Stress-Induced Neural Plasticity Mediated by Glial GPCR REMO-1 Promotes C. elegans Adaptive Behavior

Highlights

- The glial GPCR REMO-1 regulates stress-induced neural plasticity in C. elegans
- REMO-1 functions cell-autonomously to regulate glial and neuronal remodeling
- REMO-1-dependent remodeling primes animals to respond to favorable conditions

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In Brief

Lee et al. provide molecular insight into how stress-induced sensory organ remodeling affects behavior in C. elegans. They identify a glial GPCR, REMO-1, that drives structural remodeling of sensory neurons and associated glia. This alteration of a single neuron-glia pair primes animals to respond efficiently to favorable environmental conditions.
Report

Stress-Induced Neural Plasticity Mediated by Glial GPCR REMO-1 Promotes *C. elegans* Adaptive Behavior

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SUMMARY

Animal nervous systems remodel following stress. Although global stress-dependent changes are well documented, contributions of individual neuron remodeling events to animal behavior modification are challenging to study. In response to environmental insults, *C. elegans* become stress-resistant dauers. Dauer entry induces amphid sensory organ remodeling in which bilateral AMsh glial cells expand and fuse, allowing embedded AWC chemosensory neurons to extend sensory receptive endings. We show that amphid remodeling correlates with accelerated dauer exit upon exposure to favorable conditions and identify a G protein-coupled receptor, REMO-1, driving AMsh glia fusion, AWC neuron remodeling, and dauer exit. REMO-1 is expressed in and localizes to AMsh glia tips, is dispensable for other remodeling events, and promotes stress-induced expression of the remodeling receptor tyrosine kinase VER-1. Our results demonstrate how single-neuron structural changes affect animal behavior, identify key glial roles in stress-induced nervous system plasticity, and demonstrate that remodeling primes animals to respond to favorable conditions.

INTRODUCTION

Nervous systems are robust in connectivity (Goodman and Shatz, 1993), allowing reproducibility in behavior, yet malleable in function, permitting adaptation to fluctuating environments (Albert and Riddle, 1983; Bourne and Harris, 2006; Cuschieri and Bannister, 1975; Engström, 1967; Flock et al., 1999; Golden and Riddle, 1984; Goodman and Shatz, 1993; Murray, 1993; Procko et al., 2011). Such plasticity can be accompanied by structural alterations. For example, in response to dehydration or lactation, astrocytes retract their synapse-associated processes in the paraventricular and supraoptic nuclei to alter synaptic activity (Chapman et al., 1986; Gregory et al., 1980; Hatton et al., 1984; Theodosis et al., 1986; Theodosis and Poulain, 1984, 1989). Similarly, environmental stimuli remodel sensory organs. For example, in the organ of Corti of the mammalian inner ear, glia-like Deiters’ cells form a scaffold that supports outer hair cells (Rio et al., 2002). Upon exposure to high-intensity sound, Deiters’ glia are displaced toward outer hair cells to reduce cochlear sensitivity, protecting against hearing loss (Flock et al., 1999).

Changes in animal behavior have been correlated with large-scale changes in glia and neuron ensembles, and plasticity mechanisms in individual neurons have been extensively explored (Citri and Malenka, 2008). However, linking changes at single synapses or at single sensory receptive endings to animal behavior, a major goal for understanding nervous processing, is challenging in animals in which many neurons contribute to behavior.

The nematode *C. elegans* is an excellent model in which to study sensory organ remodeling and single-cell contributions to animal behavior. Upon exposure to stressful conditions, including high temperature, starvation, and crowding, animals enter an alternative developmental state called dauer (Cassada and Russell, 1975; Golden and Riddle, 1984). The sensory nervous system of dauer animals undergoes striking changes in morphology and gene expression. These changes have been studied primarily in the context of the amphid and inner labial sensilla (Albert and Riddle, 1983; Peckol et al., 2001; Schroeder et al., 2013). Amphids are the primary *C. elegans* sensory organs, detecting chemosensory, osmotic, mechanical, thermal, light, and other stimuli (Bargmann and Mori, 1997; Perkins et al., 1986; Troemel, 1999; Troemel et al., 1995). Each of the bilateral amphids consists of 12 sensory neurons and two glial cells, the AMsh and AMso glial cells, which wrap around sensory neuron receptive endings (NREs) (Ward et al., 1975). Under normal growth conditions, the NRE of each of the bilateral AWC amphid neurons, which houses receptors for volatile odorants (Troemel, 1999; Troemel et al., 1995), is individually ensheathed by processes of adjacent AMsh glial cells (Figures 1A and 1B). Upon dauer entry, AWC NREs expand along with their associated AMsh glia wrappings. In half of dauer animals, bilateral AMsh glia fuse, allowing AWC NREs to expand beyond the midline (Figure 1C) (Albert and Riddle, 1983; Golden and Riddle, 1984; Procko et al., 2011). Thus, AMsh glia delimit AWC NRE growth. Ablation studies demonstrate that AMsh glia remodel independently of AWC NREs (Procko et al., 2011).
AMsh glia is induced by cultivation at high temperature (25°C) (Procko et al., 2011, 2012). The Otd/Otx transcription factor TTX-1; and the zinc finger transcription protein AFF-1; a VEGFR-related tyrosine kinase protein, VER-1; the proteins implicated in remodeling. These include the cell fusion protein AFF-1; a VEGFR-related tyrosine kinase protein, VER-1; the proteins implicated in remodeling.

Previous studies from our lab identified several AMsh glia proteins implicated in remodeling. These include the cell fusion protein AFF-1; a VEGFR-related tyrosine kinase protein, VER-1; the Otd/Otx transcription factor TTX-1; and the zinc finger transcription factor ZTF-16 (Procko et al., 2011, 2012). ver-1 expression in AMsh glia is induced by cultivation at high temperature (25°C) or by dauer entry and requires direct binding of TTX-1 to ver-1 regulatory sequences (Procko et al., 2011).

