The mRNA Decay Factor CAR-1/LSM14 Regulates Axon Regeneration via Mitochondrial Calcium Dynamics

Graphical Abstract

Highlights

- *C. elegans* mRNA decay factors affect axon regeneration and maintenance
- The LSM14 ortholog CAR-1 is a repressor of axon regeneration
- CAR-1 represses expression of the mitochondrial calcium regulator MICU-1
- Axon injury triggers mitochondrial calcium uptake regulated by MICU-1

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In Brief
Tang et al. dissect the roles of mRNA decay factors in *C. elegans* axon regeneration. Loss of function in the LSM14 ortholog CAR-1 results in increased axon regeneration due to increased expression of MICU-1, a mitochondrial calcium uniporter regulator that is a key target of CAR-1 repression in neurons.
The mRNA Decay Factor CAR-1/LSM14 Regulates Axon Regeneration via Mitochondrial Calcium Dynamics

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SUMMARY

mRNA decay factors regulate mRNA turnover by recruiting non-translating mRNAs and targeting them for translational repression and mRNA degradation. How mRNA decay pathways regulate cellular function in vivo with specificity is poorly understood. Here, we show that C. elegans mRNA decay factors, including the translational repressors CAR-1/LSM14 and CGH-1/DDX6, and the decapping enzymes DCP1/DCP2, and mRNA degradation by the exonuclease XRN1 (reviewed in [1]). In multiple cell types, depletion of mRNA decay factors usually leads to accumulation of stabilized mRNAs. However, increasing evidence suggests that mRNA decay factors exhibit a high degree of selectivity in mRNA stability regulation [2]. Therefore, it is crucial to identify the mRNA targets of mRNA decay factors to provide a better understanding of their cellular functions.

Neurons are polarized cells that are primed for a high degree of selective and dynamic regulation of gene expression. mRNA decay factors are widely expressed in neurons of C. elegans, Drosophila, and mammals [3–6]. For example, Drosophila Me31B/DDX6 is enriched in post-synaptic dendrites [5], and its overexpression in sensory neurons reduces higher-order dendrite arborization [3], although it remains unknown whether particular mRNA targets are involved.

C. elegans expresses many conserved mRNA decay factors, whose functions have been mostly characterized in the germline and in early embryos [7–10]. For example, the CAR-1 protein family includes yeast Scd6, Drosophila Tra1 (Trailer hitch), Xenopus RAP55, and mammalian LSM14 [11]. The interaction between CAR-1/LSM14 and CGH-1/DDX6 regulates the formation of endoplasmic reticulum and anaphase spindle network; loss of function in car-1 or cgh-1 leads to sterility and embryonic lethality due to defects in germline apoptosis and embryonic cytokinesis [7, 8, 10]. Loss of function in cgh-1 has recently been shown to affect dendrite development in PVD neurons [6]. However, the neuronal mRNA targets of CAR-1 or CGH-1 have not yet been identified.

Here, we show that loss of the translational repressors CAR-1 or CGH-1 results in axon breakage and branching, and increased axon regrowth after injury, whereas loss of mRNA

INTRODUCTION

Mature mRNAs are either actively translated or exist in a translationally repressed state that can be targeted for degradation by mRNA decay factors. Extensive biochemical studies show that the recognition of mRNAs by the mRNA decay factor LSM14 and DEAD-box RNA helicase DDX6 leads to translational repression, followed by irreversible removal of the 5’ cap by decapping enzymes DCP1 and DCP2, and mRNA degradation by the exonuclease XRN1 (reviewed in [1]). In multiple cell types, depletion of mRNA decay factors usually leads to accumulation of stabilized mRNAs. However, increasing evidence suggests that mRNA decay factors exhibit a high degree of selectivity in mRNA stability regulation [2]. Therefore, it is crucial to identify the mRNA targets of mRNA decay factors to provide a better understanding of their cellular functions.

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Here, we show that loss of the translational repressors CAR-1 or CGH-1 results in axon breakage and branching, and increased axon regrowth after injury, whereas loss of mRNA
decapping factors results in aberrant axon development and decreased axon regrowth after injury. We further identified numerous genes including the mitochondrial calcium ([Ca\(^{2+}\)]\(_{mt}\)) regulator \textit{micu-1}, whose mRNAs are bound by CAR-1 in neurons. We present multiple lines of evidence that CAR-1 represses the expression and translation of \textit{micu-1} in neurons. Additionally, we describe a [Ca\(^{2+}\)]\(_{mt}\) influx induced by axon injury and show that the dynamics of this [Ca\(^{2+}\)]\(_{mt}\) is regulated by CAR-1 and MICU-1 with dose dependency. Our studies reveal a novel mechanism whereby mRNA decay pathways regulate axonal mitochondrial calcium [Ca\(^{2+}\)]\(_{mt}\) dynamics.

RESULTS

\textit{C. elegans} mRNA Decay Factors Are Expressed in Neurons and Localize to Distinct Subcellular Granules

We first asked whether key mRNA decay components were expressed in \textit{C. elegans} neurons. We performed smFISH (single-molecule fluorescence \textit{in situ} hybridization) for endogenous \textit{car-1} mRNAs and observed widespread expression in somatic cells including touch receptor neurons (TRNs) and other neurons besides the germline (Figures 1A and S1). We examined functional transgenes of full-length DCAP-1 or CGH-1 expressed under their respective promoters [12, 13] and observed expression in many neurons including TRNs and motor neurons, localizing to cytoplasmic puncta in neuronal cell bodies (Figures 1B and S2A). To determine protein localization in single neurons, we generated transgenes expressing various mRNA decay factors using the TRN-specific \textit{mec-4} promoter. We found that in TRN cell bodies, CGH-1::GFP fully co-localized with mKate2::CAR-1 puncta (Figure 1C), whereas DCAP-1::dsRed partially co-localized with GFP::CAR-1 (Figure 1D). These results suggest mRNA decay components form at least two types of cytoplasmic granules in \textit{C. elegans} neurons. The partial colocalization of CAR-1 and DCAP-1 further implies that only some CAR-1-bound mRNAs undergo mRNA decapping and decay (Figure 1E).

Assembly of mRNA decay complexes in other organisms involves direct interactions such as binding of DDX6 to LSM14 and of DCP1 to the decapping activator PATL [14, 15]. To address whether punctate localization by \textit{C. elegans} mRNA decay factors depended on similar interactions, we examined genetic null (\textit{0}) mutants. Loss of function in \textit{patr-1}/\textit{PATL} resulted in fewer DCAP-1 puncta in neurons, whereas loss of function in \textit{cgh-1} resulted in dimmer CAR-1 puncta (Figures S2B and S2C), indicating that PATR-1 and CGH-1 promote recruitment of DCAP-1 and CAR-1, respectively. In contrast, loss of function in decapping enzymes \textit{dcap-1} or \textit{dcap-2} increased the size of CAR-1 puncta (Figure S1C). In \textit{dcap-2(0)} mutants, these enlarged CAR-1 puncta showed nearly complete colocalization with DCAP-1 (Figure S1D), suggesting that the lack of decapping activity may cause accumulation of granules containing CAR-1 and its bound mRNAs.
mRNA Decay Components Have Differential Roles in Axon Morphology and Regrowth after Injury

To study the roles of mRNA decay components in neuronal development and maintenance, we focused on the posterior TRN, PLM. In wild-type animals, PLM neurons normally extend a long anterior axon and a short posterior dendrite (Figures 2A and S3A). In car-1(0) and cgh-1(0) mutant larvae, PLM neurons displayed normal axon morphology (Figure S3B), but in day 1 mutant adults, PLM axons displayed ectopic branching, breakage, and blebbing (Figures 2A, 2B, S3A, and S3B), suggesting that car-1 and cgh-1 are not essential for PLM axon development, but are necessary for maintaining healthy axon morphology in adults. In contrast, dcap-1(0), dcap-2(0), or dcap-1(0) dcap-2(0) mutants at all larval and adult stages displayed low penetrance hook-shaped axon extensions, in addition to ectopic axon branching and blebbing (Figures 2A, 2B, and S3B), suggesting that decapping enzyme activity is important for PLM development.

