# Catecholamine receptor polymorphisms affect decision-making in *C. elegans*

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Innate behaviours are flexible: they change rapidly in response to transient environmental conditions, and are modified slowly by changes in the genome. A classical flexible behaviour is the exploration–exploitation decision, which describes the time at which foraging animals choose to abandon a depleting food supply. We have used quantitative genetic analysis to examine the decision to leave a food patch in *Caenorhabditis elegans*. Here we show that patch-leaving is a multigenic trait regulated in part by naturally occurring non–coding polymorphisms in tyra-3 (tyramine receptor 3), which encodes a G-protein–coupled catecholamine receptor related to vertebrate adrenergic receptors. tyra-3 acts in sensory neurons that detect environmental cues, suggesting that the internal catecholamines detected by tyra-3 regulate responses to external conditions. These results indicate that genetic variation and environmental cues converge on common circuits to regulate behaviour, and suggest that catecholamines have an ancient role in regulating behavioural decisions.

Despite abundant evidence for heritability of behavioural traits within and between species, only a few naturally varying traits have been associated with polymorphisms in specific genes<sup>1</sup>. Foraging for food is an ecologically relevant, environmentally regulated behaviour that is suitable for genetic analysis, as it can differ between populations of a species that live in different habitats<sup>2</sup>. An essential foraging decision is the choice between exploiting existing resources and exploring other options that may provide new resources. This decision can be described by Charnov's marginal value theorem, which proposes that the optimal time for an animal to leave a foraging ground occurs when local resource levels fall below the average level in the entire habitat<sup>3</sup>. The marginal value theorem was developed for animals foraging for food in patchy environments, but has analogies with diverse decisionmaking processes in field biology, cognitive neuroscience and economics<sup>2,4–6</sup>.

Studies of patch-leaving behaviour in the nematode C. elegans have revealed innate, environmental and experience-dependent factors that affect its foraging decisions. C. elegans rarely leaves a dense lawn of high-quality bacterial food<sup>7,8</sup>, but more frequently leaves lawns of pathogenic bacteria or lawns that are spiked with chemical repellents<sup>9,10</sup>. Males will leave lawns that do not contain potential mates<sup>11</sup>, while hermaphrodites leave lawns when animal density is high<sup>12</sup>. In addition, wild-type strains vary in their propensity to leave bacterial lawns based on a genetic polymorphism that affects the G-proteincoupled neuropeptide receptor NPR-1 (refs 12-14). This npr-1 polymorphism affects many foraging behaviours; low-activity npr-1 strains aggregate into social feeding groups, move quickly on food, and have altered responses to oxygen, carbon dioxide and pheromones compared to the N2 laboratory strain<sup>15-20</sup>. The high-activity allele of *npr-1* in N2 arose in the laboratory, probably as an adaptation to laboratory conditions<sup>19</sup>, so it is not known whether genetic variation affects C. elegans foraging in natural environments.

Natural genetic variation within a species can generate diversity in foraging behaviour, as exemplified by the polymorphic *Drosophila melanogaster foraging (for)* gene, which encodes a cyclic guanosine

monophosphate (cGMP)-dependent protein kinase<sup>21</sup>. A low-activity allele of *for* is present in *Drosophila* sitter larvae, which move slowly on a food patch; a high-activity allele of *for* is present in rover larvae, which move quickly and disperse rapidly<sup>22</sup>. A *for*-related cGMP-dependent kinase affects foraging in honeybees, ants and nematodes, suggesting that diverse animals share molecular mechanisms for behavioural regulation<sup>22,23</sup>.

To gain further insight into the genetics and neurobiology of exploratory behaviour in *C. elegans*, we here use quantitative genetic analysis to examine this behaviour's genetic architecture in wild-type strains, and show that genetic variation in multiple loci, including a catecholamine receptor, interacts with environmental conditions to regulate the exploitation–exploration decision.

