

Current Biology

Early Pheromone Experience Modifies a Synaptic Activity to Influence Adult Pheromone Responses of *C. elegans*

Highlights

- Early pheromone exposure modulates behavioral responses to the pheromone as adults
- Pheromone experience is imprinted as increased synaptic activity
- The *odr-2* GPI-linked signaling protein mediates pheromone imprinting

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In Brief

Hong et al. show that early pheromone experience in *C. elegans* hermaphrodites is imprinted via alteration of activity of a single synaptic connection and, in turn, modulates behavioral responses to the pheromone as adults.



Early Pheromone Experience Modifies a Synaptic Activity to Influence Adult Pheromone Responses of *C. elegans*

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SUMMARY

Experiences during early development can influence neuronal functions and modulate adult behaviors [1, 2]. However, the molecular mechanisms underlying the long-term behavioral effects of these early experiences are not fully understood. The *C. elegans* ascr#3 (asc-ΔC9; C9) pheromone triggers avoidance behavior in adult hermaphrodites [3–7]. Here, we show that hermaphrodites that are briefly exposed to ascr#3 immediately after birth exhibit increased ascr#3-specific avoidance as adults, indicating that ascr#3-experienced animals form a long-lasting memory or imprint of this early ascr#3 exposure [8]. ascr#3 imprinting is mediated by increased synaptic activity between the ascr#3-sensing ADL neurons and their post-synaptic SMB motor neuron partners via increased expression of the *odr-2* glycosylated phosphatidylinositol (GPI)-linked signaling gene in the SMB neurons. Our study suggests that the memory for early ascr#3 experience is imprinted via alteration of activity of a single synaptic connection, which in turn shapes experience-dependent plasticity in adult ascr#3 responses.

RESULTS

Adult *C. elegans* Transiently Exposed to ascr#3 during Larval Stages Exhibits Increased ascr#3 Pheromone Avoidance

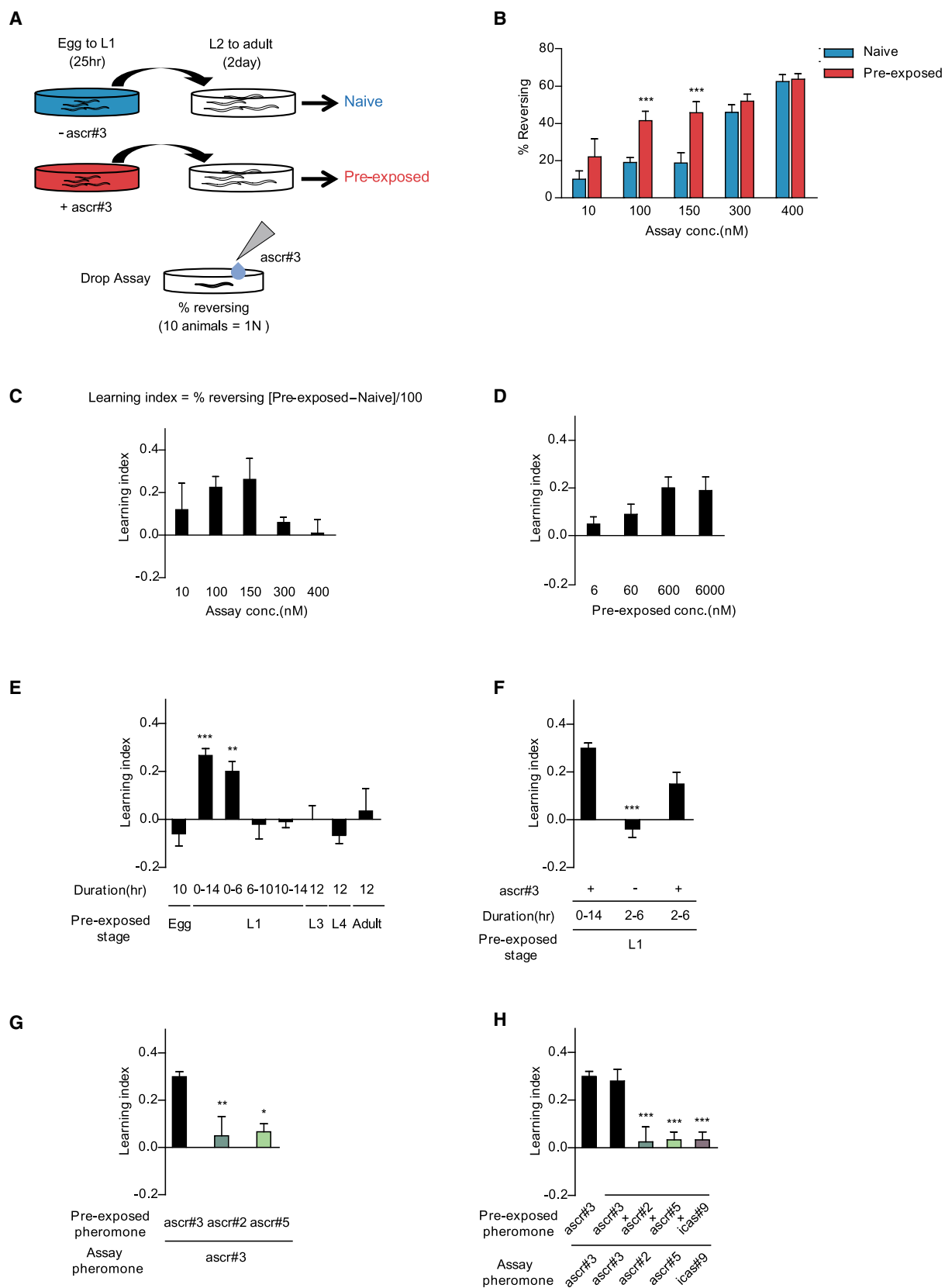
To assess whether early experience of pheromones affects ascr#3 avoidance behavior in adult worms, we transiently exposed first larval stage (L1) wild-type worms to ascr#3 and examined their responses to ascr#3 as adults (Figure 1A; see STAR Methods). To test avoidance, a pheromone diluted in buffer was applied to a freely moving animal, and the fraction

of animals that reverse was calculated (drop test assay; see STAR Methods) [9]. Pre-exposure to 600 nM ascr#3 during the L1 stage significantly improved adult avoidance of ascr#3 at concentrations that elicit only weak avoidance by adults that were not similarly pre-exposed but did not further increase avoidance at higher ascr#3 concentrations (Figure 1B). Increased ascr#3 avoidance appeared to be mediated by increased long reversals and omega turns, but not short reversals in pre-exposed animals (Figure S1A). Henceforth, we calculate increased avoidance as a learning index (LI; pre-exposed reversing rate minus naive reversing rate/100) (Figure 1C).

We next defined the optimal ascr#3 concentration that elicits the pre-exposure effect. Animals exhibited maximal adult ascr#3 avoidance when they were pre-exposed to ascr#3 concentrations higher than 600 nM (Figure 1D) [4]. Although exposure of L1 larvae to ascr#3 pheromone also promotes entry into the alternate dauer developmental stage, we note that the conditions used for imprinting are distinct from those used for dauer induction and few if any ascr#3-imprinted animals enter into the dauer stage. These results indicate that early experience of ascr#3 pheromone appears to be translated into behavioral changes in the adult stage of hermaphrodites, suggesting that ascr#3-experienced animals form a long-lasting memory or imprint for ascr#3. Wild-type *C. elegans* males exhibit neutral responses to 100 nM ascr#3 [7, 10]. We found that ascr#3-experienced adult males continue to exhibit neutral responses to 100 nM ascr#3 (Figure S1B), indicating that early ascr#3 experience appears to increase ascr#3 avoidance only by adult hermaphrodites, but not by adult males.

