RFX Transcription Factor DAF-19 Regulates 5-HT and Innate Immune Responses to Pathogenic Bacteria in Caenorhabditis elegans

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Abstract

In Caenorhabditis elegans the Toll-interleukin receptor domain adaptor protein TIR-1 via a conserved mitogen-activated protein kinase (MAPK) signaling cascade induces innate immunity and upregulates serotonin (5-HT) biosynthesis gene tph-1 in a pair of ADF chemosensory neurons in response to infection. Here, we identify transcription factors downstream of the TIR-1 signaling pathway. We show that common transcription factors control the innate immunity and 5-HT biosynthesis. We demonstrate that a cysteine to tyrosine substitution in an ARM motif of the HEAT/Arm repeat region of the TIR-1 protein confers TIR-1 hyperactivation, leading to constitutive tph-1 upregulation in the ADF neurons, increased expression of intestinal antimicrobial genes, and enhanced resistance to killing by the human opportunistic pathogen Pseudomonas aeruginosa PA14. A forward genetic screen for suppressors of the hyperactive TIR-1 led to the identification of DAF-19, an ortholog of regulatory factor X (RFX) transcription factors that are required for human adaptive immunity. We show that DAF-19 regulates TIR-1 signaling via a conserved mitogen-activated protein kinase (MAPK) signaling cascade. We show that common transcription factors control the innate immunity and 5-HT biosynthesis. Our results suggest that pathogenic cues trigger a common core-signaling pathway via tissue-specific mechanisms and demonstrate a novel role for RFX factors in neuronal and innate immune responses to infection.

Introduction

Innate immunity is an integral part of the stress response program in which the host activates a range of defense genes to enhance the chance of survival against internal and environmental threats. In mammals, signals associated with pathogenic microbes trigger Toll-like receptors to recruit Toll-interleukin receptor (TIR) domain adaptor proteins, thereby forming scaffolds with downstream signaling cascades leading to transcriptional upregulation of defense genes. A growing body of evidence indicates that classical immune proteins including Toll-like receptors and TIR domain adaptor proteins are expressed in the developing and mature brain in mammals [1,2]. It has been proposed that certain common molecular mechanisms may function in neurons and non-neuronal tissues to induce physiologically distinct responses to aversive cues [1,2,3]. Except a few cases, the gene targets of immune factors in neurons are not known and it is unclear whether those immune signaling cascades are differentially regulated in neurons and non-neuronal tissues. Consequently, identification of upstream regulators and downstream effectors of conserved core immune signaling pathways may provide insights into the regulation of the immunity as well as the regulation of neural plasticity.

Our laboratory has focused on genetic dissection of environment-dependent transcriptional regulation of the tph-1 gene, encoding the rate-limiting serotonin (5-HT) biosynthesis enzyme tryptophan hydroxylase, in the nematode Caenorhabditis elegans. Previously, we showed that tph-1 expression in a pair of ADF chemosensory neurons in the head sensory organ Amphid is modulated by two layers of transcriptional regulation according to growth conditions: signaling through the OCR-2/OSM-9 TRPV channel turns on the basal tph-1 expression under optimal growth conditions; signaling through the OCR-2/OSM-9 TRPV channel turns on the basal tph-1 expression under optimal growth conditions; and aversive growth conditions further upregulate tph-1 expression independently of OCR-2/OSM-9 [4,5]. Work from several laboratories suggests that tph-1 expression in the ADF neurons responds to pathogenic food. C. elegans feeds on bacteria and is killed by a large number of pathogenic microbes in its natural environment [6]. In an elegant study, it showed that feeding worms with the human opportunistic pathogen Pseudomonas aeruginosa PA14 triggers upregulation of tph-1 in the ADF neurons leading to aversive learning and avoidance behavior [7]. A subsequent study indicated that the TIR-domain adaptor protein

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Author Summary

Toll-interleukin receptor (TIR)-domain adaptor proteins are keys to activate signaling cascades inducing transcriptional responses to internal and external pathogenic signals in evolutionary disparate organisms. Despite lacking a homolog of the mammalian innate immunity transcriptional regulator nuclear factor-kappaB (NF-κB), the nematode Caenorhabditis elegans responds to infections by activating TIR-1 signaling targets in the innate immune system and in neurons. Through a genetic screen for factors required for TIR-1 signaling to upregulate the serotonin biosynthesis gene tph-1, we identified DAF-19, an ortholog of regulatory factor X (RFX) transcription factors that were initially discovered in human immune cells. We show that DAF-19 concerts with ATF-7, a member of the activating transcription factor (ATF)/cAMP response element-binding B (CREB) family of transcription factors, to upregulate tph-1 in the ADF chemosensory neurons and antimicrobial genes in the intestine in response to bacterial infection, reminiscent of RFX-CREB interaction in human immune cells.daf-19 mutants display heightened susceptibility to killing by the human pathogen Pseudomonas aeruginosa PA14. Our studies suggest that RFX transcriptional regulation, which is essential for human adaptive immunity, has an ancient role in controlling serotonin biosynthesis and innate immunity.

TIR-1, which was initially identified as an upstream regulator of a conserved mitogen-activated protein kinase (MAPK) signaling pathway in the innate immunity [3], is required for PA14-induced tph-1 upregulation and PA14 avoidance behavior [9]. However, the C. elegans genome lacks a homolog of nuclear factor-kappaB (NF-κB), the major transcriptional activator of the mammalian innate immunity. In addition, deletion of the sole C. elegans Toll receptor gene tol-1 did not affect the intestinal immunity or tph-1 expression [7]. These observations suggest that the TIR-1 signaling cascade may evolve more evolutionarily ancient upstream players and downstream transcription factors.

Activating transcription factor (RFX) transcription factors were first identified in human subjects of bare lymphocyte syndrome, a hereditary immunodeficiency disease, and are required for the expression of the major histocompatibility complex class II (MHC II) genes [12]. RFX proteins bind to the X-box motif on the MHC II promoters and interact with cAMP response element-binding (CREB) protein and other cofactors to form a higher order “enhanceosome”, which then recruits the non-DNA-binding transcriptional activator CIITA to turn on MHC II expression [13]. RFX factors have since been identified in diverse eukaryotic species [14,15,16] and are expressed broadly in neuronal and non-neuronal cells in animals, suggesting additional roles for RFX factors in biological processes of multiple tissues. Studies of the sole C. elegans RFX factor dat-19 have uncovered its role in the development of dendritic cilia of sensory neurons [17,18]. Subsequent studies found RFX factors regulating ciliogenesis in Drosophila [19] and mouse [20], demonstrating one aspect of RFX function conserved across phyla.

