 Paired activation of *Tph-Gal4* neurons during bends to one side resulted in a significantly higher
371 bend rate to the stimulated side relative to the unstimulated side during the test period (**Figure 4C**).
372 The probability of bending in the previously stimulated direction was also significantly elevated
373 (**Figure 4D**). In this way, activation of *Tph*-positive serotonergic neurons paired with bends to one
374 side was sufficient for the formation of a learned direction preference. Combining this result with
375 the knowledge that operant conditioning was impaired following restriction of *Ddc-Gal4 x UAS-*
376 *CsChrimson* expression to the brain and SEZ suggests that the serotonergic neurons of the VNC
377 were necessary for memory formation in this paradigm. Because *Tph-Gal4* is a broad driver line, it
378 is possible that its expression pattern contains brain or SEZ neurons outside of those in *Ddc-Gal4*.
379 The existence of these neurons could have potentially induced learning through an alternate mech-
380 anism independent from that which drove memory formation following *Ddc* neuron activation.

381 To assess whether the VNC serotonergic neurons were necessary for the observed operant
382 conditioning effect, we used *tsh-Gal80* to restrict the *Tph-Gal4* expression pattern to the brain and
383 SEZ. Paired optogenetic activation of *Tph-Gal4 x UAS-CsChrimson; tsh-LexA, LexAop-Gal80* with larval
384 bends to one side was insufficient for operant conditioning (**Figure 4C, Figure 4D**). The *Tph-Gal4*
385 expression pattern contains two neurons per VNC hemisegment (with the exception of a single
386 neuron in each A8 abdominal hemisegment), all of which are serotonergic (*Huser et al., 2012*).
387 While there are few serotonergic VNC candidates, we could not conclude from our data whether
388 the operant conditioning effect relied solely on these neurons or whether synergistic activity from
389 both the VNC and the brain or the SEZ was needed. Testing these hypotheses remains challenging

390 since, to our knowledge, no sparse driver lines exist to exclusively target VNC serotonergic neurons.
391 Under a classical conditioning paradigm, we have confirmed that there exist learning pathways
392 in *Drosophila* that rely on serotonergic neurons. We have also shown that serotonergic neurons can
393 serve as a sufficient US for operant conditioning. Notably, different circuit mechanisms underlie
394 classical and operant conditioning mediated by serotonergic neurons: activation of the brain and
395 SEZ is sufficient for classical conditioning, whereas the VNC is necessary for operant conditioning.

396 Discussion

397 Due to available genetic tools and the emerging connectome, the *Drosophila* larva is a uniquely
398 advantageous model organism for neuroscience. We have uncovered a previously unknown neu-
399 ronal mechanism of operant conditioning in the *Drosophila* larva. Serotonergic signalling can be
400 employed as a reinforcing US in both classical and operant associative learning, but we are mind-
401 ful that a single neural mechanism for learning may not exist. Distinct types of learning may share
402 neurotransmitters or circuit components, but there may remain fundamental differences in con-
403 nectivity and function. The experimental system we built was instrumental in investigating the
404 neural circuits of operant conditioning, as it combined FPGA-based real-time tracking of multiple
405 larvae with robust online behaviour detection and closed-loop stimulus presentation. This system
406 could facilitate further research in taxis (Luo *et al.*, 2010; Gomez-Marin *et al.*, 2011; Kane *et al.*,
407 2013; Jovanic *et al.*, 2019), decision-making (Krajbich, 2019; DasGupta *et al.*, 2014), and spatial nav-
408 igation and memory (Neuser *et al.*, 2008; Haberkern *et al.*, 2019). While further work is necessary,
409 our bend direction paradigm provides a strong foundation for continued study of the circuit and
410 cellular mechanisms underlying operant conditioning.

411 High-throughput operant conditioning in *Drosophila* larvae

412 We have shown that *Drosophila* larvae are capable of operant conditioning and that optogenetic
413 activation of *Ddc* neurons serves as a rewarding US during this learning process. With training,
414 larvae formed an association between the presence or absence of this US and the direction in which
415 they were bending. During testing, in the absence of any stimulation, larvae showed a significant
416 learned preference for bending towards the previously stimulated side. This learned modification
417 of future behaviour was only observed when the CS and US were paired during training; a hallmark
418 of operant conditioning. Because *Ddc-Gal4* drives expression in dopaminergic and serotonergic
419 neurons (Li *et al.*, 2000; Sitaraman *et al.*, 2008), we concluded that one or both neurotransmitters
420 are involved in memory formation under these experiment conditions.

421 Strong parallels exist between our operant learning paradigm and those employed for condi-
422 tioning direction preference in adult *Drosophila*. Consider the work of Nuwal *et al.* (2012), in which
423 tethered flies walked on a rotating ball and were rewarded with optogenetic activation of sugar-
424 sensing neurons upon turns to one direction. As a consequence of this pairing, the flies learned
425 to increase the number of turns to this side. Notably, the nature of the US remains an important
426 difference between these paradigms. Our initial attempts to operantly condition larvae using ac-
427 tivation of sugar-sensing neurons as a rewarding US were unsuccessful. These neurons, defined
428 by two different *Gr43a-Gal4* drivers, were also insufficient for memory formation when activated
429 with a paired odour in an olfactory conditioning screen. This was surprising, considering extensive
430 evidence that natural sugar can serve as a rewarding US for classical conditioning in larvae (Apos-
431 tolopoulou *et al.*, 2013; Schleyer *et al.*, 2015; Weiglein *et al.*, 2019; Honjo and Furukubo-Tokunaga,
432 2005; Neuser *et al.*, 2005; Rohwedder *et al.*, 2012; Scherer *et al.*, 2003; Schipanski *et al.*, 2008).
433 One possible explanation for these discrepancies is that multiple groups of sensory neurons must
434 be co-activated in order to relay a meaningful reward signal. Alternatively, it may be necessary to
435 adjust the temporal pattern or intensity of optogenetic stimulation.

436 It remains to be seen whether operant learning can occur by pairing roll or back-up behaviour
437 with reward or punishment. Conditioning these actions is challenging given their infrequency in
438 naïve, freely behaving animals. Rolls only occur in response to noxious stimuli (Ohyama *et al.*, 2013,

439 **2015; Robertson et al., 2013; Tracey et al., 2003**). Back-ups also occur at very low rates (**Masson**
440 **et al., 2020**). Consequently, the amount of US which larvae would receive during paired training
441 would be very small, making observable memory formation more difficult. Our high-throughput
442 tracker could potentially address this challenge with probabilistic, thermogenetic activation of roll
443 or back-up command neurons (**Ohyama et al., 2015; Carreira-Rosario et al., 2018**) and optogenetic
444 reward when performing the desired action.

445 **Neural circuits of operant conditioning**

446 From the available data, it cannot be concluded that the brain and SEZ are dispensable for operant
447 conditioning in *Drosophila* larvae. Examples from both vertebrates (**Grau et al., 1998**) and
448 invertebrates (**Horridge, 1962; Booker and Quinn, 1981**) show the spinal cord or VNC as sufficient
449 for learning, suggesting that conserved mechanisms exist for brain-independent operant condi-
450 tioning across species. This does not, however, exclude the possibility that there exist alternative
451 learning pathways using the brain. In mammals (**Redgrave et al., 2011; Balleine et al., 2009**) and
452 birds (**Fee and Goldberg, 2011**), brain correlates of operant conditioning have been identified. It is
453 unclear where such pathways would be located in the insect brain. Both our larval experiments
454 and previous adult fly studies (**Colomb and Brembs, 2016; Wolf et al., 1998; Colomb and Brembs,**
455 **2010; Booker and Quinn, 1981**) support the idea that operant conditioning can occur independently
456 of the MB, such that other learning centres might exist. To determine whether larval operant con-
457 ditioning can be fully mediated by the VNC or whether the brain or SEZ are necessary, new driver
458 lines must be created. A collection of sparse split-GAL4 lines, each specific to a distinct group of
459 serotonergic neurons, could help identify the minimum subset of neurons necessary for conveying
460 a US in our bend direction paradigm.

461 Even if the learning signal for operant conditioning can be mapped to a few serotonergic neu-
462 rons, there remain several open questions regarding the neuronal mechanisms underlying this
463 learning. Locally, neurons could drive synaptic plasticity or modulate the intrinsic excitability of
464 their postsynaptic partners. Alternatively, the learning signal could propagate further downstream,
465 yielding learning correlates elsewhere in the network. Furthermore, memory formation requires
466 integrating the US with information about the occurrence of the reinforced action. Motor feedback
467 (e. g. efference copy, **Webb, 2004; Fee, 2014**) or proprioceptive input could be used to transmit this
468 movement signal to higher-level circuits for convergence with the valence-encoding US. However,
469 if memory formation occurred at a lower level, the action-specific signal and associated valence
470 could be locally integrated inside the motor or premotor neuron without the need for feedback
471 loops.

472 **Lorenzetti et al. (2008)** proposed intracellular mechanisms for modulating the intrinsic excitabil-
473 ity of the *Aplysia* premotor neuron B51 during operant conditioning, mediated by the highly con-
474 served protein kinase C (PKC) gene. PKC signalling is also essential for operant conditioning in *Lym-*
475 *naea* (**Rosenegger and Lukowiak, 2010**) and adult *Drosophila* (**Brembs and Plendl, 2008; Colomb**
476 **and Brembs, 2016**). If the *Drosophila* larva showed evidence of PKC-induced motor neuron plas-
477 ticity, EM reconstruction of the pathways between the serotonergic neurons of the VNC and the
478 PKC-positive motor neurons could further elucidate the mechanisms of memory formation and
479 retrieval. A similar investigation of the *Drosophila* gene *FoxP* may also be informative, as its muta-
480 tion in the adult results in impaired operant self-learning (**Mendoza et al., 2014**). The vertebrate
481 homologue *FOXP2* is associated with deficits in human speech acquisition (**Lai et al., 2001**), song
482 learning in birds (**Haesler et al., 2007**), and motor learning in mice (**Groszer et al., 2008**).