The Memory for ascr#3 Is Acquired during the L1 Larval Stage

To determine whether the long-lasting memory for ascr#3 is formed during a specific developmental time window, worms were pre-exposed to ascr#3 at multiple developmental stages, and their ascr#3 responses were assessed as adults. All assays were conducted by pre-exposing animals to 600 nM pheromone



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and assessing responses to 100 nM pheromone, unless noted otherwise.

Compared to the L1 stage, *ascr#3* pre-exposure at the L3, L4, or adult stages did not enhance *ascr#3* avoidance in adults (Figure 1E), suggesting that the L1 stage is critical for acquiring the *ascr#3* memory. We could not test the L2 stage because of complexity with dauer formation [11]. We further defined the time window within the L1 stage necessary for this behavior and found that pre-exposure in a defined time period during the early L1 stage (up to 6 hr following hatching at 20°C) appears to represent the critical period for formation of the *ascr#3* memory (Figure 1E). More specifically, exposure of *ascr#3* to worms at 2–6 hr after birth appears to be necessary and sufficient for the increased *ascr#3* avoidance (Figure 1F).

C. elegans secretes additional pheromones, including *ascr#2*, *ascr#5*, and *icas#9*. Similar to *ascr#3*, these pheromones are also potent inducers of entry into the alternate dauer developmental stage under limiting food conditions [4]. Pre-exposure to 600 nM *ascr#2* or *ascr#5* alone did not affect *ascr#3* avoidance by adults, suggesting that pheromone imprinting does not reflect a memory of general unfavorable dauer-inducing conditions but specifically encodes experience of the *ascr#3* pheromone (Figure 1G). Pre-exposure to mixtures of *ascr#2*, *ascr#5*, and *icas#9* with *ascr#3* also did not affect *ascr#3* imprinting, further supporting that imprinting is specific to *ascr#3* (Figure 1H). Moreover, pre-exposure to *ascr#2* did not result in altered adult avoidance of *ascr#2* (Figure S1C), further confirming specificity of this process to *ascr#3*. In addition, pre-exposure to *ascr#3* did not affect high-osmolality glycerol avoidance (Figure S1D), suggesting that early *ascr#3* experience does not affect broad avoidance behaviors to other repulsive chemicals by adults. However, we are unable to exclude the possibility that imprinting by other ascarosides may occur at developmental stages other than the L1 stage.

Because the memory for a specific pheromone component is acquired only at the critical developmental period, we further define this altered *ascr#3* avoidance as sensory imprinting [8]. Interestingly, when animals recovered from the developmentally arrested dauer stage that was induced by limited food supply in the presence of pheromones, including *ascr#3*, at the L1 and L2 stages, their *ascr#3* avoidance was comparable to or even weaker than that of naive animals (Figure S1E) [11]. These data indicate that the *ascr#3* imprint appears to be removed following

dauer experience and/or *ascr#3* imprinting requires non-dauer-inducing conditions at the L1 stage. Moreover, it is likely that passage through the dauer stage results in genome-wide changes in gene expression patterns [12, 13], which may mask or erase *ascr#3*-imprinting phenotypes.

We next investigated perdurance of the *ascr#3* memory. We found that pre-exposed 3- or 6-day-old adults still exhibited increased *ascr#3* avoidance (Figure S1F). We also asked whether memories are transmitted to the next generation but found that progeny from imprinted mothers did not exhibit improved *ascr#3* avoidance (Figure S1G). These results indicate that the *ascr#3* imprint lasts with aging but is not inherited under these assay conditions.

***odr-2* Acts in the SMB Neurons to Increase *ascr#3* Avoidance in *ascr#3*-Imprinted Animals**

We next performed a candidate gene search to identify genes and molecules required for *ascr#3* imprinting. We first tested genes that have been shown to be required for distinct forms of learning and memory in *C. elegans*. These include the *egl-4* (cyclic guanosine monophosphate [GMP]-dependent protein kinase) gene implicated in sensory adaptation, the *casy-1* (cal-syntenin) gene involved in associative learning, and the *sra-11* (7-TM G-protein-coupled receptor), *ttx-3* (LIM homeodomain protein), or *tdc-1* (tyrosine decarboxylase) genes involved in olfactory imprinting [14–17]. However, loss of function of these genes did not affect *ascr#3* imprinting (Figure S2A).

In the course of analyzing mutants defective in odorant responses, we found that *odr-2* (*n2145*) [18, 19] mutants exhibited defects in *ascr#3* imprinting, although the ability of these mutants to avoid *ascr#3* was unaffected (Figures 2A and 2B). The *ascr#3* imprinting defects in *odr-2* mutants were fully rescued upon expression of wild-type *odr-2* cDNA driven under its upstream regulatory sequences (Figure 2C). The *odr-2* gene encodes a membrane-associated protein related to the Ly-6 (leukocyte antigen-6) superfamily of GPI (glycosylated phosphatidylinositol)-linked proteins, and *odr-2* mutants have previously been shown to exhibit decreased chemotaxis toward a set of volatile attractive chemicals [19]. *odr-2* is expressed in a set of head neurons, including RIG, RME, and SMB [19, 20]. To determine where ODR-2 acts to regulate *ascr#3* imprinting, we tested transgenic animals expressing *odr-2* wild-type sequences for

Figure 1. Transient Exposure to the *ascr#3* Pheromone after Birth Specifically Enhances *ascr#3*-Avoidance Behavior of Adults

(A) Experimental scheme of pheromone imprinting assay. The animals are exposed to dH₂O (naive: shown in blue) and *ascr#3* diluted with dH₂O (pre-exposed: shown in red) from egg to the L1 stage. The percentage of reversal is calculated by measuring avoidance frequencies to *ascr#3* exposure at the adult stage. (B and C) Percentage of reversal (B) and learning index (C) of naive and pre-exposed adult animals to 10 nM, 100 nM, 150 nM, 300 nM, and 400 nM *ascr#3*. Learning index is calculated by the percentage of the reversal rate of pre-exposed minus the reversal rate of naive divided by 100. *** indicates different *ascr#3* avoidance from naive at $p < 0.001$ by one-way ANOVA with Bonferroni's post hoc test. $n = 50$ –200 each. (D) Learning index of adult animals pre-exposed to 6 nM, 60 nM, 600 nM, and 6,000 nM *ascr#3* at L1. $n = 80$ –200 each. (E) Learning index of adult animals pre-exposed to 600 nM *ascr#3* at different developmental stages. Exposure times during the specific developmental stage are indicated as egg (10 hr), L3 (12 hr), L4 (12 hr), and adult (12 hr). For the L1 stage, exposure times are indicated as between starting and ending time of pheromone exposure. ** and *** indicate different from egg (10 hr) at $p < 0.01$ and $p < 0.001$ by one-way ANOVA with Dunnett's post hoc test, respectively. $n = 30$ –190 each. (F) Learning index of adult animals exposed to *ascr#3* for 4 hr in the L1 stage. Pre-exposure for 2–6 hr in the L1 stage is sufficient for *ascr#3* imprinting. *** indicates different from *ascr#3* pre-exposure for 14 hr at $p < 0.001$ by one-way ANOVA with Dunnett's post hoc test. $n = 80$ –120 each. (G and H) Learning index of adult animals pre-exposed and/or assayed with other pheromone components. Animals are pre-exposed to either 600 nM *ascr#3*, *ascr#2*, or *ascr#5* and assayed with 100 nM *ascr#3* (G) and pre-exposed to pheromone mixture containing 600 nM *ascr#3*, *ascr#2*, *ascr#5*, and *icas#9* and assayed with 100 nM *ascr#3*, *ascr#2*, *ascr#5*, or *icas#9* (H). *, **, and *** indicate different from the control (*ascr#3* pre-exposed and *ascr#3* assayed) at $p < 0.05$, $p < 0.01$, and $p < 0.001$, respectively, by one-way ANOVA with Dunnett's post hoc test. (G) $n = 30$ –100 each. (H) $n = 30$ –50 each. In (B)–(H), error bars represent SEM. See also Figure S1.