Adult PLM axons display robust regrowth following femtosecond laser surgery [16]. We next examined how the mRNA decay factors affected axon regrowth. We severed morphologically normal PLM axons in larval (L4) animals lacking specific mRNA decay components and imaged axon regrowth 24 h post-axotomy (hpa). car-1(0) and cgh-1(0) mutants showed increased regrowth after injury, whereas injured PLM axons in single null mutants for dcap-1, dcap-2, or patr-1, and in dcap-1(0) dcap-2(0) double mutants, all showed decreased axon regrowth (Figure 2C). This analysis implies that although these mRNA decay factors can form a protein complex and partly co-localize within the same cell, they exert differential functions in axon development, maintenance, and regrowth. Below, we focus on CAR-1, which is not required for PLM development but is a strong inhibitor of PLM axon regrowth.

CAR-1 Is a Cell-Intrinsic Inhibitor of Axon Regrowth

Three independent car-1(0) mutants all exhibited enhanced PLM axon regrowth after injury in the L4 stage, as well as adult-onset axon morphology defects (Figures 3A–3C and S3C). The enhanced regrowth in car-1(0) mutants was restored to wild-type levels by single-copy transgenes expressing car-1 using endogenous or TRN-specific promoters, but not using a muscle-specific promoter (Figure 3B), indicating that CAR-1 regulates axon regrowth cell-autonomously. Axon regrowth involves initial growth cone formation followed by axon extension [17–19]. We observed a higher rate of growth cone formation in car-1(0) mutants at multiple time points after axonal injury (Figures 3D and 3E), suggesting that CAR-1 inhibits growth cone formation throughout the process of regrowth. Moreover, overexpression of CAR-1 in TRNs resulted in severe axonal defects in adult animals and strongly impaired axon regrowth and growth cone formation after injury (Figures 3B, 3C, and S3C), indicating that axon maintenance and regrowth are sensitive to CAR-1 levels.

CAR-1 Sm and FDF Domains Are Required for Inhibition of Axon Regrowth

CAR-1 contains three conserved domains (Figure 4A): an N-terminal Sm domain, an FDF (phenylalanine-aspartate-phenylalanine) motif, and a C-terminal RGG (arginine-glycine-glycine) motif [7, 8]. The Sm domain in yeast Scd6 is essential for mRNA translational repression and stimulation of mRNA decay [20], and the FDF motif of C. elegans CAR-1 can interact with human DDX6 [14]. We next asked which domain is relevant for CAR-1 cytoplasmic puncta formation and for its function in axon regrowth. We expressed truncated GFP::CAR-1 in TRNs as single-copy insertion transgenes to ensure consistency in expression level (Figure 4A). We found that the FDF and RGG motifs, but not the Sm domain, contribute to the formation of CAR-1 puncta in...
PLM cell bodies (Figure 4A), consistent with observations in C. elegans embryos [7]. To address the function of CAR-1 mutant proteins, we performed axon injury in car-1(0) animals expressing each transgene. Expression of CAR-1(FL) or CAR-1(Sm+FDF), but not CAR-1(Sm), CAR-1(ΔSm), CAR-1(RGG), nor CAR-1(ΔFDF+RGG), in car-1(0) mutants restored axon re-growth and growth cone formation to wild-type levels (Figures 4B and 4C). These data indicate that the Sm domain and FDF motif of CAR-1, hence its role in RNA translational repression, are critical for CAR-1 function in neurons.

CAR-1 Regulates Its mRNA Expression in PLM Neurons

To identify neuronal targets of CAR-1, we constructed transgenic strains overexpressing GFP::CAR-1(FL) (juIs526) or GFP::CAR-1(ΔSm) (juIs549) in all neurons using the pan-neuronal rgef-1 promoter. In axon injury assay, we found that overexpression of GFP::CAR-1(FL), but not GFP::CAR-1(ΔSm), inhibited PLM axon regrowth (Figure 5A), confirming the importance of the Sm domain in CAR-1 function. We then performed single-end enhanced crosslinking and immunoprecipitation (seCLIP) to isolate mRNA targets of CAR-1 using these transgenic animals (see STAR Methods for details) [21, 22]. We reasoned that mRNAs repressed by CAR-1 should be more stably bound to non-functional GFP::CAR-1(ΔSm) than to GFP::CAR-1(FL). We analyzed seCLIP data to identify peaks specifically enriched in GFP::CAR-1(ΔSm) samples (STAR Methods) and selected about 30 or so potential targets (Table S1). Among them, we found that car-1 itself was specifically enriched in GFP::CAR-1(ΔSm) co-immunoprecipitates (Figure 5B), suggesting CAR-1 might repress its own expression. To test this, we performed smFISH for car-1 in animals expressing Pmec-4::GFP::CAR-1 and found that car-1 mRNA puncta partly colocalized with GFP::CAR-1 cytoplasmic puncta in wild-type and dcap-2(0) animals (Figure 5C). Since cgh-1(0) mutants display dimmer GFP::CAR-1 puncta (Figure S1C), we visualized car-1 mRNAs in cgh-1(0) mutants and observed that cgh-1(0) mutants displayed more car-1 mRNA puncta (Figures 5D and 5E), suggesting that car-1 mRNA levels are normally limited by CAR-1/CGH-1 activity.

CAR-1 Binds micu-1 Transcripts and Represses MICU-1::GFP Protein Levels in PLM Neurons

We identified 29 genes whose mRNAs showed enriched peaks in GFP::CAR-1(ΔSm) seCLIP analysis, including several calcium-related genes: cll-1, kcnl-1, micu-1, and unc-13 (Table S1). Several genes were previously analyzed for their roles in PLM axon regrowth [18, 23]. Given recent evidence for the function of mitochondria in axon regrowth [24–26], here we chose to

Figure 3. CAR-1 Is a Cell-Intrinsic Inhibitor of Axon Regrowth

(A) Map of car-1 gene showing deletion mutations. See also Table S2.

(B) PLM axon regrowth length 24 h post-axotomy in genotype indicated. Bars indicate mean ± SEM. Statistics: one-way ANOVA with Bonferroni’s post test; *p < 0.05; **p < 0.01; ns, not significant (p > 0.05). Sample size is indicated in the bar.

(C) Representative images of axon regrowth 24 h post-axotomy in wild-type, car-1(0) mutants, and Pmec-4::GFP::CAR-1(juEx7280). Anterior is to the left. Red arrows indicate site of axotomy.

(D) Representative images of re-growing axons 6 h post-axotomy in genotype indicated. PLM in car-1(0) mutants has a more persistent regenerative growth cone. Scale bar, 20 μm.

