The kpc-1 (furin) 3’UTR promotes dendritic transport and local translation of mRNAs to regulate dendrite branching and self-avoidance of a nociceptive neuron

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Summary

Mechanical stimuli on the skin of *C. elegans* are detected by the dendritic arbors of PVD nociceptive neurons, which provide uniform sensory coverage outside the head region across the entire animal. Through genetic screens, we isolate three mutants that display profound dendrite self-avoidance defects in PVD neurons. Studying dendrite self-avoidance in *C. elegans* is likely to provide new mechanistic insight into the process as the well-known self-avoidance molecule Dscam is absent from the *C. elegans* genome. Through whole genome sequencing, we identify the responsible mutations in the *kpc-1* gene. Compared to wild-type animals, a strong *kpc-1* mutant allele exhibits secondary dendrite branching defects whereas a weak *kpc-1* mutant allele displays tertiary dendrite self-avoidance defects. Here, we show that the *kpc-1* 3’UTR is required for *kpc-1*’s functions in both dendrite branching and dendrite self-avoidance. The *kpc-1* 3’UTR facilitates *kpc-1* RNA localization to branching points and contact points between sibling dendrites. Using fluorescence recovery after photoconversion, we show that the *kpc-1* 3’UTR promotes local protein synthesis in the distal segment of PVD dendrites. We identify an important secondary structural motif in the *kpc-1* 3’UTR required for tertiary dendrite self-avoidance. We demonstrate that over-expression of *kpc-1* leads to greater self-avoidance without limiting initial dendrite outgrowth, supporting a direct role of *kpc-1* in self-avoidance. Animals with *dma-1* receptor over-expression display similar secondary dendrite branching and tertiary dendrite self-avoidance defects that are suppressed with *kpc-1* over-expression, which suggests that DMA-1 is a potential KPC-1 target that is down-regulated by KPC-1. Our results support a model where KPC-1 proteins are synthesized at branching points and contact points between neighboring dendrites to
locally down-regulate DMA-1 receptors to promote dendrite branching and self-avoidance. A recently reported Schizophrenia-associated genetic variant in the 3’UTR of the human furin gene, a homolog of kpc-1, highlights the important role of the kpc-1 (furin) 3’UTR in neuronal development, which is further demonstrated by this mechanistic study.
Introduction

PVD nociceptive neurons innervate *C. elegans* skin with elaborate dendrite arbors that respond to harsh mechanical stimuli\(^1\). Dendrite arborization allows uniform sensory coverage of PVD neurons across the entire surface of the animal outside the head region. However, the mechanism underlying this process is still unknown.

The elaborate branching patterns displayed by PVD dendrites are constrained by two organizing principles: Escape from the intermediate target (the trap zone) upon arrival at it and self-avoidance\(^2-4\). Self-avoidance is a process in which self-recognition among neurites of a single neuron is followed by their repulsion from each other. This selective repulsion between dendrites of the same cell was originally observed in leech neurons and proposed to promote a uniform receptive field\(^5,6\). Recent studies using *Drosophila* as a model showed that homophilic binding of Dscam1 proteins on sister branches of the same cell (MB or da neurons) promotes repulsive interactions between them, to ensure that they diverge and grow along separate pathways\(^7-11\). Since the Dscam1 mutation does not completely disrupt dendrite self-avoidance in *Drosophila* and there is no Dscam1 homolog in *C. elegans*, studying PVD dendrite self-avoidance in *C. elegans* provides a unique opportunity to identify novel mechanisms of dendrite self-avoidance. Indeed, several known developmental pathways, including UNC-6 (netrin), MIG-14 (Wntless), and FMI-1 (Flamingo) signaling, have been implicated in PVD dendrite self-avoidance in *C. elegans*\(^4,12,13\).

It was previously reported that guidance cues SAX-7 and MNR-1 generated in the hypodermis established dendrite growth pathways and instructed dendrite arborization for PVD neurons in *C. elegans*\(^14,15\). PVD neurons first grow two longitudinally extending 1°
dendrites along the anterior-posterior (A/P) axis. Orthogonal arrays of 2°, 3°, and 4° dendritic branches emerge sequentially in a manner that alternates between the dorsal-ventral (D/V) and the A/P axes to produce an elaborate network of sensory processes\textsuperscript{3,16} (Fig. 1B). Skin cues, SAX-7 and MNR-1, and a muscle-derived cue, LECT-2, are required for dendrite pattern formation through direct interactions with the dendrite receptor DMA-1 to form a multi-protein receptor-ligand complex\textsuperscript{14,15,17–19}.

We isolated mutants through genetic screens that displayed profoundly affected dendrite self-avoidance in PVD neurons. We identified mutations in the \textit{kpc-1} gene as being responsible for this defective self-avoidance phenotype through whole genome sequencing. \textit{kpc-1} mutants displayed a significantly higher number of secondary dendrites, a significantly lower tertiary branch avoidance index, and decreased sensory coverage in the skin compared to wild-type animals. Although \textit{kpc-1} has been implicated in dendrite self-avoidance and arborization, the mechanism by which \textit{kpc-1} regulates these processes is unknown\textsuperscript{2,20,21}. KPC-1 encodes a proprotein convertase subtilisin/kexin (PCSK). Here, we show that \textit{kpc-1} functions in PVD neurons to regulate dendrite branching and self-avoidance via the intact \textit{kpc-1} 3’UTR. The \textit{kpc-1} 3’UTR is required for \textit{kpc-1} RNA localization to distal and higher-order dendrites and promotes local protein synthesis. Although over-expression of \textit{kpc-1} causes greater self-avoidance, it does not limit initial dendrite outgrowth, which supports a direct role for \textit{kpc-1} in self-avoidance. \textit{dma-1} over-expression also displays similar secondary dendrite branching and tertiary dendrite self-avoidance phenotypes that are suppressed with \textit{kpc-1} over-expression. Thus, DMA-1 is a potential KPC-1 target that becomes down-regulated with increased KPC-1 activity. Our results suggest a model in which KPC-1 is translated at branching points and contact points.
between sibling dendrites, leading to local down-regulation of DMA-1 receptors and the consequent dendrite branching and self-avoidance.
Results

Isolation of mutants that display profound self-avoidance defects in dendrites of PVD neurons

*C. elegans* PVD neurons extend a large and highly branched dendrite arbor directly beneath the hypodermal surface that envelops the worms to mediate an avoidance response to harsh mechanical force as well as to other noxious stimuli including temperature\(^1\). The elaborate dendrite branching patterns that PVD neurons display are constrained by two organizing principles: dendrite escape from the trap zone\(^2\) and dendrite self-avoidance. Self-avoidance is an important neuronal process in which dendritic branches arising from a single soma (also called isoneuronal or sister branches) turn away from one another to minimize crossing and overlap and to promote a uniform receptive field\(^2,22\).