Here we demonstrate that amphid sensory organ remodeling correlates with early dauer exit following exposure to a favorable environment. We identify a gene, remo-1 (remodeling defective-1), encoding a putative 7-transmembrane G protein-coupled receptor (GPCR) that is required for dauer-induced AMsh glia and AWC neuron remodeling. REMO-1 protein is required for expression of VER-1 in AMsh glia, functions in and is expressed in AMsh glia, and localizes to the tip of amphid sensilla. Importantly, remo-1 loss delays dauer exit behaviors. Our studies link shape changes in a single neuron-glia pair to animal behavior and demonstrate that glia-dependent nervous system remodeling following stress allows animals to efficiently detect restoration of a non-stressful environment.

RESULTS

Gliaal REMO-1 Regulates Stress-Induced VER-1/RTK Expression

To uncover genes required for AMsh glia and AWC neuron remodeling in dauer animals, we sought mutants that fail to induce ver-1 promoter::GFP transgene expression in adults grown at 25°C (Procko et al., 2011). As ver-1 is also upregulated in dauer animals and is implicated in AMsh glia remodeling (Procko et al., 2011), we reasoned that our screen might identify remodeling genes. Indeed, roles for ttx-1 and ztf-16 were discovered in this way (Procko et al., 2011, 2012). From this screen, we identified a mutant, with designated allele number ns250, that fails to turn on ver-1p::GFP in young adults cultivated at 25°C and in dauer larvae induced by starvation (Figures 2A and 2B). Genetic mapping and whole-genome sequencing showed that ns250 is not an allele of previously identified remodeling genes. Rather, ns250 mutants carry a lesion in the gene F47D2.11, which we have renamed remo-1. remo-1 is a putative Srz-type GPCR (Figure 2C) and is conserved in other nematodes (Figure S1) (Robertson and Thomas, 2006). The functions of Srz GPCRs are not well understood. The remo-1(ns250) allele is predicted to cause a glutamic acid-to-lysine change at position 278 (E278K) in the C-terminal intracellular tail of the protein (Figure 2C). This residue is part of a threonine-glutamic acid-isoleucine (TEI) motif conserved in a subfamily of Srz GPCRs (Figure S1) and lies at the terminal membrane-cytosol interface (Figures 2C and S1), suggesting that its mutation could lead to REMO-1 function deficits.

To confirm that the E278K change is the causal mutation in remo-1(ns250) mutants, we recreated the same genomic lesion in otherwise wild-type animals carrying the ver-1p::GFP transgene using CRISPR. Two independent mutants, remo-1(ns820, ns821), were isolated, and both fail to induce ver-1p::GFP expression in young adults cultivated at 25°C and in dauer larvae induced by starvation (Figures 2A and 2B). Furthermore, transgenes containing a genomic fosmid clone encompassing the remo-1 locus, or a smaller genomic fragment, restore ver-1p::GFP expression to remo-1(ns250) mutants (Figure 2D). Finally, we found that expression of remo-1 cDNA using an AMsh glia-specific promoter, derived from the F16F9.3 gene, but not an amphid sensory neuron promoter, derived from the dyf-7 gene, also restores ver-1p::GFP induction to remo-1(ns250) mutants (Figure 2D). Taken together, these results...
Figure 2. Glial REMO-1 Regulates Stress-Induced VER-1/RTK Expression

(A and B) ver-1p::GFP expression in the AMsh glia in indicated genotypes at 25°C in young adult (A) or in dauer larvae induced by starvation (B). WT, wild-type. The number of animals scored is inside bars. Error bars denote SEM of three independent scoring experiments. n.s., p > 0.05; ****p < 0.0001.

(C) Diagram of REMO-1 predicted protein structure and mutation sites generated with Protter software. ns250/820/821: glutamic acid-to-lysine change at position 278 (E278K) indicated by arrow.

(D) Histogram detail as in (A) and (B). All strains are in a remo-1(ns250) mutant background, except for wild-type animals on the left. Rescue experiments conducted with genomic fosmid rescue construct WRM0638cF06, which spans the remo-1 locus, a smaller remo-1 genomic fragment, an AMsh glia promoter (F16F9.3) driving remo-1 cDNA, or amphid sensory neuron promoter (dyf-7) driving remo-1 cDNA. All rescue constructs were injected with a co-injection marker (unc-122p::RFP). Animals were screened at young adult stage. Three independent lines were observed for each rescue construct. See also Figure S1.
strongly support a role for remo-1, acting in AMsh glia, in the induction of ver-1p::GFP expression.

**REMO-1 Functions in AMsh Glia to Regulate Dauer-Induced AMsh Glia Remodeling**

We next sought to determine whether remo-1 is required for AMsh glia remodeling in dauers. Remodeling is difficult to follow optically, as the structures in question are often separated by <20 nm. We therefore monitored remodeling functionally by assessing the frequency of fusion of the two AMsh glia, using a cytoplasmic mixing assay we previously developed (Procko et al., 2011). Briefly, daf-7(e1372) single or daf-7(e1372); remo-1(ns250) double mutants, which, independently of crowding or starvation, become dauers at 25°C as a result of the daf-7(e1372) lesion, were engineered to carry an AMsh glia::GFP transgene from an unstable extrachromosomal array (nsEx1391). First-stage larvae (L1) of these strains that express GFP in only one of the bilateral AMsh glial cells were then isolated. These mosaic animals were cultivated at 25°C for >48 h to induce dauer entry and scored for GFP redistribution to both cells, indicating fusion. Although 46% of daf-7(e1372) single mutants distribute GFP to both AMsh glia, consistent with our previous studies (Procko et al., 2011, 2012), no redistribution is seen in daf-7(e1372); remo-1(ns250) double mutants (Figure 3B). Similarly, the CRISPR-generated strains, remo-1(ns820) and remo-1(ns821), also show no GFP redistribution (Figure 3B). Consistent with these findings, daf-7(e1372) dauer animals that fail to redistribute GFP do not show fusion when examined by electron microscopy (EM) serial reconstructions (n = 3), while animals that distribute GFP do (n = 2; Figures 3C and 3D), and EM reconstructions of daf-7(e1372); remo-1(ns250) double mutants reveal a lack of fusion (n = 2; Figure 3E). Importantly, as with ver-1p::GFP rescue, expressing REMO-1 in AMsh glia, but not in AWC and other amphid neurons, fully restores AMsh glia fusion to remo-1(ns250) mutants (Figure 3B). Thus, REMO-1 is required in AMsh glia for dauer-dependent AMsh glia remodeling and functions downstream of or in parallel to DAF-7/TGFβ.

