Sucre is an attractive feeding substance and a positive reinforcer for *Drosophila*. But *Drosophila* females have been shown to robustly reject a sucrose-containing option for egg-laying when given a choice between a plain and a sucrose-containing option in specific contexts. How the sweet taste system of *Drosophila* promotes context-dependent devaluation of an egg-laying option that contains sucre, an otherwise highly appetitive tantant, is unknown. Here, we report that devaluation of sweetness/sucrose for egg-laying is executed by a sensory pathway recruited specifically by the sweet neurons on the legs of *Drosophila*. First, silencing just the leg sweet neurons caused acceptance of the sucrose option in a sucrose versus plain decision, whereas expressing the channelrhodopsin CsChrimson in them caused rejection of a plain option that was "baited" with light over another that was not. Analogous bidirectional manipulations of other sweet neurons did not produce these effects. Second, circuit tracing revealed that the leg sweet neurons receive different presynaptic neuromodulations compared to some other sweet neurons and were the only ones with postsynaptic partners that projected prominently to the superior lateral protocerebrum (SLP) in the brain. Third, silencing one specific SLP-projecting postsynaptic partner of the leg sweet neurons reduced sucrose rejection, whereas expressing CsChrimson in it promoted rejection of a light-baited option during egg-laying. These results uncover that the *Drosophila* sweet taste system exhibits a functional division that is value-based and task-specific, challenging the conventional view that the system adheres to a simple labeled-line coding scheme.

sweet neurons | egg-laying | labeled-line coding | functional division | *Drosophila*

The taste systems of many animal species are known to possess a dedicated "channel" for detecting sugars, a class of chemicals that is highly nutritious. For example, mice have been shown to encode gustatory receptors that specifically sense sugars, and the taste neurons that express these sugar receptors on their tongues generally do not express receptors that sense chemicals of another taste modality (e.g., bitterness) (1–3). Furthermore, activation of these sugar-sensing taste neurons by artificial means can drive appetitive behaviors and act as a positive reinforcer for learning (10, 13, 14), while artificial activation of bitter-sensing neurons can induce rejection behaviors and be used as a punishment for learning (10, 13, 15). Interestingly, while these results suggest that *Drosophila* sweet neurons and their mammalian counterparts have some shared properties, subsequent studies suggest that significant differences exist between them, too. First, the *Drosophila* genome appears to encode many more sweet receptors than mouse genome does (12, 16–23). Second, *Drosophila* sweet neurons appear to be able to detect some chemicals that belong to another taste modality [e.g., acetic acid (AA)] (24–27). Third, *Drosophila* sweet neurons can be found on several body parts (e.g., proboscis and legs) (8, 12, 18, 20, 23, 28–30). Interestingly, sweet neurons on different body parts of *Drosophila* do not promote identical behavioral outputs (8, 20, 23, 24, 28, 29). For example, labellar sweet neurons and esophageal sweet neurons on the proboscis have been shown to promote proboscis extension reflex (PER) and ingestion, respectively, whereas leg sweet neurons have been shown to promote PER and slowing down of locomotion (8, 12, 28, 29). Collectively, these results suggest that in contrast to the apparent homogeneity of sweet neurons in some mammals, a functional division exists among *Drosophila* sweet neurons, although the different behavioral responses promoted by different *Drosophila* sweet neurons generally appear appetitive in nature.

**Significance**

Sweet taste neurons in both *Drosophila* and mice are often thought to be hardwired to promote appetitive responses and signal the presence of reward. Here, exploiting *Drosophila* females’ robust rejection of sucrose substrates over plain ones during egg-laying in one specific context, we discovered that *Drosophila* sweet neurons can be divided into at least two anatomically and functionally distinct groups that confer positive and negative values, respectively, to options during egg-laying. This discovery reveals one design feature of the *Drosophila* sweet taste system that allows sweetness/sugars to be valued differently according to context and animals’ behavioral goal (i.e., feeding versus egg-laying), pointing to a level of flexibility and sophistication that is not seen in the system’s mammalian counterparts.