The self-avoidance mechanism in *C. elegans* is likely to provide new mechanistic insight into the process as the well-known self-avoidance molecule Dscam\(^7-11\) is absent from the *C. elegans* genome. In order to uncover new mechanisms responsible for self-avoidance in *C. elegans*, we screened approximately 9,500 genomes for mutants that displayed self-avoidance defects in secondary and tertiary dendrites. These screens uncovered three strong mutants (*xr47, xr58*, and *xr60*) that show profound self-avoidance defects in all animals examined (n>200 each). *xr47* and *xr60* mutants showed an additional secondary dendrite trapping phenotype (a defect in the secondary dendrite branching), which resulted in significant reduced number of secondary branches in *xr47* and *xr60* mutants compared to wild-type animals (Table 1). Despite displaying extensive defects in self-avoidance among secondary and tertiary sibling dendrites, *xr58* mutants showed little defect in secondary dendrite escape from the trap zone (Fig. 1, a and c; Table 1). In wild-
type animals, secondary dendrites that contact neighboring secondary branches retract in order to form a regularly spaced array of parallel branches (Fig. 1, a and b). In \textit{xr58} mutants, many secondary dendrites remain in contact with each other (Fig. 1c, arrows). In wild-type animals, tertiary dendrites initially extend but later retract once their termini come into contact with sibling tertiary branches (Supplementary S1, arrowheads). In \textit{xr58} mutants, many tertiary dendrites remain in contact with their siblings long after arborization ends (Fig. 1c and Supplementary Fig. 1, arrowheads), resulting in a significantly lower tertiary branch avoidance index (TBAI) compared to the wild-type animals (Fig. 1e; TBAI is 0.38 in \textit{xr58} mutants versus 0.93 in wild-type, p<0.0001). In wild-type animals, PVD dendrite arbors are fully extended beneath the hypodermis and cover the areas from the posterior animal body to right before the head region (Fig. 1f, Supplementary Fig. 2a, 2c, and 2d). In \textit{xr58} mutants, PVD dendrite arbors develop partially overlapping sibling menorahs, causing decreased sensory coverage of skin (Fig. 1f, Supplementary Fig. 2b, 2c, and 2d). We quantified sensory coverage of skin by PVD dendrite arbors using two measurements, receptive field index and length ratio of receptive field (Supplementary Fig. 2c,d). Both measurements show significantly reduced values in \textit{kpc-1(xr58)} mutants compared to the wild-type animals (Supplementary Fig. 2).

**Identification of responsible mutations in the \textit{kpc-1} gene**

We identified \textit{xr47}, \textit{xr58}, and \textit{xr60} mutations in the \textit{kpc-1} gene through whole genome sequencing. The \textit{xr58} allele harbors a missense mutation resulting in a P440S amino acid substitution at a conserved proline residue four amino acids away from the catalytic serine residue toward the C-terminal. It was recently shown by others that \textit{kpc-1(my24)}, a
different mutation in the same residue (P440L), caused a loss of function in the kpc-1 gene which resulted in defective secondary dendrite escape from the trap zone in PVD neurons\textsuperscript{20}. Thus, it appears that the P440 residue can be used to moderately or dramatically alter kpc-1 activity. Using the complementation test, we showed that the kpc-1(gk8) null mutation failed to complement the xr58 mutant phenotype in PVD neurons, further confirming that xr58 is a new kpc-1 allele (Fig. 6d). The xr58 mutant reveals an additional function of kpc-1 in tertiary dendrite self-avoidance as trans-heterozygous mutants (xr58/gk8) display the xr58 mutant phenotype of defective tertiary dendrite self-avoidance rather than defective secondary dendrite escape from the trap zone characteristic of gk8 (Fig. 1c, 2d; n>40). The two other kpc-1 alleles are also predicted to affect KPC-1 protein structure and function. The xr60 allele contains a missense mutation, which results in a G223S amino acid substitution in the catalytic domain, while the xr47 allele contains a G-to-A transition at the +1 position of intron 5, which abolishes the donor splice site (Fig. 1g). Compared to the gk8 null allele, xr58, xr47, and xr60 alleles are weaker in their phenotypic effects in PVD neurons as judged by the extent of menorah coverage and trapped secondary dendrites along the primary dendrite axis (Table 1). kpc-1 mRNAs can be made from xr58, xr47, and xr60 mutant alleles based on successful amplification of respective kpc-1 cDNAs from them (unpublished).

**The kpc-1 3’UTR is required for both secondary dendrite branching and tertiary dendrite self-avoidance**

kpc-1 genomic DNA containing the 5’, coding, and 3’ fragments rescued both the defective secondary dendrite escape from the trap zone and the defective tertiary dendrite self-
avoidance in the kpc-1(gk8) null allele as well as the defective tertiary dendrite self-avoidance in the kpc-1(xr58) allele (Fig. 2, a,b). To investigate the genomic requirement of the kpc-1 gene in different steps of PVD dendrite development, we first reduced the kpc-1 genomic coding region to the kpc-1 cDNA. We found that the kpc-1 cDNA without introns led to a complete rescue of the dendrite phenotypes in both kpc-1(gk8) null and kpc-1(xr58) alleles (Fig. 2, a,b). We then further reduced the kpc-1 promoter to a more PVD cell specific promoter (The F49H12.4 promoter). This Ppvd::kpc-1::kpc-1 3’UTR transgene remained able to cause a near complete phenotypic rescue, in a manner similar to the kpc-1 genomic DNA, in both kpc-1(gk8) null and kpc-1(xr58) alleles, indicating that kpc-1 acts cell-autonomously in PVD and that the kpc-1 3’UTR is required for both secondary dendrite escape (branching) from the trap zone and tertiary dendrite self-avoidance (Fig. 2a,b, and e-h). Consistent with a cell-autonomous role of kpc-1 in PVD, the expression of the kpc-1 promoter reporter was detected in PVD neurons during secondary dendrite branching and tertiary dendrite self-avoidance (Fig. 2c). When the kpc-1 3’UTR was replaced with an unrelated unc-54 3’UTR, the Ppvd::kpc-1::unc-54 3’UTR transgene was able to rescue the defective secondary dendrite escape (branching) from the trap zone but not the defective tertiary dendrite self-avoidance in the kpc-1(gk8) null allele (Fig. 2 a,b,d, arrowheads), suggesting a more stringent 3’UTR requirement for tertiary dendrite self-avoidance. In addition to tertiary dendrites, some secondary dendrites displayed self-avoidance defects in these transgenic animals (Fig. 2d, asterisks). The same Ppvd::kpc-1::unc-54 3’UTR transgene was also unable to rescue the defective tertiary dendrite self-avoidance in the kpc-1(xr58) allele (Fig. 2b). Increasing the concentration of the Ppvd::kpc-1::unc-54 3’UTR transgene from 10 ng/ml to 100 ng/ml did not alter this
observation (data not shown). Thus, these results show that both the secondary dendrite branching and the tertiary dendrite self-avoidance functions of kpc-1 are dependent on its own 3’UTR.