## Multiple loci affect leaving behaviour

Different wild-type strains of *C. elegans* vary in their tendency to leave or remain on a standardized small lawn of bacterial food (Fig. 1a). For example, adult hermaphrodites from the laboratory strain N2 leave the lawn only once every 100 min, whereas animals from the CB4856 (HW) strain isolated from pineapple fields in Hawaii leave the lawn once every 5–6 min (Fig. 1b, Supplementary Movies 1 and 2). To determine the genetic architecture of this behavioural difference between N2 and HW, we quantified leaving rates in 91 N2-HW recombinant inbred advanced intercross lines (RIAILs)<sup>24</sup>. 58 of the RIAILs had low leaving rates comparable to N2, only 6–10 had high leaving rates comparable to HW, and 23 had intermediate rates (Fig. 1c). The excess of low leaving rates and the continuous behavioural distribution in RIAILs suggest that leaving is a multigenic quantitative trait.

Quantitative trait locus (QTL) analysis of the RIAILs uncovered two regions with significant effects on leaving rates, one on the X chromosome and one on chromosome II (Fig. 1d). The X chromosome QTL overlapped with the location of the polymorphic G-protein-coupled neuropeptide receptor NPR-1, which affects many food-related behaviours<sup>12,15</sup>. The *npr-1* polymorphism has previously

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Figure 1 | Lawn-leaving behaviour varies between wild-type *C. elegans* strains. a, Lawn-leaving assays. Top, six adult HW hermaphrodites on a bacterial lawn. One animal has left the lawn and one is leaving. Bottom, track of a HW animal during 5 min of an assay. The track is colour-coded to show passage of time (colour scale at bottom). The border of the lawn is outlined. Scale bar, 6 mm. **b**, Leaving rates of six wild-type strains. **c**, Leaving rates of 91 N2-HW recombinant inbred advanced intercross lines (RIAILs)<sup>24</sup> and parental strains. **d**, QTL analysis of RIAILs shown in **c**. The horizontal line denotes the *P* < 0.01 genome-wide significance threshold. lod, log likelihood ratio. Error bars in **b** and **c**, s.e.m.

been shown to affect leaving, as well as locomotion speed on food, a behaviour that partially correlates with leaving rate<sup>12,15</sup> (Supplementary Fig. 2). Examining the *npr-1* genotype in the RIAILs revealed a strong but asymmetric correlation with leaving rates (Fig. 1c). Every strain with the N2 allele of *npr-1* had low leaving rates ( $\leq 1$  event every 20 min), but strains with the HW allele of *npr-1* could have either low or high leaving rates (Fig. 1c). The asymmetric distribution is consistent with a role for *npr-1* in leaving behaviour, but indicates that *npr-1* has epistatic interactions with other loci segregating in the RIAILs.

The involvement of npr-1 in leaving behaviour was confirmed by analysing near-isogenic lines (NILs) containing the N2 and HW npr-1alleles in the reciprocal strain background, and by examining npr-1null mutants (Supplementary Fig. 3). Specific transgenic expression of the N2 npr-1 allele in its essential site of action, the RMG motor neurons<sup>20</sup>, sharply reduced the leaving rate of HW animals (Supplementary Fig. 3). Thus npr-1 is a regulator of HW leaving rates, but not the only contributing gene.

#### tyra-3 affects leaving behaviour

Studies in yeast, flies, mice, and plants have shown that individual QTLs often resolve into several genes that contribute to phenotypic variance<sup>25–28</sup>. Similarly, fine-mapping of the  $\sim$ 1 megabase (Mb) QTL

that contained *npr-1* suggested the existence of multiple loci that affected leaving rates. A NIL with <150 kilobases (kb) of N2 DNA spanning the *npr-1* locus introgressed into HW had N2-like leaving rates (*leav-1* QTL, Fig. 2a and Supplementary Fig. 3). A second NIL with 700 kb of N2 DNA that did not cover *npr-1* introgressed into HW also had a low leaving rate, with about half the leaving rate of HW (*leav-2* QTL, Fig. 2a). These results suggest the existence of a second X-linked locus that affects leaving rates, which we called *leav-2*. The *leav-2* region did not affect leaving in the N2 genetic background (Fig. 2a), so all subsequent experiments were conducted in the HW background.

