Antagonistic serotonergic and octopaminergic neural circuits mediate food-dependent locomotory behavior in Caenorhabditis elegans

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ABSTRACT
Biogenic amines are conserved signaling molecules that link food cues to behavior and metabolism in a wide variety of organisms. In the nematode C. elegans, the biogenic amines serotonin (5-HT) and octopamine regulate a number of food-related behaviors. Using a novel method for long-term quantitative behavioral imaging, we show that 5-HT and octopamine jointly influence locomotor activity and quiescence in feeding and fasting hermaphrodites, and we define the neural circuits through which this modulation occurs. We show that 5-HT produced by the ADF neurons acts through the SER-5 receptor in muscles and neurons to suppress quiescent behavior and promote roaming in fasting worms, whereas 5-HT produced by the NSM neurons acts on the MOD-1 receptor in AIY neurons to promote low-amplitude locomotor behavior characteristic of well-fed animals. Octopamine, produced by the RIC neurons, acts through SER-3 and SER-6 receptors in SIA neurons to promote roaming behaviors characteristic of fasting animals. We find that 5-HT signaling is required for animals to assume food-appropriate behavior, whereas octopamine signaling is required for animals to assume fasting-appropriate behavior. The requirement for both neurotransmitters in both the feeding and fasting states enables increased behavioral adaptability. Our results define the molecular and neural pathways through which parallel biogenic amine signaling tunes behavior appropriately to nutrient conditions.

INTRODUCTION
The ability to alter behavior in response to nutritional cues is a crucial trait for organisms to adapt to changing environments. For example, many animals respond to scarcity of food by adopting quiescent behaviors such as hibernation or diapause (Nelson et al., 2014; van Breukelen and Martin, 2015). Hibernation is a long-term behavior characterized by suppressed movement and metabolism that occurs in response to low food availability and decreased temperature (Carey et al., 2003). How the nervous system initiates and sustains quiescent behavior during hibernation is poorly understood (Andrews, 2007).

In many organisms, the linking of behavior and metabolism (Chase and Koelle, 2007; Tecott, 2007; Roeder, 2005) to nutrition cues is performed in part through a widely conserved class of signaling molecules called biogenic amines. In mammals, biogenic amines regulate appetite, mood, weight, and other aspects of physiology (Lam, 2006). Disorders such as anorexia, obesity, and depression, whose symptoms include altered food intake, are associated with dysregulation of the biogenic amine serotonin (5-HT) (Curran and Chalasani, 2012). These disorders can be treated with drugs that increase the amount of 5-HT acting at the synapse, but how 5-HT levels relate to the complex behavioral and metabolic manifestations of these disorders is not fully understood. Thus, it is important to understand the mechanisms by which biogenic amines link environmental stimuli to behavior and metabolism.

The nematode C. elegans is a powerful model organism to dissect the neural circuits linking food, behavior, and physiology owing to its compact nervous system, manipulable genetics, and a host of food-related behaviors (Sengupta, 2013; Hart, 2006). Its nervous system signals in part through the biogenic amines 5-HT, dopamine, tyramine, and octopamine. Here we focus on the behavioral effects of 5-HT and octopamine in relation to food levels.
5-HT is involved in the modulation of multiple *C. elegans* behaviors in relation to food levels (Liang et al., 2006; Ségalat et al., 1995). During feeding, 5-HT is released and promotes locomotory slowing, thereby increasing the probability of the animal remaining on a food source (Sawin and Horvitz, 2000). In addition, 5-HT increases pharyngeal pumping (feeding) and egg laying rates on food (Song et al., 2013; Waggoner et al., 1998). Mutants lacking tryptophan hydroxylase, the rate-limiting enzyme required for 5-HT synthesis, exhibit an increased propensity to developmentally arrest, indicating that lack of the positive food signal, 5-HT, discourages development into adulthood (Sze et al., 2000).

While 5-HT is associated with food-related behaviors, multiple lines of evidence suggest that octopamine is associated with starvation-related behaviors. In *Drosophila melanogaster*, octopamine signaling is required for the hyperactivity observed in starved animals (Yang et al., 2015). Hyperactivity is thought to reflect the drive of starving animals to locate a food source. In *C. elegans*, octopamine inhibits egg-laying and feeding, and modulates response rate to aversive stimuli (Alkema et al., 2005; Mills et al., 2011; Wragg et al., 2007). However, octopamine’s role in *C. elegans* locomotion has not been well studied, and how it acts to regulate food-dependent locomotion behavior is unknown.

Recently, our laboratory investigated the dynamics of behavioral states of worms in liquid. McCloskey et al. found that food, mechanical environment, insulin signals, protein kinase G signaling, and neuropeptides regulate the incidence of three behavioral states roaming, dwelling, and quiescence (McCloskey et al., 2017; Ghosh and Emmons, 2008). While roles for acetylcholine, TGF-β, and insulin signaling have been reported in regulating these behavioral states, specific neural circuits have not been fully elucidated (Gallagher et al., 2013; You et al., 2008).

In this study, we use longitudinal imaging, pharmacology, and genetic techniques to show that the biogenic amines 5-HT and octopamine regulate behavior during feeding and fasting in *C. elegans*. We then identify the neuronal classes responsible for releasing these neurotransmitters and the receptors on which they act. We show that 5-HT and octopamine play antagonistic roles in regulating nutrient-related locomotory states.