The kpc-1 3’UTR is required for localization of kpc-1 transcripts to distal primary dendrites and higher-order tertiary dendrites in PVD neurons

To directly visualize kpc-1 mRNA distribution in PVD neurons, we utilized the MS2 tagging system, which relies on the sequence-specific interaction between the MS2 bacteriophage RNA hairpin loops and capsid proteins. In this system, the kpc-1 mRNA containing either its own 3’UTR or a control unc-54 3’UTR was fused to tandem MS2 binding sites and coexpressed with a green fluorescent MS2 capsid protein23 (Fig. 3a). The green fluorescent MS2 capsid protein, which is targeted to the nucleus by the nuclear localization signal (Fig. 3b), binds tightly to MS2 binding sites fused to the 3’UTR of the kpc-1 mRNA, allowing visualization of the kpc-1 mRNA by MS2 fluorescence in PVD neurons. Using this system, we detected significant axonal and dendritic signals for kpc-1 transcripts containing its own 3’UTR (Fig. 3c,d,g). Dendritic signals were preferentially distributed in the 1° and 3° dendrites (Fig. 3c,d, white arrows), consistent with kpc-1’s roles in regulating secondary dendrite branching from the 1° dendrites and tertiary dendrite self-avoidance. Distribution of kpc-1 transcripts in 3° dendrites was region specific. We observed enriched localization at potential branching points and retracted contact points between neighboring 3° dendrites (Fig. 3c,d, grey arrowheads). Using the mCherry reporter to label PVD dendrites, we can confirm enrichment of kpc-1 transcripts at branching points (Fig. 3E, grey arrowheads) and retracted contact points (Fig. 3E, white arrowheads). In
contrast, when the \textit{kpc-1} 3’UTR was replaced with an unrelated \textit{unc-54} 3’UTR, the \textit{kpc-1} transcripts were mainly trapped in the nucleus with some perinuclear signals detected in PVD (Fig. 3f). \textit{kpc-1} transcripts containing the \textit{unc-54} 3’UTR had a significantly lower chance to be transported to higher-order tertiary dendrites compared to \textit{kpc-1} transcripts containing its own 3’UTR (Fig. 3g,h). Together, these results indicate that the \textit{kpc-1} 3’UTR facilitates \textit{kpc-1} RNA localization to distal primary dendrites and higher-order tertiary dendrites, implicating local translation of \textit{kpc-1} in dendrites.

\textbf{The \textit{kpc-1} 3’UTR promotes local protein synthesis in the distal segment of PVD dendrites}

Our results showed that the \textit{kpc-1} 3’UTR facilitates localization of \textit{kpc-1} transcripts to the distal primary and tertiary PVD dendrites, suggesting that the \textit{kpc-1} 3’UTR promotes local translation of transcripts it regulates. To examine whether the \textit{kpc-1} 3’UTR can promote local protein synthesis, we performed fluorescence recovery after photoconversion (FRAP) of the photoconvertible Kaede protein\textsuperscript{24}, which was under control of the \textit{kpc-1} 3’UTR or a control \textit{unc-54} 3’UTR (Fig. 4). Upon photostimulation using 405 nm laser, Kaede proteins can be converted from green fluorescent proteins to red fluorescent proteins irreversibly, allowing us to distinguish the future newly synthesized green fluorescent Kaede proteins from the resident (converted) red fluorescent Kaede proteins. We carried-out photoconversion at a broad region from the mid-body to the nerve ring and measured newly synthesized green fluorescent Kaede proteins in a local area distant away from the cell body (Fig. 4a). This design ensures that the local green fluorescent signals represent newly synthesized Kaede proteins rather than resident (unconverted) Kaede proteins diffused
from the soma (Fig. 4a). Green fluorescence recovery of Kaede proteins expressed from the *kaede::kpc-1 3’UTR* is significantly faster than that from the *kaede::unc-54 3’UTR* (Fig. 4b,c), indicating that the *kpc-1 3’UTR* promotes local protein synthesis in the distal segment of PVD dendrites. Using the split GFP *rps-18* reporter to visualize ribosomes in PVD neurons\(^{25}\), we detected the protein synthesis machinery not only in the soma but also in the tertiary dendrites (Fig. 4d), further supporting protein synthesis occurs in the higher order dendrites to supply proteins and to allow for proteome remodeling.

**Identification and characterization of secondary structural motifs in the kpc-1 3’UTR required for tertiary dendrite self-avoidance**

Many studies have shown that specific elements in 5′ or 3′ untranslated regions (UTRs) of mRNA regulate mRNA stability, splicing, translation and localization\(^{26}\). microRNAs regulate mRNA translation and stability by binding to complementary sequences in the 3′UTR\(^{27}\). Other types of elements in the 3′UTR can function in either a sequence- or secondary structure-specific manner. For example, one of the earliest identified localization elements was discovered from the chick β-actin mRNA. The “zipcode” in the 3′UTR of β-actin mRNA consists of 54 nucleotides that contain tandem repeats of the conserved hexanucleotide motif ACACCC and form a stem-loop structure recognized by the Zipcode Binding Proteins (ZBPs) to regulate the localization and translation of β-actin mRNA\(^{28,29}\). In yeast, the cis-acting sequences that mediate ASH1 mRNA targeting to the bud tip provide another example of repetitive clustering and synergistic action of localization elements. Four localization elements in ASH1 mRNA were all predicted to form stem-loop structures, and each element on its own is able to localize mRNAs, but the
presence of four elements together increases the efficiency of localization\textsuperscript{30-32}. In Aplysia sensory-motor neurons, cis-elements in the 3' UTR of the neurotransmitter sensorin mRNA control the transport of sensorin mRNAs from soma to neurite\textsuperscript{33}. In mice, conditional deletion of an axon-localizing cis-element in the 3'UTR of importin β1 mRNAs in sensory neurons caused selective depletion of importin β1 mRNAs and proteins in axons, without affecting cell body levels or nuclear functions, suggesting that local translation of importin β1 mRNAs in axons enables separation of axonal and nuclear transport functions of importins, and is required for efficient retrograde signaling in injured axon\textsuperscript{34}.

In \textit{C. elegans}, it was previously reported that the \textit{cebp-1} 3’UTR is required for the localization of \textit{cebp-1} mRNAs to discrete locations in touch axons, which can be translated into proteins locally\textsuperscript{35}. What is unclear is whether those discrete locations represent synapses in touch axons. In addition, a putative stem-loop secondary structure was identified in the \textit{cebp-1} 3’UTR that inhibits \textit{cebp-1} mRNA stability or its translation\textsuperscript{36}. Thus, the \textit{cebp-1} 3’UTR appears to have both positive and negative impacts on \textit{cebp-1} gene expression.