A 100 kb minimal region for *leav-2* was identified by analysing the breakpoints of individual RIAILs (Supplementary Fig. 4 and Supplementary Methods). We characterized the genetic properties of leav-2 by crossing the leav-2 NIL strain with HW. The heterozygous F<sub>1</sub> progeny had leaving rates similar to the *leav-2* NIL (Fig. 2a), indicating that the N2 leav-2 locus was dominant to HW and suggesting that N2 transgenes covering the relevant gene should reduce the leaving rate of HW animals. Therefore, overlapping N2 genomic DNA fragments from the 100 kb minimal leav-2 region were introduced into HW animals by microinjection (Fig. 2b and Supplementary Fig. 5). A single gene in this region reduced leaving rates: tyra-3, which encodes a G-protein-coupled receptor for the invertebrate noradrenaline-like neurotransmitters tyramine and octopamine<sup>29</sup>. Tyramine and octopamine receptors are related to vertebrate adrenergic receptors, and are thought to carry out analogous functions. tyra-3 genomic fragments from the N2 strain were more active than tyra-3 fragments from the HW strain injected at the same concentration, consistent with the possibility that tyra-3 is a polymorphic gene that differs between N2 and HW (Fig. 2b).

If *leav-2* corresponds to *tyra-3*, a *tyra-3* mutation should eliminate its activity<sup>30</sup>. To test this prediction genetically, a null allele of *tyra-3* in an N2 background was introgressed into a HW background. The N2 region in the resulting NIL covered from 4.9 to 5.4 Mb of the X chromosome, the inferred position of *leav-2*. The *tyra-3(ok325)* null NIL had high (HW-like) leaving rates, suggesting that N2 *leav-2* activity was not present in the strain (Fig. 2a). Heterozygotes between HW and the near-isogenic *tyra-3(ok325)* null strain also had high leaving rates (Fig. 2a). These results are as expected if the active locus in *leav-2* is *tyra-3*; however, other genes within the introgressed regions could also contribute to the different leaving rates.

To strengthen the connection between *tyra-3* and *leav-2*, RNA interference against *tyra-3* was performed in the *leav-2* NIL that has low leaving rates due to the presence of the N2 QTL. Knockdown of *tyra-3* increased the leaving rate of the *leav-2* NIL to levels observed in HW animals, the result predicted if the *tyra-3* locus from N2 reduces leaving (Fig. 2c). Comparable experiments in a pure HW strain had minimal effects, as expected if *tyra-3* activity in HW is already low.

Further confirmation that the HW allele of *tyra-3* has reduced biological activity was provided by examining the one phenotype previously associated with *tyra-3*, avoidance of dilute octanol<sup>29</sup>. *tyra-3* null mutants avoid octanol more strongly than wild-type N2; the NIL strain with the HW *tyra-3* allele had a similar enhanced octanol response, suggesting that the HW *tyra-3* allele has reduced *tyra-3* function (Supplementary Fig. 6).

#### Non-coding changes affect *tyra-3* activity

The differential activity of N2 and HW genomic *tyra-3* fragments in the leaving assay suggested that N2 and HW alleles are functionally distinct (Fig. 2b). To identify polymorphisms between N2 and HW alleles of *tyra-3*, we sequenced ~19 kb surrounding the *tyra-3* locus in HW. There were 34 differences between HW and the N2 consensus genomic sequence (Fig. 3a): 33 non-coding changes and a single coding difference that changed a glutamic acid in the *tyra-3b* isoform to glycine.