(E) Quantitation of percentage of growth cones at 6, 24, and 48 h post-axotomy. Statistics: Fisher’s exact test; *p < 0.05; **p < 0.01.
focus on micu-1, which encodes a mitochondrial calcium uptake regulator of the MICU1 family (Figure S4A). Careful inspection of the seCLIP peak profiles revealed that GFP::CAR-1(ΔSm) samples were specifically enriched on micu-1 (Figure 6A). To further verify this finding, we performed smFISH of micu-1 mRNA. In wild type PLM neurons, micu-1 mRNA puncta partly colocalized with GFP::CAR-1 (Figure 6A), similar to car-1 mRNA. In car-1(0) mutants, we observed increased numbers of micu-1 mRNA puncta in PLM cell bodies (Figures 6B and 6C). The increased micu-1 mRNA puncta observed in car-1(0) mutant TRNs was suppressed by expressing CAR-1(FL) or CAR-1(Sm+FDF), but not CAR-1(ΔSm) (Figure 6C), suggesting that CAR-1(ΔSm) domain mediates translational repression. micu-1 mRNA puncta did not colocalize with GFP::DCAP-1 (Figure S5A), suggesting that DCAP-1 does not regulate expression of micu-1 mRNA directly.

Next, we characterized endogenous MICU-1 expression by inserting GFP into the carboxy-terminus of the endogenous micu-1 gene (see STAR Methods). MICU-1::GFP was detected at low levels in germ cells, epidermis, and some muscles (Figure S4B). MICU-1::GFP was detected at low levels in PLM neuron soma, colocalizing with a mitochondrial marker, but was below the limit of detection in axons (Figure 6D). As micu-1 transcript levels were increased in car-1(0) mutants, we examined whether MICU-1::GFP signals were different in car-1(0) mutant compared to wild type. We observed a consistent 1.2-fold increase of MICU-1::GFP intensity in PLM cell bodies in car-1(0) mutant compared to wild type (Figures 6E and 6F). Collectively, these observations indicate that CAR-1 binds micu-1 mRNA and represses its translation in PLM neurons.

CAR-1-Mediated Inhibition of Axon Regrowth Is Dependent on the Mitochondrial Calcium Import Factors MCU-1 and MICU-1

MICU1 is an EF-hand protein that forms a complex with the MCU1 unipporter to regulate $[Ca^{2+}]_{mt}$ uptake [27–31]. We therefore hypothesize that axonal mitochondrial calcium uptake might influence regrowth dynamics. To test this, we first asked whether micu-1, or the unipporter mcu-1, affected PLM axon regrowth. We analyzed axon regrowth in double mutants between car-1(0) and mcu-1(0) or mcu-1(0). Loss of mcu-1 or micu-1 fully suppressed increased axon regrowth observed in car-1(0) mutants (Figure 7A). However, loss of mcu-1 or micu-1 did not suppress the increased growth cone formation observed in car-1(0) mutants, indicating that CAR-1 acts via other targets to regulate axon extension and growth cone formation. Micu-1(0) also suppressed the adult-onset axon maintenance defects of car-1(0) mutants (Figure S5B), suggesting that MICU-1 is acting downstream of CAR-1 in axon maintenance. Loss of mcu-1 or micu-1 did not rescue axon regrowth and axon morphology defects of dcap-1(0) mutants (Figures SSC and SSD), implying that micu-1 is a specific target of CAR-1 but not of DCAP-1.

Axon Injury Triggers Mitochondrial Calcium Uptake Dependent on MCU-1 and Regulated by MICU-1

We next addressed whether axon injury might affect $[Ca^{2+}]_{mt}$ flux. We targeted the Ca$^{2+}$ sensor GCaMP5 to the mitochondrial matrix (mtGCaMP) in TRNs. micu-1(0), mcu-1(0), and car-1(0) mutants displayed normal axonal mitochondrial distribution and baseline mtGCaMP levels (Figures S6A and S6B). We confirmed that injury causes a cytosolic Ca$^{2+}$ ([Ca$^{2+}]_c$) transient (Figure 7B) [32], with no significant difference between wild-type, car-1(0), mcu-1(0), and micu-1(0) mutants (data not shown).

In wild-type PLM axons, injury triggered a rapid increase in mtGCaMP fluorescence, consistent with axonal $[Ca^{2+}]_{mt}$ uptake (Figures 7B and 7C). We measured the relative change in mtGCaMP fluorescence ($\Delta F/F_0$) and found that the increased mtGCaMP fluorescence peaked within seconds of injury followed by decay to baseline levels after 5 min (Figures 7B–7D). Loss of function in mcu-1 abolished the axotomy-triggered mtGCaMP transient (Figure 7B). In contrast, micu-1(0) PLM axons showed a higher peak mtGCaMP fluorescence immediately after injury but decayed faster, and by 5 min after injury,
had mtGCaMP levels similar to wild-type animals (Figure 7B). The increase in mtGCaMP fluorescence observed in wild-type and micu-1(0) axons after injury was dependent on MCU-1, because mcu-1(0) micu-1(0) double mutants did not display any transient mtGCaMP increase following axonal injury, resembling mcu-1 single mutants (Figure 7B). Our results show that MCU-1 is essential for the transient [Ca^{2+}]_{mt} uptake after axon injury, whereas MICU-1 appears to inhibit initial Ca^{2+} overload, consistent with the gatekeeping function of mammalian MICUs [30, 33, 34].

**car-1(0) Mutants Display More Sustained Axonal [Ca^{2+}]_{mt} Uptake**

We further addressed whether the axotomy-triggered [Ca^{2+}]_{mt} influx was regulated by CAR-1 by examining [Ca^{2+}]_{mt} uptake in car-1(0) animals. The mtGCaMP transients triggered immediately (2.4 s after axotomy) by injury in car-1(0) were similar to those in wild-type axons (Figures 7B and 7C). However, car-1(0) mutants displayed a more sustained elevation in [Ca^{2+}]_{mt} levels compared to wild type; this elevation was completely dependent on MCU-1 (Figures 7B–7D). As expression of MICU-1 was elevated in car-1(0) mutants (Figure 6), we next tested whether loss of micu-1 was able to restore wild-type [Ca^{2+}]_{mt} levels in car-1(0) mutants. We observed wild-type [Ca^{2+}]_{mt} levels in car-1(0) and micu-1(0) double mutants (Figure 7B). Further, slight elevation of micu-1 levels in the TRNs resulted in increased axon regrowth and sustained axonal [Ca^{2+}]_{mt} uptake after injury (Figures S7A–S7C). Axon regrowth was impaired when micu-1 was overexpressed to a high level (Figure S7A), suggesting axon regrowth is sensitive to the level of MICU-1. Overall, our results suggest that mild elevation of MICU-1 expression is responsible for the more sustained [Ca^{2+}]_{mt} uptake and increased axon regrowth of car-1(0) mutants upon axon injury.