Secondary structural motifs derived from the alignment of \textit{kpc-1} 3’UTRs from 4 closely-related species, including \textit{C. elegans} (Supplementary Fig. 3), as well as individual Turbofold II-predicted secondary structures\textsuperscript{37}, revealed that two stem-loop structural motifs (SLS1 and SLS2) are conserved in the 3’ UTR of all \textit{kpc-1} homologs (Fig. 5a). The \textit{kpc-1} transgene with its 3’UTR being replaced by a control \textit{unc-54} 3’UTR lost the ability to rescue tertiary dendrite self-avoidance defects in the \textit{kpc-1(gk8)} null allele (Fig. 5f). Deletion analysis showed that SLS2 but not SLS1 is required for the \textit{kpc-1} 3’UTR to mediate tertiary dendrite self-avoidance since ΔSLS2 but not ΔSLS1 in the \textit{kpc-1} 3’UTR
abolished the ability of the \textit{kpc-1} transgene to rescue tertiary dendrite self-avoidance defects in the \textit{kpc-1}(gk8) null allele (Fig. 5f). The \textit{kpc-1} transgene with mutations that disrupt base-pairing in the stem region of SLS2 in the 3’UTR significantly increased contacts between sibling tertiary dendrites (indicating losing self-avoidance functions; Fig. 5c,f) whereas the \textit{kpc-1} transgene with reciprocal mutations that preserve base-pairing in the stem region of SLS2 in the 3’UTR did not increase contacts between sibling tertiary dendrites (indicating maintaining self-avoidance functions; Fig. 5d,f).

All the \textit{kpc-1} transgenes tested here partially rescued secondary dendrite trapping defects in the \textit{kpc-1}(gk8) null allele, suggesting that the modifications we introduced to the transgenes, including the 3’UTR replacement, deletions, and mutations, did not result in degradation of the \textit{kpc-1} transcripts (Fig. 5e). Levels of the \textit{kpc-1} RNA expressed from various transgenes in \textit{kpc-1}(gk8) mutants were analyzed by RT-PCR. All \textit{kpc-1} 3’UTR modifications were made from the PF49H12.4::\textit{kpc-1}:\textit{kpc-1} 3’UTR construct, where only the short-isoform \textit{kpc-1}s RNA would be generated. We found that the \textit{kpc-1} RNA expression levels are comparable among different \textit{kpc-1} transgenes (Supplementary Fig. 4), indicating that modifications we introduced to the \textit{kpc-1} 3’UTR did not affect the \textit{kpc-1} RNA stability. Furthermore, even though the level of the endogenous \textit{kpc-1} RNA in wild-type animals (containing both long and short isoforms) is lower, it promotes dendrite self-avoidance better than a higher level of the short-isoform (\textit{kpc-1}s RNA) expressed from various transgenes in PVD neurons of \textit{kpc-1}(gk8) mutants (Supplementary Fig. 4). Together, our results have identified an important secondary structural motif in the \textit{kpc-1} 3’UTR required for tertiary dendrite self-avoidance.
KPC-1 down-regulates DMA-1 receptors on PVD tertiary dendrites

While the number of tertiary dendrites is significantly reduced in sax-7, mnr-1, lect-2, or dma-1 mutants, it is significantly increased in kpc-1(xr58) mutants\textsuperscript{14,15,17–19} (Fig. 1c). This result suggests that kpc-1 likely antagonizes the sax-7/mnr-1/lect-2/dma-1 signaling pathway. Double mutant analyses revealed that sax-7, mnr-1, lect-2, and dma-1 mutations completely suppressed the kpc-1(xr58) mutant phenotype of excessive tertiary dendrites, indicating that the kpc-1 mutant effect depends on the sax-7/mnr-1/lect-2/dma-1 signaling pathway (data not shown). The kpc-1(gk8) null allele seems to have a phenotype very similar to the sax-7, mnr-1, lect-2, and dma-1 mutants\textsuperscript{14,15,17–19} (Fig. 2d,e). However, closer examination of the kpc-1(gk8) null mutant phenotype revealed that defective secondary dendrite extension to sublateral longitudinal stripes is caused by a failure of secondary dendrites to escape the trap zone\textsuperscript{2}, a phenotype that is clearly different from the less secondary dendrite outgrowth in sax-7, mnr-1, lect-2, and dma-1 mutants. Double mutant analyses between the kpc-1(gk8) null allele and sax-7, mnr-1, and dma-1 mutations have been reported in a recent paper\textsuperscript{2}, which also supported that kpc-1 antagonizes the sax-7/mnr-1/dma-1 signaling pathway in PVD neurons. dma-1, like kpc-1, encodes a transmembrane protein and acts cell-autonomously in PVD neurons\textsuperscript{2,17} (Fig. 2a,b). Thus, DMA-1 could be a potential KPC-1 target in regulating PVD tertiary dendrite self-avoidance. Supporting this possibility, endogenous DMA-1 protein levels on PVD tertiary dendrites were up-regulated in kpc-1 mutants, suggesting that KPC-1 normally down-regulates DMA-1 on PVD tertiary dendrites (Fig. 6a,b). In addition, the kpc-1(xr47) mutant allele, which showed a greater increase in the number of overlapped tertiary dendrites compared to the kpc-1(xr58) mutant allele (Fig. 6c), caused a greater up-regulation of the
DMA-1 protein level on PVD tertiary dendrites (Fig. 6a,b). A recent paper reported a different phenotypic effect of kpc-1 mutations in PVD neurons where kpc-1 mutations caused excessive DMA-1 receptors on secondary dendrites, which results in secondary dendrites trapped in the primary dendrite zone where a high level of the cognate ligand SAX-7/L1CAM is present2.

**The kpc-1-dma-1 regulatory circuit promotes dendrite self-avoidance**

To further strengthen the conclusion of a kpc-1-dma-1 regulatory circuit in promoting PVD dendrite self-avoidance, we tested whether dma-1 over-expression (OE) mimics the kpc-1 mutation in its phenotypic effect on PVD dendrites. dma-1 OE displayed a tertiary dendrite self-avoidance defect similar to that exhibited by the kpc-1(xr58) mutant allele (Fig. 7a,b). In addition, dma-1 OE caused a secondary dendrite trapping phenotype (Fig. 7d), which was further enhanced by the kpc-1(xr58) mutation (data not shown). In contrast, kpc-1 OE suppressed the secondary dendrite trapping phenotype in dma-1 OE animals (Fig. 7c-f). Although fewer kpc-1 OE; dma-1 OE animals displayed a secondary dendrite trapping phenotype, most still exhibited a weak tertiary dendrite self-avoidance defect, that was milder than dma-1 OE or kpc-1(xr58) alone. Thus, a weaker kpc-1-dma-1 regulatory circuit causes tertiary dendrite self-avoidance defects whereas a further reduced kpc-1-dma-1 regulatory circuit results in secondary dendrite trapping. Indeed, the mild kpc-1(xr58) mutant allele preferentially affects tertiary dendrite self-avoidance rather than secondary dendrite branching. Together, our results support a model in which locally expressed KPC-1 down-regulates DMA-1 receptors on PVD dendrites to promote dendrite branching and self-avoidance (Fig. 7k).
We have made different versions of KPC-1::dzGFPm-yr and KPC-1::GFP fusion proteins, both wild-type and mutant forms that disrupted the KPC-1 self-cleavage site. However, we were unable to detect any fluorescent signal beyond the PVD cell body (unpublished), suggesting that KPC-1 proteins have very short half-life in PVD dendrites. Due to this technical limitation, we cannot determine whether KPC-1 proteins are synthesized locally in tertiary dendrites. Nor can we address whether the KPC-1 protein and its substrate DMA-1 specifically collocate to higher order dendrites. As an alternative approach, we superimposed the image of dendritic signals for kpc-1 transcripts with the image of dendritic signals for DMA-1 proteins. We found that they did not show overlapping distribution pattern (Fig. 7g,h). Instead, adjacent sites of expression can be detected for kpc-1 RNAs and DMA-1 proteins (arrowheads; Fig. 7g,h), consistent with down-regulation of DMA-1 by KPC-1 at specific sites.