In this paper, we identified DAF-19 as a key transcriptional regulator of tph-1 in the ADF neurons and intestinal antimicrobial genes in C. elegans. We found that, analogous to the RFX-CREB interaction for MHC II expression in human immune cells, DAF-19 concert with ATF-7, a member of activating transcription factor (ATF)/CREB superfamily of transcription factors, acting downstream of the TIR-1 signaling cascade to control transcriptional responses to pathogenic bacterial food in C. elegans. We show that the TIR-1-DAF-19/ATF-7 pathway is differentially regulated to induce tph-1 upregulation and intestinal immunity in response to P. aeruginosa PA14. Thus, our data suggest that pathogenic signals may trigger a common core signaling pathway via cell-specific mechanisms and a RFX transcription factor acts in an ancient host to regulate 5-HT biosynthesis and the innate immunity.

Results

Isolation of tir-1(yz68) gain-of-function mutation

We carried out a forward genetic screen to identify components underlying aversive environment-induced tph-1 upregulation in C. elegans. The levels of tph-1 expression in identified neurons in living C. elegans can be estimated by quantifying fluorescence intensity of a green fluorescence protein (GFP) driven by the tph-1 promoter (tph-1::gfp) [4]. A pair of ADF neurons is the only chemosensory neurons producing 5-HT in a hermaphrodite C. elegans [21]. Each ADF neuron projects a single dendrite to the tip of the nose where the ciliated sensory endings are exposed to the external environment and its axon extends to the nerve ring, the brain of C. elegans [22]. We started with a strain expressing a stably chromosomally integrated tph-1::gfp transgene in ocr-2(yz5) mutant background, in which tph-1::gfp expression in the ADF neurons is visible under aversive growth conditions but not under optimal growth conditions, providing a visual assay for environment-dependent changes in tph-1 expression [5]. We isolated mutagenized worms with enhanced ADF tph-1::gfp under optimal growth conditions, and analyzed the mutants in the ocr-2 background as well as after the ocr-2 mutation being outcrossed, yz68 is one of the mutants retrieved from the screen.

Through single nucleotide polymorphism-based (SNP) mapping, RNA-interference (RNAi)-mediated inactivation of candidate genes in the mapped contig, and sequencing the yz68 mutant genome, we identified a nucleic acid change predicting a substitution of cysteine326 by tyrosine (C426Y) in the fourth ARM motif of the HEAT/Arm repeat region of the TIR-1 protein (Figure 1A). Several experimental data suggest that the C426Y substitution causes TIR-1 constitutive activation. First, whole mount anti-5-HT-antibody staining showed that ADF 5-HT immunoreactivity in tir-1(yz68)ocr-2 double mutants was elevated compared to the ocr-2 single mutant (Figure 1D). As 5-HT is being secreted, 5-HT immunostaining does not fully reflect the rate of 5-HT biosynthesis. With this caveat in mind, the results suggest that tir-1(yz68) enhanced 5-HT in the ADF neurons. Second, transgenic expression of tir-1(yz68) cDNA under the gpa-13 promoter (Pyga-13::tir-1(yz68)), which is expressed in ADF, AWC and ASH sensory neurons in the head region, recapitulated tph-1::gfp upregulation (Figure 1B). Third, RNAi of tir-1 in tir-1(yz68):ocr-2 mutants blocked the tph-1 upregulation (Figure 1C). The tir-1(ok1052) mutation, which causes mixed gain- and loss-of-function tir-1 phenotypes in the AWC neuron development [23] but does not affect the innate immunity [24], caused only a modest increase in ADF tph-1::gfp (Figure 1C). Collectively, these data suggest that the C426Y substitution alters a site critical for TIR-1 activation in the ADF neurons.
regulator of basal tph-1 expression in the ADF neurons under optimal growth conditions.

TIR-1 signaling pathway selectively regulates serotonergic response to pathogenic food

Previously, we identified that a number of aversive conditions may induce tph-1 upregulation in the ADF neurons [5]. We asked whether tir-1 signaling specifically mediates the response to pathogenic bacteria, or it is responsible for tph-1 upregulation under all aversive conditions. We first analyzed the intensity of tph-1::gfp in the ADF neurons in tir-1(0f) and tir-1(gf) mutants fed with the pathogenic P. aeruginosa strain PA14. Following feeding on PA14 for 6 hr, ADF fluorescence in WT animals was ~1.4-fold higher than their sibling on OP50, but tir-1(0f) mutants failed to upregulate tph-1::gfp under the same conditions (Figure 2A), consistent with prior reports [7,9]. The tir-1(yz68gf) mutant also did not exhibit a significant increase in tph-1::gfp following PA14 feeding, suggesting that the pathogen signals cannot further enhance tir-1(yz68gf) protein activity (Figure 2A).

We next tested if TIR-1 function is required for tph-1 upregulation during dauer formation. Under the conditions of starvation, high growth temperature and high levels of phero-
mones, *C. elegans* develops into a stress-resistant dauer larva through a series of cellular and physiological remodeling and turns on a battery of stress genes [25]. We previously showed that WT and *ocr-2* mutant worms upregulated ADF *tph-1::gfp* when they entered the dauer stage [5]. We therefore induced *tir-1(tm3036lf)* and *tir-1(yz68gf)* mutants to form dauers by treating the worms with dauer pheromones. We observed ADF *tph-1::gfp* upregulation in both *tir-1(tm3036lf)* and *tir-1(yz68gf)* mutants during dauer formation (Figure 2B). Thus neither *TIR-1* deficiency nor *TIR-1* hyperactivation can block *tph-1::gfp* upregulation induced by dauer formation.

We previously showed that mutations that alter the morphology of dendritic cilia of all chemosensory neurons including ADF and cause ADF *tph-1::gfp* upregulation on its own as well as in *tir-1(lf)* background. *Tir-1* and *daf-6* mutations confer additive upregulation of *tph-1::gfp* in the ADF neurons. Data represent the average of three trials each with at least 20 animals per strain per condition ± SEM. The value of ADF GFP fluorescence in WT animals under a stress paradigm and that of mutants is normalized to the value of WT animals under optimal conditions. *p*<0.05, **p**<0.01, ***p***<0.001.

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Figure 2. TIR-1 signaling selectively regulates *tph-1::gfp* response to pathogenic bacterial food. A. PA14-induced ADF *tph-1::gfp* upregulation was impaired in *tir-1(tm3036lf)* and *tir-1(yz68gf)* mutants. B. *tir-1(tm3036lf)* and *tir-1(yz68gf)* mutations did not block *tph-1::gfp* upregulation during dauer formation. C. *TIR-1* is not required for *tph-1::gfp* upregulation caused by changes in ciliary morphology. Mutations in *daf-6/Patched* alter the morphology of dendritic cilia of all chemosensory neurons including ADF and cause ADF *tph-1::gfp* upregulation on its own as well as in *tir-1(lf)* background. D. *tir-1(yz68gf)* and *daf-6* mutations confer additive upregulation of *tph-1::gfp* in the ADF neurons. Data represent the average of three trials each with at least 20 animals per strain per condition ± SEM. The value of ADF GFP fluorescence in WT animals under a stress paradigm and that of mutants is normalized to the value of WT animals under optimal conditions. *p*<0.05, **p**<0.01, ***p***<0.001.