**DISCUSSION**

In this study, we find that CAR-1 and other mRNA decay factors play multiple roles in *C. elegans* axon development, axon maintenance, and axon regrowth. We further dissected the mechanism by which CAR-1 inhibits axon regrowth after injury. We presented multiple lines of evidence that the mitochondrial calcium regulator MICU-1 is a major CAR-1 target in axon regrowth. We also show that the regulation of MICU-1 by CAR-1...
is important for fine-tuning injury-induced \([\text{Ca}^{2+}]_{\text{mt}}\) transients. Our data reveal a previously unknown molecular pathway linking mRNA decay and axonal \([\text{Ca}^{2+}]_{\text{mt}}\) dynamics. Our analyses on \(C.\) elegans mRNA decay factors are consistent with an emerging notion that these factors have distinct \textit{in vivo} functions. Focusing on the PLM neurons, we show that mRNA decay factors localize to at least two main subcellular granules, as the decapping enzymes are present in a subset of granules containing translational repressors. These observations are consistent with reports that in \(Drosophila\) neurons, decapping enzymes partly colocalize with the translational repressor Me31B/DDX6 \([5]\). Our data further show that mRNA decapping factors DCAP-1 and DCAP-2 are required for PLM developmental axon guidance and regrowth, whereas translational repressors CAR-1 and CGH-1 are not essential for PLM axon development but inhibit PLM axon regrowth and maintain adult axon integrity. We have found that MICU-1, a member of a highly conserved family of \([\text{Ca}^{2+}]_{\text{mt}}\) regulators, is a major functionally relevant target of neuronal CAR-1. \(miku-1\) transcripts are present in CAR-1-positive, DCAP-1-negative granules, suggesting different mRNA decay factors regulate expression of distinct mRNA targets. Our observation that in

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**Figure 6. CAR-1 Regulates \(miku-1\) Transcript Levels and Represses MICU-1::GFP Protein Levels**

(A) smFISH shows colocalization of \(miku-1\) mRNA with GFP::CAR-1 in the PLM cell body. Cytoplasmic puncta of \(miku-1\) mRNAs (red arrowheads) co-localize with CAR-1 protein puncta in the PLM cell body in wild-type and \(dcap-2(0)\) mutant. See also Figures S4 and S5.

(B) smFISH of \(miku-1\) mRNAs (red arrowheads) shows \(miku-1\) mRNAs form cytoplasmic puncta in PLM neurons, labeled with Pmech-7-GFP (muIs32). More \(miku-1\) mRNA puncta in \(car-1(0)\) mutants and increased accumulation in \(dcap-2(0)\) mutants were observed.

(C) Quantification of \(miku-1\) mRNA puncta in PLM cell bodies of wild-type and \(car-1(0)\) mutants. Representative images shown in (E). \(n \geq 24\) for each strain.

(D) Top shows confocal images of mito::RFP [Pmech-7-tagRFP-mito(jsIs1073)] in the absence of MICU-1::GFP(\(ju1783)\). Bottom shows confocal images showing colocalization of MICU-1::GFP(\(ju1783)\) with mito::RFP in the PLM cell bodies.

(E) Top is an illustration of the region of interest (IntROI) and background (IntB). Bottom images show MICU-1::GFP(\(ju1783)\) in PLM cell bodies (outlined), with increased expression in \(car-1(0)\). Scale bar, 10 \(\mu m\).

(F) MICU-1::GFP intensities in wild type and \(car-1(0)\) animals, normalized to wild-type control. Statistics: unpaired Student’s \(t\) test; *\(p = 0.0194\).
Figure 7. CAR-1 Represses Axonal Mitochondrial Calcium Uptake after Axon Injury

(A) PLM regrowth length 24 h post-axotomy. Statistics: one-way ANOVA with Bonferroni’s post test; *p < 0.05; **p < 0.01. Sample size is indicated in the bar. See also Figures S5, S6, and S7.

(B) mtGCaMP fluorescence intensity (ΔF/F0) over 5 min post-axotomy. Statistics: unpaired t test against wild type; *p < 0.05; **p < 0.01.

(C) Top is an illustration of the region of interest (Int ROI) and background (Int B). Bottom images show mitochondrial calcium imaging trace in wild type, car-1(0), and micu-1(0) mutant axons post-axotomy. Animals expressing the mtGCaMP sensor [Pmec-4-mito-GCaMP5[juIs550]] were severed using femtosecond laser and imaged immediately. Red arrows show the cut site. Yellow arrowheads mark mitochondria in PLM neurons. Some mtGCaMP signal is in the neuronal cytoplasm, likely reflecting incomplete targeting of mtGCaMP due to the high copy number of juIs550. Scale bar, 20 μm.

(D) mtGCaMP fluorescence intensity (ΔF/F0) at 5 min post-axotomy in genotypes indicated. Each dot represents the 5-min mtGCaMP 5 min post-injury in a single animal. Statistics: unpaired t test; **p < 0.01; ***p < 0.001.

(E) Model for regulation of axon regrowth by CAR-1 and [Ca2+]mt uptake. Axon injury induces transient mitochondrial Ca2+ uptake, dependent on MCU-1. Loss of CAR-1 results in elevated micu-1 mRNA expression, leading to more sustained mitochondrial Ca2+ uptake after injury, correlating with increased axon regrowth.
the absence of DCAP-1 or DCAP-2, the mRNA targets of CAR-1 accumulate in larger CAR-1 granules, suggests that increased binding of CAR-1 to its mRNA targets likely leads to translational repression, potentially accounting for the axonal developmental defects and decreased axon regrowth in decapping enzyme mutants. We find that at least two domains of LSM14 family proteins are important for translational repression. The Sm domain is essential for binding to 4E-T, and the FDF domain binds to DDX6/CGH-1 and serves as a platform for other mRNA decay factors [14, 35]. We find that truncated CAR-1(Sm+FDF) protein can suppress the increased micu-1 transcript level of car-1(0) mutants, despite not forming distinct puncta in neurons. Our findings are consistent with prior studies showing the RGG motif is dispensable for translational repression [36, 37]. Moreover, translational repression has been observed in the absence of visible cytoplasmic puncta formation [38, 39]. Taken together, our results suggest that CAR-1 may bind to and repress translation of mRNA targets even in the absence of visible puncta. Through characterizing in vivo mRNA targets of CAR-1 in neurons, we have uncovered a previously unknown pathway involving mitochondria calcium regulation in axon injury. The functions of mitochondrial calcium uniporter MCU complex critically depend on its regulatory subunits, including the MICU proteins and cytoplasmic calcium (reviewed in [40, 41]), and are important for human health. Mutations in human MICU genes have been linked to neurological disorders [42–44], and MCU itself is implicated in excitotoxic neuronal cell death [45]. [Ca2+]mt dynamics affect multiple aspects of cellular metabolism, including oxidative phosphorylation, Ca2+ buffering, and reactive oxygen species production, so abnormal [Ca2+]mt dynamics could affect neuronal development and function in several ways [46]. Emerging evidence links elevated [Ca2+]mt uptake to axonal degeneration in vertebrates [47, 48] and in C. elegans [49]. Moreover, increased Ca2+ levels in traumatic brain injury can be ameliorated by the inhibition of MCU1 [50], suggesting potential therapeutic benefits of mitochondrial Ca2+ regulation in brain injury. Loss of function in Drosophila MCU or MICU causes aberrant axon morphology and impairs memory formation [51]. Interestingly, expression levels of MCU and MICU are dependent on neuronal cell type and Ca2+ signaling [52]. Our results reveal that neuronal MICU-1 levels are fine-tuned by the mRNA decay machinery, adding another level of regulation to neuronal [Ca2+]mt dynamics. Axon injury triggers a transient elevation in [Ca2+]c [17, 32, 53]. In C. elegans neurons, increased [Ca2+]c could contribute to the activation of the conserved kinase DLK-1, thereby promoting axon regrowth [54, 55]. Here, we reveal that axon injury also triggers axonal [Ca2+]mt uptake in C. elegans, reminiscent of observations in vertebrate neurons [56]. We find that this [Ca2+]mt uptake is completely dependent on MCU-1 and is regulated in complex ways by MICU-1. In cultured mammalian cells, MICU1 localizes to mitochondria [31, 57–59] and inhibits MCU function at low [Ca2+]c levels (known as “gatekeeping”), but at higher [Ca2+]c levels, Ca2+ binds to MICU1 to stimulate MCU1 opening [33, 34, 59]. MICU1 therefore can inhibit or promote MCU function depending on the [Ca2+]c level. Supporting this model, our data show that loss of function in micu-1 and mcu-1 have opposing effects on the amplitude of the initial [Ca2+]mt transient after axotomy. However, the enhanced axon regrowth of car-1(0) mutants is suppressed to a similar degree by loss of function in mcu-1 or micu-1. Our data suggest that the MCU-1-stimulating role of MICU-1 at lower [Ca2+]c is most relevant to the long-term outcome in axon regrowth. MCU function is sensitive to the stoichiometry of MICU and other regulatory subunits. For example, liver mitochondria have a higher MICU1:MCU1 ratio than heart or muscle mitochondria and display increased cooperativity of [Ca2+]mt uptake [50]. We propose that in PLM neurons, CAR-1 tightly regulates MICU-1 expression via mRNA decay machinery. car-1(0) mutants have a higher MICU-1:MCU-1 ratio, causing increased [Ca2+]mt uptake after axon injury and enhanced axon regrowth (Figure 7E). Axonal transport of mitochondria has been shown to boost regrowth in C. elegans and mammalian neurons [24–26]. Our observations suggest that more sustained [Ca2+]mt uptake also promotes axon regrowth, for example potentially through increasing ATP production required for axon repair [61]. We speculate that the mild enhancement of [Ca2+]mt uptake resulting from elevated MICU expression is insufficient to trigger deleterious consequences such as opening of the mitochondrial permeability transition pore that would lead to axonal degeneration. Precise modulation of [Ca2+]mt uptake in injured neurons may provide new therapeutic routes toward effective axon repair.