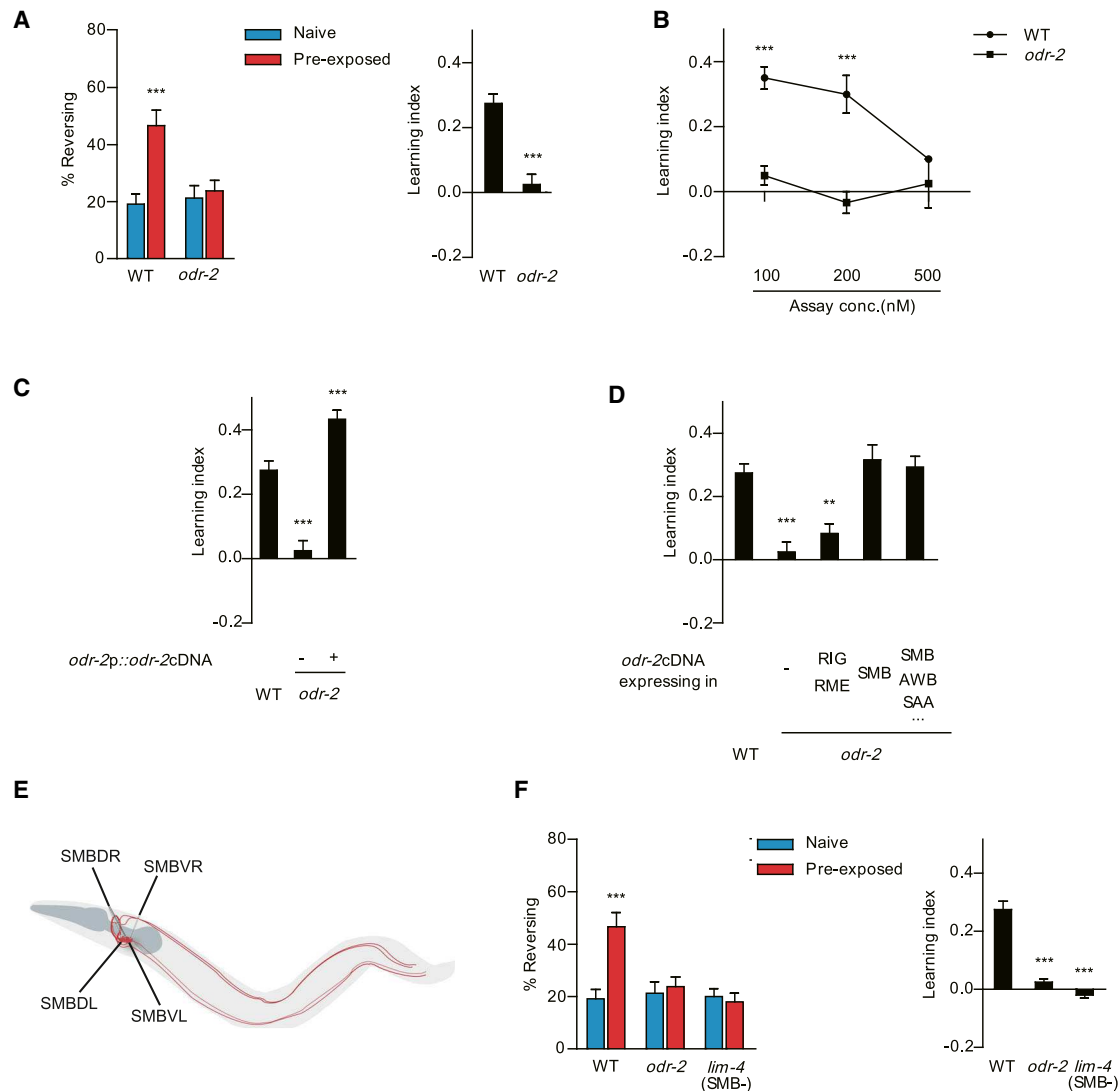


Figure 2. The Glycosylated Phosphatidylinositol-Linked Protein, ODR-2, Is Required for Increased *ascr#3* Avoidance in *ascr#3*-Imprinted Animals

(A) Percentage of reversal (left) and learning index (right) in wild-type and *odr-2* mutants. *** indicates significantly different from naive (left) and wild-type (WT) (right) at $p < 0.001$ by one-way ANOVA with Bonferroni's post hoc test (left) and by Student's *t* test (right), respectively. $n = 80$ –120 each.

(B) Learning index of wild-type and *odr-2* mutants at 100 nM, 200 nM, or 500 nM *ascr#3*. *** indicates different from wild-type at $p < 0.001$ by one-way ANOVA with Bonferroni's post hoc test. $n = 30$ –60 each.

(C and D) Learning index of wild-type and *odr-2* mutants expressing *odr-2* cDNA under the control of *odr-2* promoter (RIG, RME, and SMB; C) or cell-specific promoters, including *odr-2* (–377) (RIG and RME), *flp-12* (–339) (SMB), and *lim-4* (AWB, SAA, RID, RIV, RMD, and SMB; D). ** and *** indicate different from wild-type at $p < 0.01$ and $p < 0.001$, respectively, by one-way ANOVA with Dunnett's post hoc test. (C) $n = 80$ –120 each. (D) $n = 60$ –140 each.

(E) Schematic diagram of the SMB neurons. A pair of dorsal and ventral cell bodies (SMBDs and SMBVs) are located in the head, and their processes innervate head muscle and run posteriorly.

(F) Percentage of reversal of naive and pre-exposed animals (left) and learning index (right) in wild-type, *odr-2*, and *lim-4* mutants. *** indicates different from naive (left) and wild-type (right) at $p < 0.001$ by one-way ANOVA with Bonferroni's (left) and Dunnett's post hoc tests (right), respectively. $n = 80$ –120 each.