Author contributions: H.-L.C. and C.-H.Y. designed research; H.-L.C. and D.M. performed research; U.S. contributed new reagents/analytic tools; H.-L.C., D.M., U.S., and C.-H.Y. analyzed data; and H.-L.C. and C.-H.Y. wrote the paper.

The authors declare no competing interest.

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Edited by L. B. Vosshall, Laboratory of Neurogenetics and Behavior, Rockefeller University, New York, NY; received June 9, 2021; accepted November 7, 2021.
In this work, we report yet another striking feature of Drosophila sweet neurons that sets them apart from their mammalian counterparts, namely a functional division that is value-based and task-specific. We discovered this by taking advantage of a context-dependent but highly robust sugar rejection behavior exhibited by egg-laying females (31–34). Previous studies have shown that when selecting for egg-laying site in a small enclosure (dimension \(16 \times 10 \times 18 \text{ mm}\), Drosophila readily accept a sucrose-containing agarose for egg-laying when it is the sole option but strongly reject it when a plain option is also available (31, 32). Importantly, silencing their sweet neurons causes the females to no longer reject the sucrose option when choosing between the sucrose versus plain options (31, 32). Thus, in addition to promoting appetitive behaviors and acting as a positive reinforcer, activation of sweet neurons on an egg-laying option can also decrease the value of such an option (thereby causing its rejection over an option that does not activate sweet neurons). These observations not only suggest the existence of an apparent “antiappetitive” role of Drosophila sweet neurons when the task of animals is to select for egg-laying sites but also raise a key question as to whether such counterintuitive, value-decreasing property of sweetness detection during egg-laying may be 1) solely an emergent property of specific neurons in the brain that respond similarly to all peripheral sweet neurons but are sensitive to animals’ behavioral goal and context or 2) carried out by specific sweet neurons at the periphery and then transmitted into the brain via a unique neural pathway activated by these neurons. To disambiguate between these possibilities, we genetically targeted different subsets of sweet neurons to assess their circuit properties as well as their behavioral roles as the animals decided in either a regular or a virtual sweet versus plain decision during egg-laying, taking advantage of a high-throughput closed-loop optogenetic stimulation platform we developed recently. Our collective results support the second scenario and suggest that the value-decreasing property of sweetness/sucrose is conveyed specifically by the sweet neurons on the legs of Drosophila—and not by other sweet neurons—and the unique postsynaptic target(s) of the leg sweet neurons that send long-range projections to the superior lateral protocerebrum (SLP) in the brain. These results reveal a previously unappreciated functional and anatomical division of the Drosophila sweet taste neurons that is both task-specific and value-based, pointing to a level of complexity and sophistication that seems unmatched by their mammalian counterparts so far.

**Results**

**Sweet Neurons at Different Locations Contribute Differentially to Devaluing the Sweet Option for Egg-laying.** We first reconfirmed that Drosophila showed context-dependent sucrose rejection for egg-laying. Indeed, when tested in our high-throughput apparatus, wild-type (\(w^{1118}\)) females accepted the sucrose-containing agarose when given two sucrose-containing agaroses but rejected the sucrose option when the other option was sucrose-free (Fig. 1A and B) (31, 32). In addition, when we used either the Gr64f-GAL4 or the Gr64f-H\(^{A}\) drivers (18, 35, 36) to silence virtually all their peripheral sweet neurons, the affected females no longer rejected the sucrose agarose in the same sweet versus plain task (Figs. 1C and 2A). Thus, sweet neurons play a critical role in devaluing an agarose that contains sucrose in our decision task, thereby promoting its rejection over a plain agarose.

Previous studies have shown that cell bodies of sweet neurons can be found at different locations in a fly’s body such as its proboscis (i.e., the labellum and the esophagus), legs, and brain (8, 18, 20, 28, 30). To begin to assess whether sweet neurons at different locations contribute similarly to devaluing the sweet agarose for egg-laying, we collected several GAL4 drivers, each of which labeled a different combination of sweet neurons (Fig. 2; SI Appendix, Fig. S1 A–F). We then attempted to deduce the roles of different sweet neurons by correlating the expression patterns of these drivers with the phenotypes that they produced.