**REMO-1 Is Also Required for AWC Neuron Remodeling**

As AMsh glia can affect sensory NRE morphology (Procko et al., 2011, 2012; Singhvi et al., 2016), we assessed whether REMO-1 regulates AWC remodeling using EM. In ttx-1 mutant dauers, AWC neurons can expand within the limited space provided by each unfused AMsh glial cell by forming membrane loops to accommodate excess AWC growth (Figure 1D) (Procko et al., 2011). In these animals, AWC and AMsh glia remodeling are, therefore, uncoupled. In ztf-16 mutants, AMsh glia and AWC NREs are not well developed in non-dauer animals and remain the same in dauer animals (Figure 1E) (Procko et al., 2012). In remo-1(ns250, ns820, or ns821) mutants, we found that AWC dendrites fill the unfused glial space but do not expand further to create the membrane loops observed in ttx-1 mutants (n = 6; Figures 1D, 1F, and 3E), suggesting that remo-1 is required for both AMsh fusion and AWC expansion.

To determine whether remo-1 regulates other neuronal remodeling events, we examined IL2 sensory neurons, which exhibit dauer-dependent dendritic arborization (Schroeder et al., 2013). We found that 100% of daf-7(e1372) animals and 98% of daf-7(e1372); remo-1(ns250) double mutants exhibit remodeling (Figures S2C–S2E). Thus, although remo-1 controls dauer-induced AMsh glia and AWC remodeling, it does not control all remodeling events.

**REMO-1 Is Expressed in AMsh Glia and Localizes to the Glial Tip**

The ver-1p::GFP and AMsh glia fusion rescue studies suggest that REMO-1 is expressed in AMsh glia. To confirm this, we generated wild-type animals carrying a bicistronic transgene consisting of the remo-1 genomic locus, including its 5' intergenic region, and DNA encoding mKate2 fluorescent protein tagged with two nuclear localization signals, separated by a trans-splicing site (Figure 4A). We found that mKate2 expression is detected in AMsh glia of all larval stages, as confirmed by co-expression with a pan-glial mir-228p::myristoylGFP transgene (Figure 4B). We also detected rare expression in ASI and ASJ neurons (Figure S2), confirmed by DIO staining, but never in AWC neurons. Although ASI and ASJ control whole-animal dauer entry and exit (Bargmann and Horvitz, 1991; Cornils et al., 2011), remo-1 mutants can form dauers at normal rates, and remo-1 AMsh and AWC remodeling defects are not rescued by expression of remo-1 in these neurons (Figure 3B). The cell-autonomous functions of REMO-1, together with lack of REMO-1 expression in AWC neurons, suggest that AWC remodeling defects most likely follow from AMsh glia deficits.

To examine REMO-1 subcellular localization, we used CRISPR to knock in a transgene encoding Venus fluorescent protein sequences upstream of the remo-1 stop codon (Figure 4A) (Arribere et al., 2014). Transgenic animals show consistent Venus expression in two adjacent regions at the tip of the nose in both dauer and non-dauer larvae (Figure 4C), consistent with localization at the site of AMsh glia and AWC remodeling events.

**REMO-1 Differentially Regulates ver-1 Transcription and Amphid Remodeling**

To understand how REMO-1 functions with respect to previously identified remodeling genes, we examined expression of ttx-1 and ztf-16 and localization of AFF-1 in wild-type and remo-1(ns250) mutants. Using ttx-1 and ztf-1 transcriptional reporters, we measured the mean fluorescent intensity of GFP expression in the AMsh cell body in wild-type (n = 21 and n = 11, respectively) and remo-1(ns250) mutant (n = 21 and n = 12, respectively) backgrounds and found that ttx-1 and ztf-16 expression levels are unaffected by the remo-1 mutation. We also quantified ttx-1 and ztf-16 expression frequencies and found no difference in the fraction of animals expressing ttx-1 in AMsh glia in either wild-type (100% [n = 75]) or remo-1(ns250) mutant backgrounds (100% [n = 75]). Similarly, ztf-16 expression frequency was unchanged in wild-type (100% [n = 50]) or remo-1(ns250) mutants (100% [n = 50]).

Likewise, AMsh glia-expressed AFF-1::GFP fusion protein localizes to the nose tip in wild-type animals (100% [n = 50]) and in remo-1(ns250) mutants (100% [n = 48]). Thus, remo-1(ns250) affects ver-1p::GFP expression but not expression or localization of other genes driving remodeling.
Figure 3. REMO-1 Functions in AMsh Glia to Regulate Dauer-Induced AMsh Glia Remodeling

(A) Cytoplasmic mixing assay. Schematic of the head of the animal, which expresses an AMsh::GFP transgene from an unstable extrachromosomal array (nsEx1391). Anterior is up. See STAR Methods. Adapted from Procko et al. (2011).

(B) Percentage of animals with fused AMsh glia as scored by cytoplasmic mixing assay in indicated genotypes. All strains carry the daf-7(e1372) mutation to induce dauer when cultured at 25°C. Cell-specific expression of remo-1 in AMsh glia (F16F9.3 promoter) rescues dauer-induced AMsh glia fusion, whereas expression in the amphid neurons (dyf-7 promoter) does not. Rescue experiments were conducted in remo-1(ns250) mutant background, and three independent extrachromosomal arrays were observed per rescue construct. Number of animals scored is inside bars. Error bars denote SEM of at least three independent scoring experiments. ****p < 0.0001.