All error bars represent SEM. See also Figure S2.

rescue of the *ascr#3* imprinting defects. We found that, whereas expression of *odr-2* in RIG and RME under the control of *odr-2* (–377) promoter [20] did not rescue these behavioral defects, *ascr#3* imprinting defects were fully restored upon expression of *odr-2* exclusively in SMB under the control of *flp-12* (–339) promoter (Figure 2D) [20], indicating that ODR-2 acts in the SMB neurons to mediate *ascr#3* imprinting.

The SMB neurons consist of two left and right pairs (dorsal or ventral) of sensory/inter/motor neuron types that are located in the head and that innervate the head and neck muscles (Figure 2E) [21]. Whereas their synapse-free processes extend along the ventral or dorsal sublaterals to the tail, the SMB neurons have extensive electric and chemical synaptic contacts to other neurons in the head [21]. These neurons regulate head locomotion

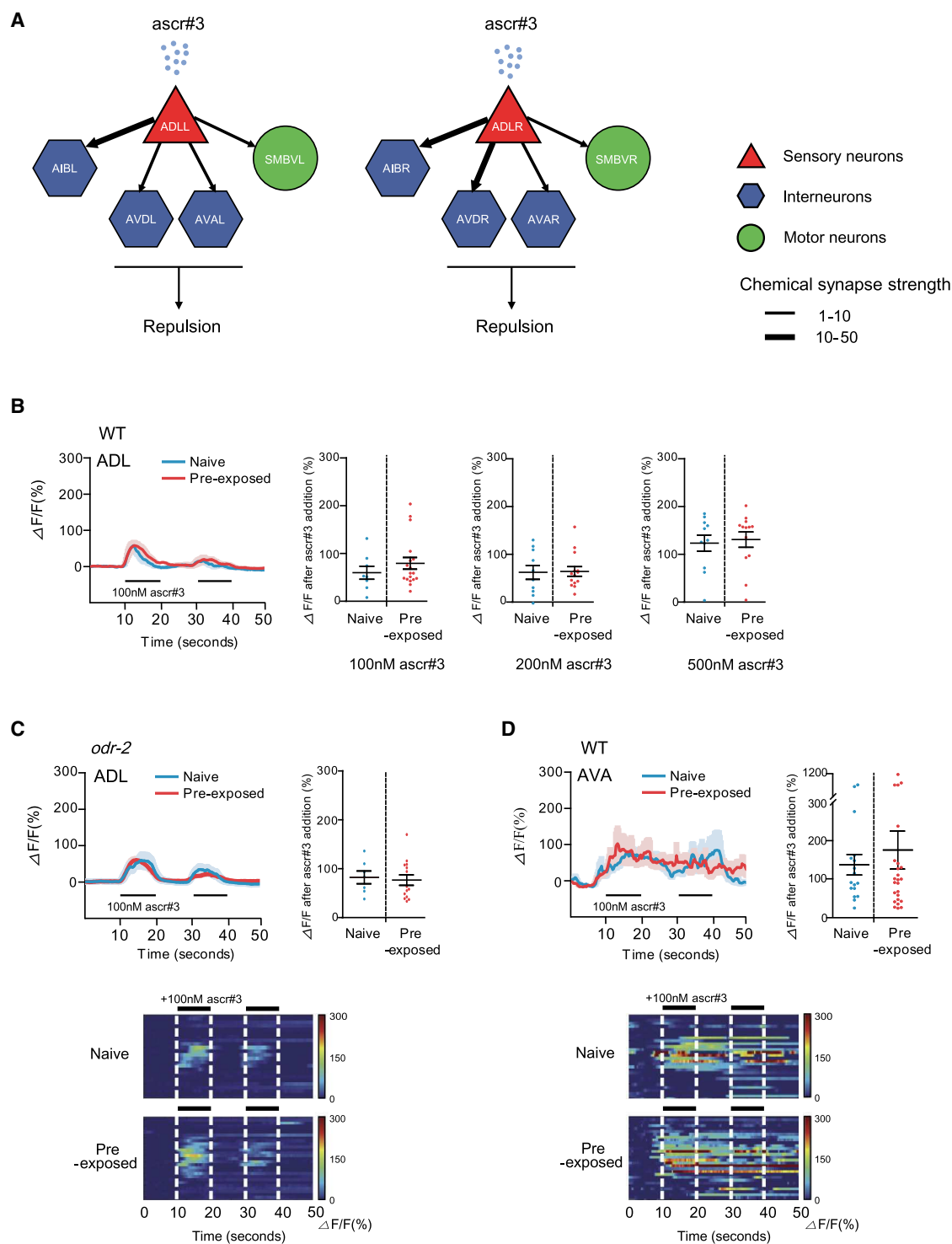


Figure 3. Ca^{2+} Response to Acute *ascr#3* Exposure Is Not Altered in the *ascr#3*-Sensing ADL Neurons and Their Downstream Command Interneurons of *ascr#3*-Imprinted Animals

(A) Post-synaptic connections of the ADLL and ADLR neurons. The AIB, AVD, AVA, and SMBV neurons are post-synaptic to ADL. Chemical synapses between sensory (triangles), inter (hexagons), and motor (circles) are indicated as arrows. Synaptic strength is indicated in thickness of lines (<http://www.wormwiring.org>). (B) Ca^{2+} transients of ADL in response to 100 nM *ascr#3* exposure. The average traces of Ca^{2+} responses during two pulses of 100 nM *ascr#3* (left) and the maximum value of Ca^{2+} responses to 100 nM, 200 nM, or 500 nM *ascr#3* (right) of naive (shown in blue) or pre-exposed animals (shown in red) in 50 s are shown. $n = 8$ (naive) and 18 (pre-exposed) each.

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[20–22]. However, their roles in chemosensory behaviors have not been explored. We next investigated that the SMB neurons are required for *ascr#3* imprinting. To address this issue, we examined *lim-4* mutants in which the SMB neurons are not fully differentiated and in which the functions of SMB are completely abolished [20]. *lim-4* mutants had been shown to exhibit additional defects, including neuronal specification [23]. Whereas *lim-4* mutants were still able to avoid *ascr#3*, pre-exposure to *ascr#3* did not enhance *ascr#3* avoidance in *lim-4* mutants (Figure 2F). These results together with *odr-2* rescue results support that the SMB neurons may play an important role in *ascr#3* imprinting.

ascr#3-Induced Responses in the ADL Chemosensory Neurons Are Unaltered in ascr#3-Imprinted Worms

In adults, *ascr#3* elicits avoidance behavior in hermaphrodites via the nociceptive ADL chemosensory neurons (Figure 3A) [7]. To describe the neuronal basis of pheromone imprinting, we first monitored intracellular Ca^{2+} dynamics in response to *ascr#3* in transgenic animals expressing the genetically encoded calcium sensor GCaMP3 in the *ascr#3*-sensing ADL neurons. The ADL neurons exhibit a rapid and transient Ca^{2+} increase upon exposure to nano-molar concentrations of *ascr#3* (Figure 3B) [7]. Both naive animals and *ascr#3*-imprinted animals displayed similar Ca^{2+} transients in adult ADL neurons upon addition of *ascr#3* (Figure 3B). These results suggest that increased *ascr#3* avoidance is not due to enhanced sensory responsiveness of ADL. Furthermore, consistent with the normal *ascr#3* avoidance behavior of *odr-2* mutants, the ADL Ca^{2+} response to *ascr#3* was not altered in *odr-2* mutants (Figure 3C).