STAR METHODS

Detailed methods are provided in the online version of this paper and include the following:

- KEY RESOURCES TABLE
- LEAD CONTACT AND MATERIALS AVAILABILITY
- EXPERIMENTAL MODEL AND SUBJECT DETAILS
- METHOD DETAILS
  - C. elegans genetics
  - Molecular cloning
  - Transgenic strain construction
  - CRISPR-mediated deletion alleles and GFP knock-in
  - Fluorescence microscopy and laser axotomy
  - Single-end enhanced crosslinking and immunoprecipitation (seCLIP)
  - Single molecule fluorescence in situ hybridization (smFISH)
- QUANTIFICATION AND STATISTICAL ANALYSIS
- DATA AND CODE AVAILABILITY

SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10.1016/j.cub.2019.12.061.

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AUTHOR CONTRIBUTIONS

N.H.T., Y.J., and A.D.C. conceived the project. N.H.T. performed all experiments unless otherwise specified. K.W.K. performed smFISH experiments in Figures 1A and S1. S.X. generated micr-1(0) mutants. S.B., B.A.Y., and G.W.Y. contributed to seCLIP analyses. N.H.T., Y.J., and A.D.C. analyzed data and wrote the manuscript, with input from all authors.

DECLARATION OF INTERESTS

G.W.Y. is a co-founder and member of the Board of Directors, on the SAB, an equity holder, and a paid consultant for Locana and Eclipse BioInnovations. G.W.Y. is a visiting professor at the National University of Singapore, with the terms of this arrangement reviewed and approved by the University of California, San Diego in accordance with its conflict of interest policies. The other authors declare no competing financial interests.

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### STAR METHODS

#### KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| Antibodies          |        |            |
| GFP Trap            | ChromoTek | Cat#gtm-20; AB_2631359 |
| **Bacterial and Virus Strains** |        |            |
| E. coli: OP50-1     | Caenorhabditis Genetics Center | RRID:WB-STRAIN:OP50-1 |

| **Chemicals, Peptides, and Recombinant Proteins** |        |            |
| Protein: Cas9-NLS purified protein | QB3 MacroLab, UC Berkeley | N/A |
| Stellaris RNA FISH Hybridization buffer | LGC BioSearch Technologies | Cat#SMF-HB1-10 |
| Stellaris RNA FISH Wash Buffer A | LGC BioSearch Technologies | Cat#SMF-WA1-60 |
| Stellaris RNA FISH Wash Buffer B | LGC BioSearch Technologies | Cat#SMF-WB1-20 |
| ProLong Antifade Mountant with DAPI | Invitrogen | Cat#P36935 |
| TRizol Reagent | Invitrogen | Cat#15596026 |
| DreamTaq DNA polymerases | Thermo Scientific | Cat#EP0705 |
| Phusion High-Fidelity DNA polymerases | Thermo Scientific | Cat#F530L |
| Protease Inhibitor Cocktail Tablet | Roche | Cat#05892970001 |

| **Critical Commercial Assays** |        |            |
| Gateway LR Clonase II Enzyme Mix | Thermo Scientific | Cat#11791-100 |
| Gibson Assembly Master Mix | New England BioLabs | Cat#E2611S |

| **Deposited Data** |        |            |
| Raw and Analyzed Data | This paper | GEO: GSE124714 |

| **Experimental Models: Organisms/Strains** |        |            |
| C. elegans: Strain wild type N2 | Caenorhabditis Genetics Center | RRID:WB-STRAIN:N2_(ancestral) |
| C. elegans: Strain Pmec-7-GFP(muls32) II | [19] | CZ10969 |
| C. elegans: Strain Pmec-4-GFP(zdl5) I | [18] | CZ10175 |
| C. elegans: Strain Pmec-17-GFP(uls31) III | [62] | CZ25265 |
| C. elegans: Strain zdl55 I; [Pdcap-1-DCAP-1::dsRed+rol-6 (su1006)])/bpl37 | This paper | CZ26592 |
| C. elegans: Strain Pmec-7-mRFP(isls973) III; Pcg-1-CGH-1::GFP(dhls1000) | This paper | CZ26244 |
| C. elegans: Strain Pcg-1-CGH-1::GFP(dhls1000); Pmc4-mKate2::CAR-1( juEx7739) | This paper | CZ25819 |
| C. elegans: Strain Pmec-4-4-GFP::CAR-1( juSi338) IV; [Pdcap-1-DCAP-1::dsRed+rol-6 (su1006)])/bpl37 | This paper | CZ25293 |
| C. elegans: Strain car-1(tm1753) I/hT2 [bli-4(e937) I let-7(q782) GFP(qls48) I;fl]; muls32 II | This paper | CZ11541 |
| C. elegans: Strain zdl55 I; dcap-2(tm2470) IV | This paper | CZ25658 |
| C. elegans: Strain zdl55 I; cgh-1( ok492) III/hT2 [bli-4(e937) I let-7(q782) GFP(qls48) I;fl] | This paper | CZ11540 |
| C. elegans: Strain zdl55 I; dcap-1( ok2139) IV | This paper | CZ11537 |
| C. elegans: Strain zdl55 I; dcap-1( ok2139) dcap-2(tm2470) IV | This paper | CZ27317 |
| C. elegans: Strain zdl55 I; patr-1(tm2402) II/ min1 [mils14 dpy-10(e128) II] | This paper | CZ11759 |
| C. elegans: Strain car-1(tm1753) I/hT2 [bli-4(e937) I let-7(q782) GFP(qls48) I;fl]; uls31 III | This paper | CZ25319 |
| C. elegans: Strain muls32 II; Pmec-4-GFP::CAR-1( juEx7280) | This paper | CZ23991 |