(C–E) Representative electron micrographs (EMs) and schematic outlines of amphid sensory organs in dauer animals of given genotypes. Dauer larvae were induced by the daf-7(e1372) mutation cultivated at 25°C. Scale bar, 1 μm.
Curiously, we found that remo-1(ns843, ns844, and ns845) predicted loss-of-function deletion alleles, as well as remo-1(ns841 and ns842) premature stop-codon insertion alleles we generated, do not perturb ver-1 expression at 25°C or in dauer larvae but still block glia remodeling (Figures S3A–S3E). These observations suggest that the effect on ver-1 expression is allele specific and demonstrate that ver-1 expression can be uncoupled from dauer remodeling. This is consistent with our previous findings that mutations in ver-1 do not fully block remodeling (Procko et al., 2011). Thus, REMO-1 likely acts upstream of additional remodeling pathways. AWC neurons also fail to remodel in these loss-of-function mutants (n = 4; Figure S3E), further suggesting that remo-1-dependent AMsh glia remodeling likely influences AWC remodeling.

**REMO-1-Dependent Remodeling Facilitates Dauer Exit**

The purpose of amphid remodeling in dauer animals has been debated. Previous studies demonstrated that amphid sensory neuron odorant receptor repertoire is altered in dauers (Peckol et al., 2001). This raises the possibility that dauers adjust sensory tuning to detect favorable environments, which would promote dauer exit behavior. We sought, therefore, to examine whether remodeling-defective animals exhibit dauer exit defects. In dauer animals, pharyngeal pumping, used to suck in bacteria, ceases (Cassada and Russell, 1975; Chow et al., 2006), and pumping resumption is the first sign of dauer exit (Albert and Riddle, 1983). To assess dauer recovery, we therefore exposed animals to GFP-expressing bacteria, which promote dauer exit, and monitored pharyngeal GFP accumulation (Figures 4D and 4E). Accumulation coincides with the onset of pharyngeal pumping, as correlated by direct observation (Figure S4A) (Chou et al., 2015; Proudfoot et al., 1993), and can be used to quickly examine many animals at a time. We found that 50% of wild-type animals exit the dauer stage at 2.9 ± 0.3 h following bacterial food introduction, while ttx-1 mutants, in which dauer-dependent AMsh glia remodeling is defective, exit dauer significantly later, with half of animals accumulating pharyngeal GFP at 4.0 ± 0.1 h (Figure 4F). Animals in which AWC neurons are ablated (Beverly et al., 2011), but in which AMsh glia fusion still occurs (Figure S4B), are even more severely delayed, with 50% recovering at 4.7 ± 0.1 h (Figure 4F). This is consistent with the observation that AWC neurons in ttx-1 mutants still expand, albeit in a disorganized fashion (Procko et al., 2011). These results suggest that remodeling of AWC neurons may drive timely dauer exit behavior. However, ttx-1 mutants also perturb neuronal temperature sensing, and AWC neuron ablations do not distinguish whether it is the presence or the specific remodeling of AWC neurons that is important. Therefore, we reasoned that remo-1 mutants, which fail to remodel both AMsh glia and AWC neurons, and which do not exhibit other defects, are better suited for testing the effects of remodeling on dauer exit.

We found that following food exposure, 50% of remo-1(ns250) animals exit dauer at 4.6 ± 0.6 h, a similar delay to that observed in AWC-ablated animals (Figure 4F), supporting the idea that dauer remodeling is indeed required for efficient dauer exit. Importantly, restoring wild-type REMO-1 to AMsh glia of remo-1(ns250) animals not only restores AMsh glia fusion, and presumably AWC remodeling, but fully rescues dauer exit delay (Figure 4F). Expression of REMO-1 in amphid neurons does not (Figure 4F). Taken together, the correlation of remodeling dynamics and dauer exit are consistent with the view that these two processes are causally linked.

To determine whether glia remodeling affects dauer exit efficiency, we examined wild-type animals that exited dauer early (2 h) or late (4 h) following exposure to GFP-tagged OP50 bacteria. We then examined AMsh glia fusion in these animals using EM. In all animals that exited dauer early, left and right AMsh glial cells fused on both dorsal and ventral sides (n = 4). However, glial fusion was more variable in animals that exited dauer late. Here, we found one animal in which AMsh glia failed to fuse on both dorsal and ventral sides, one animal in which glial fusion occurred only on one side, and one animal with glial fusion on both sides. Thus, glial remodeling may facilitate efficient dauer exit upon exposure to favorable conditions.

**DISCUSSION**

The studies presented here provide insight into how sensory neurons and their associated glia remodel in response to a changing environment and demonstrate direct behavioral consequences of such plasticity. Correlating activities of single neurons, single synapses, or single sensory endings, with animal

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**Figure 4. REMO-1 Is Expressed and Localized at the Tip of the AMsh Glia to Regulate Dauer Exit Timeline**

(A) Top: schematic of remo-1 and its upstream gene, ZC132.11. Middle: remo-1 transcriptional reporter. Bottom: remo-1 translational reporter. Black boxes represent exons, black lines indicate introns, and gray line indicates 5′ intergenic region. Scale bar, 100 bp.

(B) Fluorescence images depicting expression of remo-1 transcriptional reporter co-expressed with a pan-glial mir-228p::myristoylGFP transgene. remo-1 promoter expression in AMsh glia (left, arrow); mir-228 pan-glial promoter expression (middle left, AMsh glia arrow); merged (middle right; AMsh glia arrow); merged with DIC (right, AMsh glia arrow).

(C) Fluorescence images depicting subcellular localization of REMO-1 in AMsh glia (arrow) in adult (left) and dauer (right) animals. REMO-1 accumulates at the tip of the amphid region. Adult animals were cultivated at 25°C, and dauer state was induced by starvation. Magnified inset of the Venus accumulation at the tip of the nose. In (B) and (C), scale bar, 10 μm.

(D) Dauer recovery assay. See STAR Methods.

(E) Fluorescence images showing timeline of GFP-tagged OP50 bacteria accumulation in the pharynges of dauer animals cultivated at 15°C. Images are maximum z stack projections of the entire head volume of the animal. Outline of the animal is marked in gray. Arrow marks the accumulation of GFP-tagged OP50 bacteria in the pharynx. Anterior is up. Time in hours following exposure to GFP-tagged OP50 bacteria. Each image is of a different animal. Scale bar, 10 μm.

