Two microRNA regulatory circuits set
start and end times for dendritic arborization
of a nociceptive neuron

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Summary

Choreographic dendritic arborization takes place within a defined time frame, but the timing mechanism is currently not known. Here, we report that a precisely timed lin-4-lin-14 regulatory circuit triggers an initial dendritic growth activity whereas a precisely timed let-7-lin-41 regulatory circuit signals a subsequent developmental decline in dendritic growth ability, hence restricting dendritic arborization within a defined time frame. Loss-of-function mutations in the lin-4 microRNA gene cause limited dendritic outgrowth whereas loss-of-function mutations in its direct target, the lin-14 transcription factor gene, cause precocious and excessive outgrowth. In contrast, loss-of-function mutations in the let-7 microRNA gene prevent a developmental decline in dendritic growth ability whereas loss-of-function mutations in its direct target, the lin-41 tripartite motif protein gene, cause further decline. lin-4 and let-7 regulatory circuits are expressed at the right place and the right time to set start and end times for PVD dendritic arborization. Replacing the endogenous lin-4 promoter at the lin-4 locus with a late-onset let-7 promoter delays PVD dendrite arborization whereas replacing the endogenous let-7 promoter at the let-7 locus with an early-onset lin-4 promoter causes precocious decline in dendritic growth ability in PVD neurons. We further find that lin-28 acts upstream of let-7 in regulating developmental decline in dendritic growth ability. Our results indicate that the lin-4-lin-14 and the lin-28-let-7-lin-41 regulatory circuits control the timing of PVD dendrite arborization through antagonistic regulation of the DMA-1 receptor level on PVD dendrites.
Keywords: the *lin-4-lin-14* pathway, the *lin-28-let-7-lin-41* pathway, developmental timing genes, heterochronic genes, neuronal timers, temporal regulation, dendrite arborization, PVD neurons, *Caenorhabditis elegans*
Introduction

Studies on temporal control of cell cycle progression, circadian rhythm, and segmentation have frequently converged on the concept of biological oscillators\textsuperscript{1-14}. Biological oscillators are systems of molecules with various levels of expression and activity that act as molecular “clocks” that determine biological rhythms resilient to changes in external environments. Understanding these molecular oscillators gives a glimpse into the mechanism of timing control for the cyclical nature of these processes. However, timing control for non-cyclical biological processes is less understood. Most non-cyclical events in non-neuronal cells are transient, which makes it difficult to study their temporal regulation. In contrast, differentiation of neurons occurs in a longer timescale because of their complex structures, thereby providing us with an opportunity to study intrinsic timing mechanisms.