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| C. elegans: Strain car-1(ju1505) I/hT2 [bli-4(e937) I let-?(q782) GFP(qIs48) I;III]; mels2 II | This paper | CZ25262 |
| C. elegans: Strain car-1(ju1506) I/hT2 [bli-4(e937) I let-?(q782) GFP(qIs48) I;III]; mels2 II | This paper | CZ25264 |
| C. elegans: Strain car-1tm1753 I/hT2 [bli-4(e937) I let-?(q782) GFP(qIs48) I;III]; mels2 II; loxP-Pcar-1-GFP::car-1g+3' UTR-loxP [juSi343] IV | This paper | CZ25401 |
| C. elegans: Strain car-1tm1753 I/hT2 [bli-4(e937) I let-?(q782) GFP(qIs48) I;III]; Pmyo-3-car-1(juEx7913) | This paper | CZ26723 |
| C. elegans: Strain zdIs5 I; Prgef-1-GFP::car-1(juIs526) | This paper | CZ25296 |
| C. elegans: Strain zdIs5 I; Prgef-1-GFP::car-1(ΔSm) (juIs549) | This paper | CZ25884 |
| C. elegans: Strain dcap-2(tm2470) IV; Pmec-4-mito::GCaMP5(juIs550) | This paper | CZ25566 |
| C. elegans: Strain dcap-2(tm2470) IV; Pmec-4-mito::GCaMP5(juIs550) | This paper | CZ25567 |
| C. elegans: Strain Pmec-4-GFP::CAR-1(Sm) (juSi346) IV | This paper | CZ25444 |
| C. elegans: Strain Pmec-4-mito::GCaMP5(juIs550) | This paper | CZ25938 |
| C. elegans: Strain Pmec-7-mRFP::mito(jsIs1073) | This paper | CZ26634 |
| C. elegans: Strain Pmec-7-mRFP::mito(jsIs1073) | This paper | CZ26429 |
| C. elegans: Strain dcap-2(tm2470) IV; Pmec-4-GFP::CAR-1(ΔSm) [juSi361] IV | This paper | NM3573 |
| C. elegans: Strain Pmec-7-mRFP::mito(jsIs1073) | This paper | CZ27549 |
| C. elegans: Strain Pmec-7-mRFP::mito(jsIs1073) | This paper | CZ27519 |
| C. elegans: Strain Pmec-7-mRFP::mito(jsIs1073) | This paper | CZ27538 |
| C. elegans: Strain dcap-2(tm2470) IV; Pmec-4-GFP::CAR-1(ΔSm) [juSi361] IV | This paper | CZ20877 |
| C. elegans: Strain dcap-2(tm2470) IV; Pmec-4-GFP::CAR-1(ΔSm) [juSi361] IV | This paper | CZ26661 |
| C. elegans: Strain Pmec-7-mRFP::mito(jsIs1073) | This paper | CZ25925 |
| C. elegans: Strain muIs32; car-1(tm1753) I/hT2 [bli-4(e937) I let-?(q782) GFP(qIs48) I;III]; mels2 II; micu-1(ju1154) IV | This paper | CZ27010 |
| C. elegans: Strain Pmec-7-mRFP::mito(jsIs1073) III; Pmec-4-mito::GCaMP5(juIs550) | This paper | CZ26629 |
| C. elegans: Strain car-1(tm1753) I/hT2 [bli-4(e937) I let-?(q782) GFP(qIs48) I;III]; Pmec-7-mRFP::mito(jsIs1073) III; Pmec-4-mito::GCaMP5(juIs550) | This paper | CZ26687 |
| C. elegans: Strain Pmec-7-mRFP::mito(jsIs1073) III; Pmec-4-mito::GCaMP5(juIs550) | This paper | CZ26857 |

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| **C. elegans**: Strain Pmec-7-mRFP[sls973] III; mcu-1[Ju1154] micu-1[Ju1156] IV/nT1 [qsl51 IV;V]; Pmec-4-mito::GCaMP5[sls550] | This paper CZ26858 |
| **C. elegans**: Strain car-1(tm1753) I/hT2 [bli-4(e937) I let-7(q782) GFP[sls48] I;II]; Pmec-7-mRFP[sls973] III; mcu-1[Ju1154] IV; Pmec-4-mito::GCaMP5[sls550] | This paper CZ27060 |
| **C. elegans**: Strain Punc-17::GFP[sls48]; [Pdcap-1-DCAP-1::dsRed+rol-6(su1006)][bpl37] | This paper CZ26594 |
| **C. elegans**: Strain zdIs5 I; dcap-1(ok2139) IV; Pmec-4-GFP::CAR-1[juSi338] IV | This paper CZ27378 |
| **C. elegans**: Transgenic allele: N2 injected with 10 ng/μl of pCZGY2860 [Pmec-4-GFP::car-1(genomic)::let-858_3 UTR] and 5 ng/μl pCFJ90 [Pmyo-2-mCherry]. | This paper juEx7280 |
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| **C. elegans**: Transgenic allele: N2 injected with 10 ng/µl pCZGY372 [Pmyo-3-car-1(cDNA)::let-858_3'UTR] and 3 ng/µl pCFJ90[Pmyo-2-mCherry]. | This paper juEx7913 |
| **C. elegans**: Transgenic allele: N2 injected with 10 ng/µl pCZGY3411 [Pmec-4-GFP::dcap-1(cDNA)::let-858_3'UTR] and 3 ng/µl pCFJ90 [Pmyo-2-mCherry]. | This paper juEx8024 |
| **C. elegans**: Transgenic allele: N2 injected with 2 ng/µl pCZGY3416 [Pmec-4-micu-1-gfp] and 3 ng/µl pCFJ90 [Pmyo-2-mCherry]. | This paper juEx8048 |
| **C. elegans**: Single-copy transgenic allele: N2 injected with 50ng/µl of pCZGY3366 [Pcar-1(1062bp upstream)-GFP::car-1(genomic+768bp 3'UTR)]. | This paper juEx8050 |
| **C. elegans**: Single-copy transgenic allele: N2 injected with 50ng/µl of pCZGY3362 [Pmec-4-GFP::car-1(cDNA)::let-858_3'UTR]. | This paper juSi338 |
| **C. elegans**: Single-copy transgenic allele: N2 injected with 50ng/µl of pCZGY3367 [Pmec-4-GFP::car-1(Sm)::let-858_3'UTR]. | This paper juSi346 |
| **C. elegans**: Multi-copy transgenic allele: CZ20310 [Pmex-5-his-72::miniSOG(juSi164) III unc-119(ed3) III] injected with 10 ng/µl pCZGY3363 [Prgef-1-GFP::car-1(genomic)::let-858_3'UTR] and 3 ng/µl pCFJ90 [Pmyo-2-mCherry]. | This paper juIs526 |
| **C. elegans**: Multi-copy transgenic allele: N2 injected with 40 ng/µL of Pmec-4-mito::GCaMP5 and treated with TMP/UV. | This paper juIs550 |
| **C. elegans**: GFP knock-in allele: N2 injected with 50 ng/µl of pCZGY3412, 50 ng/µl of pCZGY3413, 10 ng/µl of pCZGY3414, 3 ng/µl Pmyo-2-mCherry and 3 ng/µl of Pmyo-3-mCherry. Strains were treated with heat shock as described in Dickinson et al., 2015. | This paper ju1783 |