Sequences that contribute to the differential activity of N2 and HW *tyra-3* alleles were localized further using transgenic assays. We fused

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N2 and HW *tyra-3b* complementary DNAs to 4.9 kb of non-coding N2 or HW sequence upstream of the *tyra-3b* start site and introduced each of the four resulting clones into the HW strain. *tyra-3* transgenes with the N2 non-coding sequence were significantly more potent than comparable transgenes with the HW sequence, regardless of whether they preceded N2 or HW *tyra-3* cDNAs (Fig. 3b), excluding the coding polymorphism and localizing a functional difference between N2 and HW *tyra-3* genes to a 4.9 kb region that harbours 5 non-coding SNPs, 1 single nucleotide insertion, and a 184 bp deletion in HW. To



narrow the relevant change down further, the 184 bp deletion was engineered into the N2 *tyra-3* genomic fragment; this clone was significantly less potent in the leaving assay than the full N2 genomic fragment (Supplementary Fig. 7). These results indicate that the 184 bp deletion represents at least part of the functional difference between N2 and HW *tyra-3* alleles.

Sequence variation in *tyra-3* non-coding regions could affect the level or location of *tyra-3* expression. Quantitative reverse transcription–PCR analysis (RT–PCR) of *tyra-3* mRNA levels in mixed-stage animals





**b**, Leaving rates of transgenic HW animals with *tyra-3b* promoters (*Ptyra-3b*) fused to *tyra-3b* cDNAs. Error bars, s.e.m. \*\*P < 0.01 by two-way ANOVA; no statistical interaction between the promoter and the cDNA. **c**, Relative amounts of *tyra-3* isoform mRNAs (**a**) in HW, N2 and *leav-2* strains (Fig. 2a). Error bars, s.d. \*\*P < 0.01 compared to HW, ANOVA with Dunnett test.

indicated that N2 expressed approximately twice as much *tyra-3* mRNA as HW, consistent with increased *tyra-3* activity in the N2 strain (Fig. 3c). The *leav-2* NIL with N2 *tyra-3* introgressed into HW also had high *tyra-3* messenger RNA levels, suggesting that *cis*-acting changes affect *tyra-3* expression (Fig. 3c).

As both N2 and HW were cultivated in the laboratory for many years before permanent cultures were frozen, we wished to exclude the possibility that the *tyra-3* polymorphisms were laboratory-derived<sup>19</sup>. Therefore, 19 kb of the *tyra-3* locus was sequenced in all wild strains tested for leaving behaviour in Fig. 1, including three strains that were frozen immediately after their isolation. Each strain represents a different *C. elegans* haplotype group<sup>24</sup>. Both N2-like and HW-like *tyra-3* sequences were represented in the wild-caught strains, confirming the wild ancestry of both alleles (Supplementary Table 1 and Supplementary Methods). Notably, the *tyra-3* locus of MY1 was identical to N2 and, correspondingly, the leaving rate of MY1 was similar to that of N2.

### tyra-3 acts in sensory neurons

The identification of *tyra-3* provided an opportunity to characterize the neuronal basis of the decision to leave or remain on a food patch. The biological activity of a transgene with 4.9 kb upstream of the *tyra-3b* start site fused to a *tyra-3* cDNA (Fig. 3b) implied that it was expressed in cells that regulate leaving behaviour. When this 4.9 kb region was fused to green fluorescent protein (GFP), it drove reliable expression in ASK, ADL, AIM, AUA, BAG, CEP, OLQ and SDQL neurons, in other unidentified neurons in the ventral ganglion and the tail, occasionally in AFD and AWC neurons, and in two non-neuronal cell types, the spermatheca and the distal tip cell (Fig. 4a and data not shown). The same set of cells was observed with reporter genes bearing either N2 or HW *tyra-3* upstream regions, and in both N2 and HW genetic backgrounds. Together with the quantitative RT–PCR data (Fig. 3c), these results suggest that different *tyra-3* expression levels, not different sites of expression, distinguish N2 and HW alleles.