483 **Serotonin as a learning signal**

484 A limited set of studies have shown that serotonergic signalling is sufficient to induce learning (**Liu**
485 **et al., 2014; Hawkins and Byrne, 2015; Ganguly et al., 2020**). Previous *Drosophila* studies highlight
486 other roles of serotonin in associative learning (**Yu et al., 2005; Keene et al., 2004, 2006; Lee et al.,**
487 **2011; Huser et al., 2017**). **Sitaraman et al. (2012)** have shown that synaptic transmission from sero-

488 tonergic neurons is essential for appetitive olfactory conditioning in the adult. Aversive olfactory
489 memory formation is impaired in flies fed with a tryptophan hydroxylase inhibitor which blocks
490 serotonin biosynthesis (*Lee et al., 2011*). Furthermore, serotonin receptor signalling is required
491 for memory formation in classical conditioning tasks (*Johnson et al., 2011*). In larvae, aversive ol-
492 factory conditioning is impaired by either ablation of serotonergic neurons during development or
493 mutations in a serotonin receptor gene (*Huser et al., 2017*).

494 Our work suggests a novel role of serotonin as a reward signal for learning in *Drosophila* larvae.
495 In our olfactory classical conditioning screen, optogenetic stimulation of serotonergic neurons in
496 the brain and SEZ was sufficient to induce strong appetitive learning. Conversely, operant con-
497 ditioning necessitated serotonergic neuron activity in the VNC. Since it remains unclear to what
498 extent serotonergic neurons in the brain and SEZ are also involved in the operant conditioning
499 effect we observed, it is possible that some neurons mediate both forms of associative learning.

500 Further investigation is necessary to better understand the function of serotonin in memory
501 formation. It is possible that even a single instance of learning leads to a variety of changes across
502 the nervous system. In the case of operant conditioning, higher brain centres, motor command
503 neurons, premotor circuits and motor neurons would all qualify as potential learning sites. In addi-
504 tion to thoroughly analysing the expression patterns of driver lines used in our classical condition-
505 ing screen, developing new, sparse driver lines targeting serotonergic neurons would be valuable
506 for identifying the minimal subset of neurons which provide the serotonergic learning signal. The
507 larval connectome could be used to subsequently trace the paths from these neurons to the MB.
508 One could then test whether learning as induced by the serotonergic US remains intact when these
509 connections are silenced. The expression pattern of serotonin receptors could also provide clues
510 about how the serotonergic signal triggers learning. One should certainly consider the possibil-
511 ity that learning is not induced by serotonin itself, but by other neurotransmitters coexpressed by
512 serotonergic neurons. This could be assessed by suppressing serotonin biosynthesis in the desired
513 neuronal subset.

514 **Materials and methods**

515 **High-throughput closed-loop tracker**

516 Hardware set-up

517 A high-resolution camera (3072 x 3200 pixels) (#TEL-G3-CM10-M5105, Teledyne DALSA, Ontario,
518 Canada) positioned above a 23 cm x 23 cm 4% agarose plate captured 8-bit greyscale images at
519 20 Hz. The agarose plate was illuminated from below by a 30 cm x 30 cm 850 nm LED backlight
520 (#SOBL-300x300-850, Smart Vision Lights, Norton Shores, Michigan) equipped with intensity con-
521 trol (#IVP-C1, Smart Vision Lights, Norton Shores, Michigan). An 800 nm longpass filter (#LP800-
522 40.5, Midwest Optical Systems, Palatine, Illinois) mounted on the camera blocked all visible wave-
523 lengths, including those used for optogenetics. When the agarose plate comprised most of the
524 camera image, each pixel corresponded to a 72.92 μm diameter section of the plate.

525 Each camera image was processed in parallel on both the host computer (#T7920, running
526 Windows 10, Dell Technologies Inc, Round Rock, Texas) and an FPGA device (#PCIe-1473R-LX110,
527 National Instruments, Austin, Texas). LabVIEW 2017 (National Instruments, Austin, Texas) software
528 extracted larval contours and interfaced with C++ software that performed real-time behaviour de-
529 tection. The LabVIEW software controlled closed-loop optogenetic and thermogenetic stimulation
530 in response to these detected behaviours.

531 Multi-animal detection and tracking

532 Raw camera images were read by the FPGA at 20 Hz and then sent to the host computer. The Lab-
533 VIEW process on the host computer then filtered out non-larval objects by combining background
534 subtraction and binary thresholding. The remaining objects were each enclosed in a rectangular
535 box of minimal size, with edges parallel to the camera image axes. We defined the following criteria
536 to detect third-instar larvae within these boxes:

- 537 • Pixel intensity range (default 25–170): the minimum and maximum brightness values for
- 538 pixels selected by binary thresholding (between 0 and 255 for an 8-bit image).
- 539 • Box side length (pixels) (default 6–100): the range of eligible values for width and height of
- 540 each box.
- 541 • Box width + height (pixels) (default 12–200): the range of eligible values for the sum of each
- 542 box's width and height.
- 543 • Box area (pixels) (default 300–900): the range of eligible values for the area of each box.

544 To track larvae over time, the host computer assigned a numerical identifier to each eligible object.
545 We used distance-based tracking with a hard threshold of 40 pixels to maintain larval ID based
546 on `centroid` position. Although identity was lost when larvae touched or reached the plate's edge,
547 new IDs were generated when larvae matched detection criteria. For each of the largest 16 objects,
548 the host computer sent a binary pixel pattern and location (defined as the centre of the box) to the
549 FPGA. Since the host computer required more than 50 ms of run time for object detection, this
550 process was not executed in every frame. On average, the FPGA received updated objects and
551 their locations every three frames.

552 The FPGA extracted object contours in three steps. Within a 2 cm² region of interest around the
553 object's centre, the FPGA first applied a user-defined binary threshold, then applied both vertical
554 and a horizontal convolution with a 2 x 1 XOR kernel, and finally generated edge pixels by com-
555 bining the results of the two convolutions using an OR operation. Contours were extracted from
556 edge images using the Moore boundary tracing algorithm (*Gonzalez and Woods, 2018*) with three
557 added error capture procedures. First, if the algorithm yielded a contour that ended prematurely
558 or contained small loops, the construction process could be reversed by up to 16 contour points to
559 find an alternative contour. Second, 10,000 FPGA clock cycles (≈ 100 us) was the maximum allotted
560 execution time, with each pixel comparison occurring within one clock cycle. In the rare event that
561 this window was exceeded, the algorithm returned the already constructed contour points. Third,
562 a contour containing fewer than 63 points was rejected and the FPGA returned the last valid con-
563 tour detected for a given larva ID. The algorithm stopped when none of the remaining neighbours
564 were edge pixels (*Figure 1–Figure Supplement 1*).

565 Contour processing and landmark detection

566 An undesired result of the FPGA contouring algorithm was the variable number of contour points
567 across larvae and frames. We aimed to detect behaviour based on a smooth contour with a fixed
568 number of 100 contour points. This contour regularization was achieved inside the Behaviour Pro-
569 gramme using Fourier decomposition and reconstruction as in *Masson et al. (2020)*.

570 The initial detection of head and tail was implemented on FPGA. The larva's head and tail were
571 defined as the contour points with the sharpest and second-sharpest curvature, respectively (*Fig-
572 ure 1–Figure Supplement 2*). While correct in most cases, this calculation sometimes led to flipped
573 detection of the two body ends. The Behaviour Programme flagged and corrected these false de-
574 tection events at run time by calculating the distance head and tail traveled between frames and
575 tracking the number of correct versus flipped detection events. The vote system correction com-
576 monly failed when the larva made large angle bends. The resulting contour was nearly-circular and
577 exhibited similar curvature across all points. The solution required resetting the vote tallies when
578 detecting these ball events (*Figure 1–Figure Supplement 2*).

579 We defined the larval spine as 11 points running along the central body axis from `head` to `tail`
580 (*Figure 1–Figure Supplement 3; Swierczek et al., 2011*). In addition to `head` and `tail`, the Behaviour
581 Programme calculated three equally distributed landmark points along the spine (`neck_top`, `neck`,
582 and `neck_down`). A fourth landmark, the `centroid`, defined the larva's location. The six landmarks
583 were collectively used to extract features for training behaviour classifiers (*Figure 1–Figure Supple-
584 ment 3*).

585 The Behaviour Programme transformed the raw contour and spine from camera coordinates

586 (in pixels) to world coordinates (in mm). If stable larval detection criteria were met, all spine points
587 were temporally smoothed using exponential smoothing (*Figure 1–Figure Supplement 3*).