Oligonucleotides

Alt-R® CRISPR-Cas9 crRNA for car-1: to make car-1(ju1505) and car-1(ju1506) deletion: /AltR1/CrArCrG rUrCrG rArCrA rUrCrC rGrUrU rArCrG rGrUrU rUrUrArCrA rGrArG rCrUrA rUrGrC rUr/AltR2/ This paper N/A

Alt-R® CRISPR-Cas9 crRNA for car-1: to make car-1(ju1505) and car-1(ju1506) deletion: /AltR1/rUrCrG rArArU rCrUrU rGrCrC rGrUrU rArCrA rGrUrA rGrUrU rUrUrArCrA rGrArG rCrUrA rUrGrC rUr/AltR2/ This paper N/A

Alt-R® CRISPR-Cas9 crRNA for micu-1: to make micu-1(ju1155) and micu-1(ju1156) deletion: /AltR1/rGrGrG rCrGrG rUrUrU rCrUrC rGrCrG rArCrG rGrUrU rUrUrArGrG rCrUrA rUrGrC rUr/AltR2/ This paper N/A

Alt-R® CRISPR-Cas9 crRNA for micu-1: to make micu-1(ju1155) and micu-1(ju1156) deletion: /AltR1/rUrCrG rGrGrG rCrGrG rUrUrU rCrUrC rGrCrG rArCrG rGrUrU rUrUrArGrG rCrUrA rUrGrC rUr/AltR2/ This paper N/A

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LEAD CONTACT AND MATERIALS AVAILABILITY

Further information and requests for reagents should be directed to and will be fulfilled by the Lead Contact, Andrew Chisholm (adchisholm@ucsd.edu). All unique/stable reagents generated in this study are available from the Lead Contact without restriction.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

The nematode *Caenorhabditis elegans* was used as the experimental model for this study. All experiments were performed with hermaphrodite animals; males were used only for crosses. *C. elegans* strains were maintained at 20°C on nematode growth medium (NGM) agar plates seeded with *E. coli* OP50.

METHOD DETAILS

*C. elegans* genetics

car-1(0), cgh-1(0) and patr-1(0) mutants display maternal effect lethality, and micu-1(0) mutants display sterility. There mutants were kept as balanced strains. Homozygous car-1(0), cgh-1(0), patr-1(0) and micu-1(0) mutants produced from heterozygous mothers were analyzed. car-1(0) and micu-1(0) mutants do not display overt behavioral or development phenotypes. We used the following transgenes: Pmec-7-GFP(muls32), Pmec-4-GFP(zdls5), Pmec-17-GFP(uls31) and Pmec-7-mRFP(jsls973) to visualize touch receptor neurons (TRNs), Punc-17-mCherry(nuls32t) and Punc-17-GFP(vsls48) to visualize cholinergic motor neurons, and

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| Alt-R CRISPR-Cas9 tracrRNA | IDT | Cat#1072532 |

Recombinant DNA

| Plasmid: Pmec-4-GFP::car-1(genomic)-let-858_3'UTR | This paper | pCZGY2860 |
| Plasmid: FRT-Hygro-FRT::car-1(cDNA)-let-858_3'UTR for CRISPR | This paper | pCZGY3362 |
| Plasmid: Pmec-4-GFP::car-1(cDNA)-let-858_3'UTR | This paper | pCZGY3363 |
| Plasmid: Pmec-4-GFP::car-1(cDNA)-let-858_3'UTR | This paper | pCZGY3364 |
| Plasmid: FRT-Hygro-FRT::car-1loxP-car-1loxP for CRISPR | This paper | pCZGY3365 |
| Plasmid: FRT-Hygro-FRT::car-1(1062bp upstream)::GFP::car-1(genomic+768bp 3'UTR) for CRISPR | This paper | pCZGY3366 |
| Plasmid: FRT-Hygro-FRT::car-1(Sm)-let-858_3'UTR for CRISPR | This paper | pCZGY3367 |
| Plasmid: FRT-Hygro-FRT::car-1(Sm)-let-858_3'UTR for CRISPR | This paper | pCZGY3368 |
| Plasmid: FRT-Hygro-FRT::car-1(JRGG)-let-858_3'UTR for CRISPR | This paper | pCZGY3369 |
| Plasmid: FRT-Hygro-FRT::car-1(RGG)-let-858_3'UTR for CRISPR | This paper | pCZGY3370 |
| Plasmid: FRT-Hygro-FRT::car-1(FDF+RGG)-let-858_3'UTR for CRISPR | This paper | pCZGY3371 |
| Plasmid: Pmec-4-GFP::dcap-1(cDNA)-let-858_3'UTR | This paper | pCZGY3411 |
| Plasmid: Peft-3::Cas9+micu-1 C1 sgRNA | This paper | pCZGY3412 |
| Plasmid: Peft-3::Cas9+micu-1 C2 sgRNA | This paper | pCZGY3413 |
| Plasmid: GFP-SEC-3xFLAG_micu-1 C1 homology_arm | This paper | pCZGY3414 |
| Plasmid: Pmec-4-micu-1 gfp | This paper | pCZGY3416 |

Software and Algorithms

| ImageJ | NIH image | RRID:SCR_003070 |
| GraphPad Prism 7 | GraphPad Software, Inc. | RRID:SCR_002798 |
| ZEN | Zeiss | https://www.zeiss.com/microscopy/us/downloads/zen.html |
Molecular cloning
To generate car-1 rescue constructs, we cloned ~3kb of car-1 genomic DNA into pCR8 vector. A full-length car-1 cDNA was generated by RT-PCR from wild type N2 total RNA. Expression constructs for tagged or truncated CAR-1 were generated from cDNA by PCR and cloned into destination vectors containing Pmec-4 or Pmyo-3 promoter using the Gateway system (Life Technologies). We expressed the Ca2+ sensor GCaMP [64] to the mitochondrial matrix (mGCaMP) in TRNs using the mec-4 TRN-specific promoter. We used Gibson assembly (New England Biolabs) to generate clones for single-copy transgenes.

Transgenic strain construction
For standard extrachromosomal transgenes, plasmid DNAs were used at 10 ng/μl, co-injection marker Ptx-3-RFP at 90 ng/μl or Pmyo-2-mCherry at 3-5 ng/μl. At least 2 independent transgenic lines were analyzed per construct. Single-copy transgenes were inserted into the transposon site exTi5882 on chromosome IV using CRISPR/Cas9. Three plasmids were injected into wild type hermaphrodites: a plasmid encoding HgK-sGFP-car-1-cDNA flanked by homology arms, a plasmid encoding Cas9-sgRNA, and a plasmid encoding co-injection marker. Animals were selected for genomic insertion based upon resistance to hygromycin (HgK), the absence of co-injection markers and transgene was confirmed by PCR genotyping. We used optogenetic mutagenesis to generate chromosomal integration of pan-neuronal GFP-CAR-1 constructs [65]. Briefly, plasmids were injected into C203010 [Pmex-5:His::miniSOG/ juIs164; unc-19(ed3)] L4 hermaphrodites, and 6 h after injection, these animals were exposed to blue light for 30 min at 4 Hz. F1 progeny with co-injection marker were singled onto seeded plates. F2 progeny from F1 plates with > 75% transmission were singled, and those showing 100% transmission of the co-injection maker in the following generation were outcrossed with wild type males to confirm chromosomal integration. We also determined by quantitative PCR the copy number of the expression construct in juIs526 and juIs549 to be 52 and 78 copies respectively. Pmec-4-mito-GCaMP5/juIs550 was integrated using UV-trimethylpsoralen and outcrossed 4 times before use.