(F) The rates of dauer exit measured by the onset of GFP-tagged OP50 bacteria accumulation in the pharynx. Black dotted line indicates the time point at which 50% of the given population recovers pumping. ttx-1(p767), AWC-ablated, and remo-1(ns250) mutants show significant dauer exit delay compared with wild-type (p < 0.0001). Dauer exit delay in remo-1(ns250) is rescued by restoring WT REMO-1 in AMsh glia (F16F9.3 promoter) but not in amphid sensory neurons (dyf-7 promoter). Dauer larvae induced by starvation. Error bars denote SEM of at least three independent scoring experiments.

See also Figures S2A, S2B, and S4A.
behavior is impossible in most model systems, as these responses require activities of large neuronal ensembles. We show here that *C. elegans* provides a unique setting that allows direct interrogation of this black box regime, in which single-neuron, neuron-glia pair, or sensory structure perturbations lead to direct behavioral output.

Our data are consistent with a model in which REMO-1 acts in AMsh glia following detection of dauer entry cues to facilitate AMsh glia remodeling. This drives expansion of AWC neuron NREs, which, in turn, appears to promote efficient dauer exit following re-introduction of *C. elegans* to a favorable environment. REMO-1 functions in parallel to or downstream of DAF-7/TGFβ signaling and may therefore directly sense environmental cues promoting dauer entry. However, it may also respond to detection of such cues by sensory neurons, respond indirectly to downstream signaling induced by neuropeptides or insulin-like peptides important for dauer entry, become active in response to internal ligands, or exhibit constitutive activity, transducing signals only when downstream signaling machinery becomes available. Identification of REMO-1 ligands may help distinguish among these possibilities. Furthermore, our results suggest that REMO-1 likely has yet to be discovered targets, which may function in both AMsh glia and AWC neurons.

REMO-1 is a member of a small family of predicted *C. elegans* GPCRs with a unique C-terminal sequence. This raises the possibility that these other GPCRs might also participate in remodeling events. Interrogating the consequences of their loss may reveal such roles.

Finally, how REMO-1-mediated expansion of AWC neurons leads to accelerated dauer exit is a fascinating area for future investigation. This study, as well as our previous studies, reveals that AMsh glia fusion occurs in only 50% of wild-type dauer animals. Why this remodeling is not fully penetrant remains unclear. However, it is possible that it represents a way for animals to hedge their bets about the quality of a dauer exit signal. If dauer animals that remodel exit more quickly, and the environmental stimulus triggering exit is only transient, these animals would not go on to reproduce. However, un-remodeled dauer animals would not initiate dauer exit and would survive until a bona fide favorable environment is detected. In such a model, the increased surface area of AWC neurons could enhance their sensitivity to favorable environmental stimuli, perhaps by increasing the number of available receptors. Enhanced sensitivity could be used to detect volatiles from further away, or to compensate for the thicker cuticle of dauer animals, which may reduce odorant engagement with receptors. Our studies show that exposure to a dauer-exit stimulus for 4 h is sufficient to induce exit of both remodeled and un-remodeled animals. Thus, if remodeling is a mechanism for testing stimulus stability, persistence of the stimulus for 4 h must be a good predictive measure of longer-term stimulus stability in natural contexts.

**STAR METHODS**

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**SUPPLEMENTAL INFORMATION**

Supplemental Information can be found online at https://doi.org/10.1016/j.celrep.2020.108607.

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**AUTHOR CONTRIBUTIONS**

I.H.L. and S.S. designed the experiments and wrote the manuscript. I.H.L. performed all experiments and analyses, except for EM, which was performed by Y.L., and the genetic screen for ver-1 expression, which was performed by C.P.

**DECLARATION OF INTERESTS**

The authors declare no competing interests.

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## STAR METHODS

### KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Bacterial and Virus Strains | | |
| **E. coli OP50** | Caenorhabditis Genetics Center (CGC); https://cbs.umn.edu/cgc/home | N/A |
| **E. coli OP50-GFP** | CGC | N/A |
| Experimental Models: Organisms/Strains | | |
| **C. elegans: N2** | CGC | N/A |
| Oligonucleotides | | |
| REMO-1[E278K] sgRNA: 5’ TCCAGTGGTTGTCATGTGGA 3’ | This study | N/A |
| REMO-1(deletion/premature stop) sgRNA #1: 5’ GCGTTGGTCAAGCAACATG 3’ | This study | N/A |
| REMO-1(deletion/premature stop) sgRNA #2: 5’ AATTCTGAGTGTGTGTTGTTGTA 3’ | This study | N/A |
| REMO-1::VENUS sgRNA: 5’ ACGGAAAATCAAAGCGAACCC 3’ | This study | N/A |
| Single stranded repair oligo REMO-1[E278K]: 5’ TATTGTGAGTATCGATTTC TTCTTGTTCAAGTGTTGTGAGTTGAAACGAT AGGAAAATCAAAGCGAACCCGG GTATTGCAGATTTCGCTTTTAAACGAT AACAAACAA 3’ | This study | N/A |
| REMO-1::VENUS homologous repair template (detail in methods) | This study | N/A |
| Recombinant DNA | | |
| pDD162 | (Dickinson et al., 2013) | Addgene |
| pHL164: remo-1[E278K] CRISPR targeting vector #1, pDD162 backbone | This study | N/A |
| pHL212: remo-1[deletion/premature stop] CRISPR targeting vector #1, pDD162 backbone | This study | N/A |
| pHL213: remo-1[deletion/premature stop] CRISPR targeting vector #2, pDD162 backbone | This study | N/A |
| pHL217: remo-1[Venus] CRISPR targeting vector, pDD162 backbone | This study | N/A |
| Fosmids WRM0638cF08 | Source BioScience https://www.sourcebioscience.com | Moerman Fosmid Library |
| nsIs22: ver-1::GFP | (Procko et al., 2011, 2012) | N/A |
| nsEx6181, -82, -83: remo-1 fosmid (WRM0638cF08) (20 ng/μl) + unc-122p::RFP (20 ng/μl) + pBluescript (60 ng/μl) | This study | N/A |
| nsEx6184, -85, -86: remo-1 gDNA (50 ng/μl) + unc-122p::RFP (20 ng/μl) + pBluescript (30 ng/μl) | This study | N/A |
| nsEx6048, 6173, 6174: F16F9.3p::remo-1 cDNA (40 ng/μl) + unc-122p::GFP (20 ng/μl) + pBluescript (40 ng/μl) | This study | N/A |

(Continued on next page)
RESOURCE AVAILABILITY

Lead Contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Shai Shaham (shaham@rockefeller.edu).