We found that inactivation of sweet neurons labeled by either Gr61a-GAL4 (36) or Gr64a\(^{GAL4}\) (18, 35) caused females to switch from rejecting to preferring the sweet agarose in the sweet versus plain task (Fig. 2B and C; SI Appendix, Fig. S1G). Because these two GAL4s labeled the same sweet neurons on the legs and the LSO (labral sense organ, on the esophagus) but differed in their labeling of neurons at other locations (Fig. 2B and C; SI Appendix, Fig. S1 B and C), these results suggest that sweet neurons on the legs, the LSO, or possibly both might be more critical for promoting the rejection of the sweet agarose.

![Fig. 1. Drosophila rely on sweet neurons to reject the sucrose option in a sucrose versus plain task.](https://doi.org/10.1073/pnas.2110158119)
However, the importance of the LSO sweet neurons was called into question as inactivating these neurons by using either Gr43aGAL4 (20, 37) or Gr64a-GAL4 (36) (a driver that differs from the “knocked-in” Gr64aGAL4)—both of which labeled LSO neurons—did not significantly impact sweet rejection (Fig. 2 D and E). Lastly, we found that the labellar sweet neurons may...
not promote sweet rejection either, as while inactivating the Gr5a-GAL4–expressing neurons reduced sweet rejection, it did so to a lesser degree than inactivating the Gr64aGAL4 neurons (Fig. 2 C and F). Gr5a-GAL4 labeled many labellar sweet neurons (16) but fewer sweet neurons on the legs (and essentially no other sweet neurons) (Fig. 2F, SI Appendix, Fig. S1F). Thus, this result is more consistent with the view that the weaker rejection exhibited by the Gr5a-GAL4 > Kir2,1 animals may be because fewer leg sweet neurons of theirs were inactivated.

Taken together, these results suggest that sweet neurons at different locations do not contribute equally to promoting rejection of the sweet agarose in the sweet versus plain decision and that those on the legs likely play more significantly a role than the rest.

Sweet Neurons on the Legs Are Solely Necessary for Devaluing the Sweet Option for Egg-Laying. To confirm that the sweet neurons on the legs are essential for promoting sweet rejection in the sweet versus plain task, we next used three intersectional approaches to manipulate different subsets of the neurons labeled by Gr64aGAL4. Gr64aGAL4 labeled the leg sweet neurons strongly and a few sweet neurons on the LSO and in the brain and, importantly, produced a very strong lack-of-sweet-rejection phenotype when used to inactivate neurons (Fig. 2C, SI Appendix, Fig. S1 C and G). First, we sought corroboration that the inability of Gr64aGAL4 > Kir2,1 animals to reject sweet option was due to inactivation of their leg sweet neurons. To that end, we found a transgene combination, Gr43a-LexA>Gal80, that can block GAL4-dependent expression in the sweet neurons on the legs (and on the LSO), but not on the labellum (SI Appendix, Fig. S2A and C; Fig. 6D). Introducing this combination into the Gr64aGAL4 > Kir2,1 animals reverted them from not rejecting the sweet agarose to clearly rejecting it again in the sweet versus plain decision (SI Appendix, Fig. S2B). Moreover, the same combination also reverted the Gr64f-GAL4 > Kir2,1 and the Gr5a-GAL4 > Kir2,1 animals from not rejecting to clearly rejecting the sweet option, too (SI Appendix, Fig. S2D). These results support that the leg, but not labellar, sweet neurons are essential for devaluing the sweet agarose.