588 Feature extraction

589 We developed a machine learning approach to address the high deformability of the larva shape,
590 ensure live execution, reduce overfitting, and limit the volume of data tagging. What follows is a
591 brief summary of larval features describing motion direction, body shape, and velocity that were
592 calculated from the contour and spine data inside the Behaviour Programme. Features were de-
593 signed as in *Masson et al. (2020)*, with notable modifications required to run the inference live:

594 1. Motion Direction (*Figure 1–Figure Supplement 4*)

- 595 • `direction_vector`: normalised vector describing the main body axis
- 596 • `direction_head_vector`: normalised vector describing the head axis
- 597 • `direction_tail_vector`: normalised vector describing the tail axis

598 2. Body Shape (*Figure 1–Figure Supplement 5*)

- 599 • `skeleton_length`: summed distances between consecutive spine points
- 600 • `perimeter`: summed distances between neighbouring contour points
- 601 • `larva_arc_ratio`: ratio of contour perimeter to convex hull perimeter (`larva_arc_ratio` \geq
602 1 and was close to 1 when larva was in either straight or ball-like shape)
- 603 • `larva_area_ratio`: ratio of the areas enclosed by the contour and its convex hull ($0 \leq$
604 `larva_area_ratio` ≤ 1 and was close to 1 when the larva was in either straight, heavily
605 curved, or ball-like shape)
- 606 • `eig_reduced`: $\text{eig_reduced} = \frac{|\lambda_1 - \lambda_2|}{\lambda_1 + \lambda_2}$ where λ_1, λ_2 were the eigenvalues of the structure
607 tensor of the larval contour with respect to the neck ($0 \leq \text{eig_reduced} \leq 1$ and `eig_-`
608 `reduced` decreased as the bend amplitude of the larva increased)
- 609 • `s`: normalised angle along the body ($-0.5 \leq s \leq 1$, was close to 1 when larva was straight,
610 and decreased with increasing bend amplitude)
- 611 • `asymmetry`: sine of the angle between `direction_vector` and `direction_head_vector`
612 (`asymmetry` > 0 when larva bent left and `asymmetry` < 0 when larva bent right)
- 613 • `angle_upper_lower`: absolute angle between `direction_vector` and `direction_head_-`
614 `vector` (despite similarity to `asymmetry`, this develops different dynamics following tem-
615 poral smoothing, which are valuable for stable left and right bend detection)

616 3. Velocity (*Figure 1–Figure Supplement 6*)

- 617 • Velocity of all six landmark points (`head_speed`, `neck_top_speed`, `neck_speed`, `neck_down_-`
618 `speed`, `tail_speed`, and `v_centroid`) in mm/s over interval $dt = 0.2\text{ s}$ (four frames)
- 619 • `v_norm`: arithmetic mean of `neck_top_speed`, `neck_speed`, and `neck_down_speed`, passed
620 through a hyperbolic tangent activation function to suppress excessively large values
- 621 • `speed_reduced`: relative contribution of `neck_top_speed` to `v_norm`, passed through a
622 hyperbolic tangent activation function to suppress excessively large values (`speed_re-`
623 `duced` increased when the anterior larval body moved quickly compared to the posterior,
624 e. g. when a bend was initiated)
- 625 • `damped_distance`: distance (mm) travelled by neck, giving greater weight to recent over
626 past events
- 627 • `crab_speed`: lateral velocity (mm/s), defined as the component of `neck_speed` orthogonal
628 to `direction_vector_filtered`
- 629 • `parallel_speed`: forward velocity (mm/s), defined as the component of `neck_speed_-`
630 `filtered` parallel to `direction_vector_filtered`
- 631 • `parallel_speed_tail_raw`: tail's forward velocity (mm/s), defined as the component of
632 `tail_speed_filtered` parallel to `direction_tail_vector_filtered`

633 • `parallel_speed_tail`: similar to `parallel_speed_tail_raw`, with the difference that `tail_`
634 `speed_filtered` was normalised prior to calculating the dot product (i. e. a measure of
635 tail movement direction which took values between -1 (backward) and +1 (forward))

To extract features in real time and address various sources of noise, we implemented exponential smoothing defined as follows for a given feature f (**Figure 1–Figure Supplement 7**):

$$f_{\text{filtered},t} = (1 - \alpha) \cdot f_{\text{filtered},t-\Delta t} + \alpha \cdot f_t$$

636 where t is unitless, but derived from the experiment time in seconds, $\alpha = \frac{\Delta t}{\tau}$ with $\Delta t = 0.05$ s and
637 $\tau = 0.25$ s. Features that had the potential to exhibit large value deviations (e. g. `v_norm`) were
638 instead bounded using a hyperbolic tangent function. Additionally, some features were exponentially
639 smoothed over a longer time window (where $\alpha_{\text{long}} = \frac{\Delta t}{\tau_{\text{long}}}$ with $\Delta t = 0.05$ s and $\tau_{\text{long}} = 5$ s)
640 (**Figure 1–Figure Supplement 7**).

Convolution was used to approximate a smoothed squared derivative for each feature (**Figure 1–Figure Supplement 8**); useful for integrating information over time without needing to further expand the feature space. The underlying mathematical concepts were motivated by **Masson et al. (2012)**. For a given feature f at time t , `f_convolved_squared` was calculated as follows:

$$f1_t = (1 - \lambda \Delta t) \cdot f1_{t-\Delta t} + \frac{1}{2} \Delta t \cdot (f_{t-\Delta t} + f_t)$$

$$f2_t = \lambda \Delta t \cdot f1_{t-\Delta t} + (1 - \lambda \Delta t) \cdot f2_{t-n\Delta t}$$

$$f_{\text{convolved_squared},t} = k \cdot (f1_t - f2_t)^2,$$

641 where $\Delta t = 0.05$ s, $\lambda = \frac{1}{\tau}$, $\tau = 0.25$ s, and $n = 5$ s. k values were empirically chosen for each feature.

642 Behaviour classifiers

643 Behaviour classifiers were developed using a user interface similar to JAABA (**Kabra et al., 2013**).
644 The underlying algorithms combined trained neural networks and empirically determined linear
645 thresholds. We developed a MATLAB (MathWorks, Natick, Massachusetts) user interface with func-
646 tions for data visualisation, manual annotation, and machine learning using the Neural Network
647 Toolbox, the Deep Learning Toolbox, and the Statistics and Machine Learning Toolbox. Here we
648 briefly describe the behaviour classifiers and provide performance results based on manual vali-
649 dation (**Table 1**).

650 The `bend` classifier was based on predefined thresholds for temporally smoothed body shape
651 features and was itself exponentially smoothed over time. Independent `left` and `right` classifiers
652 were used to initially detect bend direction. To detect left and right bends, these classifiers were
653 combined with the smoothed `bend` classifier using an AND conjunction. The raw time series of left
654 and right bends was further smoothed post-acquisition using a custom MATLAB script: two bends
655 to the same side separated by less than 200 ms were combined into a single long bend, and short
656 bends of less than 200 ms were removed from analysis.

657 To improve `left` and `right` detection performance, we developed a classifier for circular larval
658 contours. This `ball` classifier used a feed-forward neural network with a single fully connected
659 hidden layer whose inputs were normalised values of `eig_reduced`, `larva_arc_ratio`, and `larva_`
660 `area_ratio`. The hidden layer consisted of five neurons with a hyperbolic tangent activation func-
661 tion. The output layer contained a single neuron and used a sigmoid activation function. The
662 neural network was trained in MATLAB on a manually annotated data set for 500 epochs using a
663 cross-entropy loss function and scaled conjugate gradient backpropagation. If a `ball` was detected
664 within the previous 1.5 s, `left` and `right` classifiers were overwritten to match the last detected
665 bend direction prior to the beginning of the `ball`.

666 The `back` classifier detected individual backward peristaltic waves based on thresholds for smoothed
667 tail velocity features combined with no `ball` detection within the previous 1.5 s.

668 Two different classifiers were used to detect crawling. `forward` detected longer forward crawl
669 periods based on thresholds for smoothed tail velocity features combined with no `ball` detection

Table 1. Manual quantification of behaviour detection performance.

back (268 events from 24 larvae in 60 minutes of video data)	
Precision	86.5%
Recall	88.4%
bend (714 events from 24 larvae in 60 minutes of video data)	
Precision	95.6%
Recall	96.4%
Accuracy of left and right detection (true-positive bends)	97.3%
forward (425 events from 24 larvae in 60 minutes of video data)	
Precision	97.8%
Recall	94.1%
forward_peristaltic (2954 events from 24 larvae in 60 minutes of video data)	
Precision	99.5%
Recall	93.6%
Events which are falsely combined with another event	10.7%
Events which are detected as more than one event	1.2%
roll (240 events from 24 larvae in 60 minutes of video data)	
Precision (rolls and roll-like events)	96.6%
Recall (rolls)	86.7%
Recall (roll-like events)	25.8%

670 within the previous 1.5 s. `forward_peristaltic` detected individual forward peristaltic waves based
671 on the `forward` classifier and a threshold on forward tail velocity.

672 The `roll` classifier was based on thresholds for body shape and velocity combined with no ball
673 detection and was exponentially smoothed over time. If a `roll` was detected within the previous
674 1.5 s, `forward`, `forward_peristaltic`, and `back` classifier values were reset to reduce false-positive
675 detection for these classifiers. Unusual behaviour patterns such as rapid bending or twitching
676 could be observed in addition to true larval rolling. These behaviours were considered "roll-like"
677 events during manual validation of the `roll` classifier's performance.

678 Optogenetic stimulation

679 Optogenetic stimulation was achieved using two digital micromirror devices DMDs to project light
680 patterns onto larvae on the agarose plate. During the hardware design process, two different DMD
681 models were tested. One contained an integrated 613 nm LED (#CEL-5500-LED, Digital Light Inno-
682 vations, Austin, Texas) and the other (#CEL-5500-FIBER, Digital Light Innovations, Austin, Texas)
683 received input from an external 625 nm LED (#BLS-GCS-0625-38-A0710, Mightex Systems, Ontario,
684 Canada) controlled by a BioLED light source control module (#BLS-13000-1, Mightex Systems, On-
685 tario, Canada) and fed through an optic fibre (#LLG-05-59-420-2000-1, Mightex Systems, Ontario,
686 Canada). Both DMDs operated like a 768 x 1024 pixel monochrome red light projector with nu-
687 merous rotatable micromirrors used to modulate the intensity of individual pixels. Although both
688 achieved similar light intensities, each DMD on its own was insufficient for optogenetic stimulation
689 of larvae. We installed both devices on the system such that their projections each covered the
690 entire agarose plate. In this way, the summed light intensities of the two DMDs could be achieved
691 at all locations. Accurately aiming light at crawling larvae required spatial calibration of each DMD.
692 Calibration was performed by projecting square spots at fixed DMD pixel locations and linearly
693 fitting the corresponding camera coordinates.