The neurons whose activity is regulated by *tyra-3* were localized further by expressing *tyra-3* cDNAs from cell-type-specific promoters. *tyra-3* expression in ASK or BAG sensory neurons significantly reduced leaving, but expression in the CEP or ADL sensory neurons

did not (Fig. 4b). The ASK neurons sense attractive food-derived amino acids<sup>31</sup> and regulate search behaviours after animals are removed from food<sup>32,33</sup>. The BAG neurons sense  $CO_2$  and  $O_2$ , two cues associated with bacterial metabolism<sup>34,35</sup>. Lowering  $O_2$  to levels that activate BAG reduced leaving rates (Supplementary Fig. 8).

To investigate whether the *tyra-3* non-coding polymorphism affects expression in relevant neurons, single-copy N2 or HW *tyra-3b* promoters driving GFP were inserted into a single, defined chromosomal location using the MosSCI technique<sup>36</sup>. GFP levels in ASK neurons were significantly higher for transgenes containing the N2 promoter compared to those containing the HW promoter (Fig. 4c). These results suggest that the N2 *tyra-3* locus is associated with higher *tyra-3* expression in ASK, as well as higher *tyra-3* mRNA expression at a whole-animal level; expression in BAG was not examined.

The behavioural functions of ASK and BAG, and the effect of *tyra-3* on those functions, were assessed by killing the neurons in different genetic backgrounds. Killing the ASK neurons reduced the leaving rate of HW animals, indicating that ASK can promote leaving (Fig. 4d). The ablation resembled the effect of the ASK::*tyra-3* transgene, suggesting that *tyra-3* reduces ASK activity. In agreement with this idea, killing the ASK neurons in a strain with the N2 high-activity *tyra-3* allele did not reduce their leaving rates further. The effect of *tyra-3* on ASK was selective for this assay; *tyra-3* did not reduce lysine chemotaxis, a second ASK-dependent behaviour (Supplementary Fig. 9).

Killing the BAG neurons increased leaving rates in the strain with the N2 *tyra-3* allele, demonstrating that BAG neurons prevent leaving (Fig. 4d). However, killing BAG had no effect in the strain with the HW *tyra-3* allele, suggesting that BAG activity is already low in this strain under the assay conditions. The ablation and genetic results suggest that the N2 *tyra-3* allele decreases ASK activity and increases BAG activity, two changes that act together to prevent leaving (Supplementary Fig. 1).

#### **Gene-gene-environment interactions**

Like most natural behaviours, the decision to leave a food patch is regulated by multiple genes and the environment; it responds to genetic variation in *tyra-3*, *npr-1*, and additional genes on the autosomes





*tyra-3b* in specific cells. **c**, Left, GFP fluorescence intensity in ASK of HW animals with a MosSCI insertion of N2 or HW 4.9 kb *tyra-3b* promoter::GFP. Right, schematic of MosSCI technique<sup>36</sup>. **d**, Leaving rates after killing ASK or BAG in HW and *leav-2* strains (Fig. 2a). Error bars, s.e.m. \*P < 0.05, \*\*P < 0.01, \*\*\*P < 0.001 by *t*-test or ANOVA with Dunnett test.

(Fig. 1) as well as food quality and quantity<sup>7,8</sup>. Our results suggested that the N2 *npr-1* allele was epistatic to *tyra-3*; animals with the N2 *npr-1* allele had low leaving rates regardless of the *tyra-3* genotype (Fig. 2a). However, N2 *npr-1* reduced the leaving rate to almost zero, making it difficult to detect any further reduction. To make the assay more powerful, leaving was assayed on bacterial lawns of different densities. Leaving rates of all genotypes increased on thinner lawns and decreased on thicker lawns (Supplementary Fig. 10), but the thickness of the lawn changed the genetic interaction between *tyra-3* and *npr-1*. In the standard leaving assay, *tyra-3* polymorphisms had different effects only in the presence of the HW *npr-1* allele; on a thinner lawn, only in the presence of the N2 *npr-1* allele (Supplementary Fig. 10). Thus the epistatic relationship between *npr-1* and *tyra-3* is defined by the specific environment, not by an intrinsic regulatory relationship between the genes.