**MATERIALS AND METHODS**

**Strains**

*C. elegans* hermaphrodites were cultured as previously described (Brenner, 1974). All worms were hermaphrodites grown on standard agar plates with OP50 *E. coli* until just prior to experiments. N2 Bristol was used as the wild-type reference strain. The following strains were used in this study: MT15434: *tph-1(mg280) II*, DA1814: *ser-1(ok345) X*, CX12800: *ser-3(ad1774) I*, AQ866: *ser-4(ok512) III*, RB2277: *ser-5(ok3087) I*, *ser-5(tm2654) I*, *ser-5(tm2647) I*, KQ1048: *ser-6(tm2146) IV*, DA2100: *ser-7(tm1325) X*, MT9668: *mod-1(ok103) V*, MT9180: *mod-1(n3034) V*, MT9667: *mod-1(nr2043) V*, MT13113: *tdc-1(n3419) II*, MT9455: *tbh-1(n3247) X*, CX13079: *oct-1(ok371) X*, DA2289: *tph-1(mg280) II*, kyEx947[pceh-2::*tph-1(+):gfp punc-122::gfp(+)], DA2290: *tph-1(mg280) II*, kyEx949[psrh-142::*tph-1(+):gfp punc-122::gfp(+)], SSR664: *mod-1(ok103) V*, [Pmod-1::mod-1], SSR644: *mod-1(ok103) V*, [Pttx-
3::mod-1], SSR642: mod-1(_ok103) V; [Pgcy-36::mod-1], SSR665: mod-1(_ok103) V; [Pfpl-1::mod-1], SSR662: mod-1(_ok103) V; [Pfpl-8::mod-1], VN289: ser-6(tm2104); tzIs3[cre::gfp, lin-15(+)], VN441: ser-6(tm2104); tzIs3; vnEx142[ceh-17::ser-6, lin-44::gfp], YX179: ser-5(ok3087) I; [Pser-5::ser-5, Pmyo-2::mCherry], YX180: ser-5(ok3087) I; [Pmyo-3::ser-5, Pmyo-2::mCherry], YX181: ser-3(ad1774) I; [Pceh-17::ser-3, Pmyo-2::mCherry], YX184: ser-5(ok3087) I; [Psra-6::ser-5, Pmyo-2::mCherry], YX185: ser-5(ok3087) I; [Punc-119::ser-5, Pmyo-2::mCherry].

WorMotel Design and Fabrication
To fabricate the WorMotel, we developed a 3D-printing based molding method (Shepherd et al., 2011). We designed a chip containing a rectangular array of either 48 or 240 rounded wells with 3 mm diameter, 3 mm depth, and center-to-center spacing of 4.5 mm (Fig. 1). Each well was surrounded by a 0.5 mm wide and 3 mm deep channel, which served as a moat. Designs of the WorMotel masters were created using MATLAB. We printed a master corresponding to the negative of this shape with an Objet30 photopolymer 3D printer using the material VeroBlack. To mold the WM devices, we mixed Dow Corning Sylgard 184 PDMS according to the manufacturer’s instructions and poured 35 g or 5 g of PDMS into the 240-well or 48-well masters, respectively. We then degassed the poured PDMS in a vacuum chamber for 1 hour. Devices were cured overnight at 40°C and then removed from molds using a spatula.

Liquid Behavioral Assays
9 µl of liquid NGM buffer (NGB), which consists of the same components as NGM but without agar or peptone (Stiernagle, 2006; McCloskey et al., 2017), was added to each well of the WorMotel just prior to adding worms. Young adult animals were added manually to the liquid contained in each well of the WorMotel. Where indicated, the NGM was supplemented with either drugs or food. As a food source we used E. coli DA837 (Davis et al., 1995), a streptomycin resistant derivative of OP50 (Brenner, 1974). For experiments where food was added to wells, bacteria was added to NGM to a final OD of approximately 1.

For experiments with added exogenous biogenic amines, we added 5-HT hydrochloride, tyramine hydrochloride, or octopamine hydrochloride (Sigma Aldrich) in the indicated concentrations to liquid NGM prior to filling the WorMotel. Multiple genotypes or conditions were assessed on the same WorMotel. Behavioral comparisons were made only between conditions that were assayed simultaneously on the same WorMotel.

PDMS devices were placed inside a 90 mm diameter petri dish. To maintain humidity inside the dishes, we used water-absorbing polyacrylate crystals. Sterile water was added to the crystals in a ratio of 150:1 (water:crystals) by weight. Approximately 5 g of hydrated crystals were added around the WorMotel. We placed lids on all dishes. To prevent accumulation of water condensation, lids were prepared by coating with a 30% solution of Tween 20 (Sigma-Aldrich) in water, which was allowed to dry before use.

Finally, the filled WorMotel was placed upside down on the bottom of the petri plate and the lid was added. The petri plate was placed on a glass platform inside a custom imaging rig as previously described (Churgin and Fang-Yen, 2015). A digital camera (see Image Acquisition) was placed beneath the glass platform and images were acquired every second until 60,000
images had been acquired, corresponding to about 16.6 hours of recording time (Churgin and Fang-Yen 2015).

**Image Acquisition**
Images were captured with an Imaging Source DMK 23GP031 camera (2592 x 1944 pixels) equipped with a Fujinon lens (HF12.5SA-1, focal length 12.5 mm). We used IC Capture imaging software (Imaging Source) to acquire time lapse images through a gigabit Ethernet connection. All experiments were carried out under dark-field illumination using four 4.7” red LED strips (Oznium) positioned on the glass platform surrounding the WorMotel. Images were saved and processed by a 64-bit computer with a 3.40 GHz Intel Core i3 processor and 4 GB of RAM. Images were processed and analyzed using custom-written MATLAB software (see Image Processing).

The camera’s field of view was adjusted such that 48 wells of the WorMotel were visible on the image. This field of view corresponded to a spatial resolution of approximately 15 μm.

**Image Processing and data analysis**
All data processing and analysis was carried out using MATLAB (Mathworks). Subsequent images were subtracted to generate difference images of pixel value intensity change. The difference image was normalized by average pixel intensity across the two subtracted images. A Gaussian smoothing filter with standard deviation equal to one pixel was applied to the delta image in order to reduce noise. A binary threshold of 0.35 was then applied to the filtered delta image in order to score whether or not movement occurred at each pixel location. All pixels in which movement occurred were summed up and the resulting value was called the ‘activity’ between the two frames.

To calculate the time spent quiescent, we first time-averaged raw activity data, with time resolution one second, with a smoothing kernel of five seconds. For each animal, we then calculated the number of frames where the activity was equal to zero. The resultant value was divided by the duration of the assay to calculate the fraction of time spent quiescent.

**Histogram Fitting**
We generated a histogram of the five-second time-averaged activity for each worm. We then used a non-linear least squares fitting algorithm to fit each individual worm’s activity histogram to the sum of two exponential terms (with two unknown parameters each) and a Gaussian term (with three unknown parameters). We therefore fit seven parameters. The zero bin on the activity histogram, corresponding to quiescent frames, was excluded from the histogram before the fitting process as the contribution of quiescence had already been calculated.

To calculate the fraction of time spent roaming, we calculated the area under the Gaussian curve component of the fit. To calculate the fraction of time spent dwelling, we calculated the area under the two exponential curve components of the fit.

**Manual Validation of Behavior**
We picked at random three fasting and three feeding worms from a previously recorded experiment to manually score roaming and dwelling behavior. We examined videos recorded at
8 frames per second rather than one frame per second because we found it easier for a manual user to score behavior when data was recorded at a higher frame rate. Nevertheless, activity values were still calculated using frames captured one second apart as in all other experiments.