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ADF neurons can detect and discriminate multiple aversive cues, and indicate that TIR-1 is selectively involved in the pathogen signaling transduction pathway.

**daf-19 is a suppressor of tir-1(yz68gf)**

To identify the effectors of TIR-1 signaling, we carried out a suppressor screen for mutants that abrogate tph-1 expression in tir-1(yz68gf) mutants. Using a combination of SNP mapping, non-complementation tests and sequencing the mutant genomes, we identified that two mutations, yz60 and yz70, are allelic to the daf-19 gene, encoding the sole C. elegans ortholog of the RFX transcription factors (Figure 3A). Subsequent experiments with our alleles and the previously existing daf-19(m86)-null mutation revealed that DAF-19 function is required not only for tir-1(yz60gf) to upregulate tph-1::gfp, but also for ADF tph-1::gfp expression under optimal growth conditions (Figure 3B), during dauer formation (data not shown) and in aberrant cilia backgrounds (Figure 3E; Figure S2). The reduced tph-1::gfp in daf-19 mutants was fully rescued by transgenic expression of WT daf-19 genomic sequence (Figure 3B) or daf-19 cDNA driven by the gpa-13 promoter (Figure 3D).

An implicit concern was that daf-19 deficiency alters ADF cell fates. To directly analyze the effect of daf-19 deficiency on tph-1 expression, we inactivated daf-19 by RNAi after 5-HT phenotypes established. Figure 3C shows that 100% of larval stage 1 (L1) worms expressed tph-1::gfp prior to RNAi treatment, 33% of the animals lost ADF GFP after 24 hr RNAi treatment, and by 48 hr, 100% of the animals showed no GFP in the ADF neurons. DAF-19 is required for the expression of cilia IFT components [17]. Although we showed that aberrant cilia induced ADF tph-1::gfp upregulation [5] (Figure S2), complete lacking cilia could inhibit tph-1 expression. To rule out this possibility, we used lipophilic dye Dil staining to examine the dendritic cilia morphology of the chemosensory neurons in worms treated with daf-19 RNAi. We observed that RNAi of daf-19 eliminated ADF tph-1::gfp prior to a detectable change in the cilia (Figure 3C).

We investigated whether the reduced tph-1 expression is a secondary consequence of reduced ocr-2 and tir-1 expression in daf-19 mutants. daf-19(yz69) (Figure 3Hi) and daf-19(m86)-null (not shown) mutants expressed a GFP reporter for ocr-2 (ocr-2::gfp) in ADF and other chemosensory neurons indistinguishable from WT animals. Diminished tph-1::gfp expression also cannot be ascribed to reduced tir-1 expression, as transgenic expressing tir-1(yz68) cDNA by the gpa-13 promoter failed to increase ADF tph-1::gfp in daf-19 mutants (Figure 3G), although the Pgpa-13::tir-1(yz68c) transgene conferred a significantly increase of ADF tph-1::gfp as tested in ocr-2 mutant background but not in ocr-2;daf-7(gf) double mutants (Figure 4A). Thus, tph-1 upregulation induced by PA14 and tir-1(yz60gf) requires both DAF-19 and ATF-7.

**daf-19 deficiency suppresses the enhanced innate immunity of tir-1(yz68gf) mutants**

daf-19 is expressed broadly in neurons but also in the intestine (Figure S3) [31,32]. We used epistasis analysis to investigate the role of DAF-19 in TIR-1-mediated innate immunity. When incubated on a lawn of PA14, tir-1(yz60gf) mutants exhibited enhanced resistance to killing by PA14, in contrast to the heightened susceptibility of tir-1(lf) mutants (Figure 5A). daf-19(m86) and daf-19(yz70) mutants exhibited heightened susceptibility to PA14 compared to WT (Figure 5B). Similarly, RNAi of daf-19 enhanced the susceptibility to PA14 compared to mock RNAi (Figure 5C). Transgenic expression of daf-19 cDNA in the intestine partially rescued PA14 resistance in daf-19 mutants (Figure 5D). daf-19(m86)-null mutation suppressed enhanced immune resistance of the tir-1(yz60gf) mutants, showing that the enhanced pathogen resistance of tir-1(yz60gf) mutants also requires DAF-19 function (Figure 5E). However, the daf-19(m86);tir-1(yz68gf) double mutants did not display heightened pathogen susceptibility as seen in daf-19(m86) single mutants, suggesting additional transcriptional regulators involved in the immunity induced by TIR-1 activation.

We tested the functional relationship between DAF-19 and ATF-7 in the innate immunity. Both atf-7(gf) and atf-7(lf) mutants exhibited heightened susceptibility to killing by PA14, although the immunodeficiency phenotype of atf-7(gf) mutants is weaker [30]. If ATF-7 and DAF-19 function in parallel, we could expect a stronger immunodeficiency phenotype in a double mutant of atf-7(gf) and daf-19 relative to the single mutants. Contrary to the prediction, atf-7(gf)/daf-19 double mutants displayed a survival rate on PA14 comparable to the atf-7(lf) single mutant (Figure 5E). This result is more consistent with the model in which DAF-19 regulates ATF-7 targets in the immune system.

**TIR-1- and ATF-7-regulated immune reporter genes are also regulated by DAF-19**

The exact detoxification mechanisms of the C. elegans immunity are not known. To validate the role of DAF-19 in the innate immunity, C. elegans, an ortholog of the mammalian ATF2/ATF7/CREB5 family of transcription factors [29], has been implicated as a transcriptional repressor of antimicrobial genes in the intestine; activation of FMK-1 p38 MAPK de-represses ATF-7, hence upregulating the antimicrobial genes [30]. We therefore analyzed tph-1::gfp expression in atf-7(gf) and atf-7(lf) mutants. atf-7(gf) mutation presumably constitutively represses TIR-1 signaling targets. atf-7(gf) did not downregulate ADF tph-1::gfp under optimal growth conditions (Figure 4A), as seen in mutants with reduced tir-1, TIR-1 downstream mapk-1 MAPKKK and skk-1 MAPKK (Figure 1C, Figure S1) [9]. Like tir-1(lf) mutants, atf-7(gf), as well as daf-19 mutants, failed to upregulate ADF tph-1::gfp following 6 hr feeding on PA14 (Figure 2A, Figure 4B). By contrast, both atf-7(gf) and atf-7(lf) mutants upregulated ADF tph-1::gfp during dauer formation (Figure 4C). These data suggest that ATF-7 confers the specificity to upregulate tph-1 in response to pathogenic bacteria.