694 We determined that DMD illumination using the default light output was not uniform at plate
695 level, which could have resulted in variable optogenetic stimulation depending on larval location.
696 The maximum achievable light intensity at the plate's edge was approximately 40% of the peak
697 value at its centre. We therefore normalised the pixel intensity of the DMD image to the highest
698 intensity uniformly achievable at all plate locations. A look-up table containing the normalisation
699 factor for each DMD pixel was then calculated using bi-linear interpolation with approximately
700 100 light intensity values measured across the plate. To accommodate for possible differences in
701 non-uniformity between the two DMDs, this intensity calibration was performed for both DMDs
702 simultaneously following spatial calibration. When fully calibrated, the system could achieve a
703 uniform light intensity of 285 $\mu\text{W}/\text{cm}^2$.

704 A user-defined Behaviour Programme protocol operated on the behaviour detection output
705 and sent 8-bit optogenetic stimulation instructions to the LabVIEW application. Because the Lab-
706 VIEW application updated DMD projections at 20 Hz, the delay between behaviour detection and
707 closed-loop optogenetic stimulation of individual larvae did not exceed 50 ms. Furthermore, if two
708 or more larvae were close enough such that their corresponding stimulation areas overlapped, the
709 light intensity in the overlapping region was set to the smallest of those values to avoid undesired
710 stimulation.

711 Thermogenetic stimulation

712 Thermogenetic stimulation was achieved by heating up larvae with a custom IR laser set-up. A
713 1490 nm laser diode beam (#2CM-101, SemiNex, Peabody, Massachusetts) was fed into a two-
714 axis galvanometer system (#GVSM002, Thorlabs, Newton, New Jersey), both controlled by an ana-
715 logue output device (#PCIe-6738, National Instruments, Austin, Texas). Two mirrors inside the
716 galvanometer were rotated around orthogonal axes to target the beam spot to any user-defined
717 location on the agarose plate. The beam spot measured approximately 5 mm in diameter, de-
718 pending on the beam's angle of incidence to the plate. Mirror positions were controlled by two
719 integrated motors that received voltage inputs. Each voltage pair clearly defined the laser beam's
720 position.

721 Spatially calibrating the galvanometer was necessary to obtain a map between larval locations
722 in world coordinates and the mirror motor input voltages. A visible aiming beam was scanned
723 across the agarose plate using a fixed set of voltage pair inputs to the galvanometer. With the
724 optical filter removed from the camera, the aiming beam's location in camera coordinates was au-
725 tomatically extracted from the image using binary thresholding. Two voltage-to-camera look-up
726 tables were generated through bi-linear interpolation of these measured coordinates. For accu-
727 rately targeted thermogenetic stimulation, the location of the larval centroid was first converted
728 to camera coordinates using the existing world-to-camera transform and was then mapped to a
729 pair of galvanometer input voltages using the look-up tables.

730 Laser intensity calibration was also necessary to ensure that all larvae received the same stimu-
731 lation regardless of their position on the agarose plate. A larva's location changed the laser beam's
732 angle of incidence, causing the illuminated spot at plate level to take an elliptical shape with variable
733 size. Although laser beam power was constant, the changing spot area generated inconsistencies
734 in the amount of IR light covering each larva. Calibration was used to normalise the desired laser
735 intensity to achieve constant power per unit area. A visible aiming beam was scanned across the
736 plate and the camera image automatically measured the beam's spot size at various locations. Bi-
737 linear interpolation was then used to generate a pixel-wise look-up table containing the necessary
738 scaling factors for the laser power. At the location where the laser spot area was smallest, the
739 maximum power was reduced to 67.3%. We also accounted for a nonlinear relationship between
740 the laser source input voltage and the laser's total power output by generating a voltage-to-power
741 map from manual measurements. With these transformations, the system could calculate the
742 laser source input voltage necessary to produce uniform, 5.26 W stimulation at any location.

743 A user-defined Behaviour Programme protocol operated on the 20 Hz behaviour detection out-

744 put and sent thermogenetic stimulation instructions to the LabVIEW application which controlled
745 the galvanometer and laser. Four centroid locations were specified on every frame, enabling a
746 single galvanometer to cycle the laser beam between four individual larvae at 20 Hz. Within the
747 available 50 ms time window, each larva was heated for 11 ms. Switching off the laser input for
748 1.5 ms between larvae accounted for small time fluctuations surrounding each new galvanometer
749 position update and helped avoid undesired stimulation of other plate areas (**Figure 2D**). If fewer
750 than four objects were detected in a given frame, the remaining galvanometer target locations
751 were set to the plate's centre and the corresponding laser intensity was set to zero. This temporal
752 pattern of galvanometer position updates yielded no more than 100 ms delay between behaviour
753 detection and closed-loop thermogenetic stimulation.

754 Three parameters influenced larval temperature increase following thermogenetic stimulation
755 with the IR beam: i) the laser power, ii) the total duration of the stimulus, and iii) the order in which
756 the galvanometer cycles between locations in its 80 Hz movement. Preliminary experiments sug-
757 gested that these parameters could be adjusted to simultaneously stimulate eight or twelve larvae
758 using a single galvanometer. This could potentially eliminate the need to install three additional
759 laser sources to target all 16 larvae.

760 **Software availability**

761 All software code is available upon request.

762 **Fly strains and larval rearing**

763 We used the following fly strains: *58E02-Gal4* (Bloomington stock 41347), *69F06-Gal4* (Bloomington
764 stock 39497), *72F11-Gal4* (Bloomington stock 39786), *attP2* (**Pfeiffer et al., 2008**), *Ddc-Gal4* (**Li et al.,**
765 **2000**), *SS01989* (own stock), *TH-Gal4* (**Friggi-Grelin et al., 2003**), *Tph-Gal4* (**Park et al., 2006**), *Trh-Gal4*
766 (**Alekseyenko et al., 2010**), *UAS-CsChrimson* (Bloomington stock 55134), *UAS-CsChrimson; tsh-LexA,*
767 *LexAop-Gal80* (Dr Stefan Pulver, Dr Yoshinori Aso), *UAS-dTrpA1* (Dr Paul Garrity), *UAS-GFP* (**Nern et al.,**
768 **2015**), and *w¹¹¹⁸* (**Hazelrigg et al., 1984**).

769 Fly stocks were maintained in vials filled with standard cornmeal food (**Wirtz and Semey, 1982;**
770 49.2 ml of molasses, 19.9 g of yeast, 82.2 g of cornmeal, 7.4 g of agarose, 9.8 ml of 20% Tegosept
771 solution in 95% ethanol and 5.2 ml of propionic acid in 1 litre of water). For proof-of-principle
772 and operant and classical learning experiments, eggs were collected overnight for approximately
773 12–18 hours on standard cornmeal food plates with additional dry yeast to increase laying. These
774 experiments were performed using foraging-stage third-instar larvae (72–96 hours after egg laying)
775 reared at 25°C and 65% humidity (**Ohyama et al., 2013, 2015; Jovanic et al., 2016, 2019; Eschbach**
776 **et al., 2020b**). Specifically for optogenetics experiments, larvae were raised in the dark and a 1:200
777 retinal solution (diluting 1 g of powdered all-*trans*-retinal (#R240000, Toronto Research Chemicals,
778 Ontario, Canada) in 35.2 ml of 95% ethanol) was added to the food unless indicated otherwise.
779 For immunohistochemistry, eggs were collected during daytime for approximately four hours on
780 standard cornmeal food plates with added yeast. Dissections were performed using wandering-
781 stage third-instar larvae (118–122 hours after egg laying).

782 **Immunohistochemistry and confocal imaging**

783 All dissections, immunohistochemical stainings, and confocal imaging were done using a proce-
784 dure adapted from **Jenett et al. (2012)** and **Li et al. (2014)**. Larval CNSs were dissected in cold 1x
785 phosphate buffer saline (PBS, Corning Cellgro, #21-040) and transferred to tubes filled with cold
786 4% paraformaldehyde (Electron Microscopy Sciences, #15713-S) in 1x PBS. Tubes were incubated
787 for one hour at room temperature. The tissue was then washed four times in 1x PBS with 1%
788 Triton X-100 (#X100, Sigma Aldrich St. Louis, Missouri) (PBT) and incubated in 1:20 donkey serum
789 (#017-000-121, Jackson Immuno Research, West Grove, Pennsylvania) in PBT for two hours at room
790 temperature.

791 The tissue was then incubated in the primary antibody solution, first for four hours at room
792 temperature and then for two nights at 4°C. This solution contained mouse anti-Neuroglial (1:50,
793 #BP104 anti-Neuroglial, Developmental Studies Hybridoma Bank, Iowa City, Iowa), rabbit anti-
794 green fluorescent protein (GFP) (1:500, #A11122, Life Technologies, Waltham, Massachusetts) and
795 rat anti-N-Cadherin (1:50, #DN-Ex #8, Developmental Studies Hybridoma Bank, Iowa City, Iowa) in
796 PBT. This solution was then removed and the tissue washed four times in PBT. The tissue was then
797 incubated in the secondary antibody solution, first for four hours at room temperature and then
798 for two nights at 4°C. This solution contained Alexa Fluor 568 donkey anti-mouse (1:500, #A10037,
799 Invitrogen, Waltham, Massachusetts), FITC donkey anti-rabbit (1:500, #711-095-152, Jackson Im-
800 muno Research West Grove, Pennsylvania) and Alexa Fluor 647 donkey anti-rat (1:500, #712-605-
801 153, Jackson Immuno Research West Grove, Pennsylvania) in PBT. After removal of the secondary
802 solution, the tissue was washed in PBT four times and mounted on a coverslip coated with poly-L-
803 lysine (#P1524-25MG, Sigma Aldrich, St. Louis, Missouri).