Materials Availability
All unique/stable reagents generated in this study are available from the Lead Contact without restriction.

Data and Code Availability
This study did not generate any unique datasets or code.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

C. elegans
C. elegans were cultivated using standard methods (Brenner, 1974). All animals were cultivated at 15°C or 25°C, unless otherwise noted. Bristol N2 strain was used as wild-type. Mutants recovered by Ethyl methanesulfonate (EMS) mutagenesis were outcrossed at least three times before use. Most strains had the nsls22 (ver-1p::GFP) IV reporter transgene. Co-injection markers used were unc-122p::GFP, unc-122p::RFP, or unc-122p::DsRed expressed in coelomocytes, or plasmids as otherwise noted. Integrated transgenic

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| nsEx6045, -46, -47: dyf-7p::remo-1 cDNA (40 ng/µl) + unc-122p::GFP (20 ng/µl) + pBluescript (40 ng/µl) | This study | N/A |
| nsls391: mir-228p::myr-GFP + lin-15(+) | This study | N/A |
| nsls6249: pHIL28[remo-1p::SL2-NLSx2-mKate2] (80 ng/µl) + unc-122p::RFP (10 ng/µl) | This study | N/A |
| nsEx1391: F16F9.3p::GFP + pRF4[rol-6( suicide1006)] | (Procko et al., 2011, 2012) | N/A |
| nsls855: F28A12.3p::myr-GFP | This study | N/A |
| nsEx1942: ttx-1p::GFP + pRF4[rol-6( suicide1006)] | (Procko et al., 2011) | N/A |
| nsEx3001: R08E3.4p::GFP + pRF4[rol-6( suicide1006)] | (Procko et al., 2012) | N/A |
| nsEx2727: F16F9.3p::aff-1 cDNA-GFP + pRF4[rol-6( suicide1006)] | (Procko et al., 2011) | N/A |
| nsEx6027: unc-122p::RFP (20 ng/µl) + pBluescript (80 ng/µl) | This study | N/A |
| nsEx6125, -26, -27: pIL20 [F16F9.3p::REMO-1 E278K] (10 ng/µl) + unc-122p::GFP (20 ng/µl) + pBluescript (70 ng/µl) | CGC; (Beverly et al., 2011) | N/A |
| oysIs85: ceh-36p::TU813 + ceh-36p::TU814 + srxT-1p::GFP + unc-122p::DsRed]. TU813 and TU814 are split caspase vectors (Chalfie Lab) subcloned downstream of the ceh-36 promoter. | | |

Software and Algorithms

| Software | Source | URL |
|----------|--------|-----|
| Prism 8 | GraphPad | N/A |
| ImageJ | (Schneider et al., 2012) | https://imagej.nih.gov/ij/ |
| Protter | (Omasits et al., 2014) | http://wlab.ethz.ch/protter/start/ |
| Jalview | (Waterhouse et al., 2009) | https://www.jalview.org/ |

RESOURCES

RESOURCES

Continued
strains were generated with UV/trioxalen treatment (Mello et al., 1991) (Sigma, T6137). AWC neurons were ablated via cell-specific expression of caspases under ceh-36 promoter sequences (Beverly et al., 2011). Some strains listed below were sourced from the CGC, funded by NIH Office of Research Infrastructure Programs (P40 OD010440).

Alleles used in this work are:

LGII: daf-7(e1372), unc-119(ed3)
LGV: ttx-1(p767), remo-1(ns250, ns820, ns821, ns841, ns842, ns843, ns844, ns845, ns846[Venus])
LGX: ztf-16(ns171)

METHOD DETAILS

Forward Genetic Screen
Fourth-stage larvae carrying nsIs22 in the N2 strain background were mutagenized with ethylmethanesulfonate (EMS) for 4 hours. Individual P0s were picked to separate 9cm NGM agar plates with seeded OP50. F2 animals were screened under a fluorescent dissecting scope (Leica) for mutants that failed to express ver-1p::GFP in the AMsh glia (Procko et al., 2011, 2012).

Gene Identification
A combination of Hawaiian Snip-SNP mapping (Wicks et al., 2001), and whole genome sequencing (Zuryn et al., 2010) using galign (Shaham, 2009) to align reads, was used to identify remo-1(ns250). The remo-1 mutation was confirmed by fosmid rescue (WRM0638cF08); remo-1 genomic transgene rescue; and re-introduction of E278K CRISPR alleles.

Germline Transformation
Germline transformation was carried out as previously described (Mello and Fire, 1995). Plasmid mixes containing the plasmid of interest, co-injection markers, and pBluescript were injected into one or both gonads of young adult hermaphrodites (Mello and Fire, 1995). pBluescript was used to adjust the DNA concentration of injection mixes as necessary. Injected animals were singled onto NGM plates and allowed to grow for two generations. Transformed animals were picked onto single plates based on co-injection marker expression, and screened for stable inheritance of the extrachromosomal array. Only lines from different P0 hermaphrodites were considered independent. Isolated strains were staged based on morphology under a dissecting microscope and were screened for ver-1p::GFP expression, amphid remodeling, IL2 remodeling and/or dauer recovery.

Plasmid Construction
Plasmids were constructed using Gibson cloning into the applicable backbone plasmid (Gibson et al., 2009). remo-1 cDNA was constructed by cloning each exon from the fosmid WRM0638cF08 and fusing in sequential order using Gibson cloning. remo-1 promoter region includes the entire remo-1 genomic locus, including its 5' intergenic region. CRISPR-related vectors were generated using a site-directed mutagenesis protocol on plasmid pDD162 (Dickinson et al., 2013; Liu and Naismith, 2008).