We tested further whether constitutive repression function of atf-7(gf) could suppress tph-1::gfp upregulation by tir-1(yz60gf). We crossed the Pga-13::tir-1(yz68c) transgene into atf-7(gf) mutants. The Pga-13::tir-1(yz68c) transgene conferred a significantly increase of ADF tph-1::gfp as tested in ocr-2 mutant background but not in ocr-2;daf-7(gf) double mutants (Figure 4A). Thus, tph-1 upregulation induced by PA14 and tir-1(yz60gf) requires both DAF-19 and ATF-7.
immunity, we made use of the fact that bacterial infections cause the intestine to induce the transcription of a battery of secretory proteins that are thought to produce antimicrobial effects [33,34]. Transcriptional regulation of those candidate antimicrobial genes has been used as an assay for genetic delineation of C. elegans immune pathways [35,36]. For example, the atf-7(gf) allele, as well as a number of tir-1(If) alleles, were identified based on the diminished intestinal expression levels of a GFP reporter for the ShK-like toxin peptide gene T24B8.5 (T24B8.5::gfp), and af-7(If) alleles were identified as suppressors of the diminished T24B8.5::gfp of af-7(gf) [30]. We therefore analyzed the same integrated T24B8.5::gfp reporter in tir-1 and daf-19 mutants. On a
lawn of standard bacterial food *E. coli* OP50, *T24B8.5::gfp* intensity in the intestine of *tir-1(yz68)* was substantially enhanced relative to WT animals (Figure 6B, 6I), further validating the constitutive activity of *tir-1(yz68gf)*. By contrast, intestinal *T24B8.5::gfp* was markedly reduced in *daf-19* mutants as in *atf-7(gf)* and *tir-1(lf)* mutants (Figure 6C, 6E, 6I). Thus, DAF-19 deficiency results in downregulation of an immune gene marker regulated by TIR-1 and ATF-7.

Similar to the effect of the *daf-19* mutation on *tir-1(yz68gf)* immunity (Figure 5E), the *daf-19(n868)* and *daf-19(yz69)* mutations diminished the increased *T24B8.5::gfp* expression in *tir-1(yz68gf)* mutants, although the GFP level in the *daf-19; tir-1(yz68gf)* double mutants was higher compared to the *daf-19* single mutants (Figure 6D, 6I). We also detected an increase in *T24B8.5::gfp* in the *atf-7(gf)* intestine relative to WT animals; two tested *daf-19* alleles both reversed the increased *T24B8.5::gfp* in *atf-7(gf)* mutants (Figure 6F, 6I, 6D, 6I).

To confirm the role for DAF-19 in the regulation of candidate antimicrobial genes, we used quantitative real-time RT-PCR (qPCR) to measure the expression of endogenous *T24B8.5* as well as three other ATF-7-regulated candidate antimicrobial genes. Every tested antimicrobial gene was reduced in two tested *daf-19* alleles compared to WT animals (Figure 7B). In contrast, the message levels of these antimicrobial genes were increased in *tir-1(yz68gf)* mutants (Figure 7B). *daf-19* mutation diminished the increases of those antimicrobial genes in *tir-1(yz68gf)* mutants (Figure 7B), similar to that seen with *T24B8.5::gfp* in the intestine. As the controls, we analyzed *atf-7(gf)* and *atf-7(lf)* mutants. We observed that the message levels of *K08D8.5* and *F35E12.5* were significantly reduced, and *C17H12.8* elevated in the *atf-7(lf)* mutant (Figure 7A), as previously reported [30]. Consistent with our observation of enhanced *T24B8.5::gfp* in *atf-7(lf)* intestine, we found *T24B8.5* message level increased in *atf-7(lf)* mutants (Figure 7A, 7B). We did observe dramatically reduced message levels of all tested genes in *atf-7(gf)* and *tir-1(lf)* mutants (Figure 7A, 7B), as previously reported [30,36].

Our data thus far indicated that DAF-19 is required for TIR-1 signaling to upregulate *tph-1* in the ADF neurons and candidate antimicrobial genes in the intestine. We wished to determine whether DAF-19 regulates every TIR-1 target, or it selectively mediates TIR-1 regulation of pathogen inducible genes. It has been well established that, TIR-1 specifies asymmetrical expression of the olfactory receptor STR-2 in one of two AWC olfactory neurons during the development [23]. We observed that neither AWC neuron expressed *str-2::gfp* in the *tir-1(yz68gf)* mutant, as
Figure 5. DAF-19 regulation of TIR-1-mediated innate immunity. A. Survival rates of WT and loss- and gain-of-function mutants of tir-1 fed PA14 under standard slow killing pathogenesis assay conditions. B. daf-19 mutants displayed elevated susceptibility to killing by PA14. C. Survival rates of vector control and daf-19 RNAi-treated worms fed PA14. D. Transgenic expression of daf-19 in the intestine by the ges-1 promoter partially rescued the elevated PA14 susceptibility of daf-19 mutants. E. daf-19 deficiency suppressed the enhanced PA14 resistance of tir-1(yz68gf) mutants. F. daf-19;atf-7(lf) mutants displayed elevated susceptibility to killing by PA14 similarly to atf-7(lf) mutants. The fraction of live animals was determined at each indicated time points. The experiments were performed in triplicates. Results are representative of at least two independent experiments.

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seen in mutants with excessive TIR-1 activity [23]. However, daf-19 mutants did not exhibit the AWC phenotype seen in tir-1(lf) mutants (Figure S4). Thus, DAF-19 is critical for TIR-1 signaling to induce transcriptional responses to pathogenic bacteria in the 5-HT neurons and the intestine, but is not required for TIR-1 to regulate neural development.

Figure 6. Reduced immune gene marker in the intestine of daf-19 mutants. A–H. Representative photomicrographs showing two-day-old adult WT and indicated mutants expressing a GFP reporter for the ShK-like toxin peptide gene T24B8.5 (T24B8.5::gfp) in the intestine, all the animals shown with the anterior to the left. I. Quantification of T24B8.5::gfp intensity in the intestine of two-day-old adults. Fluorescence was quantified by measuring pixel intensity of three areas along the body (as depicted in B). T24B8.5::gfp intensity was reduced in atf-7(gf) and two daf-19 alleles, but increased in tir-1(yz68gf) and atf-7(lf) mutants. daf-19 deficiency diminished the increased T24B8.5::gfp in tir-1(yz68gf) as well as atf-7(lf) mutants. Data represent the average of three trials each with 20 animals per strain ± SEM, and the value of GFP fluorescence in mutants is normalized to that of WT animals analyzed on the same day. Statistics between WT and mutants is marked on the top of each bar, and that between two indicated groups is marked on the top of lines, * p<0.05, ** p<0.01, *** p<0.001.