804 The coverslip with the CNSs was dehydrated by moving it through a series of jars containing
805 ethanol at increasing concentrations (30%, 50%, 75%, 95%, 100%, 100%, 100%) for ten minutes
806 each. The tissue was then cleared by soaking the coverslip with xylene (#X5-500, Fisher Scientific,
807 Waltham, Massachusetts) three times for five minutes each. Finally, the coverslips were mounted
808 in dibutyl phthalate in xylene (DPX, #13512, Electron Microscopy Sciences, Hatfield, Pennsylvania)
809 with the tissue facing down on a microscope slide with spacers. The DPX was allowed to dry for at
810 least two nights prior to confocal imaging with an LSM 710 microscope (Zeiss).

811 Details on the confocal imaging settings are provided in the respective figure captions. Confocal
812 images were analysed using Fiji (ImageJ). Neurons were counted by specifying regions of interest
813 around the cell bodies using raw image stacks.

814 **Verification of optogenetic and thermogenetic stimulation efficiency**

815 We assessed the multi-larva tracker's optogenetic and thermogenetic stimulation efficiency through
816 open-loop experiments. The behavioural readout was rolling upon exposure to stimulation. All lar-
817 val handling and experiments were performed in the dark to avoid unintended optogenetic stimu-
818 lation. The one-minute experiment protocol began with a 15 s initialisation period in which larvae
819 acclimated to the agarose plate and the *rol1* behaviour classifier stabilised. In three subsequent
820 15 s stimulation cycles, larvae received 5 s of open-loop stimulation followed by 10 s without stim-
821 ulation (**Figure 2B**, **Figure 2E**). Optogenetics were performed with the maximum available red light
822 intensity of 285 $\mu\text{W}/\text{cm}^2$. Thermogenetics were performed with 40% of the maximum available
823 laser intensity.

824 We analysed both optogenetic and thermogenetic experiment data using identical assessment
825 and exclusion criteria. For each larva, the criterion for a single *rol1* was detection of the behaviour
826 for at least 300 ms during a given 15 s stimulation cycle. This threshold ensured true rolls were
827 counted, as opposed to rapid larval bends characteristic of aversion to light.

828 **High-throughput operant conditioning**

829 Experiment procedures

830 We performed high-throughput operant conditioning using our multi-larva closed-loop tracker. All
831 larval handling and experiments were performed in the dark to avoid unintended optogenetic
832 stimulation. We used water to wash approximately 10–12 larvae out of their food. Using a brush,
833 we immediately placed these larvae into the centre of the agarose plate in such a way that they were
834 not touching each other. We placed the agarose plate inside the tracker on top of the backlight and
835 then shut the tracker door. Larvae were given at least 30 s to accustom to their new environment
836 before we started the experiment.

837 The experiment protocol began and ended with a one-minute test period without optogenetic
838 stimulation. Between these test periods were four, three-minute training sessions during which
839 larvae received red light stimulation of 285 $\mu\text{W}/\text{cm}^2$ for the entire duration of the detected bend.

840 Which side received stimulation was randomized across trials such that approximately 50% of lar-
841 vae were trained to develop a right bend preference and 50% a left bend preference. No stimulus
842 was triggered when the larva was bending right or when its body was straight. The test periods
843 were each separated by three-minute periods without stimulation. After the first minute of this
844 period, a brush was used to gently move all larvae back to the centre of the plate and larvae were
845 given time to recover before the beginning of the next training session. This recentring addresses
846 problems encountered when performing prolonged experiments with freely behaving larvae on a
847 small agarose plate. The longer larvae are left undisturbed, the more likely they are to touch the
848 plate's edge, causing tracking disruption and temporary loss of valid objects. This shrinks sample
849 size and reduces training efficiency by decreasing the proportion of animals which are receiving
850 the stimulus.

851 Control experiments were designed so that valid objects received optogenetic stimulation un-
852 correlated with behaviour. These control experiments were split into 60 s time bins, during which
853 each valid object was randomly assigned a stimulus train from this same time bin, pulled from a
854 prior experiment where stimulation correlated with behaviour.

855 Data analysis

856 Data analysis was conducted using custom MATLAB software. To ensure high quality data, it was
857 necessary to remove invalid objects from the data set prior to behavioural analysis. These included
858 corrupted objects (e.g. scratches on the plate or residual food) that the software mistook for larvae.
859 They also included larvae that lost their object identity and were consequently detected for only
860 part of the experiment (e.g. after temporarily reaching the plate's edge or touching other larvae).
861 After equally splitting each experiment into 60 s time bins, we retained objects for analysis that
862 fulfilled strict criteria: i) the object must have been detected in every frame of the bin; ii) the object's
863 initial detection must have occurred at least 20 s prior to the start of the bin; iii) at no point during
864 the bin did the smoothed velocity of the larval centroid exceed 1.5 mm/s; and iv) the mean of the
865 smoothed centroid velocity across the object's detection period in the bin was at least 0.5 mm/s.
866 To quantify the accuracy of this method, we manually reviewed 350 videos of objects flagged as
867 valid for a given 60 s bin. In this group, we observed no severely corrupted objects. In one case
868 (0.3%), a larva briefly touched another larva. In another case (0.3%), head and tail of a larva were
869 falsely detected the majority of the time, leading to flipped detection of left and right bends.

870 When analysing valid bin data for operant conditioning of bend direction preference, we counted,
871 for each larva, the numbers of left and right bends initiated within the bin. This was defined as the
872 bend rate towards the respective direction. The probability of the larva bending towards the side
873 paired with the optogenetic stimulus was defined as the ratio of the number of bends towards this
874 side to the total number of bends initiated in the bin. We pooled together all larval data within
875 each bin because bends to the left and right were each paired with the optogenetic stimulus for
876 approximately half of the larvae. Mean and standard error were calculated for bend rate to the
877 stimulated side, bend rate to the unstimulated side, and probability of bending towards the stim-
878 ulated side. For the control condition in which larvae received random stimulation during 50% of
879 bends regardless of direction, we calculated mean and standard error for bend rates to the left
880 and right and the probability for bending towards the left. Bend rates to either side were com-
881 pared to each other using a two-sided Wilcoxon signed-rank test. The probability for bending to
882 a given side was compared to chance level (0.5) using a two-sided Wilcoxon signed-rank test. The
883 behaviour characteristics of experimental animals were compared to the control group using a
884 two-sided Mann-Whitney *U* test.

885 Classical conditioning

886 Experiment procedures

887 *CsChrimson* (Klapoetke *et al.*, 2014) was expressed under the control of driver lines targeting can-
888 didate valence-conveying neurons. These driver lines were classified based on expression pattern

889 and previous functional data and are known to drive expression in larvae. Optogenetic activation
890 of neurons (US) was paired with odour presentation (CS) to induce olfactory memory (**Figure 4A**).
891 For each driver line, data was acquired from at least two separate crosses.

892 Classical conditioning followed a procedure similar to those described in *Gerber and Hendel*
893 *(2006)*, *Saumweber et al. (2011)* and *Eschbach et al. (2020b)*. Approximately 40 third-instar larvae
894 were transferred onto a 4% agarose petri dish. Larvae were presented with an odour (1:10⁴ ethyl
895 acetate in ddH₂O) pipetted onto two pieces of filter paper attached to the lid of the dish. This en-
896 closed dish was exposed to red light (630 nm, 350 μW/cm²) for three minutes. Larvae were then
897 transferred to a new agarose-filled petri dish with no odour on its lid (“air”) and placed in the dark
898 for three minutes. This training procedure was repeated three times, with alternating presentation
899 of odour/light and air/dark (paired group). An unpaired group receiving reciprocal stimulus presen-
900 tation (odour/dark, air/light) was trained simultaneously. This ensured that any observed effects
901 were attributable to learning rather than innate odour preference or avoidance. The training trial
902 order was reversed in half of the experiments, starting with air instead of odour presentation.

903 After training, larvae of both paired and unpaired groups were immediately transferred to a
904 1 cm middle zone in the centre of fresh agarose-filled petri dishes. A lid was placed on each dish,
905 with odour presented on one side (odour side) but not the other (air side). After a three-minute
906 test period in the dark, the number of larvae on the odour side, the air side, and in the middle zone
907 were manually counted and entered into an Excel spreadsheet (Microsoft Corporation, Remond,
908 Washington).

909 Data analysis

All data was manually entered into MATLAB and analysed using custom software. For each experi-
ment, a performance index (PI) was calculated as follows:

$$\text{Pref}_{\text{paired}} = \frac{\#(\text{larvae on odour side}) - \#(\text{larvae on air side})}{\#(\text{larvae on plate})} \quad (\text{paired dish})$$

$$\text{Pref}_{\text{unpaired}} = \frac{\#(\text{larvae on odour side}) - \#(\text{larvae on air side})}{\#(\text{larvae on plate})} \quad (\text{unpaired dish})$$

$$\text{PI} = \frac{\text{Pref}_{\text{paired}} - \text{Pref}_{\text{unpaired}}}{2} \quad (\text{combined})$$

910 PIs take values between -1 and +1, where a positive PI reflects appetitive learning and a neg-
911 ative PI reflects aversive learning. Mean and standard error were calculated for each condition.
912 Statistical differences between two groups were tested using a two-sided Mann-Whitney *U* test
913 with Bonferroni correction. Significance compared to zero was tested with a two-sided Wilcoxon
914 signed-rank test with Bonferroni correction.