In our second approach, we directly ruled out a significant contribution from the LSO and the brain-intrinsic sweet neurons labeled by Gr64aGAL4. We created flies that contained the following transgenes: Otd-nls::FLP, UAS-FRT-mCherry-FRT-Kir2,1, and Gr64aGAL4. Because Otd-nls::FLP (38) is expressed in only the brain-intrinsic neurons and a few subsets of sensory neurons that project into the brain (e.g., LSO sweet neurons) but not in any sensory neurons on the legs, this transgene combination allowed Kir2,1 to be expressed in only the Gr64aGAL4–expressing sweet neurons in the brain and LSO (Fig. 3A). These animals clearly rejected the sweet agarose in the sweet versus plain task (Fig. 3B), in stark contrast to the lack of sweet rejection exhibited by the Gr64aGAL4 > Kir2,1 animals (Fig. 2C; SI Appendix, Fig. S1G), suggesting that neither the brain-intrinsic nor the LSO sweet neurons labeled by Gr64aGAL4 are essential for devaluing the sweet agarose.

In our third approach, we restricted Kir2,1 to be expressed in only the Gr64aGAL4–expressing sweet neurons on the legs by putting Otd-nls::FLP together with Tub-FRT-stop-FRT-Gal80, UAS-Kir2,1, and Gr64aGAL4 (Fig. 3C). These animals no longer rejected the sweet agarose for egg-laying (Fig. 3D), directly
demonstrating the requirement of the leg sweet neurons in promoting sweet rejection in the sweet versus plain decision task. We obtained a similar result when we used a different transgene combination to inhibit just the Gr64a-GAL4-expressing neurons on the legs (Fig. 3E).

Thus, results from these intersectional approaches strongly suggest that sweet neurons on the legs, but not in other locations, are solely necessary for females to devalue a sucrose-containing agarose in our sweet versus plain task.

A Closed-Loop Platform to Examine the Impact of Artificial Stimulation of Sweet Neurons During Egg-Laying. Next, we wished to determine whether artificial activation of the leg sweet neurons on a plain agarose is sufficient to decrease its value for egg-laying. One approach is to express the channelrhodopsin CsChrimson (39) in the leg sweet neurons and assess how the flies would choose between two plain agaroses, one of which is “baited” with light. We designed and built SkinnerSys, a high-throughput platform that can illuminate flies in closed loop during egg-laying (Fig. 4A and B; Movies S1 and S2). Briefly, SkinnerSys consists of three major components: 1) SkinnerTrax, code we previously developed for real-time tracking and delivering light to multiple individual animals in closed loop (15, 40); 2) an apparatus that contains 40 individual two-choice arenas; and 3) a custom printed circuit board (PCB) that allows LED illumination of different arenas to be independently controlled (Fig. 4A). Thus, SkinnerSys can assay in a highly parallel manner how individual females respond to activation of specific neurons of interest as they explore different options for egg-laying. Importantly, because SkinnerSys can deliver light pulses to animals according to their position in real time, we can control optogenetic activation of neurons of interest with desired spatial precision (Fig. 4B).

As a proof-of-concept experiment, we first used Gr64f-GAL4 to express CsChrimson in all peripheral sweet neurons and examined how the females would choose when given a plain agarose on which light is consistently off versus a plain agarose on which light is turned on only when the fly is on the agarose (Fig. 4B). We found that Gr64f-GAL4 > CsChrimson animals not fed with retinal showed no clear biases for either option (Fig. 4C and D). In contrast, retinal-fed animals robustly rejected the light-on agarose over the light-off agarose (Fig. 4C and D). This is a demonstration that artificial activation of sweet neurons on a plain option was sufficient to drive its rejection for egg-laying over another plain option on which such activation did not occur. Positional heatmaps revealed that...
retinal-fed Gr64a\text{GAL4} > CsChrimson females had a slight but statistically significant positional preference for the light-on substrate (Fig. 4C and E), suggesting that while activation of all sweet neurons on an option can decrease its value for egg-laying, it still confers a mild appetitive quality to flies when they are not laying eggs.