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PA14 triggers TIR-1 signaling via tissue-specific mechanisms

The finding of the shared transcriptional effectors of TIR-1 signaling in the ADF neurons and intestine raised an intriguing question as to whether pathogenic bacterial signals trigger the neurons and immune cells in the same manner in C. elegans.
Although little is known about how the worm senses the presence of pathogens and relays the signals to TIR-1, the protein kinase D DKF-2 is thought to promote transcriptional responses to PA14 by activating the TIR-1 signaling pathway [37]. However, we found that the \( \text{dkf-2(pr3)} \)-null mutation did not suppress \( \text{tph-1::gfp} \) upregulation by \( \text{tir-1(yz68gf)} \) or PA14 (Figure 8A).

Previously, we showed that a gain-of-function mutation in the calcium/calmodulin-dependent protein kinase II (CaMKII) UNC-43 upregulates ADF \( \text{tph-1::gfp} \) [4]. We hypothesized that UNC-43 could be a component of the TIR-1 signaling pathway in the ADF neurons, similar to its involvement in TIR-1-mediated AWC development [23]. To test this hypothesis, we analyzed \( \text{tph-1::gfp} \) in \( \text{unc-43(lf)};\text{tir-1(yz68)} \) double mutants. We observed that \( \text{unc-43(lf)} \) abrogated ADF \( \text{tph-1::gfp} \) upregulation by \( \text{tir-1(yz68gf)} \) or PA14 (Figure 8A).

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In cultured mammalian neurons, CaMKII activation alters the subcellular localization of signaling components to initiate cellular responses [38]. UNC-43 has been shown enriched in postsynaptic sites of a number of neuronal types and was co-immunoprecipitated with TIR-1 when co-expressed in cultured mammalian cells [23,39]. If UNC-43 regulates TIR-1 subcellular distribution, then it is possible that the subcellular distribution of the TIR-1(yz68gf) protein also depends on UNC-43. We tested this idea by comparing the subcellular distribution of GFP-tagged TIR-1 and TIR-1(yz68gf) proteins expressed in chemo sensory neurons by the \( \text{gpa-13} \) promoter in WT and \( \text{unc-43(lf)} \) backgrounds. In WT...
animals, TIR-1(WT)::GFP was observed in punctate structures in the axons around nerve ring, with more diffused fluorescence seen in the cell bodies (Figure 8C). By contrast, TIR-1(yz68gf)::GFP displayed increased punctate structures in the axons as well as in the cell bodies (Figure 8D). Importantly, TIR-1(WT)::GFP and TIR-1(yz68gf)::GFP punctate structures were reduced in unc-43(lf) mutants (Figure 8C, 8D). Based on these observations, we speculated that the C426Y substitution of TIR-1(yz68gf) alters the protein conformation, thereby enhancing its interaction with UNC-43 in particular cellular compartments where TIR-1 interacts with MAPK signaling components, but that TIR-1(yz68gf) cannot efficiently interact with its downstream components in the absence of UNC-43. Collectively, these results suggest that PA14 triggers distinct mechanisms to activate TIR-1 signaling to induce DAF-19/ATF-7 targets in the ADF neurons and the intestine. Our data indicate that UNC-43 is required for TIR-1 signaling in ADF, and DKF-2 is not.

Discussion

Here, we used an unbiased genetic approach to identify molecular mechanisms underlying transcriptional responses to pathogenic bacterial infection in *C. elegans*. We identified DAF-19, the ortholog of RFX transcription factors. We showed that DAF-19 acts downstream in the TIR-1 pathway to induce *tph-1* in the ADF neurons and antimicrobial genes in the intestine. Our genetic analyses suggest that bacterial infection may trigger the TIR-1 signaling cascade via cell-specific mechanisms, but common TIR-1
downstream transcription factors regulate neuronal and immune responses to infection. Our data demonstrate that RFX gene function, which is required for human adaptive immunity, regulates 5-HT biosynthesis and innate immunity in C. elegans (Figure 9).

DAF-19 is an ancient transcriptional regulator of innate immunity

Parallel to the C. elegans NSY-1/MAPKK-SEK-1/MAPKK-PMK-1/p38 MAPK pathway, the ASK-1/MAPKKK to p38 MAPK pathway regulates mouse innate immunity [40]. While members of the NF-κB family of transcription factors are the major effectors in mammalian innate immune responses, the ASK-1-p38 innate immune pathway is independent of NF-κB [40]. Since C. elegans lacks NF-κB, this MAPK pathway was proposed to act via effectors evolutionarily more ancient than NF-κB in the host defense systems [40]. In this study, we identify that DAF-19 RFX acts as a transcription factor downstream of this core innate immune pathway in C. elegans. Our genetic analyses suggest that DAF-19 interacts with ATF-7, a member of the ATF/CREB family of transcription factors, to regulate the 5-HT biosynthesis pathway in response to pathogenic bacterial infection.

Figure 9. DAF-19 regulation of 5-HT biosynthesis and innate immunity. A. DAF-19 is engaged in multiple independent signaling pathways that regulate tph-1 expression in ADF neurons. Based on the genetic interaction of DAF-19 and ATF-7 in the TIR-1 signaling pathway, we speculate that DAF-19 interacts with different cofactors at different cis-elements of the tph-1 promoter in response to different environmental cues. DAF-19 may simultaneously interact with multiple cofactors to confer additive upregulation of tph-1 in response to multiple aversive cues. B. DAF-19 regulates TIR-1 targets in response to pathogenic bacterial infection, but is not required for TIR-1 to regulate AWC neuron development. However, pathogenic bacterial infection may trigger the TIR-1-DAF-19/ATF-7 pathway via cell-specific mechanisms.

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superfamily of transcription factors, to upregulate \( tpk-1 \) in the ADF neurons and antimicrobial genes in the intestine in response to the pathogenic \( P. aeruginosa \) strain PA14, reminiscent of the RFX-CREB partnership for human MHC II gene expression.

Several lines of evidence point to RFX factors as an ancient mechanism for enhancing survival under aversive conditions, one predicting the divergence of stress responses and immunity. First, RFX is a regulator of cell cycle in a nutrient sensing pathway in \( Schizosaccharomyces pombe \) [41] and an effecter of the DNA damage and replication checkpoint pathway essential for \( Saccharomyces cerevisiae \) survival under replicative stress [42]. Second, biochemical experiments have identified a role for RFX factors in RAS signaling-regulated transcription in mammalian epithelial cells [43]. Third, in addition to mediate transcriptional responses to signaling-regulated transcription in mammalian epithelial cells experiments have identified a role for RFX factors in RAS survival under replicative stress [42]. Second, biochemical CREB partnership for human MHC II gene expression.