The ADL sensory neurons drive *ascr#3* avoidance through their chemical synapses [7]. Inspection of the anatomical wiring data from the *C. elegans* connectome suggests that *ascr#3* signals from the ADL neurons are transmitted to a few major post-synaptic target neurons, including AIB (1st layer interneurons), and the AVA and AVD backward command interneurons (Figure 3A). Previous studies have shown that activity in AIB, AVA, and AVD is increased during backward movement or reversals [24]. We found that, whereas the AIB or AVD neurons exhibited noisy but low levels of Ca^{2+} responses upon exposure to *ascr#3* (Figures S3A and S3B), the AVA cell bodies responded strongly and consistently to repeated *ascr#3* addition with increased Ca^{2+} levels (Figure 3D). However, these responses were unaltered upon *ascr#3* imprinting (Figures 3D, S3A, and S3B). These results imply that *ascr#3* signals in the ADL neurons may be transmitted mainly via AVA but that imprinting does not enhance *ascr#3* avoidance via this ADL-AVA circuit.

The SMB Neurons Mediate Increased ascr#3 Avoidance in ascr#3-Imprinted Animals

The *C. elegans* connectome data indicate that the SMBV neurons are additional post-synaptic partners of the ADL sensory

neurons (Figure 3A), although the synaptic strength appears to be weak. Because the SMB neurons appear to play a role in *ascr#3* imprinting, we then examined how SMB is involved in *ascr#3*-avoidance behavior.

We first expressed GCaMP3 in SMB under the control of *flp-12* (–339) promoter that drives transgene expression in all four SMB neurons [20]. We note that a SMBV-specific promoter is not currently available. Moreover, because the cell bodies of SMBD and SMBV are located in close proximity, differentiating among these neurons is challenging [20]. We thus examined *ascr#3* responses in all SMB neurons together. Acute *ascr#3* exposure did not elicit Ca^{2+} transients in the SMB neurons of naive control animals (Figure 4A; Movie S1). However, we detected consistent and robust Ca^{2+} transients in response to acute *ascr#3* exposure in *ascr#3*-imprinted animals (Figure 4A; Movie S2). Because SMB may detect body muscle contractions that could be responsible for the observed Ca^{2+} responses in SMB, we paralyzed worms with the nicotinic acetylcholine receptor agonist levamisole. Levamisole-treated *ascr#3*-imprinted animals still exhibited similar or even enhanced Ca^{2+} dynamics upon *ascr#3* exposure (Figure S4A).

To test whether the observed Ca^{2+} transients in SMB in *ascr#3*-imprinted animals are transmitted from ADL, we examined Ca^{2+} responses in transgenic animals expressing TeTx (tetanus toxin light chain) specifically in the ADL neurons to block synaptic transmission from ADL [7]. Ca^{2+} transients of SMB in *ascr#3*-imprinted animals were strongly suppressed by blocking ADL synaptic transmission (Figures 4B and S4B), suggesting that *ascr#3* imprinting results in enhanced *ascr#3* signal transmission from ADL to SMB. These results indicate that *ascr#3* imprinting specifically alters the ADL-SMB synaptic transmission.

Moreover, the *ascr#3*-imprinting-dependent Ca^{2+} response in SMB was completely abolished in *odr-2* mutants (Figure 4C), indicating that *odr-2* elicits increased activity of SMB to *ascr#3* exposure in imprinted animals by enhancing the ADL-SMB synaptic transmission.

Upregulation of odr-2 Expression of L1 Larvae Is Sufficient for Increased ascr#3 Avoidance in Adults

To further investigate the role of *odr-2* in mediating increased SMB activity of *ascr#3*-imprinted animals, we monitored *odr-2* expression in SMB and found that *odr-2* expression in SMB was increased after transient exposure to *ascr#3* at the L1 stage and was maintained through adulthood (Figure 4D). In contrast, *odr-2p::gfp* expression was unaffected in other *odr-2*-expressing cells, including RIG and RME (Figure 4D). To examine whether upregulation of *odr-2* at the L1 stage is sufficient for the increased *ascr#3* avoidance in adults, we expressed *odr-2* cDNA in wild-type animals with the inducible expressed heat-shock promoter. Compared to animals with no heat-shock

(C) Ca^{2+} transients of ADL in response to 100 nM *ascr#3* exposure in *odr-2* mutants. The average traces of Ca^{2+} responses during two pulses of 100 nM *ascr#3* (left), heatmap of individual Ca^{2+} responses to *ascr#3* (bottom), and the maximum value of Ca^{2+} responses to 100 nM *ascr#3* (right) in naive (shown in blue) or pre-exposed (shown in red) *odr-2* mutants are shown. $n = 7$ (naive) and 14 (pre-exposed) each.

(D) Ca^{2+} transients of AVA in response to 100 nM *ascr#3* exposure. The average traces of Ca^{2+} responses during two pulses of 100 nM *ascr#3* (left), heatmap of individual Ca^{2+} responses to *ascr#3* (bottom), and the maximum value of Ca^{2+} responses to 100 nM *ascr#3* (right) in naive (shown in blue) or pre-exposed animals (shown in red) are shown. $n = 7$ (naive) and 8 (pre-exposed) each.

All error bars represent SEM. See also Figure S3.