#### Discussion

Our results show that natural variation in *tyra-3* affects patch leaving, a behaviour representative of the exploration–exploitation decision. *tyra-3* encodes a G-protein-coupled receptor activated by the invertebrate transmitters tyramine and octopamine<sup>29</sup>, which are structurally related to vertebrate adrenaline and noradrenaline. Catecholamines are known to regulate arousal systems that affect many behaviours and behavioural decisions. In *C. elegans*, octopamine drives sensory, molecular and behavioural responses to starvation, and tyramine affects specific aspects of locomotion<sup>37–40</sup>. In insects, octopamine affects locomotory activity, arousal and aggression<sup>41–43</sup>. Mammalian noradrenaline is generally implicated in arousal behaviours, and noradrenaline release from the locus coeruleus is associated with switching between different tasks, a cognitive function with analogies to the exploration–exploitation decision<sup>6</sup>.

Relatively few natural behavioural variations have been mapped to the single-gene level in any animal, and it is interesting that several of these variations affect G-protein-coupled receptor signalling systems<sup>44,45</sup>. We speculate that these receptor pathways may serve as common substrates of behavioural variation. All animal genomes encode many G-protein-coupled receptors with different expression patterns. These receptors may provide a reservoir for genetic changes, as alteration in an individual receptor could cause relatively discrete effects without disrupting the entire system.

QTL mapping in rodents and in *Drosophila* indicates that most behavioural traits are polygenic, with widespread epistatic effects<sup>1,46</sup>. In agreement with this conclusion, our analysis suggests the existence of epistatic interactions between *tyra-3*, *npr-1* and at least one additional locus. Importantly, the non-additive interactions between *tyra-3* and *npr-1* are not stable, but vary based on the genetic background and the environment, similar to what has been found with yeast sporulation QTLs<sup>47</sup>.

By integrating genetic studies of *C. elegans* foraging with neuronal analysis, we can provide a first-level description of underlying mechanisms. The sensory neurons that express *tyra-3* detect food-related cues; we suggest that they integrate these external cues with internal arousal states detected by *tyra-3*, and that different *tyra-3* alleles confer differential sensitivity to these arousal states (Supplementary Fig. 1). Thus variation in *tyra-3* lies at the intersection of many forms of behavioural flexibility: rapid responses to environmental cues, short-term modulation by internal state fluctuations, and long-term genetic changes that lead to adaptive changes in innate behaviours.

#### METHODS SUMMARY

Standardized leaving assays were conducted by videotaping seven adult hermaphrodite animals for 30 min on a bacterial lawn that was low-density compared to their growth lawn. The number of leaving events was recorded manually by examining the video recordings, and further behavioural analysis was conducted with a Matlab code adapted from the Parallel Worm Tracker<sup>48</sup>. A leaving event was defined as an episode in which the whole body of an animal left the bacterial lawn and the animal did not reverse immediately to return to the lawn. The leaving rate was calculated as the number of leaving events per worm minute spent inside the bacterial lawn. Experiments on each strain were repeated at least three times.

QTL analysis was performed on the mean leaving rates of N2-HW recombinant inbred advanced intercross lines (RIAILs) by non-parametric interval mapping in R/qtl<sup>24,49</sup>. Significance levels were estimated from 10,000 permutations of the data. Near-isogenic lines (NILs) were created by backcrossing a chromosomal region or allele into the desired genetic background at least 9 times. Extrachromosomal transgenes were made by injection of DNA clones into the gonads of young adult hermaphrodites together with a fluorescent coinjection marker<sup>50</sup>. To control for variation between transgenes, two to five independent lines from each injection were characterized. Single-copy insertion of transgenes was performed using the direct MosSCI transposition technique, targeting the *ttTi5605* Mos allele on chromosome II<sup>36</sup>.

Full Methods and any associated references are available in the online version of the paper at www.nature.com/nature.

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**Supplementary Information** is linked to the online version of the paper at www.nature.com/nature.