We examined behavior for approximately 10 minutes at each of three time points: 2 hours, 3 hours, and 4.5 hours from the start of the experiment. The purpose of using three time points was to ensure all three types of behavior were observed for each worm, not to assess temporal variations in behavior. Indeed, all three behaviors including roaming, dwelling, and quiescence were observed for each worm manually scored. A user blind to the results of the fit analysis scored behavior as roaming, dwelling, or quiescent for each worm for each frame.

Next, the activity histogram for frames the user scored as roaming and dwelling were individually generated and compared to the Gaussian or exponential components, respectively, of the full-data fit. To quantify the agreement between user-scored and computer-scored behavior, we calculated the $R^2$ value.

**Visualizing Data in Roaming-Dwelling Space**

After fitting, the roaming and dwelling fraction was calculated for each worm of a given genotype and experimental condition. For each population, the mean roaming and dwelling fraction were calculated, along with the standard deviation and orientation of principal components in roaming-dwelling space. Ellipses with dimensions corresponding to the standard error of the mean for each experimental condition were plotted with their orientation in the direction of greatest variation for that experimental condition as determined by principal component analysis.

**Cloning and Transgenic Strain Construction**

Rescue constructs for SER-5 were cloned using Gateway Technology (Life Technologies). The desired promoters, ser-5 cDNA, and unc-54 3′ UTR were acquired in entry slots 1, 2, and 3, respectively. The following plasmids in Gateway entry vectors were gifts of Zheng-Xing Wu: sra-6p, ser-5p, and ser-5 cDNA (Guo et al., 2015). The LR recombination reaction was performed to produce each rescue construct. Each rescue plasmid was confirmed via sequencing. The rescue plasmid ceh-17::ser-3 was a gift from Satoshi Suo (Yoshida et al., 2014). Transgenic strains were constructed by microinjection in the *C. elegans* germline. SER-5 rescue constructs were injected into RB2277: ser-5(ok3087) mutants. SER-3 rescue constructs were injected into CX12800: ser-3(ad1774). For microinjections, 30 ng/µl of the desired plasmid was injected with 2 ng/µl of myo-2::mCherry coinjection marker and 100 ng/µl of 1 kb DNA ladder.

**Experimental Design and Statistical Analysis**

Differences in population behavior distributions were carried out using a two-tailed t-test. Error bars throughout the manuscript are presented as standard error of the mean, with N taken to be the number of biological replicates per condition.

**RESULTS**

**Quantification of locomotion behavior in swimming *C. elegans* with longitudinal imaging**
We first sought to better characterize *C. elegans* behavior in feeding and fasting conditions. We monitored locomotion behavior of young adult worms using the WorMotel, a custom microplate device consisting of an array of individual wells that can be used to longitudinally image animals under solid or liquid media conditions (Churgin *et al.*, unpublished manuscript). We assayed worms under liquid conditions and quantified locomotion behavior using custom machine vision software (see Methods).

We first monitored young adult N2 worms for 16 hours in liquid NGB (see Methods) grown with or without food (Figure 1). As previously reported, after a period of a few hours of continuous high activity, fasting worms in liquid began to cycle between high activity and behavioral quiescence (Figure 1) (Ghosh and Emmons, 2008; McCloskey *et al.*, 2017). The behavior of feeding and fasting worms was qualitatively very different. Fasting worms oscillated between extended high activity periods and short low activity periods, whereas feeding worms exhibited much more intermediate activity. We then plotted activity histograms for individual worms. From these histograms, we confirmed that fasting worms exhibit a more bimodal activity distribution, whereas feeding worms exhibit a mostly monotonically decreasing activity distribution.

To quantify behavior, we applied an empirical fit to the activity histogram on an individual worm basis (Figure 2a-b). We used a nonlinear least squares algorithm to fit each worm’s activity histogram to a sum of two exponential terms and one Gaussian term (see Methods). Worm locomotion is usually classified into one of three behavior states: roaming, dwelling, or quiescence (McCloskey *et al.*, 2017; Flavell *et al.*, 2013). Roaming is characterized by rapid movement and propagation of body waves down the length of the worm’s body. Dwelling is characterized by lower amount of movement and propagation of body waves in the anterior but not posterior of the worm’s body. Quiescence is characterized by complete cessation of movement and is observed both in high food environments, where it is termed satiety, and in fasting conditions, where it is termed fasting quiescence (You *et al.*, 2008; Ghosh and Emmons, 2008). We therefore wanted to classify worm activity measurements according to these commonly applied labels.

We hypothesized that the exponential and Gaussian terms of the fit corresponded to dwelling and roaming behavior, respectively. To test this idea, we performed manual assessment of behavioral states from recorded videos. We found a strong overlap between the Gaussian component of the fit with the histogram of frames in which a worm’s behavior was manually scored as roaming (Figure 2c). Similarly, we found a strong overlap between the exponential components of the fit and the histogram of frames in which a worm’s behavior was manually scored as dwelling (Figure 2d). Combining both frames scored as dwelling and roaming together largely matched the overall fit (Figure 2e, Table 1).

These results indicate that the exponential components of the histogram fit correspond well with frames in which worms are dwelling, and the Gaussian component corresponds with frames in which worms are roaming. Therefore, for each worm we simply summed up the area under each of these two components to calculate the fraction of time individuals spent dwelling and roaming. Quiescence was calculated as the fraction of time each worm’s activity level was zero.
We found large quantitative differences in locomotion behavior between feeding and starved worms. We found that feeding caused an elevation in behavioral quiescence, a reduction in roaming, and an increase in dwelling in comparison to fasting (Figure 3j-n). Therefore, food status modulates multiple locomotion parameters in young adult worms.

5-HT signaling is required for worms to appropriately adapt behavior from fasting to feeding

Having established that food levels affect locomotion behaviors, we sought to determine which neurotransmitters were responsible for these changes. Since serotonin is a transmitter known to regulate behavior in response to food, we first examined the behavior of tph-1 mutants, which lack endogenous 5-HT, in response to feeding and fasting. These mutants exhibit increased fasting quiescence, decreased feeding quiescence, and increased roaming while feeding (Figure 3). Therefore, endogenous 5-HT signaling is required for both wild-type fasting and feeding behavior.