Although abundant knowledge has been learnt in past decades on temporal control of cell fate specification of neurons\textsuperscript{15-17}, less is known for timing of their wiring to give rise to complex neuronal circuits and for timing of their plasticity. In the nervous system, establishment of neuronal connectivity during development and decline in neuronal plasticity during aging are controlled with temporal precision, but the timing mechanisms are largely unknown. The heterochronic pathways are important temporal regulators of animal development and involve a number of microRNA-regulated post-transcriptional genetic circuits, including important interactions between the \textit{lin-4} microRNA and its direct target, the \textit{lin-14} transcription factor gene\textsuperscript{18,19} and between the \textit{let-7} microRNA and its direct target, the \textit{lin-41} tripartite motif (TRIM) protein gene\textsuperscript{20,21}. Since the discovery of
the lin-4-lin-14 and the let-7-lin-41 regulatory circuits broadly expressed in the nervous system\textsuperscript{22,23}, evidences indicating a widespread role of the lin-4-lin-14 and the let-7-lin-41 regulatory circuits in timing neuronal assembly and plasticity start to accumulate. Distinct microRNA regulatory circuits were recently shown to control orderly neuronal connectivity and regeneration potential decline\textsuperscript{22-25}. The lin-4 microRNA and the LIN-14 transcription factor regulate transition of sequential events in AVM neuronal connectivity. Up-regulation of lin-4 and down-regulation of lin-14 signal the end of netrin-mediated axon pathfinding to allow synapse formation in AVM neurons\textsuperscript{22,24}. Similar functions of lin-4 and lin-14 in temporal regulation of axon pathfinding have been reported in HSN and PLM neurons\textsuperscript{26,27}. The lin-4-lin-14 regulatory circuit also temporally controls synaptic rewiring of postmitotic DD motor neurons\textsuperscript{28,29}. In postmitotic PVT interneurons, lin-14 temporally regulates onset of zig gene expression, which is required for maintenance of ventral nerve cord structure\textsuperscript{30}. The let-7 microRNA and the LIN-41 tripartite motif protein control timing of a post-differentiation event in AVM neurons\textsuperscript{23,25}. The progressive increase of let-7 and the progressive decrease of lin-41 contribute to a normal developmental decline in AVM axon regeneration. lin-4 and let-7 microRNAs also regulate the establishment of axodendritic polarity in the DA9 motor neuron early in development and its maintenance at the adult stage, respectively\textsuperscript{31}. Despite advances in understanding timing of neuronal connectivity and regeneration decline, intrinsic timing mechanisms that control choreographic dendritic arborization, an integral part of establishing a functional neural circuit, are still elusive.
Dendritic arborization of the PVD nociceptive neurons takes place within a defined time period starting at the L2 stage and ending at the young adult stage during *C. elegans* development. The PVD dendritic arbor is established by a complex but well-ordered array of non-overlapping sister dendrites. The creation of this structure involves a stereotypical series of branching decisions, which makes PVD neurons an ideal model system to study timing mechanisms of dendritic arborization. A 1° dendrite extends from the PVD cell body along the A/P axis at the location of the lateral nerve fascicle. Orthogonal arrays of 2°, 3°, and 4° dendritic branches envelop the animal in a manner that alternates between the D/V and the A/P axes to produce an elaborate network of sensory processes. Growth of these dendrite branches depends on the DMA-1 dendrite receptor in PVD neurons. What timing mechanisms initiate and terminate the arborization of this complex dendritic network? To answer these questions, we search for molecules in PVD neurons that may provide intrinsic temporal controls for dendritic arborization. We identify two timing regulatory circuits that restrain the growth of PVD dendrites within a set time frame. The initial dendritic outgrowth in PVD neurons is actuated by a precisely timed *lin-4-lin-14* regulatory circuit, which positively regulates the DMA-1 dendrite receptor level. The dendritic growth is subsequently slowed down by a precisely timed *lin-28-let-7-lin-41* regulatory circuit, which negatively regulates the DMA-1 dendrite receptor level, as dendritic arborization comes to an end.
Results

**lin-4 and let-7 are temporally expressed in PVD neurons during dendrite arborization**

To identify timing mechanisms that restrict PVD dendritic arborization within a defined time period, we sought to identify molecules that are expressed in PVD neurons and whose expression coincides with start and end times of PVD dendritic arborization. We discovered lin-4 and let-7 microRNAs to be excellent candidates that fit both criteria. To understand the temporal expression of lin-4 and let-7 genes in PVD neurons, we developed a 1.9-kb lin-4 promoter reporter and a 2.9-kb let-7 promoter reporter that were stably expressed in PVD (Figure 1; Table S2). The expression levels of these two promoter reporters in the whole animal at different stages correlated strongly with the whole animal stem-loop reverse transcription polymerase chain reaction (RT-PCR) quantification of lin-4 and let-7 microRNAs, indicating that these two reporters are reliable. Using these reporters, we determined the timing of lin-4 and let-7 expression in PVD during dendritic arborization (Figures 1J and 1K). The lin-4 reporter was expressed highly in PVD at the late L2-early L3 stage, when the secondary dendrites start elaborating (Figures 1G, 1H, and 1J). In contrast, the let-7 reporter was expressed at relatively low levels in PVD at the L3 stage, but was significantly elevated from the L4 stage onward, when growth of the terminal quaternary dendrites comes to an end (Figures 1G, 1I, and 1K). These results indicate that lin-4 and let-7 are expressed at the right place and the right time to initiate and terminate PVD dendritic arborization, respectively.