CRISPR-mediated deletion alleles and GFP knock-in
We generated car-1 deletions using two CRISPR RNA (crRNAs), 5’-CTAATCTTGGTCTCGATGTA-3’ and 5’-CAAGCTCGAC ATCGTATTACG-3’ (Integrated DNA Technologies) targeting the coding sequence in the first exon of car-1. The crRNAs were injected into wild type hermaphrodites with purified Cas9 (MacroLabs, University of California, Berkeley), trans-activating crRNA (tracrRNA) and dpy-10 crRNA, as described [66]. Among 147 Dpy F1 worms, we identified two worms heterozygous for car-1 deletions ju1505 and ju1506. Like car-1(tm1753), eggs laid by homozygous car-1(ju1505) and car-1(ju1506) worms failed to hatch, indicating all three alleles are null mutants. car-1 alleles were balanced over hT2 I; III [bii-4(e937) I let-7(q782) GFP/qs48] III; [67].

We generated micu-1 deletion alleles as described [67]. We designed two subgenomic RNAs (sgRNAs): GTGG GCCGTTTCTCCGCAAGAC and ATCGATCCTAACGAAGAC. pU6::micu-1 sgRNAs were generated by primer annealing and injected into wild type or mcu-1(ju1154) worms as mixtures of 40 ng/μl of each pU6::micu-1 sgRNA, 100 ng/μl of Pelt-3-Cas9-SV40 NLS::tbb-2’3’UTR, and 20 ng/μl of Pcol-19-GFP as coinjection marker. We obtained worms heterozygous for mcu-1(ju1155) in N2 background and mcu-1(ju1156) in mcu-1(ju1154) background. mcu-1 alleles were balanced over n71 IV; V [qs151] or tmC9 [ln(gib::19 lgc:52 ln(mec-3 unc-31) i) IV [68].

We inserted GFP at the C terminus of mcu-1 using the self-excising drug selection cassette [69]. We designed two subgenomic RNAs (sgRNAs): CTATAAAAATGGAAGAGGAC and ATTGCCACTGTGAAATCTTG. We injected 50 ng/μl of each mcu-1 sgRNA, 10 ng/μl of homology arm repair template, 3 ng/μl Pmyo-2-mCherry and 3 ng/μl of Pmyo-3-mCherry into wild type worms. New alleles are in Table S2.

Fluorescence microscopy and laser axotomy
Fluorescence images were collected using Zeiss LSM510, LSM710, or LSM800 confocal microscopes, unless specified. Laser axotomy was performed as described [16, 18]. Images shown were projected as z stack. For live imaging of GCaMP fluorescence, we collected images every 240 ms using the spinning-disk confocal. Femtosecond laser power was 140 to 180 mW with a shutter time of 1.2–1.5 ms. The mitochondrion closest to the axotomy site in the proximal axon was set as the region of interest (ROI). We outlined the mitochondrion and measured its integrated density using ImageJ and subtracted the integrated density from background using an identical outline within the axon to obtain net fluorescence (F). Baseline fluorescence (F0) was obtained from the ROI one frame before axotomy. The change in fluorescence ΔF was expressed as the ratio of change with respect to the baseline.

Single-end enhanced crosslinking and immunoprecipitation (seCLIP)
We performed single-end enhanced crosslinking and immunoprecipitation (seCLIP) as previously described [21] with a few modifications. Worms expressing pan-neuronal GFP::CAR-1 [Prgef-1-GFP::CAR-1(juIs527)] or pan-neuronal GFP::CAR-1(ΔSm) [Prgef-1-GFP::CAR-1(ΔSm)/juIs549] were UV-crosslinked at 3 kJ/m² then resuspended in lysis buffer [50 mM Tris-HCl pH7.4, 100 mM NaCl, 1% NP-40, 0.1% SDS, 0.5% sodium deoxycholate, supplemented with one tablet of protease inhibitor cocktail tablet (Roche) in 20 ml]. Subsequent steps were performed as described [21]. Immunoprecipitation was performed using GFP-Trap (ChromoTek). Washes were performed using lysis buffer instead of high salt wash buffer. The quality of sequencing reads from fastq
files was evaluated by FastQC (http://www.bioinformatics.babraham.ac.uk/projects/fastqc). Bioinformatic analysis of sequencing reads was performed as previously described [22].

43,905,923 and 64,702,729 reads were obtained from GFP::CAR-1(FL) and GFP::CAR-1(ΔSm) sample respectively, of which 4,022,150 [CAR-1(FL)] and 5,470,606 [CAR-1(ΔSm)] reads uniquely mapped to C. elegans genome, after removing PCR duplicates and repeats. We identified 1190 genes with peaks > 2x enriched in CAR-1(ΔSm) over CAR-1(FL) (Table S1) using CLIPper peak calling [70]. Fold enrichment was calculated using the number of eCLIP reads overlapping CLIPper-identified peaks, after normalizing to total usable read counts in each sample. We also analyzed fastq files using HISAT2 [71], followed by transcript assembly using Cufflinks [72] and differential analysis using Cuffdiff (p value < 0.05) [73]. 29 genes were enriched in CLIPper and in Cufflinks-Cuffdiff analyses (Table S1). We inspected all genes enriched in CAR-1(ΔSm) and focused on micu-1, which showed 2x Peak enrichment in exon 7 at ChrIV:13299691-13299814.

Single molecule fluorescence in situ hybridization (smFISH)

Forty-eight 20-nt probes for car-1 or micu-1 mRNAs were designed and purchased from LGC BioSearch Technologies and conjugated to Cal Fluor 610. We performed smFISH using mixed stage animals according to a modified Stellaris protocol [74]. We used Pmec-7-GFP(muIs32) or Pmec-4-GFP(zdIs5) animals expressing GFP in TRNs as a reference for mRNA localization in PLM neurons. For imaging we used Leica DMi8 microscope, Andor spinning disk confocal (CSU-W1) and iXon ultra 888 EMCCD camera. All images shown are z stack projections.

QUANTIFICATION AND STATISTICAL ANALYSIS

Statistical tests were performed using GraphPad Prism. Two-way comparisons of categorical data were performed using Fisher’s exact test. For multiple comparisons, we used one-way ANOVA with Bonferroni’s post test. All data compared using unpaired Student’s t test passed the D’Agostino & Pearson omnibus normality test. To compare regrowth between experiments with different control means, we normalized each experimental data point by dividing it by its control means. Sample sizes are indicated in Figures (dot plots) or in Figure legends.

DATA AND CODE AVAILABILITY

The raw and analyzed seCLIP datasets have been deposited at the Gene Expression Omnibus (accession number: GSE124714).