We found that tph-1 worms exhibited more roaming compared to N2 worms when assayed with food (Figure 3). This is consistent with previous reports that tph-1 worms roam more on a bacterial lawn than N2 (Flavell et al., 2013). Moreover, we found that food treatment in tph-1 animals had a lesser effect on the fraction of time spent quiescent. These results indicate that tph-1 mutants are unable to fully adopt wild-type feeding behaviors in response to food. Our findings are consistent with the idea that 5-HT is released in response to food and is required for adapting locomotion behaviors to a feeding-appropriate state.

Differential effects of 5-HT produced by the ADF and NSM neurons

We asked in what neurons 5-HT is produced to regulate locomotory fasting quiescence. 5-HT is produced primarily by two neuron pairs in the head, ADFL/ADFR and NSML/NSMR, and one neuron pair near the vulva, HSNL/HSNR, as determined by expression of a Ptph-1::GFP transgene as well as antibody staining (Zheng et al., 2005; Sze et al., 2002; Liang et al., 2006). Previous reports showed that either fasting (Cunningham et al., 2012) or daf-2 mutation (Estevez et al., 2006) caused a reduction in tph-1 reporter expression only in the ADF neuron, implying that 5-HT production by ADF is reduced in response to a reduction nutrient signals.

We hypothesized that 5-HT in the ADF neurons plays a role in fasting quiescence. To test this idea, we examined tph-1 mutant worms in which wild-type TPH-1 was rescued in the ADF or NSM neurons with the cell-specific promoters srh-142 or ceh-2, respectively (Song et al., 2013). As before, we noticed that tph-1 mutants exhibit less roaming and less dwelling compared to wild type (Figure 4a). We found that tph-1 rescue in ADF, but not in NSM, was sufficient to rescue roaming and dwelling to wild type levels, whereas rescue of tph-1 in NSM caused a drastic reduction in roaming (Figure 4a). Therefore, this data suggests that 5-HT produced by the ADF neuron is required for wild-type fasting behavior, whereas 5-HT produced by the NSM neuron antagonizes wild-type fasting behavior.

As before, tph-1 mutants exhibited slightly but significantly more quiescence than wild type, and this increase was restored to wild-type levels by rescue of TPH-1 in ADF (Figure 4b). By contrast, rescue of TPH-1 in NSM resulted in a large increase in fraction of time spent quiescent (Figure 4b). As before, fasting tph-1 mutants exhibited an insignificant reduction in roaming, and
this reduction was reversed by rescue of TPH-1 in ADF (Figure 4a). By contrast, rescue of TPH-1 in NSM exacerbated the reduction of roaming, suggesting that these two neurons act antagonistically to regulate the amount of roaming.

These results show that 5-HT produced by the ADF neurons is necessary and sufficient to restore wild-type fasting quiescence and behavior. Furthermore, these findings show that ADF and NSM act antagonistically with respect to roaming behavior.

Exogenous 5-HT suppresses fasting quiescence and increases dwelling
Having established that endogenous 5-HT signaling regulates fasting quiescence, we sought to determine the effect of exogenous 5-HT on the behavior of fasting worms. When we added exogenous 5-HT to the liquid media, we observed in wild-type worms an increase in dwelling and reduction in roaming (Figure 5a-b). Consistent with our results that tph-1 mutants lacking 5-HT-signaling exhibit higher fasting quiescence (Figure 3-4), treatment with exogenous 5-HT suppressed wild-type fasting quiescence (Figure 5c).

Although exogenous 5-HT suppressed quiescence, it also reduced roaming (Figure 5e). Therefore, exogenous 5-HT both reduces overall movement while also reducing the time spent completely immobile. Accordingly, exogenous 5-HT causes a dramatic increase in dwelling (Figure 5d). These results are consistent with reports that 5-HT slows locomotion (Ranganathan et al., 2000; Flavell et al., 2013). Therefore, exogenous 5-HT acts to promote dwelling behavior and suppress both quiescence and roaming.

5-HT promotes roaming via SER-5 and dwelling via MOD-1
We sought to determine the receptors through which exogenous 5-HT modulate locomotion. In C. elegans, 5-HT acts strongly through four G-protein coupled receptors and one 5-HT-gated chloride channel to regulate locomotion, feeding, fat storage, egg-laying, and other phenotypes (Ranganathan et al., 2000; Carre-Pierrat et al., 2006; Gürel et al., 2012). To determine the receptors on which 5-HT acts, we tested the available single 5-HT receptor mutants with and without exogenous 5-HT under fasting conditions.

We found that exogenous 5-HT treatment had similar effects on quiescence, roaming, and dwelling in wild type animals as in ser-1, ser-4, and ser-7 mutants (Figure 5a), indicating that these receptors do not contribute significantly to locomotion behavior. By contrast, exogenous 5-HT had a drastic effect on behavior of both ser-5 and mod-1 mutants (Figure 5b). We found that 5-HT suppressed quiescence in all 5-HT receptor mutants tested except for ser-5 (Figure 5c). Instead of suppressing quiescence in ser-5 mutants, applying exogenous 5-HT resulted in greatly increased quiescence in three different alleles compared to untreated worms. Furthermore, roaming was strongly suppressed in 5-HT-treated ser-5 mutants, indicating that SER-5 promotes movement in response to 5-HT.

Applying exogenous 5-HT to mod-1 mutants, by contrast, resulted in quiescence suppression in three different alleles to a greater degree than for wild type worms (Figure 5c). Furthermore, roaming increased and dwelling decreased in 5-HT-treated mod-1 mutants (Figure 5d-e), indicating that 5-HT acts through MOD-1 to suppress movement. These results are consistent with reports showing that MOD-1 promotes dwelling behavior (Ranganathan et al., 2000; Flavell...
et al., 2013). Together, our results indicate that 5-HT acts antagonistically through SER-5 to increase roaming and suppress quiescence and through MOD-1 to increase dwelling and suppress roaming (Figure 10).

**MOD-1 acts in AIY neurons and SER-5 acts in body wall muscles and neurons to mediate behavioral effects of 5-HT**

We next sought to determine in what cells or tissues the MOD-1 and SER-5 receptors act to mediate 5-HT's effects on behavior. MOD-1 is a 5-HT-gated chloride channel (Ranganathan, Cannon, and Horvitz 2000) that has been shown to act in the nervous system to mediate locomotion (Flavell et al., 2013), learning (Zhang et al., 2005), and fat levels (Noble et al., 2013). SER-5 is a G-protein coupled receptor that has been shown to act in the ASH neurons to mediate the behavioral response to octanol and in muscles to mediate egg-laying (Harris et al., 2009; Hapiak et al., 2009). We therefore performed rescue experiments for MOD-1 and SER-5 in neurons, muscle, and specific cells to identify where these receptors act to mediate behavioral responses to 5-HT.