The pathogenic microarray analysis [36] with the database of several independent datasets identified hundreds of candidate genes. Thus, DAF-19 is likely to function in the predicted dimerization (DIM) domain, suggestive for a possible role in protein-protein interaction in DAF-19 function.

What could be the mechanism by which DAF-19 regulates \( tpk-1 \) and those antimicrobial genes? Earlier studies of RFX regulation of human MHC II genes provided evidence that RFX factors regulate gene expression indirectly through recruiting additional transcriptional regulators to the promoters [13]. In a recent study, RFX factors were shown to protect promoters against epigenetic silencing by DNA methylation through recruiting chromatin-remodeling factors [47]. Thus, one plausible mechanism could be that DAF-19 and ATF-7 bind to a promoter element shared in common among the pathogen-inducible genes, and infection induces the TIR-1 signaling cascade leading to phosphorylation of \( ATF-7 \) [30] and transcriptional activation of the targets. However, genomewide X-box motif search identified hundreds of candidate genes, only a few antimicrobial genes are among them [48] and no X-box can be recognized in the promoter of the \( tph-1 \), \( T24B8.5 \). C17H12.8, K00D0.5 or F35E12.5 genes. Thus, DAF-19 is likely to modulate non-consensus X-boxes on the promoters via co-regulators. Alternatively, DAF-19 may modulate chromatin structure, facilitating the binding of other transcriptional regulators to the promoters. We favor the binding-via-cofactor model because our data thus far suggest that DAF-19 is involved in multiple environment-dependent regulations of \( tpk-1 \) expression, whereas \( ATF-7 \) selectively regulates the response to PA14. We speculate that DAF-19 interacts with distinct cofactors that are regulated by distinct environmental cues (Figure 9A). Further experiments are required to determine whether DAF-19 directly interacts with \( ATF-7 \). In addition, the possibility that DAF-19 and ATF-7 regulate yet unidentified transcription factor(s), which in turn regulate pathogen-inducible genes, cannot be excluded. It is perhaps interesting to note that our two \( daf-19 \) alleles both are located in the predicted dimerization (DIM) domain, suggestive for the importance of protein-protein interaction in DAF-19 function.

While the exact mechanism of DAF-19 on the target gene promoters remains to be elucidated, our data showed that DAF-19 and \( ATF-7 \) regulate common immune gene markers. Remarkably, by comparing the list of TIR-1 signaling targets identified by microarray analysis [36] with the database of several independent expression profiling of \( daf-19 \) mutants [49,50,51], we found that 102 out of 215 TIR-1 gene targets are among those differentially expressed in \( daf-19 \) mutants; one of them is \( C17H12.8 \), which was confirmed by our qPCR. We showed that \( daf-19 \) mutations can suppress increased expression of immune gene markers in \( tir-1(yz56gf) \) mutants. However, \( daf-19\text{-null} \text{, } tir-1(yz56gf) \) double mutants did not display reduced expression of those immune genes as seen in the \( daf-19 \) single mutants, judging by \( T24B8.5::gfp \) in living worms and qPCR. Consistent with the gene expression analyses, the \( daf-19\text{-null} \) mutation blocked the enhanced resistance to killing by PA14 in \( tir-1(yz56gf) \) mutant but the \( daf-19\text{-null} \) mutant did not exhibit heightened susceptibility as \( daf-19 \) single mutants did. These observations may be consistent with the model in which DAF-19 and ATF-7 interact with additional transcription factor(s) that also contribute to the regulation of those antimicrobial genes.

Multiple mechanisms relay pathogen signals to the TIR-1-DAF-19 signaling pathway

There is converging evidence from neurobiology and immunology suggesting that the brain immune privilege is not absolute [52]. Internal and external pathogenic products can infiltrate into the CNS and there is extensive bi-directional communications between the CNS and immune systems [53,54]. The finding of a large number of classical immune proteins in the CNS has led to the idea that common molecular mechanisms may be involved in neuronal and immune responses to pathogenic signals [1,2,53]. Our identification of the TIR-1-DAF-19/ATF-7 pathway in regulating \( tph-1 \) and antimicrobial genes supports this idea. Moreover, our genetic analysis suggests that activation mechanisms of this core signaling pathway in the neurons and intestine during infection differ. A prior work showed that DKE-2 is required for TIR-1 signaling cascade to induce intestinal immunity [37]. We found that \( dsf-2\text{-null} \) did not prevent PA14-induced \( tph-1::gfp \) upregulation. Instead, we identified a requirement of UNC-43 CaMKII for \( tph-1::gfp \) upregulation induced by PA14 and \( tir-1(yz56gf) \). Analysis of TIR-1(\( WT\text{-::GFP} \)) and TIR-1(\( yz68gf \)) Gfp suggests that UNC-43 regulates subcellular distribution of TIR-1. While several mammalian TIR domain adaptor proteins have been shown to translocate to the plasma membrane following Toll-like receptor activation, SARM, the ortholog of TIR-1, is activated in the brain by neural toxicity via a yet unidentified mechanism [55,56]. Our results raise the possibility that the cue associated with pathogenic bacterial infection triggers Ca\textsuperscript{2+} signaling to activate TIR-1 in the ADF neurons. It may be sensible that neuronal and immune cells detect distinct molecular cues associated with infection thereby coordinating neuronal and physiological responses.

Materials and Methods

Strains

\( C. elegans \) strains were maintained at 20°C on NGM agar plates seeded with a lawn of \( Escherichia coli \) OP50 as a food source. WT animals were Bristol strain N2. The Hawaiian isolate CB4856 was used in genetic mapping of the \( daf-19 \) and \( tir-1 \) mutations. The following existing mutant strains were used in this study: \( alf-7(qd22gg) \), \( alf-7(qd22 qd130) \), \( alf-6(e1757) \), \( alf-19(m66) \), \( daf-2(\text{ps}) \), \( dfy-1(yz66) \), \( eri-1(mg189) \), \( lin-15(b74) \), \( ocr-2(yz3) \), \( sek-1(\text{ls4}) \), \( tir-1(yz105) \), \( unc-43(e408) \). Transgenic strains used in this study were: \( agl219 \text{, } I\text{s}[\text{T24B8.5::gfp} \text{, } ttx-3::gfp] \text{[30]} \), \( CX3505 \text{, } k\text{y1468(}\text{A}2\text{::}\text{gfp} \text{, } lin-44(yz52)] \text{[57]} \), \( GR1333 \text{, } y\text{z171(tph-1::gfp} \text{, } Rol-6(d) \text{[21]} \), \( I\text{[cat-1::gfp]} \text{[58]} \), \( J\text{1222 \text{, } Es2942[ocar-2::gfp} \text{, } Rol-6(d) \text{[4]} \), \( JY449 \text{, } Es1002(\text{e2::gfp} \text{, } Rol-6[d]) \text{[39]} \).
Identification of tir-1(yz68), daf-19(yz69), and daf-19(yz70) mutations