Activation of the Leg Sweet Neurons Is Sufficient to Decrease the Value of a Plain Option for Egg-Laying. To test whether optogenetic activation of the leg sweet neurons on a plain option can cause the animal to devalue it for egg-laying, we first stimulated Gr64a\text{GAL4} expressing neurons in closed loop. As described earlier, Gr64a\text{GAL4} labeled sweet neurons on the legs strongly, as well some LSO and brain-intrinsic sweet neurons, but no labellar sweet neurons (Fig. 2C; SI Appendix, Fig. S1C). Similar to Gr64f:GAL4 > CsChrimson females, retinal-fed Gr64a\text{GAL4} > CsChrimson females strongly rejected the light-baited plain agarose in the light-on versus light-off task (Fig. 5A and B). Curiously, however, positional heatmaps showed that these animals preferred to spend time away from the light-on agarose, too (Fig. 5A and C).

We next restricted CsChrimson to be expressed in only the leg sweet neurons by combining the following transgenes: Gr64a\text{GAL4}, UAS-CsChrimson, Otd-nls::FLP, and Tub-FRT-stop-FRT-GAL80 (Fig. 3C). When fed with retinal, these flies rejected the light-option over the light-off one for egg-laying (Fig. 5D and E), demonstrating the sufficiency of optogenetic activation of just the leg sweet neurons in promoting rejection of an option on which such activation occurs over an option on which it does not. Interestingly, these animals no longer avoided spending time on the illuminated option (Fig. 5D and F). Further, males, virgins, and non-egg-laying mated females of the same genotype all showed similar positional indifference between the light-on versus the light-off options (SI Appendix, Fig. S3), so we observed devaluing of the light-baited option only in egg-laying females and only for egg-laying.

Next, we examined how females with CsChrimson expressed in only their leg sweet neurons behaved when both options were baited with light (Fig. 5G and H). This experiment is important because optogenetic activation of the leg sweet neurons may simply shut down egg-laying as opposed to devaluing an option. We found that females readily laid eggs on both...
Some of the Labellar Sweet Neurons Have a Role in Increasing the Value of the Sweet Agarose for Egg-Laying. We have so far focused on assigning the “value-decreasing function” of sweet neurons during egg-laying to those on the legs. Here, we address the peculiar results that whereas inhibiting neurons using different pan-sweet-neuron drivers caused females to become indifferent between the sweet and plain options (Figs. 1C and 2A), inhibiting neurons using either Gr64a-GAL4 or Gr61a-GAL4 caused preference for the sweet option over the plain one (Fig. 2B and C; SI Appendix, Fig. S1G). These findings raise the intriguing possibility that some of the sweet neurons that are not labeled by these two GAL4 drivers (e.g., labellar and some pharyngeal sweet neurons) may act to increase the value of a sweet option over a plain one when activated; however, their impact on the sweet versus plain decision may normally be suppressed or obscured by the dominant value-decreasing function of the leg sweet neurons.

To test this idea, we first asked whether some of the pharyngeal neurons may be responsible for promoting the sweet preference exhibited by the Gr64a-GAL4 > Kir2.1 flies. We simultaneously silenced Gr64a-GAL4- and Gr43a-GAL4-expressing neurons and found that these “double inhibition” animals still exhibited a preference for the sweet option (Fig. 6A). Because Gr43a-GAL4 labeled virtually all pharyngeal sweet neurons (20, 37) (Fig. 2E), this result rules out a significant role of these neurons in promoting the peculiar sweet preference we observed. Next, we simultaneously silenced Gr64a-GAL4- and Gr64f-GAL4-expressing neurons and found that the sweet preference of these animals reduced significantly (Fig. 6B), suggesting that the labellar sweet neurons may play a role in promoting sweet preference as they were clearly...
labeled by Gr64f-GAL4 (Fig. 2A). Indeed, simultaneously inhibiting Gr64a-GAL4, and Gr5a-GAL4-expressing neurons or simultaneously inhibiting Gr64d-GAL4, and Gr5d-GAL4-expressing neurons significantly reduced the sweet preference exhibited by the Gr64a-GAL4 > Kir2.1 animals (Fig. 6C), too. Like Gr5a-GAL4, Gr5d-GAL4 also labeled many labellar sweet neurons (18, 35).