We first tested strains with a mod-1 mutant background in which wild type MOD-1 was expressed under its endogenous promoter as well as under four neuron-specific promoters: Pttx-3, which is expressed in AIY, AIA, and AIZ neurons, Pflp-8, which is expressed in AUA and URX neurons, Pflp-1, which is expressed in RIG, AIA, and AIY neurons, and Pgcy-36, which is expressed in URX neurons (Noble et al., 2013). We found that rescuing MOD-1 under its endogenous promoter as well under the TTX-3 promoter resulted in partial rescue of wild type behavior when treated with 5-HT (Figure 6a-e). By contrast, rescuing MOD-1 under the FLP-1, FLP-8, or GCY-36 promoters did not result in behavior distinguishable from mod-1 mutants (Figure 6a-e). The TTX-3 promoter has been previously reported to be drived expression strongly in the AIY interneurons (Flavell et al., 2013). Therefore our results show that MOD-1 expressed in the AIY neurons is necessary and sufficient to restore wild type response to 5-HT.

Next, we generated rescue constructs for the SER-5 receptor under its endogenous promoter, a neuron-specific promoter (Punc-119), a body wall muscle-specific promoter (Pmyo-3), and an ASH-specific promoter (Psra-6) (see Methods), and injected them into a ser-5 mutant background. We found that rescuing SER-5 under its endogenous promoter fully restored the wild type response to 5-HT (Figure 6f-j). We found that rescuing SER-5 in either muscles or neurons rescued the behavioral response to 5-HT, whereas rescuing SER-5 in only the ASH neuron did not (Figure 6f-j). Therefore, SER-5 can act in both muscle or neurons to promote movement in response to 5-HT.

**Octopamine and tyramine have opposite effects on locomotion**

We next turned our attention toward neurotransmitters typically associated with fasting. The biogenic amines tyramine and octopamine modulate multiple behaviors and appear to signal starvation and promote starvation-related behaviors, such as suppressed pumping and egg-laying (Alkema et al., 2005; Rex and Komuniecki, 2002; Rex et al., 2004; Suo et al., 2006). We sought to determine the roles, if any, of these neurotransmitters in locomotion behavior mediated in relation to food.

We first tested worms defective for TDC-1, the tyrosine decarboxylase enzyme required for
synthesis of tyramine and its downstream product, octopamine. We found that tdc-1 mutants exhibited increased fasting quiescence, increased dwelling, and reduced roaming relative to wild type (Figure 7a). These characteristics of fasting tdc-1 mutants largely overlap with those of feeding N2 worms (Figure 3), indicating that animals lacking tyramine and octopamine behave as if feeding even in the absence of food.

Loss of TDC-1 abolishes the worm’s ability to produce both tyramine and octopamine. To determine whether lack of tyramine or octopamine was responsible for the altered behavior in tdc-1 mutants, we tested behavior after either exogenous tyramine or octopamine was applied. Addition of exogenous octopamine suppressed the increased fasting quiescence in tdc-1 mutants as well as in wild type. By contrast, exogenous tyramine slightly increased fasting quiescence in wild type, but not in tdc-1 mutants (Figure 7b). These results suggest that octopamine is required for wild-type levels of fasting quiescence, and that octopamine acts to suppress quiescence.

We also found that treatment of both wild type and tdc-1 worms with tyramine resulted in increased dwelling and reduced roaming, whereas octopamine treatment slightly reduced dwelling and slightly increased roaming (Figure 7c-d). These results indicate that octopamine and tyramine have largely opposing roles on locomotion, with tyramine promoting reduced locomotion typical of feeding worms and octopamine promoting increased locomotion typical of fasting worms.

**Octopamine signaling is required for worms to appropriately adapt behavior to fasting**

Next, we examined the behavior of tbh-1 mutants, which lack only octopamine. We found that - tbh-1 mutants, like tdc-1 mutants, exhibited elevated quiescence, elevated dwelling, and decreased roaming under fasting conditions in comparison to wild type animals. The defects in quiescence and roaming, but not dwelling, were reversed by exogenous treatment with octopamine (Figure 8a, c-e). These results show that in worms, as in flies, octopamine is responsible for the hyperactivity observed during fasting (Yang et al., 2015).

The increased dwelling, decreased roaming, and increased quiescence evident in fasting tbh-1 worms again recalls, like that of fasting tdc-1 worms, the behavior of feeding wild type animals (Figure 3a, c, j, k). The finding that fasting tbh-1 worms resemble feeding N2 worms suggests that octopamine, but not tyramine, is required for worms to adopt locomotion behaviors associated with fasting.

We next considered the behavior of tbh-1 mutants in the presence of food. Treatment of tbh-1 mutants with food caused a reduction in roaming and increasing in dwelling compared with fasting tbh-1 worms. This trend is similar to that observed in wild type worms but occurred to a greater extent in tbh-1 animals (Figure 3k, l-n). Feeding tbh-1 worms exhibit greatly increased dwelling, greatly reduced roaming, and no change in quiescence compared to both fasting tbh-1 worms. Feeding tbh-1 worms also exhibit more dwelling and less roaming than feeding wild-type worms, suggesting that these animals overcompensate their locomotion behavior to food compared to wild type.

The finding that feeding tbh-1 worms exhibit a large fraction of dwelling is consistent with a qualitative view of their behavior, which shows a fairly homogenous, low level of activity.
throughout the 16 hours of the assay, compared with the relatively more heterogeneous behavior of both feeding wild type and tph-1 worms (Figure 3a-c, e, g, i). These results reinforce the notion that octopamine acts to promote movement, but also implies that this pathway acts during feeding as well as fasting.

Both SER-3 and SER-6 are required for exogenous octopamine’s effects on locomotion
We next sought to determine the receptor(s) through which exogenous octopamine acts to excite roaming and suppress quiescence. We tested mutants for the three known octopamine receptors (Sun et al., 2011; Komuniecki et al., 2004; Komuniecki et al., 2012). We found that response to octopamine was abolished in both ser-3 and ser-6 mutants but remained largely intact in octr-1 mutants (Figure 8a-e). Treatment with exogenous octopamine significantly increased roaming and decreased quiescence in wild type and octr-1 mutants, but was unable to do so in either ser-3 or ser-6 mutants. These results indicate that octopamine requires both SER-3 and SER-6 receptors to affect locomotion.