yz68 is a dominant mutation isolated from a genetic screen for mutants with enhanced GFP expression in ADF chemosensory neurons after ethyl methane sulfonate (EMS) mutagenesis of ocr-2(yz5) mutant carrying an integrated tph-1::gfp transgene as described previously [5]. Genetic mapping using single-nucleotide polymorphisms (SNP) of C. briggsae localized yz68 to a contig of 1.43 map on the chromosome III. To identify the mutant gene, 174 genes located in the contig were individually inactivated in yz68 mutants by RNAi, and the clone F13B10.1 expressing double stranded(ds)-RNA of tir-1 suppressed the tph-1::gfp upregulation of the yz68 mutant. Sequencing yz68 genomic DNA revealed a G to A transition resulting in a cysteine426 to tyrosine substitution in the fourth ARM motif of the HEAT/Arm repeat region of TIR-1. The amino acid altered in yz68 is in reference of the tir-1a isoform. yz69 and yz70 mutants are recessive mutations isolated from an EMS mutagenesis screen for mutants with dramatically reduced/absence of ADF tph-1::gfp. Analysis of the amphid morphology with fluorescence dye Dil revealed that ciliated neurons in yz69 and yz70 mutants were dye filling defective. CB4856 SNP mapping localized yz69 to a contig of 1.28 map units between the polymorphisms in the clones C18D1 and F44F4 on the chromosome II. Non-complementation assays with dye-filling mutants within the region indicated that both yz69 and yz70 were allelic to daf-19. Sequencing the daf-19 gene of the mutants revealed in yz69 a G to A transition predicting an opal mutation in the 9-UTR of the daf-19 gene of the mutants isolated from an EMS mutagenesis screen for mutants with dramatically reduced/absence of ADF tph-1::gfp. Analysis of the amphid morphology with fluorescence dye Dil revealed that ciliated neurons in yz69 and yz70 mutants were dye filling defective. CB4856 SNP mapping localized yz69 to a contig of 1.28 map units between the polymorphisms in the clones C18D1 and F44F4 on the chromosome II. Non-complementation assays with dye-filling mutants within the region indicated that both yz69 and yz70 were allelic to daf-19. Sequencing the daf-19 gene of the mutants revealed in yz69 a G to A transition predicting a cysteine to tyrosine substitution at the conserved dimerization domain, and in yz70 a G to A transition predicting an opal mutation in the dimerization domain. The amino acid changes depicted in Figure 3A are in reference of daf-19a isoform.

Generation of transgenic animals

All constructs were generated by PCR. daf-19(g) was a ~14.8 kb genomic fragment amplified from the WT genome encompassing 2.9 kb 3’-upstream promoter sequence, exons/introns, and 574 bp 3’-UTR of the daf-19 gene. To express tir-1 and daf-19 in specific neurons, we fused tir-1 and daf-19 cDNA sequences individually to the gpa-13 promoter, which is expressed in three pairs of amphidial ciliated sensory neurons ADF, ASH and AWC. The gpa-13 promoter is expressed additionally in PHA and PHB phasmid neurons located in the tail [60]. tir-1 promoter is expressed additionally in PHA and PHB phasmid neurons located in the tail [60]. tir-1/WT and tir-1(yz68) cDNA were amplified from cDNA mixture prepared from total RNA of WT and tir-1(yz68) animals, respectively, using primers corresponding to the tir-1a isoform. The cDNA of the daf-19c isoform was amplified from the plasmid PS0243 (kindly provided by P. Swoboda). The 2.6 kb gpa-13 promoter sequence amplified from the plasmid PS0243 was fused to the sequences in the order of a cDNA, GFP and unc-54 3’UTR or a cDNA, mCherry and unc-54 3’UTR. To express DAF-19 in the intestine, the cDNA sequence of daf-19a isoform was fused to the 2.9 kb ges-1 promoter. For each construct, products from three independent PCR reactions were pooled to reduce potential PCR errors. The pooled PCR products were purified (Qiagen) and microinjected at the concentration of 50 ng/μl into worms. The plasmid containing either a dominant rol-6 gene (rol-6(d)], ebf-2::gfp or unc-122::RFP was co-injected as a transgenic marker.

RNAi

All RNA interference (RNAi) experiments were done in the background of eri-1(dauer-15B), which enhances sensitivity to RNA interference in neurons [61]. RNAi assays were carried out by feeding worms E. coli HT1115 expressing dsRNA of a target gene or the control empty L4440 vector (Addgene RNAi library, University of Cambridge, England). RNAi clones were individually cultured overnight in Luria broth containing 100 μg/ml ampicillin, 500 μl of the bacterial culture were seeded onto agar plates containing NGM supplemented with 1 mM IPTG and 25 μg/ml carbenicillin to induce dsRNA expression, and incubated overnight at room temperature. For RNAi of tir-1 and yz-1, about 60 eggs were placed onto each plate and allowed to hatch, grow to adults and lay eggs. F1 progeny were transferred to a fresh plate, and tph-1::gfp in the ADF neurons of L4-stage worms of F2 generation quantified. For developmental RNAi of daf-19, synchronized L1 worms were transferred to the plates (day 0), the worms were transferred to freshly prepared plates every day, and the expression of tph-1::gfp in ADF neurons or DiI staining of cilia morphology were analyzed on indicated days. DiI staining was done as previously described [5]. For each RNAi experiment, three independent trials each with three replicates were performed, and data from one representative trial presented.

Indirect immunofluorescence microscopy

Whole-mount staining of worms with anti-5-HT antibody was performed as described previously [21]. The staining patterns were visualized via Alexa Fluor 594 or 488 conjugated secondary antibodies (Molecular Probe) under an AxioImager Z1 microscope equipped with proper filters, and images were captured using AxioCam MR digital camera (Zeiss, Northwood, NY). To quantify the intensity of 5-HT immunoreactivity, images of ADF neurons in individual worms were captured under a 40× objective lens at a fixed exposure time of 3 ms with 100% UV exposure level. For each image, fluorescence intensity of a circular 10 pixels area within the ADF cell body was quantified using the ImageJ software (National Institute of Health, Bethesda, Maryland). To exclude the background, fluorescence intensity over a circular 10 pixels area posterior to the ADF cell body in the same image was quantified, and the value of the background was subtracted from the value of the ADF area.