Next, we asked whether animals may prefer a plain agarose on which their labellar sweet neurons are optogenetically activated. To restrict CsChrimson to be present only in their labellar sweet neurons, we again introduced Gr3a-LexA>GAL80, a transgene combination that suppressed GAL4-dependent expression in virtually all leg sweet neurons but spared the labellar ones, into Gr5a-GAL4 > CsChrimson animals (Fig. 6D). Interestingly, retinal-fed animals that carried all these transgenes indeed showed a statistically significant preference to lay eggs, as well as to spend time, on the illuminated option over the unilluminated one (Fig. 6E and F).

Collectively, these results support the idea that activation of some of the labellar sweet neurons on an option increases its value for egg-laying, but such value-increasing function has little behavioral impact when the leg sweet neurons are activated on the same option. This result is consistent with the recent findings that some of the labellar sweet neurons can be activated by AA and promote egg-laying preference for AA in an AA versus plain task (24).

The Leg Sweet Neurons Receive Different Presynaptic Modulations from the Labellar Sweet Neurons. Having found that the leg sweet neurons were uniquely critical for sweet rejection during egg-laying, we next explored whether information collected by the leg sweet neurons might be processed differently from that collected by the rest of the sweet neurons. We first assess specific presynaptic modulations received by sweet neurons by using GFP Reconstitution Across Synaptic Partners (GRASP), a tool that can detect whether two groups of neurons have direct contacts as well as the directionality of such contacts (41, 42). Previous studies have reported that axons of sweet neurons labeled by Gr5a-LexA—a driver that labels both the labellar and the leg sweet neurons (29)—receive direct presynaptic inputs from the dopaminergic (DA), octopaminergic (OA), and GABAergic neurons in the subesophageal zone (SEZ) and that these inputs can modify sugar-activated feeding responses (43–46). We replicated these experiments by using Gr64f-GAL4 to label virtually all peripheral sweet neurons and indeed detected GRASP signals between axons of Gr64f-GAL4-expressing neurons and processes from the DA, OA, and GABAergic neurons in the SEZ (SI Appendix, Fig. S4).

In contrast, axons of the leg sweet neurons (labeled by Gr64a-GAL4) did not appear to directly contact either the DA or the OA neurons (SI Appendix, Fig. S5), although they did form bidirectional contacts with the GABAergic neurons in the ventral nerve cords (VNCs) and possibly also the SEZ (SI Appendix, Fig. S5). These results suggest that the DA and the OA systems do not directly modulate the output of—or do they receive any direct input from—the leg sweet neurons, revealing one difference in how information collected by the leg versus the labellar sweet neurons is processed at the first stage of information relay.

The Leg Sweet Neurons Have a Direct Postsynaptic Target That Is Not Shared by the Rest of the Sweet Neurons. To examine whether information collected by the leg sweet neurons and the rest of the sweet neurons might be routed differently into higher brain areas, we next used trans-Tango to assess their postsynaptic targets. trans-Tango is a circuit-tracing technique that can label the direct postsynaptic targets of specific GAL4-expressing neurons of interest (47). In fact, the developers of trans-Tango were among the first to show that the Gr64f-GAL4—expressing sweet neurons have numerous direct postsynaptic partners in the brain (47). We replicated this experiment and found that trans-Tango tracing of Gr64f-GAL4 indeed labeled neurons that elaborated processes in the SEZ as well as neurons that projected to the superior medial protocerebrum (SMP) and the SLP in the brain (Fig. 7A).