The tertiary dendrite self-avoidance defect could result from either excessive dendrite growth ability that overcomes a self-avoidance mechanism or insufficient self-avoidance. To distinguish between these two possibilities, we performed laser dendritomy in PVD neurons at the early L4 stage when tertiary dendrites underwent self-avoidance and assessed the dendrite growth ability 24h after surgery. Dendrites were cut at the primary branch 20 μm anterior to the cell body such that the entire anterior dendrite arbor was disconnected from the cell body after surgery. We found that PVD dendrite growth ability was not significantly enhanced in kpc-1(xr58) mutants compared to wild-type animals (Fig. 7i). This result indicated that the dendrite self-avoidance defect in kpc-1(xr58) mutants cannot be attributed to excessive dendrite growth ability.
In contrast, over-expression of \textit{kpc-1} significantly increased the distance (gap) between neighboring tertiary dendrites (Fig. 7c,j). This result suggested that \textit{kpc-1} either limited tertiary branch outgrowth or promoted tertiary dendrite self-avoidance. Time-lapse imaging revealed that neighboring tertiary dendrites in \textit{kpc-1} over-expression animals actually contacted at first and retracted subsequently, suggesting that \textit{kpc-1} over-expression did not limit tertiary branch outgrowth (data not shown). Together, these results support that \textit{kpc-1} promotes dendrite self-avoidance rather than limiting dendrite outgrowth.
Discussion

Our results support a model where locally synthesized KPC-1 at branching points and contact points between sibling dendrites down-regulates DMA-1 receptors to promote dendrite branching and self-avoidance (Fig. 7k). A recent paper showed that KPC-1 targeted DMA-1 to endosomes for degradation in PVD neurons. It is likely that a similar mechanism is utilized in a site-specific manner for the secondary dendrite branching and the tertiary dendrite self-avoidance. KPC-1 encodes a proprotein convertase subtilisin/kexin (PCSK). At a functional level, there are two types of mammalian PCSKs. PCSK3/furin activates substrates by cleavage of precursor proteins, such as TGF-β, whereas PCSK9 inactivates substrates by induced degradation of proteins, such as LDL receptors. Our results indicate that KPC-1 is more closely related to PCSK9 at a functional level in its inactivation of DMA-1. It has been shown that PCSK9 is expressed in the vertebrate CNS, including the cerebellum, and is important for CNS development. However, it remains to be seen whether PCSK9 plays a similar role to KPC-1 in dendrite branching and self-avoidance by down-regulating guidance receptors in dendrites.

It is foreseeable that the level of DMA-1 receptors on tertiary dendrites is tightly controlled during dendrite outgrowth and self-avoidance (Fig. 6). For example, the DMA-1 receptor level should be high on tertiary dendrites during dendrite outgrowth. However, the DMA-1 receptor level needs to be down-regulated upon contact between sibling tertiary dendrites, resulting in dendrite retraction. An important question that remains to be addressed in the future is how the KPC-1 activity is dynamically controlled to regulate a corresponding DMA-1 level. There are two possible regulatory mechanisms that could be responsible: expression versus activation of KPC-1 at the contact points between sibling
dendrites. Our observation of the enriched distribution of \textit{kpc-1} transcripts at the contact points appears to support the former case (Fig. 3e). However, it is not known how \textit{kpc-1} transcripts are targeted to contact points and whether contact-mediated activation of RNA transport may be responsible.

It was recently reported that 12 conserved RNA-binding protein (RBP)-encoding genes are involved in PVD dendrite development in \textit{C. elegans} \cite{43}. Among them, loss of function in \textit{cgh-1} and \textit{cpb-3} resulted in an increase in the number of secondary and tertiary dendrite branches. The phenotype of increased number of secondary and tertiary dendrite branches is also shared by \textit{kpc-1}(\textit{xr58}) mutants (1c,d). Like \textit{kpc-1}, \textit{cgh-1} and \textit{cpb-3} act cell-autonomously in PVD neurons to control dendrite development. Also, like \textit{kpc-1} transcripts, CGH-1 and CPB-3 proteins are localized to PVD dendrites, raising the possibility that CGH-1 and CPB-3 may regulate the \textit{kpc-1} function in dendrites by binding to the 3′UTR of \textit{kpc-1} transcripts, an interesting question to be studied. A Schizophrenia-associated genetic variant in the 3′UTR of the human furin gene, a \textit{kpc-1} homologous gene, was recently shown to result in downregulation of furin expression by acquiring a miR-338-3p binding site, leading to reduced BDNF production. It remains to be seen whether this human furin 3′UTR genetic variant disrupts transport and local translation of the furin mRNA in dendrites and whether it affects dendrite patterning in Schizophrenia-causing neurons \cite{44,45}.  

\[\text{Equation}\]
Experimental Procedures

Strains and plasmids

*C. elegans* strains were cultured using standard methods\(^{36}\). All strains were grown at 20°C. Standard protocol was used for the plasmid constructions. Strains and plasmids are listed in Supplementary Tables S1 and S2.

Transgenic animals

Germline transformation of *C. elegans* was performed using standard techniques\(^ {47}\). For example, the *Pkpc-1(5K)::kpc-1* transgene was injected at 10 ng/ml along with the coinjection marker *ofm-1::rfp* at 50 ng/µl. Transgenic lines were maintained by following the *ofm-1::rfp* fluorescence.

Tertiary branch avoidance index (TBAI)

TBAI was calculated by dividing the number of visible gaps between tertiary branches by the number of menorahs. A menorah-like structure is composed of a collection of 2° (“stem”), 3° (“base”) and 4° (“candles”) branches. In the case where discrete menorahs cannot be easily differentiated due to self-avoidance defects, we counted each menorah as a single 2° (“stem”) branch that extended 3° (“base”) branches.

Scatter plots of quaternary dendrite termini

To map out quaternary dendrite termini, we used a set of customized MatLab scripts (MatLAB, Mathworks, Natick, MA) to effectively unroll the worm’s cylindrical surface, quantifying anterior-posterior distances using the coordinate parallel to the body centerline.
and dorsal-ventral distances along the cylindrical surface. In order to consolidate data from different worms, all scatter plots are scaled to each worm’s circumference. In each scatter plot, the top line indicates the dorsal nerve cord, the bottom line indicates the ventral nerve cord, and the wild-type morphology of the PVD primary dendrite and axon is drawn in green. The distance between the top and bottom lines corresponds to 1/2 of total worm circumference, and the horizontal axis shows a portion of body length equivalent to approximately 3 circumferences.