Cytoplasmic Mixing Assay
AMsh glia fusion was observed using a fluorescence redistribution assay described in detail in Procko et al. (2011, 2012). First-stage mosaic larvae carrying the nsEx1391 array in one of the two AMsh glia were picked onto fresh seeded plates. Mosaic animals were cultivated for at least 48 hours at 25°C to induce dauer entry due facilitated by daf-7(e1372) temperature-sensitive allele. Animals were only scored if they were dauer larvae by morphology at the end of the assay period. Mosaic animals were scored for redistribution of GFP to both cells, indicating fusion.

Dauer Selection
To induce dauer formation by starvation, embryos were collected by hypochlorite treatment and washed extensively in M9. Approximately 200 embryos were placed onto an NGM plate containing a small lawn of OP50 (~10 μL of saturated OP50, 24 hr before use). Embryos were allowed to hatch and develop at 15°C or 25°C into dauer larva. Failure to include OP50 on the plate resulted in animals that arrested as starved L1 stage larvae. Dauer animals were selected by treatment with 1% SDS in M9 solution for 15 min. Alternatively, young animals carrying the daf-7(e1372) mutation were induced to form dauers by incubation at 25°C.

Dauer Recovery Assay
Pharyngeal Pumping Assay
Dauer animals induced by starvation were individually selected onto a plate with a thick lawn of OP50 and incubated at 15°C. Dauers were examined every 30-60 min for consistent grinder movement, indicative of pharyngeal pumping, under the dissecting scope.

Uptake of GFP-Expressing Bacteria
OP50-GFP bacteria (CGC) were cultivated in LB media. 24 hr before the assay, approximately 40 μL of bacteria were plated evenly on the NGM plates and dried overnight in the dark at 15°C. Dauer animals induced by starvation were singled out on the seeded plate
and cultivated at 15°C in the dark. Every 30-60 min, dauer animals were screened for GFP accumulation in the pharynx under the dissecting scope.

**Dye Filling Assay**
DiO (3,3′-diodocyanine) (Sigma) stock solution was made at 5 mg/ml in N,N-dimethylformamide. Animals were stained with DiO (stock solution diluted 1:10000 in M9 buffer) for 30–60 min, followed by three M9 washes to remove excess dye.

**Generation of remo-1 Alleles and Reporter Using CRISPR-Cas9 Genome Editing**
Alleles of remo-1 were generated using the co-CRISPR-based genome editing method as previously described (Arribere et al., 2014). pDD162 was used as a vector backbone and the following sgRNA sequences were added for each individual CRISPR attempt (E278K: 5′ TCCAGTGTTGCTCATGGGA 3′; deletion and premature stop alleles: #1 (5′ GGCTTTGTCAAGCATACTG 3′) and #2 (5′ AATTCTAGTGTATTCTTTGTGA 3′)) to generate remo-1 targeting vectors. A dpy-10 sgRNA-pDD162-based vector was also generated (5′ GCCACCATGGCACCCAGGAG 3′). Single-stranded repair oligos were ordered from Sigma (E278K: 5′ ATTGTGTAGTAC GATTTCCCTGTGGTGTGCTAGTGAGCAGaaATACAAAGCGAACCCGTTGTTATTCGAGATTGTCATGGAACAAA 3′; dpy-10(cn64): 5′ CATTGAACCTCAATCATGAGAAATGACTGAAACCCGTACGGCTATGGTACGG GAGCTTCCATGCGTCAACACAGCCTAT 3′) (Arribere et al., 2014). N2 animals were injected with the following mix: 50 ng/μl dpy-10 sgRNA, 50 ng/μl remo-1[E278K] targeting vector (or a mix of 25 ng/μl remo-1[deletion/premature stop] targeting vector #1 and 25 ng/μl remo-1[deletion/premature stop] targeting vector #2), 20 ng/μl dpy-10(cn64) repair oligo, and 20 ng/μl remo-1[E278K] repair oligo (when appropriate) in 1x injection buffer (20 mM potassium phosphate, 3 mM potassium citrate, 2% PEG, pH 7.5). F1 animals with Dpy or Rol phenotypes were picked to individual plates, indicating a CRISPR-based editing event had occurred. F1 animals were allowed to lay eggs, and then genotyped for successful co-conversion of the remo-1 locus using PCR and restriction enzyme screening or Cel1 digestion of heteroduplex DNA (Ward, 2015). Non-Rol, Non-Dpy F2 animals were then singled and homozygosed for the remo-1 mutation or deletion.

The Venus reporter was generating by using Cas9-triggered homologous recombination (Dickinson et al., 2013). Venus sgRNA sequence (5′ ACAGGAATCAAGGCCAACCC 3′) and homologous Venus repair template (5′ 1.49kb homolog arm of the 3′ end of remo-1 coding sequence)-loxP-reversed unc-119(+) cassette-loxP-Venus transgene-1.5kb remo-1 downstream homology arm 3′) were used to insert the Venus reporter transgene at the 3′ end of remo-1 sequence. DP38(unc-119(ed3)) animals were injected with the following mix: 10 ng/μl pG8H (rab-3p::mCherry, Addgene #19359), 5 ng/μl pCFJ104 (myo-2p::mCherry, Addgene #19328), 2.5 ng/μl pCFJ90 (mve-2p::mCherry, Addgene #19327), 50 ng/μl REMO-1::VENUS targeting vector, 10 ng/μl Venus repair oligo in 1x injection buffer (20 mM potassium phosphate, 3 mM potassium citrate, 2% PEG, pH 7.5). Three injected worms per plate were left to starve for 2 weeks at 25°C. Non-red animals were picked individually and allowed to lay eggs, and then genotyped for successful insertion. The loxP-unc-119(+)-loxP cassette was subsequently removed by injecting the following mix: 50 ng/μl pDD104 (eft-3p::Cre::tibb-2 3’UTR; Addgene #47551), 2.5 ng/μl pCFJ90 (mve-2p::mCherry, Addgene #19327), 47.5 ng/μl pBluescript in EB buffer. The animals in which the unc-119(+) gene was excised were confirmed and isolated by DNA sequence analysis. The unc-119(ed3) mutation was then crossed out. Removing the unc-119(+) cassette did not change Venus expression pattern.