We then compared trans-Tango–traced targets of the Gr64a-GAL4, versus Gr64e-GAL4—expressing neurons (Fig. 7B and C). We chose these two GAL4s for comparison because while they labeled the same brain-intrinsic and LSO sweet neurons, only Gr64e-GAL4 labeled the leg sweet neurons (Fig. 2C and D; SI Appendix, Figs. 6C–D and S6a). These GAL4s produced quite similar trans-Tango patterns (and both had targets that were not seen when we trans-Tango traced Gr64f-GAL4, as Gr64f-GAL4 did not label brain-intrinsic sweet neurons). However, one major difference was evident: whereas the long-range SLP projection was present among the targets of Gr64a-GAL4—expressing neurons, it was absent among the targets of Gr64e-GAL4—expressing neurons (Fig. 7B and C). This result suggests that the SLP projection may be unique to the postsynaptic targets of the leg sweet neurons. Consistent with this idea, the SLP projection was also present when we traced the targets of Gr5a-GAL4—expressing sweet neurons (Fig. 7D), like Gr64a-GAL4, Gr61a-GAL4 also labeled many leg sweet neurons (Fig. 2B; SI Appendix, Fig. S1B).

To assess whether the SLP projection might indeed be unique to the postsynaptic partners of the leg sweet neurons, we trans-Tango traced targets of Gr64f-GAL4—expressing neurons in intact versus leg-amputated animals (Fig. 7E and E′). A previous study has shown that leg amputation can cause axons of leg taste neurons to degenerate (8), and we thus reasoned it may also prevent the targets of leg taste neurons from being traced. Indeed, we found that axons of the leg sweet neurons were barely visible in the leg-amputated Gr64f-GAL4 > trans-Tango animals (Fig. 7E′), and, importantly, the SLP projection traced by trans-Tango was eliminated, while the SMP projection remained intact (Fig. 7E′). We observed a similar absence of the SLP projection when we amputated the legs of Gr5a-GAL4 > trans-Tango flies (SI Appendix, Fig. S6b). These results suggest that among all sweet neurons, only the leg neurons have postsynaptic partners that send prominent projections to the SLP.

To rule out that the trans-Tango–traced SLP projection may be more sensitive to leg amputation for nonspecific reasons, we attempted tracing by restricting GAL4 activity to only the leg sweet neurons (Fig. 7F) and confirmed that postsynaptic partners of the leg sweet neurons sent a clear long-range projection to SLP (Fig. 7F), as well as to the SMP and the SEZ, two areas that were targeted by postsynaptic partners of some other sweet neurons (Fig. 7F).

In our last set of experiments, we addressed the potential functional significance of the SLP-projecting targets of the leg sweet neurons. A group of projection neurons known as TPN2 has recently been shown—which we confirmed (SI Appendix, Fig. S7A–C and Movie S3)—to be a direct postsynaptic target of the leg sweet neurons (13). Cell bodies and dendrites of TPN2 are located in the VNCs, but their long axons ascend and target the SLP in the brain prominently (Fig. 8; SI Appendix, Fig. S7A) (13). To assess the role of these SLP-projecting TPN2 neurons in egg-laying decisions, we selectively 1) inhibited them and assessed how the animals chose in the sucrose versus plain task and 2) expressed CsChrimson in them and assessed how the animals chose in the light-on versus light-off task. We found that females with reduced activities of TPN2 showed reduced rejection of sucrose in the sucrose versus plain task, while retinal-fed animals with CsChrimson expressed in their TPN2 clearly rejected the light-baited agarose for egg-laying, despite having a positional
preference for it, in the light-on versus light-off task (Fig. 8).

[Curiously, while virgins and non–egg-laying mated females showed positional preference for light, too, males did not (SI Appendix, Fig. S7D).] Thus, the egg-laying decision phenotypes of TPN2 parallel that of bidirectional manipulations of the leg sweet neurons.

Fig. 7. Postsynaptic partners of the leg sweet neurons have a unique long-range projection in the brain. (A–D) Representative images showing sweet neurons labeled by different GAL4s (green) and their traced postsynaptic partners (red). (A) Gr64f-GAL4-expressing neurons and their partners, (B) Gr64a-GAL4-expressing neurons and partners, (C) Gr64a-GAL4-expressing neurons and partners, (D) Gr61a-GAL4-expressing neurons and partners. Pink arrows: processes of brain-intrinsic sweet neurons not labeled by Gr64f-GAL4. White arrows: SLP-targeting projections. (E and E') Representative images showing Gr64f-GAL4-expressing neurons and their traced postsynaptic partners in intact (E) and leg-amputated flies (E'). (F) Representative images showing specifically the Gr64aGAL4-expressing leg neurons and their partners (scale bars, 60 μm).
Collectively, our results support the view that information collected by the leg sweet neurons is processed differently from that by the rest of the sweet neurons and that the SLP is a candidate area for housing neurons that process option values during egg-laying site selection.