SER-3 and SER-6 act in the SIA neurons to mediate octopamine's effect on locomotion
We next sought to determine where the SER-3 and SER-6 receptors acted to mediate octopamine's effect on behavior. A previous report showed that SER-3 and SER-6 acted in the SIA neurons to mediate the transcription of the starvation-responsive transcription factor, CREB (Yoshida et al., 2014). We therefore tested whether rescuing SER-3 and SER-6 in the SIA neurons was sufficient to restore wild type response to octopamine.

We found that expressing SER-3 or SER-6 under the Pceh-17 promoter, which drives expression specifically in the SIA and ALA neurons, was sufficient to restore wild type response to octopamine in SER-3 and SER-6 mutants, respectively (Figure 9a-e). A previous study found that octopamine modulated CREB levels in the SIA neurons, suggesting that SER-3 and SER-6 function in the SIA neurons and not ALA (Suo et al., 2006; Yoshida et al., 2014). Therefore, our results suggest that SER-3 and SER-6 also function in the SIA neurons to mediate octopamine's effect on locomotion.

DISCUSSION

Quantitative, high-throughput imaging enables characterization of complicated behavioral phenotypes
In this study, we have used the WorMotel, a novel microplate designed for long-term behavioral imaging of C. elegans, to elucidate the molecular and neural pathways underlying the locomotion behavioral effects of the biogenic amines 5-HT and octopamine. By acquiring high-throughput behavioral measurements, we quantified the time spent quiescent, dwelling, and roaming for up to 48 worms simultaneously over the first 16 hours of young adulthood. Automated behavioral analyses like those described here will enable quantification of complicated behavioral phenotypes and thus allow for the dissection of how neural, genetic, and environmental variables act together to generate behavior.

As evidenced here and described elsewhere, animals utilize the same basic behaviors under different experimental conditions. In C. elegans, those basic behaviors are quiescence, roaming, and dwelling. However, the probability of each respective behavior is modulated as necessary to
achieve an organism's goals appropriate to the current environmental state. Therefore, the ability to automatically detect and annotate different behavioral phenotypes in a quantitative way is crucial for us to understand the nuanced way the relative weights of different behavioral states are modulated in response to the environment.

Our findings support the prevailing view that 5-HT and octopamine signaling are required for appropriately adapting behavior to the presence and absence of food, respectively. We also describe the receptors and sites of action of these signalling molecules. Together, our results delineate the neural circuit and molecular mechanisms that appropriately govern locomotion behavior in relation to food levels (Figure 10).

Two 5-HT-producing neurons exert opposing effects on locomotion
Our results suggest that ADF-produced 5-HT acts through SER-5 to promote roaming, whereas NSM-produced 5-HT acts through MOD-1 to promote dwelling. Therefore, serotonin acts both to increase and decrease locomotion, depending on its source. The fact that serotonin alone can both excite and suppress locomotion is supported by the fact that fasting thb-1 mutants, which lack the roaming-promoting signal octopamine, are able to generate periods of both dwelling and roaming (Figure 3c, h). This variation in behavior could be produced by the alternate release of 5-HT from ADF and NSM to produce periods of high and low locomotion, respectively.

5-HT acts antagonistically through SER-5 and MOD-1 to regulate locomotion
Previous results have shown that other 5-HT receptors act to slow locomotion in C. elegans (Ranganathan et al., 2000). Our result suggests that SER-5 is the sole receptor through which 5-HT can stimulate roaming. Similarly, MOD-1 appears to be the primary receptor through which 5-HT promotes dwelling (Figure 5). Therefore, we confirm that 5-HT promotes dwelling via MOD-1 and show for the first time that 5-HT promotes roaming via SER-5 (Figure 10).

Our results suggest that ADF is an activity promoting neuron and NSM is activity-suppressing neuron. Therefore, it is plausible that ADF-produced 5-HT acts through SER-5 to regulate locomotion, as previously found in the context of copper avoidance behavior (Guo et al., 2015). Our results would then suggest that NSM-produced 5-HT therefore acts through MOD-1 to suppress locomotion.

MOD-1 acts in the AIY neurons and SER-5 acts in muscles and neurons to regulate locomotion
We found that rescuing MOD-1 under either its endogenous promoter or the TTX-3 promoter restored wild type response to 5-HT. Previous reports have also found that expressing MOD-1 under the TTX-3 promoter, which is strongly expressed in the AIY neurons, is required to promote dwelling in feeding wild type worms (Flavell et al., 2013; Hapiak et al., 2009). We confirm this role for promotion of dwelling by MOD-1 in the AIY neurons. Given that MOD-1 is a 5-HT-gated chloride channel (Ranganathan et al., 2000) and optogenetic activation of AIY promotes roaming (Flavell et al., 2013), it is likely that MOD-1 acts to silence AIY in response to 5-HT. Therefore, MOD-1 likely acts to suppress activity in the AIY neurons in order to promote dwelling in response to 5-HT.

We found that SER-5 could act in either muscle or neurons to promote movement in response to
5-HT. A previous study found that SER-5 acted in vulval muscle to regulate serotonergic stimulation of egg-laying (Hapiak et al., 2009). However, this study did not describe a role for SER-5 in regulating movement. Another study found that SER-5 acts in the nervous system specifically in the ASH neurons to mediate response times to octanol (Harris et al., 2009). Therefore, SER-5 occupies multiple tissue sites of action to mediate diverse behaviors. Egg-laying is associated with a transient increase in movement (McCloskey et al., 2017; Waggoner et al., 2000). Therefore, it is possible that SER-5 acts in the muscle to increase egg-laying rate, which secondarily acts to increase movement. An alternate possibility is that SER-5 acts directly on body wall muscles to promote movement. We can, however, rule out a role for the ASH neuron in regulating locomotion behavior in response to 5-HT (Figure 6f-j). More work is required to identify whether SER-5 acts endogenously in neurons, muscles, or both to regulate locomotion behavior.

**Octopamine promotes roaming behavior via SER-3 and SER-6 in the SIA neurons**

Both fasting and treatment with exogenous octopamine excite activity the cAMP response element-binding protein (CREB) transcription factor in the SIA neurons (Yoshida et al., 2014; Suo et al., 2006). This effect requires the presence of both SER-3 and SER-6. Therefore, it has been hypothesized that these receptors act non-redundanty in the SIA neurons to mediate octopamine's effects in response to starvation, perhaps via a threshold effect or dimerization.