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#### **METHODS**

Analysis of behaviour in the leaving assay. Six-centimetre NGM agar plates were seeded with 70  $\mu$ l (conditioning plate) or with 10  $\mu$ l (assay plate) of a fresh overnight culture of *Escherichia coli* HB101 diluted in LB to  $A_{600nm} = 2.0.90$  min after seeding the plates, ten young adult hermaphrodites were picked onto the conditioning plate. 30 min after being placed on the conditioning plates, seven of the animals were transferred onto the lawn of the assay plate. The 30 min leaving assay began 1 h after placing the seven animals on the assay plate. The number of leaving events was recorded manually by examining the video recordings, and further behavioural analysis was conducted with a Matlab code adapted from the Parallel Worm Tracker<sup>48</sup>. A leaving event was defined as an episode in which the whole body of an animal left the bacterial lawn and the animal did not reverse immediately to return to the lawn. The leaving rate was calculated as the number of leaving events per worm minute spent inside the bacterial lawn. Experiments on each strain were repeated at least three times.

**Quantitative trait locus analysis.** The N2-HW recombinant inbred advanced intercross lines (RIAILs) used in this study represent the terminal generation of a 20-generation pedigree founded by reciprocal crosses between N2 and HW. The lines were constructed through 10 generations of intercrossing followed by 10 generations of selfing<sup>24</sup>. They have been genotyped at 1,454 nuclear and one mitochondrial markers and have a 5.3-fold expansion of the F<sub>2</sub> genetic map<sup>24</sup>. QTL analysis was performed on the mean leaving rates of N2-HW RIAILs by non-parametric interval mapping in R/qtl<sup>49</sup>. Significance levels were estimated from 10,000 permutations of the data.

**Quantitative RT-PCR.** Total RNA from mixed stage worms was isolated with Trizol. 1.5 µg of RNA and oligo-dT were used for reverse transcription using SuperScript III First-Strand Synthesis (Invitrogen) according to the manufacturer's instructions. Real-time PCR was performed with Fast SYBR Green Master Mix (Applied Biosystems) on a 7900HT Real-Time PCR System (Applied Biosystems). *act-3* was used as the calibrator for relative quantitation. 5' primers corresponded to upstream exons that distinguished *tyra-3* isoforms, and 3' primers corresponded to shared exon sequence. Primers used were: *tyra-3a*  $\Leftrightarrow$  *tyra-3c.2\_F*, ccacttgcaatagcagcag; *tyra-3b\_F*, ggctatttggtggtggtttg; *tyra-3a*  $\Leftrightarrow$  *tyra-3b\_R*, tccttctggcgtcgaaatac; *act-3\_F*, tcacgatcatgagaccattcaaa; *act-3\_R*, gcaaattgtagtgggtcttcttatg.

tyra-3 expression pattern. The N2 and HW 4.9 kb tyra-3b promoters were amplified using primers: tcaacctaaccactaactaaggg and cGatgaagcaagatgtcaggt, which overlaps the coding region by 4 bp. The ATG start codon is mutated to ATC (mutation is uppercase in primer). These promoters were individually fused by PCR to a fragment containing GFP followed by the unc-54 3'-UTR, as described  $^{\scriptscriptstyle 51}$  . These PCR products were injected individually into both HW and N2 animals at 20 ng µl<sup>-1</sup>. Cells expressing GFP were identified by Nomarski microscopy in both L1 and adult hermaphrodites. The identification of some cells was aided by injecting Ptyra-3b::GFP-expressing animals with promotermCherry fusions with established expression patterns. In this manner, the AIM neurons were identified as Ptyra-3b::GFP-expressing cells based on their position and the absence of co-localization with Pttx-3::mCherry. The BAG neurons coexpressed Ptyra-3b::GFP and Pflp-17::mCherry. The CEP neurons co-expressed Ptyra-3b::GFP and Pdat-1::mCherry. The ASK neurons co-expressed Ptyra-3b::GFP and Psra-9::mCherry. The ADL neurons co-expressed Ptyra-3b::GFP and Psri-51::mCherry.