**Laser dendritomy**

For the femtosecond laser surgery, we used a cavity-dumped Ti:sapphire laser oscillator\textsuperscript{48,49} to generate laser pulses ~100 fs in duration and 200 kHz in repetition rate. The laser pulses were tightly-focused onto targeted primary dendrites using a Nikon 100x, 1.4 NA oil-immersion objective. The vaporization threshold corresponds to pulse energies of 5-15 nJ. Successful laser dendritomy was confirmed by visualizing the targeted area immediately after surgery. Dendrites were cut at the primary branch 50 µm anterior to the cell body in the early L4 stage when tertiary dendrites undergo self-avoidance. 1°, 2°, 3°, and 4° dendrites in the anterior dendrite arbor were disconnected from the cell body after primary dendritomy. Newly regenerated dendrites can be easily differentiated from the old dendrites 24 hr after surgery due to difference in the fluorescent intensity of the \textit{Pser-2::GFP} dendrite marker.

**Imaging and quantifying dendrite arbors**
The morphology of neuronal cell bodies and dendrites was based on high-magnification Z-stacks using a Zeiss 60x, 1.4 NA oil-immersion objective. For laser dendritomy, we mounted individual animals on 2% agar pads and anaesthetized them with 3 mM sodium azide, the lowest possible concentration to keep adult animals immobilized. Laser dendritomy was performed and worms were recovered within 10 minutes of sodium azide treatment. Recovered worms were placed on fresh plates with bacterial foods and imaged 24 hours after dendritomy using a Hamamatsu ORCA AG camera. For imaging and quantifying uninjured dendrite arbors, young adult animals were mounted on 2% agar pads and anaesthetized with 20 mM sodium azide.

The dendrite length of regenerating neurons was quantified 24 hours after surgery. Dendrite lengths were calculated as the actual contour length between the injury site and dendrite termini measured along the cylindrical surface of each worm, by tracing dendrites through a 3-dimensional image stack. P values for the length measurements were calculated using a Student’s t-Test.

**Fluorescence microscopy and confocal imaging**

Images of DMA-1-YFP were captured in live animals using a Plan-Apochromat 40X/1.3NA objective on a Zeiss LSM 880 confocal microscope (Carl Zeiss).

**Fluorescence recovery after photoconversion**

Worms were mounted on 2% agarose pad and anaesthetized with 7.5 mM tetramisole. Photoconversion of Kaede was performed under 405 nm laser on the region of interest (ROI). Fluorescence quantification of ROI intensity was measured every 5 seconds over 6
minutes following photoconversion (Zeiss Imaging Browser). At each time point, percentage green fluorescence signal recovery was determined by \((F_n - F_0)/(F_p - F_0)\), where \(F_n = \) fluorescence intensity \(n\) seconds after photoconversion; \(F_0 = \) fluorescence intensity immediately after photoconversion; \(F_p = \) fluorescence intensity right before photoconversion. All images were taken in live animals using a 40x, 1.3 NA objective and 488 nm and 561 nm laser on a Zeiss LSM 880 confocal microscope.

**Statistics**

Average data of dendrite number, dendrite length, TBAI, receptive field index, length ratio of receptive field, and reporter expression intensity are presented as means ± SEM. Data of % PVD neurons with trapped secondary dendrites, defective dendrite self-avoidance, and dendritic signals are presented as proportions ± SEP. Statistical analyses were carried out with the Primer of Biostatistics software for the Student’s t-Test or the two-proportion Z-test. \(P < 0.01\) or \(P < 0.001\) was considered statistically significant.

**Author Contributions**

Y.Z. conceived, designed, performed, and analyzed experiments, and made constructs. M.S. conceived, designed, performed, and analyzed experiments, made constructs, and drafted the article. H.C. conceived, designed, performed, and analyzed experiments, and drafted the article. T.F. performed genetic screens, identified dendrite self-avoidance mutants, and analyzed experiments. N.Z. imaged and quantified green fluorescent MS2 capsid proteins in PVD dendrites. W.Z. made crispr line and contributed unpublished
essential data and reagents. C.F.C. conceived, designed, and analyzed experiments. C.C. conceived, designed, analyzed and interpreted data, and drafted the article.

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Figure legends

**Figure 1.** *kpc-1(xr58)* mutations preferentially affect PVD dendrite self-avoidance. (a, c) Images of PVD dendrites in wild type (a) and *xr58* mutants (c). Scale bar, 20 μm. (b) Schematic of dendrites (green) and axon (grey) extended from a PVD neuron. PVD projected primary (1°) dendritic processes anteriorly and posteriorly from the soma. Higher order dendritic branching (secondary (2°), tertiary (3°), quaternary (4°)) alternated between anterior-posterior and dorsal-ventral directions to generate a regularly spaced array of parallel branches. *xr58* mutants, while showing effective secondary dendrite escape from the trap zone, displayed extensive defects in self-avoidance of sibling dendrites. Arrows point to contacts between neighboring secondary dendrites. Arrowheads point to contacts between neighboring tertiary dendrites. (d) Quantification of secondary dendrites in wild type versus *kpc-1(xr58)* mutants. (e) Tertiary branch avoidance index (TBAI) in wild type versus *kpc-1(xr58)* mutants. *kpc-1(xr58)* mutants displayed significantly lower TBAI than wild-type animals. (f) Scatter plots showing the positions of 4° dendrite termini in wild-type young adults and *kpc-1(xr58)* young adults. In each scatter plot, the top line (dorsal nerve cord), the bottom line (ventral nerve cord), and the wild-type PVD axon and primary dendrite morphology (green) are shown. (g) Schematic of the KPC-1 protein motifs showing locations of *xr47* and *xr60* mutations.

**Figure 2.** *kpc-1* is required for both the secondary dendrite escape from the trap zone and the tertiary dendrite self-avoidance but the requirements of two functions are qualitatively different. (a) *kpc-1* functions in PVD neurons to promote secondary dendrite escape from the trap zone independent of the *kpc-1* 3’UTR. Percentage PVD neurons with
secondary dendrite trapping in kpc-1(gk8) mutants carrying various transgenes. (b) The kpc-1’s function in PVD for tertiary dendrite self-avoidance is dependent on the kpc-1 3’UTR. Percentage PVD neurons with defective tertiary dendrite self-avoidance in kpc-1(gk8) and kpc-1(xr58) mutants carrying various transgenes. ND, not determined. (c) Identification of the kpc-1 promoter reporter expression in PVD neurons. (d) Images of PVD neurons in kpc-1(gk8) mutants with or without the Ppvd::kpc-1::unc-54 3’UTR transgene. The Ppvd::kpc-1::unc-54 3’UTR transgene rescued the secondary dendrite trapping but not the tertiary dendrite self-avoidance defect in kpc-1(gk8) mutants. Arrowheads point to contacts between neighboring tertiary dendrites. Asterisks indicate contacts between neighboring secondary dendrites. (e, f) Images of PVD neurons in kpc-1(gk8) mutants without or with the Ppvd::kpc-1::kpc-1 3’UTR transgene. The Ppvd::kpc-1::kpc-1 3’UTR transgene rescued both the secondary dendrite trapping and the tertiary dendrite self-avoidance defect in kpc-1(gk8) mutants. (g, h) Images of PVD neurons in kpc-1(xr58) mutants without or with the Ppvd::kpc-1::kpc-1 3’UTR transgene. The Ppvd::kpc-1::kpc-1 3’UTR transgene rescued the tertiary dendrite self-avoidance defect in kpc-1(xr58) mutants. Arrowheads point to contacts between neighboring tertiary dendrites. Scale bar, 20 μm.