We found that SER-3 and SER-6 also mediate octopamine’s effect on behavior, and that both receptors function in SIA to mediate this effect. Therefore, our results further support the notion that both SER-3 and SER-6 are required for octopamine’s effect on the response to fasting by modulating quiescent behavior (Figure 8-9). Our results suggest that the CREB transcription factor may play important roles in regulating fasting locomotion behavior.

**5-HT and octopamine promote behaviors associated with feeding and fasting, respectively**

Our results suggest that food initiates release of 5-HT, but that in wild type animals a reserve of octopamine signaling antagonizes this food-induced 5-HT release to antagonize the effects of 5-HT on behavior and promote roaming even when food is present. This explains why the behavior of feeding N2 worms is different from N2 worms treated with exogenous 5-HT. The dose of exogenous 5-HT permeating the cuticle in these experiments may be much larger than that released endogenously, so exogenously supplied 5-HT dominates the endogenous octopamine signal that normally counteracts 5-HT’s effect. By contrast, no octopamine is present to counteract the 5-HT released in tbh-1 mutants supplied with food, so the behavior of these animals resembles that observed in wild type treated with exogenous 5-HT (cf. Figures 3k and 7a-b).

Despite the finding that 5-HT treatment primarily promotes feeding-associated locomotion phenotypes and octopamine treatment primarily promotes fasting-associated phenotypes, we find that both transmitters are required for wild type behavior under both feeding and fasting conditions. We find that although well-fed wild type animals mostly make small movements, they also exhibit periods of high locomotion activity. This is consistent with previously published reports showing that on food, worms spend time primarily in dwelling state but also exhibit short-lived roaming states (Flavell et al., 2013).
Our results suggest that by continuing to release lower levels of the non-dominant biogenic amine—octopamine under feeding conditions and 5-HT under fasting conditions—worms continue to maintain access to a variety of behaviors, some of which are normally associated with an environmental state they do not currently find themselves in. By reducing, but not completely eliminating, behaviors associated with environmental conditions not currently being experienced, animals maintain a high degree of adaptability.

**Conserved roles of 5-HT and octopamine**

5-HT and octopamine have conserved roles over diverse phyla. 5-HT is found in the protozoan Tetrahymena as an intracellular regulator of the cyclic AMP (cAMP) pathway (Turlejski, 1996), a known regulator of behavior. The majority of vertebrate 5-HT receptor families also modulate cAMP levels through their effects on adenylate cyclase, indicating a remarkable degree of conservation of the molecular role of this signaling molecule over two billion years of evolution (Turlejski, 1996). The evolutionary conservation of these receptors' downstream signaling pathways suggests that understanding the molecular roles of these receptors in model organisms may contribute to our understanding of these receptors' roles in mammals.

Octopamine is found primarily in invertebrates, but also in trace amounts in mammals, and is structurally similar to the fight-or-flight hormones adrenaline and noradrenaline. Here we find that octopamine’s role in worms is similar to that in flies, where it induces hyperactivity upon fasting as well wake-promotion (Crocker and Sehgal, 2008). However, it was not previously known which receptors octopamine acted through to induce hyperactivity in *Drosophila*. The conservation of behavioral effects of biogenic amines suggests that roles for specific receptors may also be preserved.

Understanding the role of these signals in invertebrates may inform our understanding of the molecular bases for more complex manifestations of heightened activity based on these signals in mammals, such as aggression and anxiety.
REFERENCES


factor UNC-86 regulates the tph-1 tryptophan hydroxylase gene and neurite outgrowth in specific serotonergic neurons.’, *Development*, 129(16), pp. 3901–3911.


Figure 1. Experimental setup and data analysis

a) Experimental setup b) Example image. Individual *C. elegans* are visible in each well. c) Detail of three wells. d) Locomotion activity map for worms in liquid incubated with (top half) or without (bottom half) bacterial food.
Figure 2. Quantitative analysis and manual validation

a) Activity histogram for a fasting worm (black curve) with fit overlaid (red dashed curve).
b) Activity histogram for a feeding worm with fit overlaid.
c) Activity histogram for frames during which a single worm’s behavior was manually scored as roaming (black curve). The Gaussian component of the fit is overlaid (red dashed curve).
d) Activity histogram for frames during which the same worm’s behavior as in c was manually scored as dwelling (black curve). The exponential component of the fit is overlaid (red dashed curve).
e) Activity histogram for frames in the same worm’s behavior as in c and d was manually scored. The full fit, consisting of both exponential and Gaussian components, is overlaid (red dashed curve).
Figure 3. Serotonin and octopamine signaling are required for normal fasting and feeding behavior.

a-c) Locomotion activity maps for N2, tph-1, and tbh-1, respectively.

d) Example activity trace for a fasting N2 worm.
e) Example activity trace for a feeding N2 worm.
f) Example activity trace for a fasting tph-1 worm.
g) Example activity trace for a feeding tph-1 worm.
h) Example activity trace for a fasting tbh-1 worm.
i) Example activity trace for a feeding tbh-1 worm.
j) Roaming fraction versus dwelling fraction for individual fasting (black circles) or feeding (blue triangles) N2 worms. Principal component ellipses are plotted to indicate population averages and standard errors.
k) Principal component ellipses illustrating fraction of time spent roaming versus dwelling for N2 (black), tph-1 (green), and tbh-1 (red) under fasting (solid ellipses) or feeding (dotted ellipses) conditions. Ellipse dimensions correspond to standard error of the mean.
l) Quantification of fraction of time spent quiescent for N2 (n=132), tph-1 (n=132), and tbh-1 (n=48) worms under fasting and feeding conditions.
m) Fraction of time spent dwelling.
n) Fraction of time spent roaming. *, p<0.05; **, p<0.01; ***, p<0.005
Figure 4. Antagonistic roles for two serotonergic neurons

a) Principal component ellipses illustrating fraction of time spent roaming versus dwelling for N2 (solid black), tph-1 (green x’s), and tph-1 neuron-specific rescues in ADF (red dots) or NSM (purple circles).  
b) Fraction of time spent quiescent.  
c) Fraction of time spent dwelling.  
d) Fraction of time spent roaming.  
In all graphs, genotypes represented are N2 (n=84), tph-1(mg280) (n=84), tph-1; Ex[Psrh-142::tph-1 (+)] (n=84), and tph-1; Ex[Pceh-2::tph-1 (+)] (n=80).  * p<0.05; **, p<0.01; ***, p<0.005
Figure 5. Exogenous serotonin acts through SER-5 to promote roaming and through MOD-1 to suppress roaming