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921 Competing interests

922 The authors declare that no competing interests exist.

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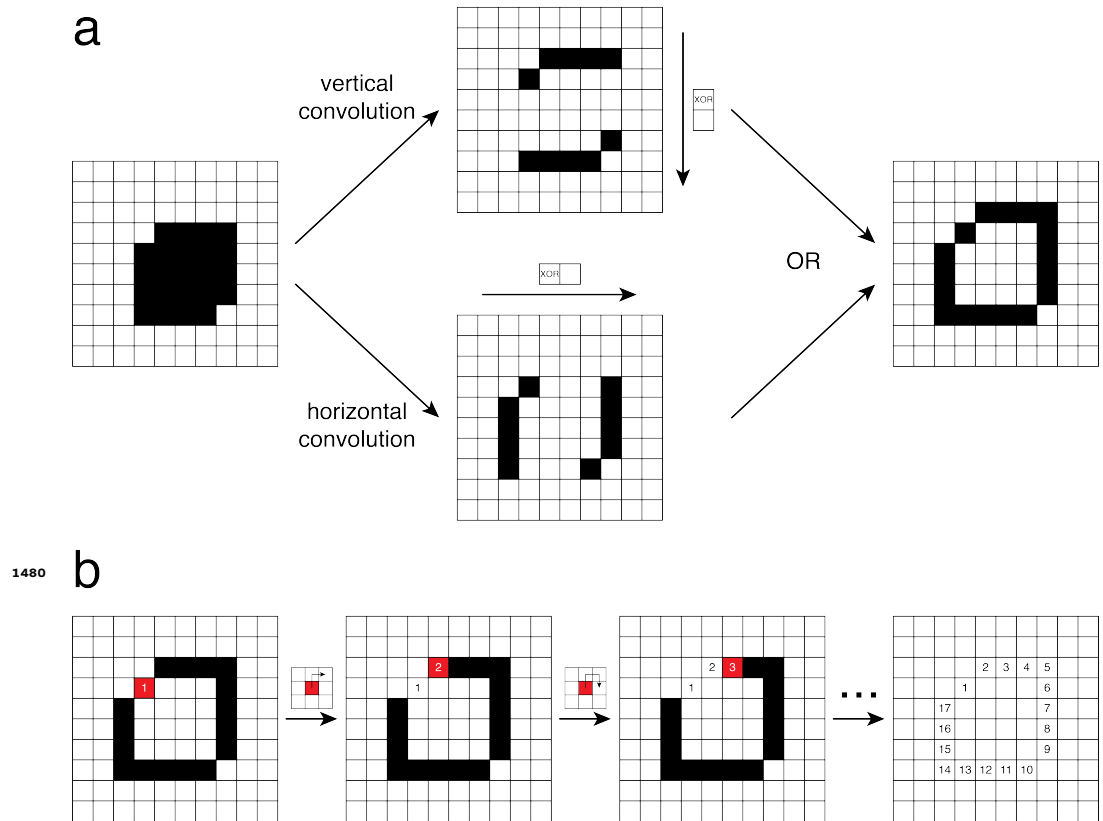
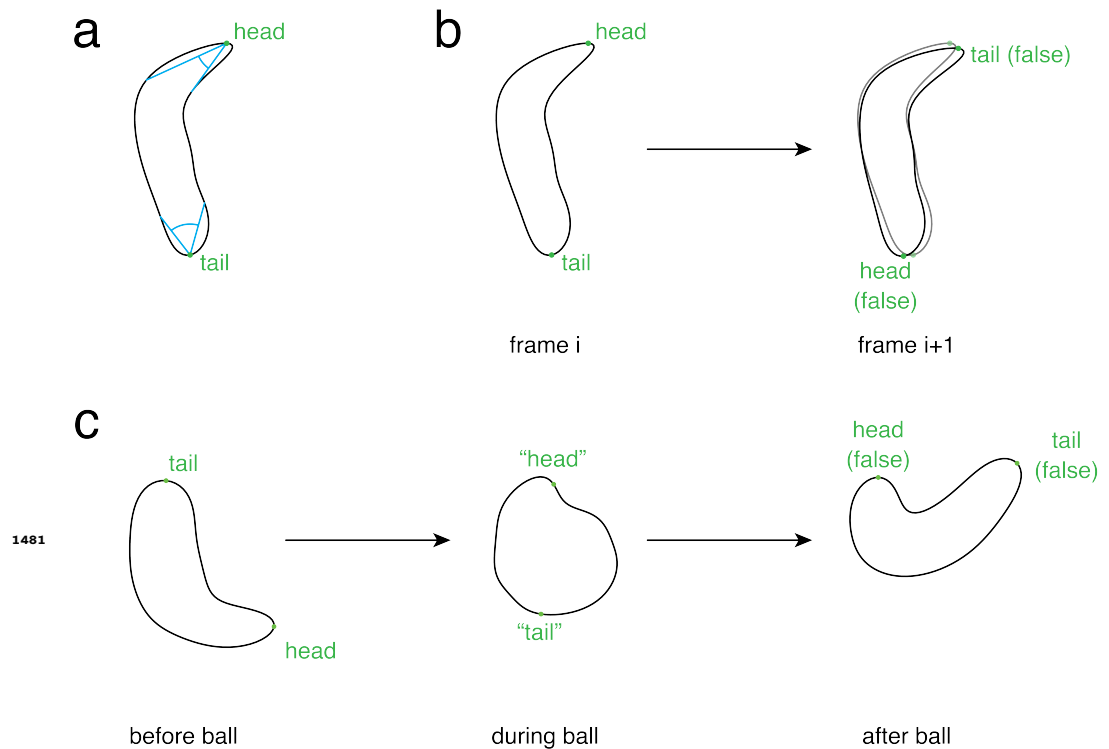


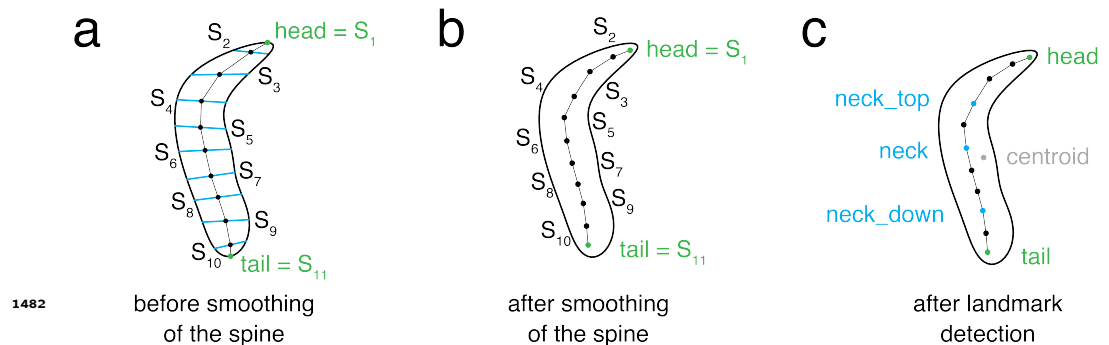
Figure 1-Figure supplement 1. Contour calculation on field-programmable gate array (FPGA).

A simplified example is shown using a 10 x 10 pixel box containing a small object. **a.** The object (black) was detected against the background (white) using binary thresholding. Edge pixels were detected by combining the results of vertical and horizontal image convolution with a 2 x 1 XOR kernel using an OR operator. **b.** The contour points were reconstructed in an iterative process, starting with the edge pixel closest to the centre of the box. The next contour point was defined as the first neighbouring pixel that was found to be an edge pixel. Neighbouring pixels were assessed clockwise from the pixel directly above the contour point. The process ended when no eligible edge pixels could be found.



1481

Figure 1-Figure supplement 2. Detecting head and tail. The larval contour (black outline) and head and tail (green) are shown. **a.** Initial detection of head and tail. The head was the contour point with the sharpest curvature. The tail was the contour point with the next-sharpest curvature which did not lie in close proximity to the head. **b.** The initial detection of head and tail was incorrect in some cases. False detection could be corrected by swapping head and tail, thereby minimising the distances from head and tail in the current frame (solid contour) to head and tail in the previous frame (transparent contour). **c.** The correction described in **b** failed if larvae curled up such that the contour appeared circular ("ball"). To eliminate this source of false head and tail detection, these events were detected using a ball classifier.



1482

Figure 1-Figure supplement 3. Calculating a smooth spine and landmark points. The larval contour is shown (black outline). The spine S was comprised of eleven points (black), including head and tail (green). **a.** The raw spine points were obtained by finding the centres between equally spaced contour points on either half of the contour as defined by head and tail. The first spine point was the head, the last spine point was the tail. **b.** The smooth spine was obtained by exponentially smoothing the raw spine. **c.** Four additional landmark points, `neck_top`, `neck`, and `neck_down` (blue), and the contour `centroid` (grey), were calculated.

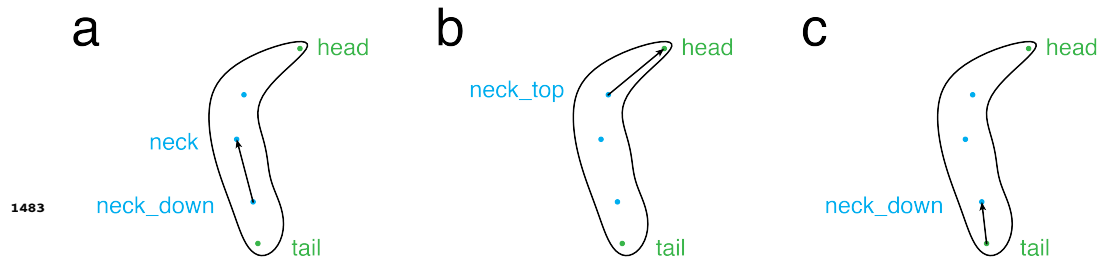


Figure 1-Figure supplement 4. Calculating direction vectors. Three direction vectors were calculated based on head, tail, and the landmark points. **a.** `direction_vector` was the normalised vector from `neck_down` to `neck`. **b.** `direction_head_vector` was the normalised vector from `neck_top` to `head`. **c.** `direction_tail_vector` was the normalised vector from `tail` to `neck_down`.