Assessment of GFP reporter levels and statistics

The expression of a chromosomally integrated tph-1::gfp reporter in ADF neurons in living WT or mutant worms was evaluated by measuring GFP fluorescence intensity. Images of ADF neurons in individual animals were captured at a fixed exposure time. The external contour of each ADF neuron was delineated, and fluorescence intensity within the entire neuron was quantified. For developmental RNAi of tph-1::gfp, the worms were transferred to freshly prepared plates every day, and the expression of tph-1::gfp in ADF neurons or DiI staining of cilia morphology were analyzed on indicated days. DiI staining was done as previously described [5]. For each RNAi experiment, three independent trials each with three replicates were performed, and data from one representative trial presented.

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lay eggs for 2–3 hr in a 25°C incubator, the adults were then removed from the plates, and dauer developed from hatched eggs on the plates were analyzed 72 hr later. For each strain, the value of dauers was compared to that of L4 grown on the plates without pheromone and assayed on the same day.

The expression of T24BR.5::gfp in the intestine was analyzed in two-day-old adult worms carrying the integrated agl219/T24BR.5::gfp; ttx-3::gfp transgene cultured on OP50 at 20°C. Images of the intestine were captured under a 10× objective lens at a fixed exposure time, and fluorescence intensity was quantified by measuring pixel intensity of three areas along the body of each animal as depicted in Figure 6B.

Data represent the average of at least three trials unless specified otherwise. For each trial, 15–20 animals per strain per condition/treatment were analyzed and compared to the controls assayed on the same day. WT animals under the same conditions and treatments were analyzed and compared for every experiment. Unpaired Student’s t-test was used for comparisons between a mutant and the same day. WT animals under the same conditions and treatments were analyzed and compared to the controls assayed on the same day. WT animals under the same conditions and treatments were analyzed and compared for every experiment. Unpaired Student’s t-test was used for comparisons between a mutant and WT and between two mutants or two treatments.

P. aeruginosa PA14 pathogenesis assays

The standard PA14 slow killing assays were performed as previously described [62]. Briefly, PA14 was cultured in King’s broth overnight and the culture was seeded at the center of 3.5 cm diameter assay plates and incubated at 37°C for 20 hr followed by 20 hr incubation at room temperature. 40–50 L4 worms per strain were transferred onto each assay plate, incubated at 25°C and scored for dead or live every 8 hr. Worms were scored as dead as no response was detected after prodding with a platinum wire. daf-19 mutants tend to claw off the plate. So for each assay, more than 300 L4 daf-19 mutants were transferred to each plate, and live and dead animals on the agar surface were scored at indicated time points; dead animals on the wall of the plate not counted. For each strain, three replicates were analyzed for each experiment. To test the effect of RNAi of daf-19, 4 to 6 gravid animals were grown on the RNAi plates as described above seed with either E.coli HT1115 harboring empty control plasmid L4440 or plasmids expressing RNAi against daf-19. G1 progeny at the L4 stage were used to test the susceptibility to killing by PA14.

Quantitative real-time PCR (qPCR) analysis

Total RNA from 100 one-day-old adults of WT and indicated mutant strains was extracted using Trizol (Invitrogen), reserve transcribed to cDNA using the SuperScript III system (Invitrogen), and the cDNA was used for qPCR analyses using the StepOnePlus machine (Applied Biosystems) and SYBR Green detection system (Applied Biosystems) in triplicated reactions. The primers for qPCR were designed using Primer Premier 5.0 (Premier Biosoft) (Figure S5). Values were normalized against the reference gene act-1 [37]. gpd-2 was analyzed as a second control showing no change relative to act-1. Fold change was calculated using the delta Ct method [61]. qPCR analysis of L4 worms of those strains showed comparable results; data of the adults are presented.

Supporting Information

Figure S1 Regulation of tph-1::gfp expression in the ADF neurons by TIR-1 downstream ny-1 MAPKKK and sek-1 MAPKK RNAi of ny-1 suppressed tph-1::gfp upregulation by tir-1(yz68gf). RNAi of ny-1 and loss-of-function mutation of sek-1 did not confer tph-1::gfp reduction, indicating that the TIR-1 signaling pathway is designated primarily to upregulate tph-1 expression in response to pathogen infection. Data represent the average of three trials each with at least 15 animals per strain ± SEM. The value of GFP fluorescence in mutants was normalized to that of WT animals, and the value of RNAi-treated animals is normalized to that of mock RNAi with an empty vector. Statistics between WT and individual mutants and RNAi-treated animals is marked on the top of each bar, and that between two indicated groups is marked on the top of lines, *** p<0.001, unpaired student’s t test. (TIF)

Figure S2 tph-1::gfp in the ADF neurons of the IFT mutant dyf-1. dyf-1 mutant animals were treated with a mock RNAi with an empty vector or a vector expressing RNAi against tir-1 or daf-19. RNAi of tir-1 did not reduce tph-1::gfp in the ADF neurons, compared to mock RNAi. By contrast, RNAi of daf-19 abolished ADF tph-1::gfp. Data represent the average of three trials each with at least 13 animals per strain ± SEM. The value of RNAi of tir-1 and daf-19 is normalized to that of dyf-1 mutants treated with the empty vector. *** p<0.001, unpaired student’s t test. (TIF)

Figure S3 daf-19 is expressed in the intestine. Top, an image of L4 worm expressing GFP driven by a genomic fragment encompassing 2.9 kb 5’-upstream sequence to exon 8 of daf-19. Bottom, a bright field image showing the position of the same worm. Anterior is to the left, and arrows point to the nuclei of the intestinal cells. (TIF)

Figure S4 DAF-19 is not required for TIR-1 regulation of AWC cell fates, while atf-7 is. A–D. Photomicrographs showing L4 animals expressing an integrated GFP reporter for the olfactory receptor str-2 (str-2::gfp); daf-19 and WT animals expressed str-2::gfp stochastically in one of two AWC neurons. Neither AWC expressed str-2::gfp in tir-1(zc60gf) mutants. E. Quantification of str-2::gfp expression in AWC neurons in daf-19, atf-7 and tir-1 signaling mutants. atf-7(gf) mutants exhibited str-2::gfp expression pattern as seen in tir-1 mutants, but atf-7(gf) showed mixed str-2::gfp patterns of tir-1(gf) and tir-1(zc60gf). daf-19 did not display the str-2::gfp phenotype of tir-1(gf), nor suppressed the str-2::gfp phenotype of tir-1(zc60gf). Data represent the percentage of the animals of a strain showing each AWC phenotype. n, number of animals analyzed. (TIF)

Figure S5 Primer sequences used for qPCR analysis of candidate antimicrobial genes. (TIF)

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Author Contributions

Conceived and designed the experiments: YX MM SC JYS. Performed the experiments: YX MM SC LX JYS. Analyzed the data: YX MM SC LX JYS. Wrote the paper: YX MM SC JYS.

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