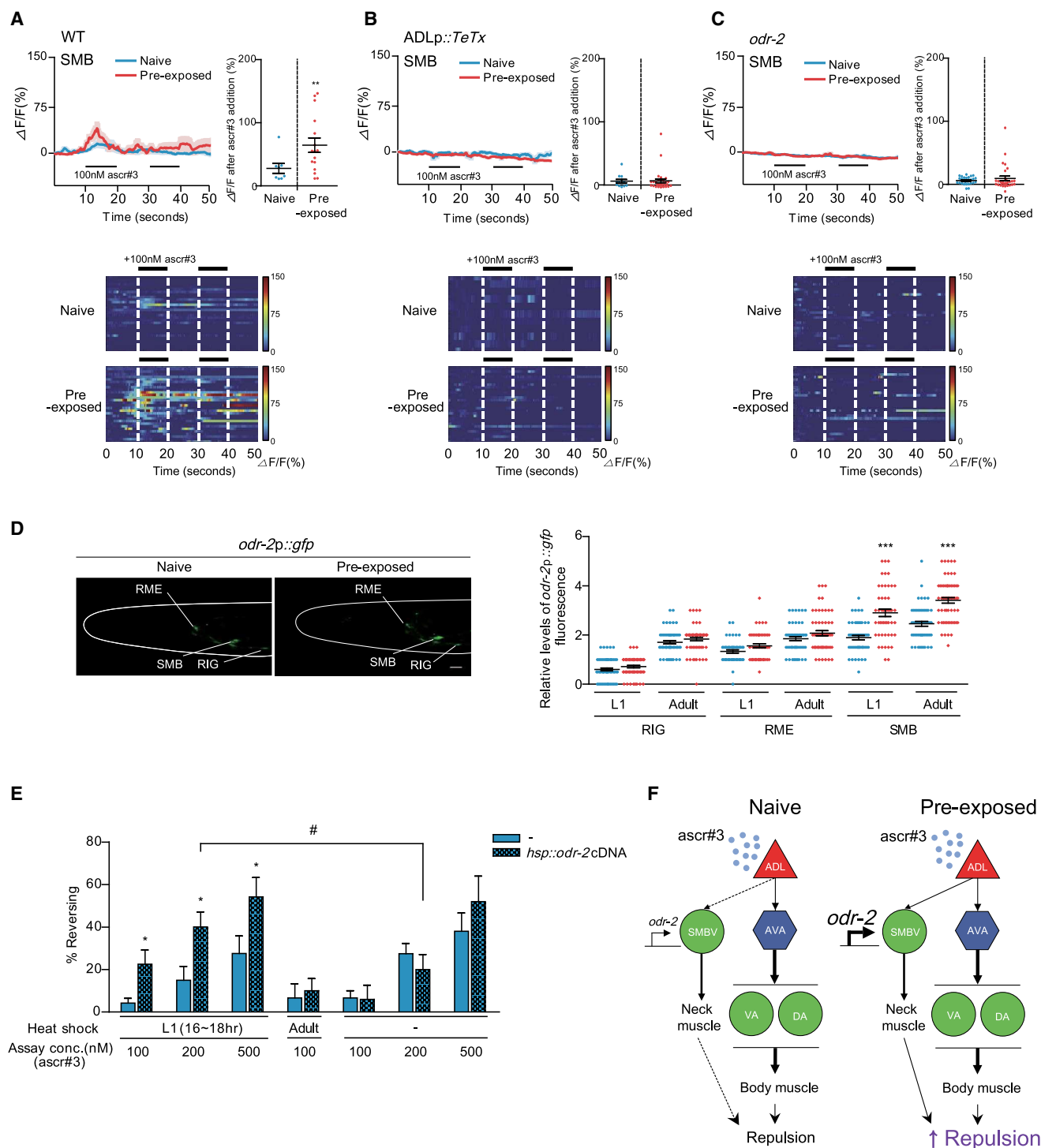


Figure 4. The SMB Motor Neurons Mediate Enhanced *ascr#3* Avoidance in *ascr#3*-Imprinted Animals

(A–C) Ca^{2+} transients of the SMB neurons in response to 100 nM *ascr#3* in wild-type animals (A), wild-type animals expressing ADLp::TeTx (B), or *odr-2* mutants (C). The average traces of Ca^{2+} responses during two pulses of 100 nM *ascr#3* (left), heatmap of individual Ca^{2+} responses to *ascr#3* (bottom), and the maximum value of Ca^{2+} responses to 100 nM *ascr#3* (right) in naive (blue) or pre-exposed animals [10] are shown. ** indicates different from naive at $p < 0.01$ by Student's *t* test. $n = 8$ –30 each.

(D) Representative images (left) and GFP quantification (right) of *odr-2p::gfp* expression in RIG, RME, and SMB of naive (shown in blue) and pre-exposed (shown in red) animals at the L1 or adult stages are shown. The images were taken of adults. *** indicates different from wild-type at $p < 0.001$ by one-way ANOVA with Bonferroni's post hoc test. $n = 50$ –60 each. The scale bar represents 10 μm .

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treated or those heat-shocked only at the adult stage, transient induction of *odr-2* gene activity at the L1 larval stage was sufficient to increase *ascr#3* avoidance in adults (Figure 4E). Thus far, our results are consistent with a model in which the increased *odr-2* expression in SMB of the *ascr#3*-imprinted animals causes functional changes in the ADL-SMB synaptic activities.

DISCUSSION

The well-defined nervous system of *C. elegans* with only 302 neurons mediates a broad spectrum of behaviors. However, these behaviors are plastic and can be modulated by the animal's experience [25, 26]. For example, *C. elegans* is able to constantly monitor and adapt to their environment by forming short-term and/or long-term memories in their hard-wired neural circuits (see a review by [27]). *C. elegans* has also been shown to imprint long-term memories of environments, including odor or pathogen exposure experienced during development, via largely uncharacterized mechanisms [15, 17]. Here, we show a novel imprinting paradigm in *C. elegans*, where exposure of newly hatched animals to the *ascr#3* pheromone increases adult neuronal activity in response to *ascr#3*, resulting in altered behavior. Because the long-lasting memory for *ascr#3* pheromone is formed during a critical period of early development and shapes the *ascr#3* avoidance behavior of adults, this learning and long-term memory process for the pheromone could be referred to as sensory imprinting [8]. Sensory imprinting is a form of long-term memory that occurs in many animals: for example, homing of salmon and neonatal attachment of rodents [28, 29]. This type of behavioral plasticity could be essential for animal survival in natural environments. *ascr#3* is the most abundant and potent aversive pheromone in *C. elegans* hermaphrodites and may represent conditions of overcrowding [7, 30, 31]. Thus, newly hatched worms may pair this specific *ascr#3* pheromone memory to their neuronal activity to promote searching behavior for more favorable environments as adults or to shape currently unidentified behavioral or physiological states for optimal fitness.

We previously showed that the ADL sensory neurons detect *C. elegans* pheromone *ascr#3* and drive *ascr#3* avoidance through their chemical synapses [7]. The ADL neurons directly connect to more than twenty post-synaptic neurons, and inspection of the anatomical wiring data, including the number of synapses, led us to hypothesize that *ascr#3* signals from ADL are transmitted to the three major post-synaptic target neurons, including the AIB, AVA, and AVD interneurons. In this work, we have found that, under non-*ascr#3* pre-exposed (naïve) conditions, the *ascr#3* signals are transmitted via the ADL-AVA synapses (Figure 4F). However, whereas transient exposure to *ascr#3* at the L1 larval stage does not appear to

affect this ADL-AVA synaptic transmission, this experience instead increases the ADL-SMB synaptic activities that are normally inactive in naïve animals and may account for the observed enhanced *ascr#3*-avoidance behavior in adult worms (Figure 4F).

The conclusion that the SMB neurons are the critical site for mediating *ascr#3* imprinting is supported by these findings: (1) the SMB neurons respond to *ascr#3* exposure in *ascr#3*-imprinted, but not in naïve, animals; (2) expression of *odr-2* that is necessary and partly sufficient for *ascr#3* imprinting is increased specifically in SMB of *ascr#3*-imprinted animals; and (3) *lim-4* mutants in which SMB functions are abolished fail to imprint *ascr#3* experience. Laser ablation of the SMB motor neurons causes increased reversal frequency and wave amplitude of forward locomotion [20, 22]. Because the SMB neurons innervate head and neck muscles and *ascr#3* exposure now activates the SMB neurons in *ascr#3*-imprinted animals, our results suggest that early *ascr#3* experience causes the SMB neurons to sensitize to subsequent *ascr#3* exposure and modulate backward movement via the alteration of head/neck muscle activity (Figure 4F). For example, in addition to body muscle contraction triggered by ADL-AVA synaptic transmission, head/neck muscle contraction due to enhanced SMB neuronal activities upon *ascr#3* imprinting may increase reversal rates further [24, 32] or the SMB neurons could directly activate backward motor circuits via interneurons, including the SAA neurons [21].