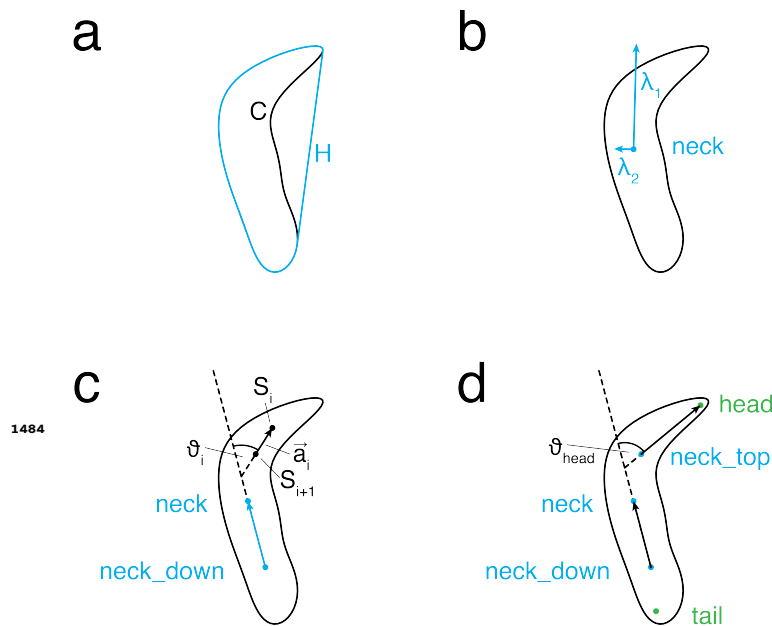
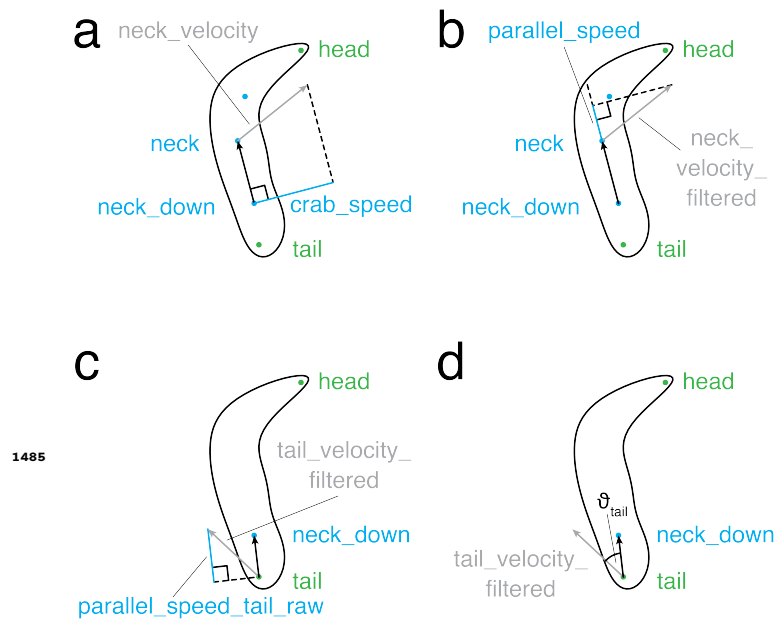


Figure 1-Figure supplement 5. Features describing body shape. **a.** Outline of a larva with contour C (black) and its convex hull H (blue). **b.** Shown here are the eigenvectors (blue) of the larval contour (black) structure tensor with respect to `neck` and their corresponding eigenvalues λ_1 and λ_2 . **c.** ϑ_i was defined as the angle between `direction_vector` (blue) and the vector \vec{a}_i that passed through spine points S_i and S_{i+1} (black). **d.** ϑ_{head} was defined as the angle between `direction_vector` and `direction_head_vector`. `head` and `tail` are shown in green.



1485

Figure 1-Figure supplement 6. Velocity features. The larval contour is shown in black while head and tail are shown in green. **a.** crab_speed (blue) was defined as the component of neck_speed (grey) that was orthogonal to direction_vector_filtered (black). **b.** parallel_speed (blue) was defined as the component of neck_speed_filtered (grey) that was parallel to direction_vector_filtered (black). **c.** parallel_speed_tail_raw (blue) was defined as the component of tail_speed_filtered (grey) that was parallel to direction_tail_vector_filtered (black). **d.** ϑ_{tail} was defined as the angle between tail_speed_filtered (grey) and direction_tail_vector_filtered (black).

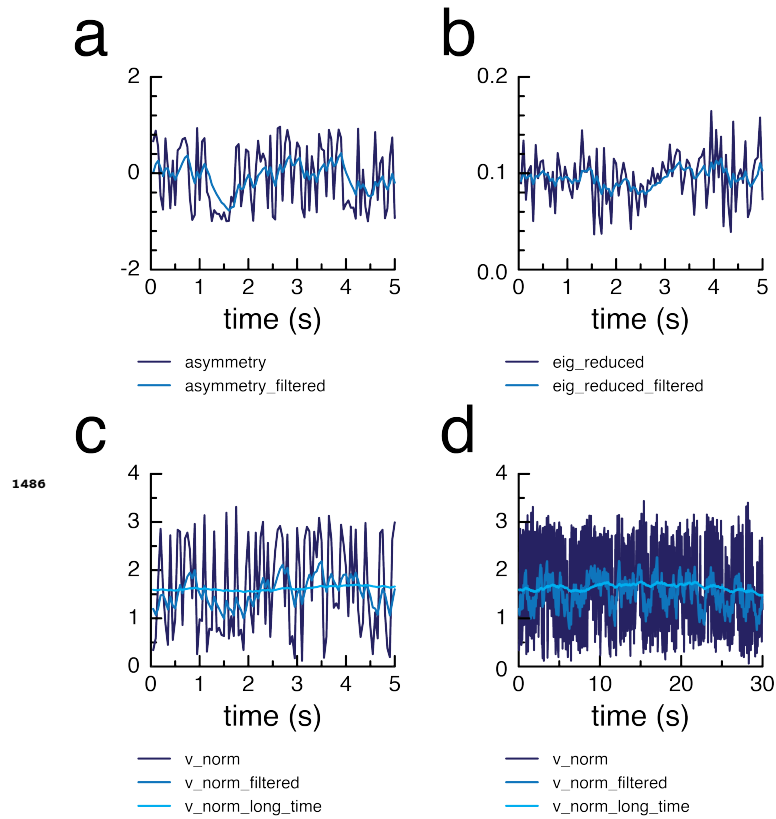


Figure 1-Figure supplement 7. Temporal smoothing of features. a-b. Example graphs of raw (dark blue) and filtered (mid blue) *asymmetry* (a) and *eig_reduced* (b) values over time. c-d. Example graphs of raw (dark blue), filtered (mid blue), and long-time filtered (light blue) *v_norm* values over a short (c) and a long (d) period of time.

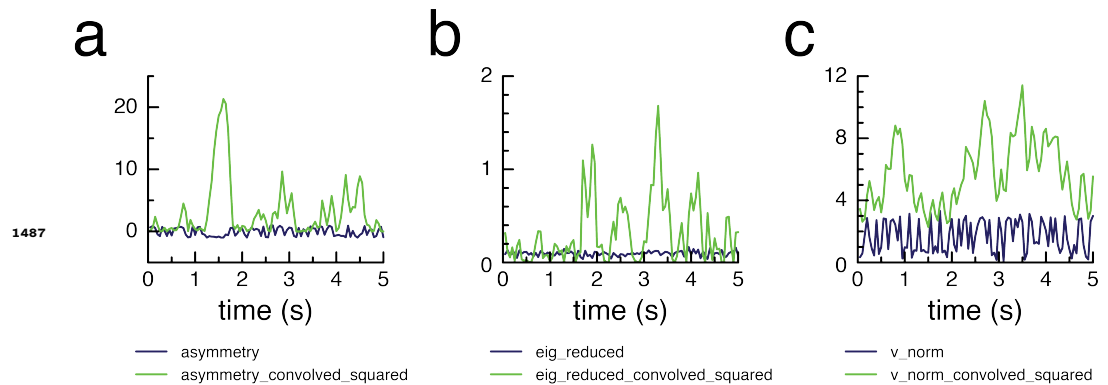
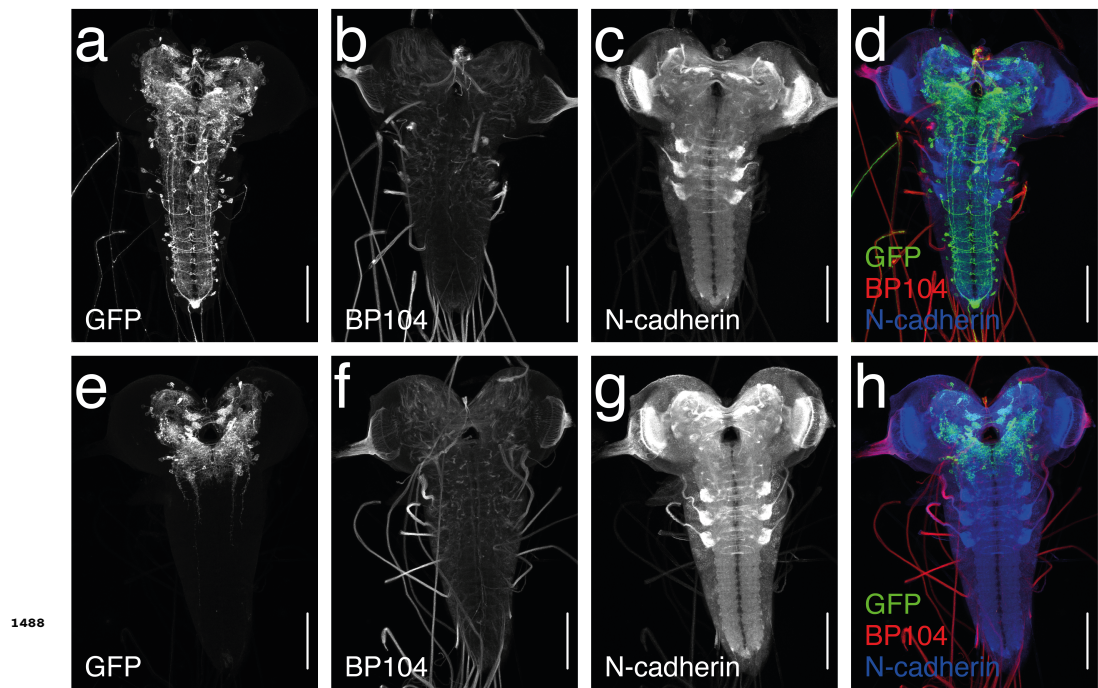


Figure 1-Figure supplement 8. Differentiation by convolution. Example graphs of raw (dark blue) and convolved squared (green) *asymmetry* (a), *eig_reduced* (b) and *v_norm* (c) values over time.