**Figure 3. The kpc-1 3’UTR targets the kpc-1 RNA to higher-order dendrites.** (a) The MS2 tagging system is composed of the kpc-1 mRNA containing its own 3’UTR or a control unc-54 3’UTR fused to 24xMS2 binding sites co-expressed with a green fluorescent MS2 capsid protein. The nuclear localization signal (NLS) is used to initially target green fluorescent MS2 proteins to the nucleus. (b) Image of the PVD neuron showing...
enriched nuclear signals for the nuclear localized green fluorescent MS2 capsid protein. Asterisk indicates soma. Scale bar, 20 µm. (c) Image of the PVD neuron showing axonal and dendritic signals for kpc-1 transcripts containing the kpc-1 3’UTR and MS2 binding sites. The white arrow points to dendritic signals in the primary dendrite and the grey arrow marks axonal signals. Grey arrowheads indicate the potential branching points at 3° dendrites and contact points between neighboring 3° dendrites. The dashed box area in 3c was blown up and shown in 3d. Anterior is to the left, dorsal is up. Scale bar, 20 µm. (e) Image of the PVD neuron showing axonal and dendritic signals for kpc-1 transcripts containing the kpc-1 3’UTR and MS2 binding sites. Low level of the mCherry protein was used to label PVD. Open arrowheads indicate branching points and white arrowheads indicate 3° dendrite termini (retracted contact points between neighboring 3° dendrites). Scale bar, 20 µm. (f) Image of the PVD neuron showing enriched soma signals for kpc-1 transcripts containing the control unc-54 3’UTR and MS2 binding sites. Asterisk indicates soma. Scale bar, 20 µm. (g) Average number of punctate MS2 fluorescence signals on 3° dendrites in transgenic animals expressing kpc-1 transcripts, containing either its own 3’UTR or the control unc-54 3’UTR. (h) Percentage PVD neurons with punctate MS2 fluorescence signals on 3° dendrites in transgenic animals expressing kpc-1 transcripts, containing either its own 3’UTR or the control unc-54 3’UTR.

**Figure 4. The kpc-1 3’UTR promotes local protein synthesis in the distal segment of PVD dendrites.** (a) Representative images of Kaeda-labelled PVD dendrites pre- and post-photoconversion. The dashed white box indicates a wide photoconverted region. The solid yellow box denotes a targeted area of analysis, which is distal to the soma within the wide
photoconverted region. Scale bar, 20 µm. (b) Pre-photoconversion of resident Kaede proteins and post-photoconversion of newly synthesized Kaede proteins. Kaeda expression in PVD dendrites is controlled by either a control unc-54 3’UTR or a kpc-1 3’UTR. Scale bar, 20 µm. (c) The kaeda::kpc-1 3’UTR transcript is expressed at a higher rate compared to the control kaeda::unc-54 3’UTR transcript. Quantification of green signal recovery in a targeted distal area after photoconversion of a wider region of PVD dendrites. Statistical analysis using 2-way ANOVA. Average data of Kaeda expression intensity are presented as means ± SEM. (d) Pan-neuronal expression of a split GFP reporter allows visualization of ribosomes in PVD neurons. The smaller part of split GFP (GFP11) is tagged to the endogenous rps-18 (GFP11::rps-18). The larger part of split GFP (GFP1-10) is overexpressed from a pan-neuronal promoter regf-1 using a high-copy transgene. GFP1-10 binds to the GFP11::RPS-18 in neurons and illuminates RPS-18. The Pser-2::mCherry marker labels PVD dendrites. Ribosomes are mostly restricted in the PVD soma, which is marked by an asterisk. Arrowheads indicate RPS-18 signals in tertiary dendrites and dendrite branching points. Scale bar, 20 µm.

**Figure 5. A putative stem-loop secondary structure in the kpc-1 3’UTR is required for tertiary dendrite self-avoidance.** (a) Illustration of putative stem-loop secondary structural motifs and specific mutations in the kpc-1 3’UTR. (b-d) Representative images of the kpc-1(gk8) mutant, the kpc-1(gk8) mutant expressing the Ppvd::kpc-1::kpc-1 3’UTR transgene (SLS2 mutated base pairing disrupted), and the kpc-1(gk8) mutant expressing the Ppvd::kpc-1::kpc-1 3’UTR transgene (SLS2 mutated base pairing preserved). Arrowheads point to contacts between neighboring tertiary dendrites. Scale bars, 20 µm.
(e) Percentage PVD neurons with secondary dendrite trapping in kpc-1(gk8) mutants and kpc-1(gk8) mutants carrying various transgenes. (f) Quantification of contact points between sibling tertiary dendrites in kpc-1(gk8) mutants and kpc-1(gk8) mutants carrying various transgenes. ND, not determined. ***p<0.0001 by one-way ANOVA.

Figure 6. Differential mutant effects displayed by different kpc-1 alleles. (a) Representative images showing DMA-1::YFP expression on PVD dendrites in wild type, kpc-1(xr58), and kpc-1(xr47) mutants. Scale bars, 20 µm. (b) Average fluorescent intensity of the DMA-1::YFP on PVD tertiary dendrites in wild type [dma-1(wy996)], kpc-1(xr58) mutants [kpc-1(xr58); dma-1(wy996)], and kpc-1(xr47) mutants [kpc-1(xr47); dma-1(wy996)]. dma-1(wy996) marker strain was created using the Crispr technology to knock in the YFP reporter gene to the 3' end of the endogenous dma-1 locus to generate a dma-1::yfp fusion gene, which allows us to quantify the endogenous level of DMA-1 proteins based on fluorescent intensity. (c) Quantification of tertiary dendrites in wild type versus kpc-1 mutants. (d) Image of the trans-heterozygous kpc-1(xr58)/kpc-1(gk8) mutant displaying self-avoidance defects similar to those displayed by homozygous kpc-1(xr58) mutants. Scale bars, 20 µm. Confirmation of the kpc-1 mutant phenotype in PVD dendrite self-avoidance by a non-complementation test.