**Extrachromosomal transgenes.** Transgenes were made by injection of DNA clones into the gonads of young adult hermaphrodites together with a fluorescent coinjection marker<sup>50</sup>. To control for variation between transgenes, two to five independent lines from each injection were characterized.

Generation of MosSCI lines and quantitation of GFP fluorescence in ASK. Single-copy insertion of transgenes was performed using the direct MosSCI technique targeting the *ttTi5605* Mos allele on chromosome II, as described<sup>36</sup>. A schematic of the mechanism underlying MosSCI is shown in Fig. 4c.

The pCFJ151 targeting vector was modified by the introduction of an FseI restriction site into the multiple cloning site by site-directed mutagenesis using the primers gtaatacgactcacttaaggccggccctaaggggtaccagagctcacc and ggtgagctctggt accctctagggccggccttaagtgagtcgtattac to make pAB1. An FseI-SpeI fragment from a pSM vector containing N2-Ptyra-3b::N2-tyra-3b::SL2 GFP::unc-54 3'-UTR or HW-Ptyra-3b::N2-tyra-3b::SL2 GFP::unc-54 3'-UTR was cloned into pAB1.

For each *tyra-3*-containing test plasmid, about 50 EG4322 animals were injected with a mixture of *tyra-3* plasmid, pGH8, pCFJ90, pCFJ104 and pJL43.1. After positive and negative selection and full sequencing of the insert, two inserted transgenes each of N2-*Ptyra-3b* and HW-*Ptyra-3b* were backcrossed to HW males seven times, selecting GFP-fluorescent hermaphrodites each generation. The transgene-containing chromosome was then homozygosed.

The strains containing the single-copy transgene in a HW background were injected with *Psra-9*::mCherry to identify ASK. Young adult hermaphrodites were examined on a Zeiss Imager Z.1 with a ×60 objective focused on ASK using mCherry to prevent bleaching of GFP signal. Fluorescence signals were acquired with fixed acquisition times (30–50 ms for mCherry, 100 ms for GFP). Background mean fluorescence intensity adjacent to ASK was subtracted from the ASK signal.

**RNAi.** RNA interference was performed essentially as described<sup>52</sup>. A fragment common to all *tyra-3* isoforms was amplified. The following primers were used, which include the T7 sequence (underlined): <u>taatacgactcactatagggagagaaaatggc</u> agcaggacttt; <u>taatacgactcactatagggagagaaatccccgcagtctgtggagt</u>. *in vitro* transcription was performed with RiboMAX kit (Promega). dsRNA was injected at 1.2 µg µl<sup>-1</sup> into the gonads of adult hermaphrodites. Eggs laid 24 and 48 h after injection were used for the behavioural assays.

**Octanol avoidance assay.** Avoidance assays were conducted essentially as described<sup>53</sup>. In brief, ~20 three-day-old animals were picked off of their growth plates food into a transfer plate without bacteria where they were allowed to crawl and rid themselves of bacteria. Animals were then transferred onto an NGM plate without food. After 40 min, a microcapillary with 30% octanol (v/v diluted fresh every day in ethanol) was presented in front of the animal's nose. The time to reverse was recorded. If animals did not reverse within 20 s, the assay was stopped. Animals were presented with odour 1–3 times per experiment, with at least 3 min of rest interval. We replicated published results demonstrating that *tyra-3* null mutants had more rapid responses than N2 in the presence of exogenous serotonin and tyramine<sup>29</sup> but also observed more rapid responses in the absence of exogenous neuromodulators, as shown in Supplementary Fig. 6.

**Cell ablations.** For leaving behaviour assays, ASK was ablated with a laser microbeam as described<sup>54</sup>. BAG was killed using split human caspase 3 fragments<sup>55</sup> expressed from *flp-17* and *glb-5* promoters that overlapped only in BAG. For lysine chemotaxis assays, ASK was killed using a mouse caspase 1 gene expressed from the *sra-9* promoter<sup>56</sup>. The ASK strain was a gift from R. Shingai.

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