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Figure 3-Figure supplement 1. *Ddc-Gal4* expression pattern without and with *tsh-Gal80* restriction. Maximum intensity projections of confocal images obtained after immunohistochemical staining. **a, e**; green in **d** and **h**. Targeting a green fluorescent protein (GFP) antibody to the *mVenus* tag of *CsChrimson*. **b, f**; red in **d** and **h**. Staining against BP104. **c, g**; blue in **d** and **h**. Staining against N-cadherin. **a-d**. *Ddc-Gal4* x *UAS-CsChrimson* larvae. Manually counting the cell bodies in the image stacks revealed more than 200 GFP-positive neurons located in the brain, subesophageal zone (SEZ), and ventral nerve cord (VNC), including the PAM cluster dopaminergic neurons innervating the mushroom body ($n = 2$). This confirmed that *Ddc-Gal4* drives broad expression across the central nervous system (CNS) (Lundell and Hirsh, 1994; Li et al., 2000). **e-h**. *Ddc-Gal4* x *UAS-CsChrimson*; *tsh-LexA*, *LexAop-Gal80* larvae. As expected, no GFP-positive neurons were found in the VNC ($n = 6$). *Ddc-Gal4* brain and SEZ expression remained largely unaffected by GAL80, as the GFP-positive neurons in both areas that could be consistently identified in *Ddc-Gal4* x *UAS-CsChrimson* larvae ($n = 3$) were also present in *Ddc-Gal4* x *UAS-CsChrimson*; *tsh-LexA*, *LexAop-Gal80* larvae ($n = 3$). **a-h**. Plan-Apochromat 20x objective, resolution: 592 x 800 pixels, scale bar: 100 μ m. Images courtesy of the HHMI Janelia FlyLight team.

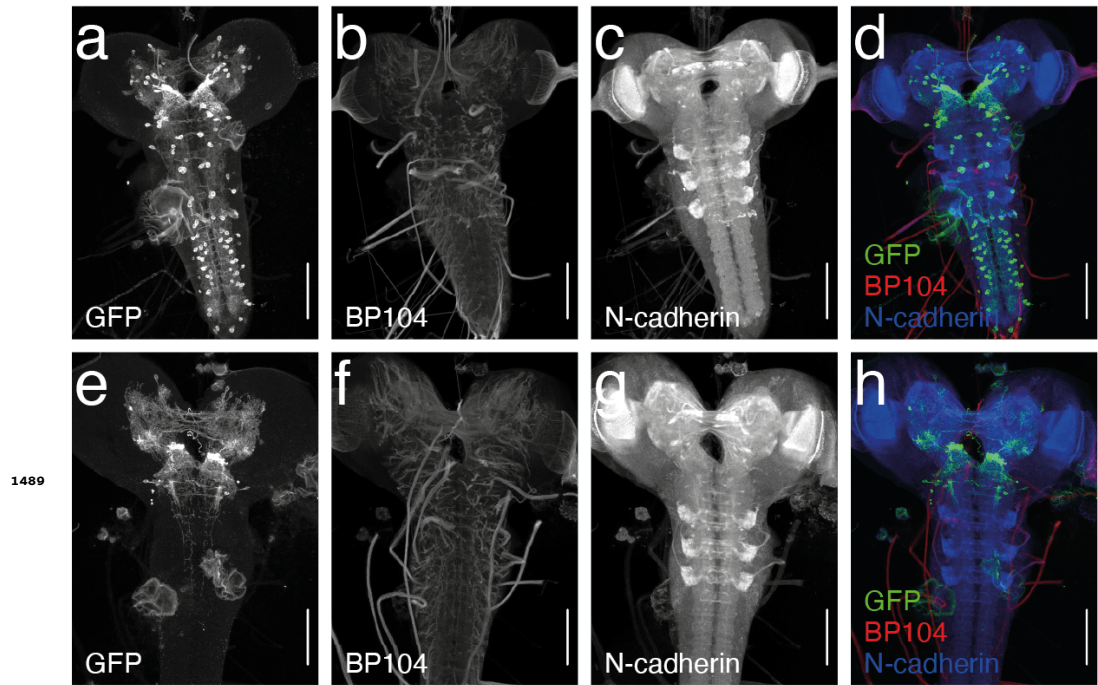


Figure 4-Figure supplement 1. *Tph-Gal4* expression pattern without and with *tsh-Gal80* restriction. Maximum intensity projections of confocal images obtained after immunohistochemistry. **a-d.** *Tph-Gal4* x *UAS-CsChrimson* larvae, **e-h.** *Tph-Gal4* x *UAS-CsChrimson*; *tsh-LexA*, *LexAop-Gal80* larvae. **a, e;** green in **d** and **h.** Staining against green fluorescent protein (GFP) antibody targeting the *mVenus* tag of *CsChrimson*. **b, f;** red in **d** and **h.** Staining against BP104. **c, g;** blue in **d** and **h.** Staining against N-cadherin. **a-h.** Plan-Apochromat 20x objective, resolution: 592 x 800 pixels, scale bar: 100 μ m. Image courtesy of the HHMI Janelia FlyLight team.

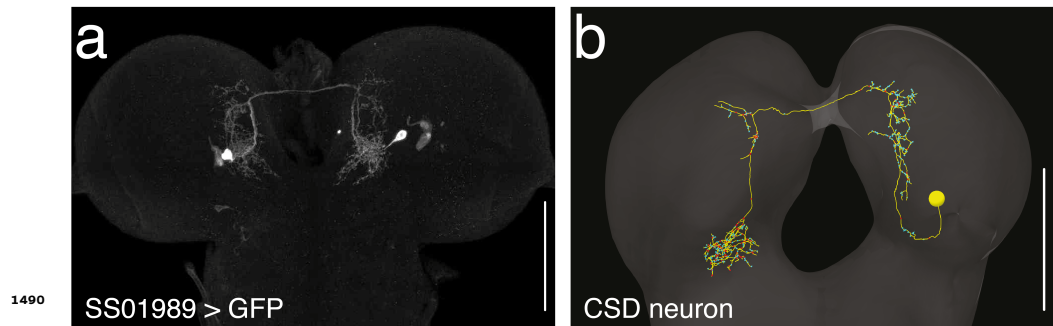


Figure 4-Figure supplement 2. *SS01989* exclusively drives expression in the CSD neuron. **a.** Confocal image of a third-instar *SS01989* x *UAS-GFP* larva CNS, derived from maximum intensity projections, obtained after immunohistochemical staining against *GFP*. C-Apochromat 40x objective, resolution: 975 x 651 pixels, scale bar: 100 μ m. Image courtesy of the HHMI Janelia FlyLight team. **b.** Electron microscopy reconstruction of the contralaterally projecting serotonergic-immunoreactive deutocerebral (CSD) neuron from the central nervous system of a first-instar larva (Berck et al., 2016), scale bar: 50 μ m.

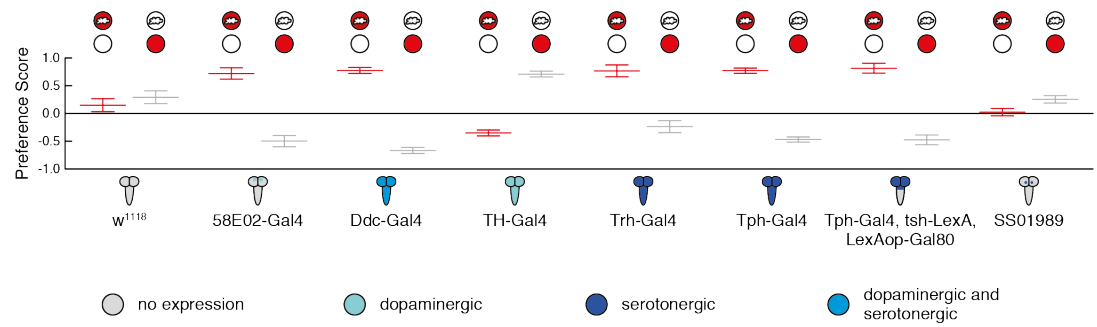


Figure 4-Figure supplement 3. Paired and unpaired group data for olfactory conditioning experiments. The data shown here underlies the performance indices depicted in **Figure 4B**. Gal4 expression depicted as color-coded central nervous system. Preference scores for paired (light/odour, dark/air) and unpaired (dark/odour, light/air) groups are shown in red and grey, respectively.