Figure 7. dma-1 over-expression phenocopied the kpc-1(xr58) mutant allele. (a, b) Images of PVD dendrites in kpc-1(xr58) mutants and dma-1 over-expression animals. Arrowheads point to contacts between neighboring tertiary dendrites. Scale bars, 20 µm. (c-e) Images of PVD dendrites in kpc-1 over-expression animals, dma-1 over-expression
animals, and animals over-expressing both kpc-1 and dma-1. Scale bars, 20 µm. (f) kpc-1 over-expression suppressed the secondary dendrite trapping phenotype in dma-1 over-expression animals. (g, h) Images of PVD neurons showing dendritic signals for kpc-1 transcripts and DMA-1 proteins. kpc-1 transcripts containing kpc-1 3′UTR and MS2 binding sites were visualized by green fluorescent MS2 capsid proteins. DMA-1 proteins were visualized by tagged mCherry fluorescent proteins. White arrowheads point to dendritic signals for kpc-1 transcripts and grey arrowheads indicate dendritic signals for DMA-1 proteins. Scale bars, 15 µm. (i) Bar chart of average PVD dendrite length in wild type and kpc-1(xr58) mutants. Dendritomy was carried out at the early L4 stage when tertiary dendrites underwent self-avoidance. Newly grown PVD dendrite length was measured 24 hours after dendritomy. Error bars indicate SEM. (j) Bar chart of average distances between neighboring tertiary dendrites in wild type and kpc-1 over-expression animals at the young adult stage. For each animal, average distances of the gaps between neighboring tertiary dendrites were measured per 150 µm anterior to the cell body. Asterisks indicate that kpc-1 over-expression animals differ from wild-type at P<0.01 (**). Error bars indicate SEM. (k) Model of the kpc-1-dma-1 regulatory circuit in dendrite branching and self-avoidance. kpc-1 is expressed at branching points and contact points between the neighboring 3° dendrites, which down-regulates DMA-1 receptors locally, leading to dendrite branching and retraction of sibling dendrites. SAX-7 cues are distributed highly in the 1° and the 3° dendrite growth pathways.
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Figure 1

(a) WT
(b) Pro domain
(c) xr58
(d) Number of secondary branches
(e) Tertiary branch avoidance index
(f) WT YA
(g) gk8 deletion

WT kpc-1(xr58)
Figure 2

(a) Bar chart showing the percentage of PVD neurons with trapped secondary dendrites. The x-axis represents different conditions, and the y-axis represents the percentage of neurons. The bars indicate the number of neurons per condition.

(b) Bar chart showing the percentage of PVD neurons with defective dendrite self-avoidance. The x-axis represents different conditions, and the y-axis represents the percentage of neurons.

(c) Images showing the expression of Pkpc-1::GFP and Ppvd::mCherry in wild type and mutant conditions. The merged image shows the co-expression of both proteins.

(d) Image showing the expression pattern of kpc-1 in wild type and mutant conditions. The arrowheads indicate the location of the expression.

(e) Image of kpc-1(gk8) expression pattern.

(f) Image of kpc-1(gk8); Ex[Ppvd::kpc-1::kpc-1 3'UTR] expression pattern.

(g) Image of kpc-1(xr58) expression pattern.

(h) Image of kpc-1(xr58); Ex[Ppvd::kpc-1::kpc-1 3'UTR] expression pattern.

Legend:
- Pkpc-1::GFP
- Ppvd::mCherry
- Merged
- kpc-1(gk8)
- kpc-1(xr58)
- Ex[Ppvd::kpc-1::kpc-1 3'UTR]
Figure 3
Figure 4
Figure 5

- a: Diagram of kpc-1 3'UTR and its base pairing with SLS1 and SLS2.
  - Mutated paired nucleotides
  - Un-paired nucleotides
  - Mutated un-paired nucleotides

- b: Image of kpc-1(gk8) showing secondary dendrite trapping defects.

- c: Image of kpc-1(gk8); [PVDp::kpc-1 cDNA::kpc-1 3'UTR] with SLS2 mutated base pairing disrupted.
  - Severe: 23
  - Moderate: 18
  - Mild: 23
  - No: 34

- d: Image of kpc-1(gk8); [PVDp::kpc-1 cDNA::kpc-1 3'UTR] with SLS2 mutated base pairing preserved.

- e: Bar chart showing secondary dendrite trapping defects.
  - WT: 65
  - kpc-1(gk8): 49
  - unc-54 3'UTR: 50
  - kpc-1 3'UTR ASLS1: 54
  - kpc-1 3'UTR ASLS2: 57
  - kpc-1(gk8); Ex[PVDp::kpc-1 cDNA::kpc-1 3'UTR]: 52

- f: Bar chart showing no. of contact points between sibling tertiary dendrites.
  - WT: 21
  - kpc-1(gk8): 34
  - unc-54 3'UTR: 43
  - kpc-1 3'UTR ASLS1: 47
  - kpc-1 3'UTR ASLS2: 52
Figure 6

**a**
- *dma-1(wy996)*
- *dma-1(wy996); kpc-1(xr58)*
- *dma-1(wy996); kpc-1(xr47)*

**b**
- Relative DMA-1::YFP intensity on tertiary branches.
- DMA-1(wy996)
- DMA-1(wy996)***
- KPC-1(xr58)
- KPC-1(xr47)

**c**
- Number of overlapped tertiary dendrites 100µm anterior to soma.
- WT
- KPC-1(xr58)
- KPC-1(xr47)***

N=32

N=39

N=34

N=15

N=17

N=16

N=39

N=34

N=32

Figure 6
Figure 7

**Figure 7**

**Panel a**
- *kpc-1(xr58)*

**Panel b**
- *dma-1 OE (defective 3° dendrite self-avoidance)*

**Panel c**
- *kpc-1 OE*

**Panel d**
- *dma-1 OE (trapped 2° dendrites)*

**Panel e**
- *kpc-1 OE; dma-1 OE*

**Panel f**
- Trapped 2° dendrites
- Defective 3° dendrite self-avoidance
- Wild type

**Panel g**
- White arrow: *kpc-1* mRNAs
- Grey arrow: DMA-1 proteins

**Panel h**
- White arrows: *kpc-1* mRNAs
- Grey arrows: DMA-1 proteins

**Panel i**
- Average dendrite length (μm)
- Wild type: N=18
- *kpc-1(xr58)*: N=15
- No significant difference (ns)

**Panel j**
- Average distances between neighboring 3° dendrites (μm)
- Wild type: N=183
- *kpc-1 OE*: N=210
- Significant difference (**)**

**Panel k**
- 3° attraction
  - (during outgrowth)
- 3° repulsion
  - (during self-avoidance)

**Legend**
- KPC-1
- DMA-1
- PVD dendrites
- SAX-7
- 1°
- 2°
- 3°
- 4°

**Graph**
- 0% PVD neurons
- 100% PVD neurons
- Trapped 2° dendrites
- Defective 3° dendrite self-avoidance
- Wild type

**Table**

|     | N=15 | N=18 |
|-----|------|------|
| Wild type |     |      |
| *kpc-1(xr58)* |     |      |

**Statistical Significance**
- ns: No significant difference
- **: Significant difference
| Genotype     | N  | Menorah coverage zone per 200 μm of 1° dendrite | Trapped 2° dendrite zone per 200 μm of 1° dendrite |
|--------------|----|-----------------------------------------------|--------------------------------------------------|
| Wild type    | 32 | 183.99                                        | 0                                                |
| kpc-1(xr58)  | 39 | 182.88                                        | 38.51                                            |
| kpc-1(xr47)  | 32 | 143.46                                        | 73.18                                            |
| kpc-1(xr60)  | 27 | 58.73                                         | 163.90                                           |
| kpc-1(gk8)   | 32 | 3.14                                          | 196.86                                           |