Our data show that *odr-2* is essential for *ascr#3* imprinting. *odr-2* encodes a novel protein distantly related to murine Ly6 lymphocyte antigen protein that is tethered to membrane by a GPI anchor [19]. In vertebrates, GPI-anchored proteins have roles in development and neurogenesis by regulating several signaling pathways, including Notch, Wnt, or transforming growth factor β (TGF- β) [33]. Moreover, a set of Ly6-related genes, including *Lynx1* and *Lynx2*, have been shown to be expressed in the nervous system and appear to be accessory components of nicotinic acetylcholine receptors, thus modulating their functions [34, 35]. Our results indicate that the ODR-2 GPI-anchored protein changes synaptic activity between ADL and SMB. ODR-2 proteins are localized on cell bodies and neuronal processes [36]. Thus, it is possible that ODR-2 may be co-localized with and regulate functions of neurotransmitter receptors in SMB.

Understanding how individual synapses are functionally or anatomically altered upon sensory imprinting is the essential first steps toward being able to dissect molecular and neuronal mechanisms underlying sensory imprinting and other forms of behavioral plasticity. Given that structures and functions of neural circuits are evolutionarily conserved, we expect that our work will lead to a general framework for understanding how circuits are modulated in higher animals, including humans.

(E) Percentage of reversal of adult transgenic animals expressing *hsp::odr-2* cDNA under heat shock at the L1 and adult stages. * indicates different from *hsp::odr-2* cDNA (–), and # indicates different from no heat shock at $p < 0.05$ by Student's *t* test. $n = 50$ –70 each.

(F) Model for circuit mechanisms underlying *ascr#3* avoidance in naïve or *ascr#3*-imprinted animals. In naïve animals, ADL detects *ascr#3* and transmits the signals to the post-synaptic AVA command interneurons and downstream VA/DA motor neurons. In pre-exposed animals, the increased expression of *odr-2* in SMB initiates synaptic transmission from ADL to SMB that mediate *ascr#3* signal transmission from ADL not only to AVA but SMB. SMB activation may serve to sensitize background for backward movement via excitation of head muscles.

In (A)–(E), error bars represent SEM. See also Figure S4.

STAR★METHODS

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

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SUPPLEMENTAL INFORMATION

Supplemental Information includes four figures and two movies and can be found with this article online at <http://dx.doi.org/10.1016/j.cub.2017.08.068>.

AUTHOR CONTRIBUTIONS

M.H., L.R., M.C.O., and A.R.J. performed the experiments; J.K., S.C., R.A.B., and H.C. provided reagents; M.H., L.R., M.C.O., Y.H.H., K.J.L., P.S., S.E.H., and K.K. analyzed and interpreted data; and M.H., L.R., P.S., S.E.H., and K.K. wrote the manuscript.

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STAR★METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Chemicals, Peptides, and Recombinant Proteins		
Levamisole	Sigma	Cat# 16595-80-5
ascr#3	This paper	
Critical Commercial Assays		
PDMS Sylgard 184 Silicone Elastomer Kit	Dow Corning	Product code: 1064291
Experimental Models: Organisms/Strains		
<i>odr-2(n2145)</i> V	<i>Caenorhabditis</i> Genetics Center	RRID: WB-STRAIN: CX2304
<i>odr-2(n2145)</i> V ; <i>IskEx606[odr-2Δ3p::odr-2cDNA]</i>	This paper	N/A
<i>odr-2(n2145)</i> V ; <i>IskEx607[flp-12Δ3p::odr-2cDNA]</i>	This paper	N/A
<i>odr-2(n2145)</i> V ; <i>IskEx608[lim-4p::odr-2cDNA]</i>	This paper	N/A
<i>lim-4(ky403)</i> X	<i>Caenorhabditis</i> Genetics Center	RRID: WB-STRAIN: CX3937
<i>IskEx218[sre-1p::GCaMP3; unc-122p::dsRed]</i>	This paper	N/A
<i>odr-2(n2145)</i> V ; <i>IskEx218[sre-1p::GCaMP3; unc-122p::dsRed]</i>	This paper	N/A
<i>IskEx597[nmr-1p::GCaMP3; unc-122p::dsRed]</i>	This paper	N/A
<i>IskEx113[flp-12Δ3p::GCaMP3; unc-122p::dsRed]</i>	This paper	N/A
<i>IskEx611[sre-1p::TeTx; unc-122p::gfp]; IskEx113[flp-12Δ3p::GCaMP3]</i>	This paper	N/A
<i>odr-2(n2145)</i> V ; <i>IskEx113[flp-12Δ3p::GCaMP3]</i>	This paper	N/A
<i>IskEx11[odr-2p::gfp; unc-122p::dsRed]</i>	This paper	N/A
<i>IskEx418[hsp::odr-2cDNA; unc-122p::gfp]</i>	This paper	N/A
<i>IskEx895 [avr-14p::gfp; unc-122p::dsRed]; IskEx28 [flp-12p::mcherry; unc-122p::gfp]</i>	This paper	N/A
<i>odr-2(n2145); IskEx895 [avr-14p::gfp; unc-122p::dsRed]; IskEx28 [flp-12p::mcherry; unc-122p::gfp]</i>	This paper	N/A
<i>egl-4(ks60)</i> IV	<i>Caenorhabditis</i> Genetics Center	RRID: WB-STRAIN: FK223
<i>sra-11(ok630)</i> II	<i>Caenorhabditis</i> Genetics Center	RRID: WB-STRAIN: RB816
<i>odr-3(n2150)</i> V	<i>Caenorhabditis</i> Genetics Center	RRID: WB-STRAIN: CX2205
<i>odr-7(ky4)</i> X	<i>Caenorhabditis</i> Genetics Center	RRID: WB-STRAIN: CX4
<i>odr-10(ky32)</i> X	<i>Caenorhabditis</i> Genetics Center	RRID: WB-STRAIN: CX32
<i>casy-1(tm718)</i> II	National BioResource Project	N/A
<i>casy-1(pe401)</i> II	<i>Caenorhabditis</i> Genetics Center	RRID: WB-STRAIN: JN414
<i>ttx-3(ks5)</i> X	<i>Caenorhabditis</i> Genetics Center	RRID: WB-STRAIN: FK134
<i>tdc-1(n3419)</i> II	<i>Caenorhabditis</i> Genetics Center	RRID: WB-STRAIN: MT13113
<i>IskEx610[npr-9p::GCaMP3; unc-122p::dsRed]</i>	This paper	N/A
Software and Algorithms		
AxioVision software	Zeiss	RRID: SCR_002677
ZEN 2010 Lite Edition software	Zeiss	RRID: SCR_013672
ImageJ	https://imagej.nih.gov/ij/	RRID: SCR_003070
MATLAB	MathWorks	RRID: SCR_001622
Prism 5.0	GraphPad Software	RRID: SCR_002798
Adobe Illustrator CS6	Adobe	RRID: SCR_014198

CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Dr. Kyuhyung Kim (khkim@dgist.ac.kr).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

C. elegans N2 strain was used as wild-type. Wild-type and transgenic animals were maintained at 20°C with abundant *E. coli* OP50 as food following standard conditions [37]. Day 1 (~24 hr after mid-L4 larval stage) young adult, hermaphrodite animals were used for experimentation, with the exception of the *odr-2p::gfp* quantification study in which the L1 stage animals were also assayed (Figure 4D).