a) Principal component ellipses illustrating fraction of time spent roaming versus dwelling for N2 (black), ser-1 (brown), ser-4 (blue), and ser-7 (yellow) under fasting (solid ellipses) or fasting + 10 mM 5-HT (dotted ellipses) conditions. b) Principal component ellipses illustrating fraction of time spent roaming versus dwelling for N2 (black), ser-5 (purple), and mod-1 (cyan) under fasting or fasting + 10 mM 5-HT conditions. c) Fraction of time spent quiescent. Number of animals tested were equal for both with and without 5-HT conditions: N2 (n=124), ser-1 (n=26), ser-4 (n=18), ser-5(ok3087) (n=54), ser-5(tm2647) (n=24), ser-5(tm2654) (n=24), ser-7 (n=38), mod-1(ok103) (n=28), mod-1(nr2043) (n=24), mod-1(n3034) (n=16). d) Fraction of time spent dwelling. e) Fraction of time spent roaming. *, p<0.05; **, p<0.01; ***, p<0.005
Figure 6. MOD-1 acts in AIY neurons and SER-5 acts in muscle and neurons to mediate 5-HT’s effects on locomotor behavior

a) Principal component ellipses illustrating fraction of time spent roaming versus dwelling for N2 (black), mod-1 (green), mod-1;Pmod-1::mod-1 (red), and mod-1;Pttx-3::mod-1 (blue) under fasting (solid ellipses) or fasting + 10 mM 5-HT (dotted ellipses) conditions. b) Principal component ellipses illustrating fraction of time spent roaming versus dwelling for N2 (black), mod-1 (green), mod-1;Pflp-1::mod-1 (gray), mod-1;Pflp-8::mod-1 (magenta), and mod-1;PgcY-36::mod-1 (blue) under fasting or fasting + 10 mM 5-HT conditions. c) Fraction of time spent quiescent. Number of animals tested were equal for both with and without 5-HT conditions: N2 (n=148), mod-1 (n=148), mod-1;Pmod-1::mod-1 (n=50), mod-1;Pttx-3::mod-1 (n=38), mod-1;Pflp-8::mod-1 (n=22), mod-1;Pflp-1::mod-1 (n=24), mod-1;PgcY36::mod-1 (n=24). Significance markings are relative to mod-1 worms treated with 5-HT (arrow). d) Fraction of time spent dwelling. e) Fraction of time spent roaming. f) Principal component ellipses illustrating fraction of time spent roaming versus dwelling for N2 (black), ser-5 (green), ser-5;Pser-5::ser-5 (red) under fasting (solid ellipses) or fasting + 10 mM 5-HT (dotted ellipses) conditions. g) Principal component ellipses illustrating fraction of time spent roaming versus dwelling for N2 (black), ser-5 (green), ser-5;Pmyo-3::ser-5 (blue), ser-5;Punc-119::ser-5 (light blue), and ser-5;Psra-6::ser-5 (magenta) under fasting (solid ellipses) or fasting + 10 mM 5-HT (dotted ellipses) conditions. h) Fraction of time spent quiescent. Number of animals tested were equal for both with and without 5-HT conditions: N2 (n=72), ser-5 (n=72), ser-5;Pser-5::ser-5 (n=16), ser-5;Pmyo-3::ser-5 (n=16), ser-5;Punc-119::mod-1 (n=24), ser-5;Psra-6::mod-1 (n=16). i) Fraction of time spent dwelling. j) Fraction of time spent roaming. *, p<0.05; **, p<0.01; ***, p<0.005
Figure 7. Octopamine and tyramine exert opposing effects on locomotory behavior

a) Principal component ellipses illustrating fraction of time spent roaming versus dwelling for N2 (black) and tdc-1 (gray) under fasting (solid ellipses), fasting + 10 mM tyramine (dashed ellipses), or fasting + 10 mM octopamine (dotted ellipses) conditions. b) Fraction of time spent quiescent. c) Fraction of time spent dwelling. Number of animals tested: N2 control (n=103), N2 with tyramine (n=52), N2 with octopamine (n=60), tdc-1 control (n=104), tdc-1 with tyramine (n=52), tdc-1 with octopamine (n=60). d) Fraction of time spent roaming. *, p<0.05; **, p<0.01; ***, p<0.005
Figure 8. Exogenous octopamine promotes roaming and suppresses quiescence through both SER-6 and SER-3

a) Principal component ellipses illustrating fraction of time spent roaming versus dwelling for N2 (black), tbh-1 (red), or octr-1 (teal) under fasting (solid ellipses) or fasting + 10 mM octopamine (dotted ellipses) conditions. b) Principal component ellipses illustrating fraction of time spent roaming versus dwelling for N2 (black), ser-3 (purple), and ser-6 (orange) under fasting or fasting + 10 mM octopamine conditions. c) Fraction of time spent quiescent. Number of animals tested were equal for both with and without octopamine conditions: N2 (n=80), tbh-1 (n=54), octr-1 (n=26), ser-3 (n=56), ser-6 (n=72). d) Fraction of time spent dwelling. e) Fraction of time spent roaming. *, p<0.05; **, p<0.01; ***, p<0.005
Figure 9. SER-6 and SER-3 function in the SIA neurons to mediate octopamine’s effects on behavior

a) Principal component ellipses illustrating fraction of time spent roaming versus dwelling for N2 (black), ser-6 (purple) and ser-6;ceh-17::ser-6 (red) under fasting (solid ellipses) or fasting + 10 mM octopamine (dotted ellipses) conditions. b) Principal component ellipses illustrating fraction of time spent roaming versus dwelling for N2 (black), ser-3 (green), and ser-3;ceh-17::ser-3 (blue) under fasting or fasting + 10 mM octopamine conditions. c) Fraction of time spent quiescent. Number of animals tested were equal for both with and without octopamine conditions: N2 (n=144), ser-6 (n=48), ser-6;ceh-17::ser-6 (n=48), ser-3 (n=96), ser-3;ceh-17::ser-3 (n=64) *, p<0.05; **, p<0.01; ***, p<0.005
Figure 10. Model for biogenic amine locomotion effects
### Tables

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Table 1. Manual behavior score versus automatic fit $R^2$ values