**Microscopy and Image Processing**
ver-1::p::GFP (nisls22) expression was assayed using a fluorescence dissecting microscope (Leica). Young adult hermaphrodites or dauer animals were scored. Compound microscope images were taken on an Axioplan II microscope using an AxioCam CCD camera (Zeiss) and analyzed using the Axiovision software (Zeiss). Some images were collected on a DeltaVision Core imaging system (Applied Precision) with a PlanApo 60x/1.42 na or UPlanSApo 60x/1.3 oil-immersion objective and a Photometrics CoolSnap HQ camera (Roper Scientific). Images were deconvolved using softWoRx program (GE Healthcare).

ttx-1 and ztf-16 expression levels were assayed using transcriptional reporter transgenes nsEx1942 and nsEx3001, respectively. Using ImageJ, we hand-traced the AMsh cell body at the focal point section and measured the mean fluorescent intensity in wild-type and remo-1(ns250) mutant backgrounds. For AFF-1, we examined the nsEx2727 transgene, an AMsh glia-expressed AFF-1::GFP fusion protein, and examined protein localization at the tip of the nose.

**Electron Microscopy**
Dauer animals for electron microscopy were grown at 25°C and selected for AMsh glial fusion or no fusion based on GFP fluorescence (see cytoplasmic mixing assay for more details). Animals were prepped and sectioned using standard methods (Lundquist et al., 2001). Imaging was acquired with a FEI Tecnai G2 Spirit BioTwin transmission electron microscope equipped with a 4K digital camera. The image acquisition, processing, and analysis was performed by using the SerialEM and IMOD software (Schorb et al., 2019).

**QUANTIFICATION AND STATISTICAL ANALYSIS**
Statistical analysis was performed using GraphPad Prism. Sample sizes and statistical parameters are reported in the main text, figures, and figure legends. When comparing a transgenic line to the parental strain, a minimum of three independent lines were scored.
Data is judged to be statistically significant when $p < 0.05$ by unpaired t test or one-way ANOVA followed by Tukey’s test where appropriate. The rates of the onset of GFP-tagged bacteria accumulation in the pharynx were plotted as survival from dauer and were compared by log-rank test using GraphPad Prism 8.

**Protein Structure Prediction**
Prediction of protein structure generated using Protter software (Omasits et al., 2014)

**Multiple Sequence Alignment**
A subfamily of Srz predicted GPCR amino acid sequences were aligned using Jalview v.2.11 (Waterhouse et al., 2009)
Figure S1. Multiple sequence alignment of the nematode Srz predicted GPCR subfamily. Related to Figure 2. Sequence alignment using Jalview of remo-1 paralogs and orthologs in the Srz predicted GPCR subfamily. Paralogs: ZC132.9 and ZK218.4. Orthologs: CBN07202, CBN17678 (C. brenneri); CRE15966, CRE15968, CRE15970, CRE15967, CRE15960 (C. remanei); CBG06292, CBG06289, CBG06291 (C. briggsae). remo-1(ns250) mutation site is marked by blue arrow. The conserved threonine-glutamatic acid-isoleucine (TEI) motif is colored in pink.
Figure S2

A  remo-1p::SL2::NLSx2::mKate2
Merged
DIO

B

C

D  remo-1(+) dauer
remo-1(ns250) dauer

E  remo-1(ns250) dauer
Figure S2. Characterization of remo-1 expression pattern and effects on other remodeling events in dauer. Related to Figure 4. (A,B) Fluorescence images of young adult animals carrying remo-1 transcriptional reporter stained with DiO (right). DiO dye filling assay was conducted to stain amphid sensory neurons in vivo. White outlines trace the worm. (A) ASI expression (left); merged (middle). (B) ASJ expression (left); merged (middle). Scale bar, 10μm. (C) Percentage of dauers with IL2 dendritic arborization in indicated genotypes. 50 animals scored for each genotype. (D,E) Fluorescence images depicting expression of IL2 reporter (F28A12.3p::myr-GFP) in remo-1(+) (D) and remo-1(ns250) (E) dauer animals. Dauer larvae were induced by the daf-7(e1372) mutation cultivated at 25°C. (D-E) Scale bar, 10μm.
Figure S3

**A**

remo-1

ns843 (4TM deletion)
ns844/845 (4TM deletion)
ns841/842 (early stop)

100bp

**B**

% dauers with fused AMsh glia

remo-1(-)

daf-7(e1372)

**C**

25°C

% animals with GFP in AMsh

remo-1(-)

**D**

% animals with GFP in AMsh

remo-1(-)

**E**

remo-1(ns842)

1 μm
**Figure S3. remo-1 allele genetics. Related to Figure 4.** (A) *remo-1* gene structure and mutation sites. Boxes represent exons and lines indicate introns. Point mutations and deletions relative to the *remo-1* start codon: *ns841/842*: leucine-leucine at positions 185-186 change to alanine-stop codon indicated in orange; *ns843*: 15 amino-acid deletions at positions 157-171 indicated in green; *ns844/845*: 21 amino-acid deletions at positions 154-174 indicated in pink. (B) Histogram detail as in Figure 3B. (C) Histogram detail as in Figure 2A. (D) Histogram detail as in Figure 2B. (E) Representative electron micrograph (EM) and schematic outline of an amphid sensory organ in dauer animals of the indicated genotype. Scale bar, 1μm.
Figure S4. Dauer exit measured by onset of OP50-GFP accumulation and effects of AWC ablation on remodeling. Related to Figures 1 and 4 (A) OP50-GFP accumulation in the pharynx coincides with the onset of pharyngeal pumping measured by direct observation of consistent grinder movement. Histogram detail as in Figure 4F. (B) Histogram detail as in Figure 3B. n.s., p > 0.05.