METHOD DETAILS

Generation of *C. elegans* transgenic lines

odr-2 18a cDNA was amplified with a set of primers containing restriction sites, 5'-*AgeI* and 3'-*NotI* using PCR and inserted into pMC10 for performing the *odr-2* rescue experiment [19]. Upstream regulatory sequences, including *odr-2* (RIG, RME and SMB), *odr-2* (−377) (RIG and RME), *flp-12* (−339) (SMB), and *lim-4* (AWB, SAA, RID, RIV, RMD and SMB), were fused with *odr-2*cDNA to drive its expression in specific neurons [20]. To generate the *hsp::odr-2*cDNA transgene, the *hsp16.2* promoter was amplified from the *hsp16.2p::unc-3*cDNA transgene [38].

For Ca²⁺ imaging experiments of the AIB, AVD and AVA neurons, the promoter region of *sre-1p::GCaMP3* transgene was exchanged with *npr-9* (AIB) or *nmr-1* (AVD and AVA) promoters [7, 39]. *nmr-1p* was amplified from pCZGY1553 (a kind gift from Yishi Jin), and *npr-9p* was 2076 bp of upstream regulatory sequence amplified from genomic DNA by PCR. These promoters were ligated with GCaMP3 using the restriction sites, *HindIII* and *BamHI*.

Each transgenic *C. elegans* strain was generated by microinjection of the rescue construct (10 ng/μl) with *unc-122p::dsRed* (50 ng/μl). The *hsp::odr-2*cDNA construct was injected with the *unc-122p::gfp* marker (50 ng/μl).

Pheromone imprinting

For pheromone imprinting, two types of 35 mm noble agar plates seeded with *E. coli* OP50 were made: the negative control plate contained distilled water (naive) and the experimental plate contained pheromones (pre-exposed). Figures 1G and 1H utilized experimental plates containing 600 nM *ascr#3*, *ascr#2* and *ascr#5*, and 600 nM *ascr#3*, *ascr#2*, *ascr#5* and *icas#9*, respectively. All other assays in this study used experimental plates containing 600 nM *ascr#3* for imprinting. To pre-expose animals to *ascr#3*, five 1-day-old young adult hermaphrodites were placed onto each control and experimental plate for 2~3 hr until 30 to 50 eggs were laid. After 1 day (~26 hr), the naive or pre-exposed L1 stage animals were transferred onto 60 mm nematode growth media plates seeded with *E. coli* OP50 food and grown for 2.5 days until the young adult stage. For male imprinting experiments, we added ~20 males with L4 hermaphrodites for 1 day to mate and picked 5 adult hermaphrodites onto each naive and pre-exposed plates. For imprinting analysis of the next generation, 5 adults from naive or pre-exposed group (P0) were placed onto new control plates and allowed to lay 30~50 eggs (F1). The adults were then subsequently removed and the plates were incubated for 2.5 days at 20°C.

Post-dauer assay

Dauer formation assay was based on the protocol in [40]. 20 μL heat-killed OP50 was seeded onto 3.5 mm 600 nM *ascr#3* noble agar plates. 5~10 young adults were placed and discarded when over 50 eggs were obtained. The eggs were grown at 25°C for 68~72hr. Animals in dauer formation were picked onto new OP50 seeded 60 mm plates and grown at 20°C for 2~3 days. We assayed *ascr#3* avoidance with adult animals which were grown at 20°C as control.

Behavioral assay

The drop assay was performed as described with modifications [7]. All behavioral assays were performed in the absence of food and used 100 nM *ascr#3* diluted in M13 buffer (30 mM Tris-HCl [pH 7.0], 100 mM NaCl, 10 mM KCl), except Figure S1D which 200 mM glycerol was used [41]. To measure the percent of animals reversing upon *ascr#3* exposure, a young adult was transferred onto a plate without food and M13 buffer was dropped near the head of worm. If the worm failed to reverse in response to M13 after 10 s, *ascr#3* was dropped onto the front of the head and avoidance behaviors (reversal) were monitored for 10 s. Short or long reversals were defined as reversals with fewer than two head bends or more than two head bends, respectively [42]. Long reversals were counted as repulsion in this study. The imprinting index was calculated as the percentage of reversals of pre-exposed animals minus percentage of reversals of naive animals. Each assay tested the avoidance response of 10 young adult hermaphrodite animals, and at least 3 independent trials were performed.

In vivo calcium imaging

Calcium imaging experiments were performed as described previously [7], using microfluidics chips that were produced in-house with a custom made chrome mask and master mold [43]. Briefly, PDMS Sylgard 184 Silicone Elastomer Kit (Dow Corning) was solidified on the master mold at 70°C for 2 hr, and then was attached on a 24 × 24 mm glass coverslip using CUTE plasma equipment (Air) (FEMTOSCIENCE). The *C. elegans* transgenic strains used for calcium imaging experiments included *sre-1p::GCaMP3* (ADL), *flp-12* (−339)*p::GCaMP3* (SMB), *npr-9p::GCaMP3* (AIB) and *nmr-1p::GCaMP3* (AVA and AVD) were used [7]. Each animal was

exposed to fluorescent light for 1~2 min and the images were captured under fixed exposure time (100ms) for 1 min with Zeiss Axioplan microscope using a 40X objective and a Zeiss AxioCam HR. The images were analyzed by AxioVision software, ImageJ, and custom-written scripts in MATLAB [7].

Levamisole treatment

25 mM levamisole (Sigma) was used to paralyze animals. 25 mM levamisole was diluted in M9 buffer from 0.25 M levamisole. The animals were placed into the solution for about 1 min, during which pharyngeal pumping ceased, indicating paralysis.

Heat shock treatment

Heat shock treatment was performed as previously described [44]. After 16 hr from egg laying, animals were twice exposed to a 33°C heat shock for 30 min with a 1 hr recovery at 20°C in between. After the second heat shock treatment, the worms were incubated at 20°C until the adult stage and behavioral assays were performed (see above).

QUANTIFICATION AND STATISTICAL ANALYSIS

Representative images

Images in Figures 4D and 4G were acquired using a Zeiss LSM700 Confocal microscope and Zeiss LSM780 Confocal microscope, respectively. The images were taken and analyzed through ZEN 2010 and ImageJ.

GFP quantification

For GFP quantification, the animals were anaesthetized by using 1 M NaN₃ and placed on 2% agarose pad with a 24 × 24 mm coverslip. *odr-2p::gfp* strain was integrated using UV crosslinker (UVP). A minimum of 30 *odr-2p::gfp* animals were exposed to 100 mM ascr#3 or control plates at the L1 larval stage following the imprinting protocol described above and quantified for GFP fluorescence. GFP expression was scored under 40X objective Zeiss Axioplan microscopy by observing the brightness of the neuron cell body and process. The animals were observed under 40X objective Zeiss Axioplan microscopy and LSM700 Confocal microscope. The images were taken and acquired by AxioVision or Zen 2010. All strains were assayed in parallel in two independent experiments.

Statistical tests

All of statistics was analyzed using Prism 5.0. When only two groups were compared, two tailed unpaired Student's t test was used. One-way ANOVA test was utilized to evaluate variation among more than two groups of datasets. The post hoc tests used include Dunnett's test for multiple comparisons obtained at the same experimental condition, and Bonferroni's test when the multiple comparisons were made from data produced at different conditions. All error bars indicate SEM. Detailed information of each statistical analysis is described in the figure legends.