**Discussion**

In this work, we present results suggesting that sweet neurons on different body parts of *Drosophila* do not contribute equally to determining how a sweet option should be valued during egg-laying. First, whereas inactivating their leg sweet neurons abolished females’ plain preference in the sweet versus plain decision, inactivating the rest of the sweet neurons did not. Second, optogenetic activation of just the leg sweet neurons on a plain agarose was sufficient to cause its rejection over another plain agarose on which such activation was withheld. In contrast, selective activation of just the labellar sweet neurons promoted preference for the plain agarose on which it occurred. Third, bidirectional manipulations of the activities of TPN2—a postsynaptic partner unique to the leg sweet neurons—produced egg-laying decision phenotypes that were qualitatively similar to those of the leg sweet neurons. These findings suggest strongly that the *Drosophila* sweet taste system possesses a functional and anatomical division that is valued-based and task-specific.

How do our findings add to our understanding of the *Drosophila* sweet taste system in general? First, our findings suggest that sweetness is not always appetitive; rather, it is a cue that can bidirectionally modify the value of an egg-laying option. Second, our findings suggest that sweet neurons on the legs can engage a value-modifying circuit may allow animals more flexibility in adjusting the value of a sweet option according to decision contexts. For example, while flies prefer the plain option in certain two-choice contexts (31, 32, 48), they prefer the sweet option when laying eggs in a significantly larger enclosure (49). Perhaps larger enclosures promote sweet preference either by enhancing the output of the value-increasing pathway potentially mediated by the labellar sweet neurons or by dampening the value-decreasing one mediated by the leg sweet neurons.

Finally, where might the sweet versus plain egg-laying decision be made in the *Drosophila* brain? While egg-laying preference has been frequently used for studying the function of different sensory systems (24, 50–61), the central circuit that assigns, retains, and compares values of egg-laying options has yet to be elucidated. Two recent studies suggest that the descending egg-laying command neurons (oviDNs) must be an integral component of the decision circuit as they not only are capable of triggering egg-laying when directly activated but also express a [Ca^{2+}] signal that tracks the relative value of an egg-laying option (33, 48). On the other hand, while successful rejection of the otherwise acceptable sweet option in the sweet versus plain task requires that the animals hold in memory their recent encounters focusing on a limited set of behavioral tests. For example, our findings suggest that some of the sweet neurons can even promote genuine avoidance, albeit in an artificial setting (Fig. 5 A–C), hinting at the potential existence of another function of sweet taste neurons that has yet to be explored.
with the preferred plain option, our previous studies have ruled out a critical role of the learning-and-memory center mushroom bodies in this decision (32). Our current results put forward the SLP as a candidate for housing the neurons that signal values to the decision-maker (Figs. 7 and 8). Given that dendrites of oVd1Ns arborize in the SMP (33), scrutinizing the electron microscopy (EM) connectome (62) for neurons that relay information from the SLP to SMP may help uncover some of the components that convert sweetness into a value-modifying signal during egg-laying.

Methods

In this work, we used various fly strains we obtained from colleagues and the Bloomington Stock Center, several published methods, and a closed-loop stimulation setup (SkinnerSys) we developed to assess the behavioral roles of neurons of interest during egg-laying site selection. Origins of the fly strains and more detailed descriptions of the published methods as well as SkinnerSys can be found in the SI Appendix. The code we used for closed-loop stimulation and the chamber and PCB design files for SkinnerSys can be found at https://github.com/ulrichstern/SkinnerTrax.

Data Availability

All study data are included in the article and/or SI Appendix.

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