**The lin-4-lin-14 regulatory circuit initiates dendritic outgrowth in PVD neurons**
We previously reported that the *lin-4* microRNA represses the expression of the LIN-14 transcription factor to inhibit AVM axon attraction\(^{22,24}\). Further analysis of *lin-4* and *lin-14* reporters revealed overlapping expression of two genes in many other neurons, including PVD, at the early L3 stage (Figures 1A-1C and 1H), when PVD neurons are sending out the secondary dendrites. We showed that *lin-4*(e912) loss-of-function (*lf*) and *lin-14*(n355) gain-of-function (*gf*) mutants displayed a similar phenotype of limited dendritic outgrowth in PVD neurons (Figures 2A-2C and 2E), suggesting that the *lin-4* microRNA may inhibit the expression of the LIN-14 transcription factor to initiate dendrite outgrowth. Consistent with this interpretation, reduced *lin-14* activity caused opposite effects. In wild-type animals at the mid L3 stage, PVD dendrites can only grow up to the tertiary branch (Figure 2F). However, in *lin-14*(n179) reduction-of-function (*rf*) mutants at the same developmental stage, the mid L3 stage, PVD dendrites can grow up to the quaternary branch and complete the menorah organization (Figure 2F), suggesting precocious dendrite outgrowth. In addition, the number of overlapped tertiary dendrites caused by excessive tertiary dendrite growth was significantly higher in *lin-14*(n179rf) mutants than in wild-type animals at the young adult stage (Figures 2A, 2D, 2G, and S1). Thus, a reduction of function in the *lin-14* gene causes precocious and excessive dendritic outgrowth. Precocious PVD dendritic outgrowth in *lin-14*(n179rf) mutants was not caused by precocious PVD cell fate specification since PVD cells were not specified prematurely in *lin-14*(n179rf) mutants (data not shown). The *lin-14*(n179rf) mutant phenotype of excessive PVD tertiary dendrite growth can be rescued by re-expressing the *lin-14* gene in the PVD neuron, suggesting that *lin-14* acts cell-autonomously in PVD (Figure 2G). To further strengthen the statement that the *lin-4* microRNA targets the *lin-14* transcription
factor to initiate dendritic outgrowth in PVD neurons, we tested whether *lin-14(n179rf)* mutations suppress the *lin-4(e912lf)* mutant phenotype of limited dendritic outgrowth. We found that this is indeed the case (Figure 2F). *lin-4(lf); lin-14(rf)* double mutants displayed precocious dendritic outgrowth phenotypes of the *lin-14(n179rf)* single mutant rather than limited dendritic outgrowth phenotypes of the *lin-4(lf)* single mutant (Figure 2F). Taken together, our results indicate that the *lin-4-lin-14* regulatory circuit initiates dendritic outgrowth in PVD neurons (Figure 7E).

The *let-7-lin-41* regulatory circuit slows down dendrite growth in the final stage of dendrite arborization

We previously reported that the *let-7* microRNA represses the expression of the LIN-41 tripartite motif protein to inhibit AVM axon regeneration in older neurons\(^{23,25}\). Here, we further studied the expression of *let-7* and *lin-41* reporters and found overlapping expression of two genes in many other neurons, including PVD, at the L4 stage (Figures 1D-1F and 1I), when growth of the terminal quaternary dendrites in PVD comes to an end. We performed laser dendritomy on the primary dendrite in PVD neurons at different developmental stages and found that PVD dendritic growth ability is significantly lower at the adult stage than at the L3 stage (Figures 3A, 3B, and 3E), suggesting that PVD neurons undergo a developmental decline in dendritic growth ability. Our expression analysis showed that the *let-7* expression in PVD was at relatively low levels at the L3 stage, but was significantly elevated from the L4 stage onward, which implicates its contribution to the developmental decline in PVD dendritic growth ability (Figure 1K). Indeed, the dendritic growth ability in adult *let-7* mutants was indistinguishable from that seen in wild-
type animals at an earlier developmental stage, the L3 stage (Figures 3A-3E), suggesting that let-7 mutations may retard a normal developmental decline in dendritic growth ability. In the young adult stage, while let-7(n2853rf) mutations significantly enhanced, lin-41(n2914lf) mutations significantly reduced dendritic growth ability in PVD neurons (Figure 3F). The lin-41(n2914lf) mutant phenotype in PVD dendrites can be rescued by re-expressing the lin-41 gene in the PVD neuron, suggesting that lin-41 acts cell-autonomously in PVD (data not shown). In addition, lin-41 mutations suppressed the let-7(n2853rf) mutant phenotype of enhancing dendrite growth ability (Figure 3F). Together, these results indicate that the let-7-lin-41 regulatory circuit slows down dendrite growth in the final stage of dendrite arborization (Figure 7E).

**lin-28 inhibits the let-7-lin-41 circuit to regulate the timing of dendrite arborization**

Previous studies have shown that the LIN-28 RNA-binding protein blocks maturation of the let-7 microRNA in both invertebrates and vertebrates33-36. We found that a lin-28::GFP fosmid-based reporter is expressed in PVD neurons, which contains the lin-28 upstream cis-regulatory, exonic, intronic, and downstream cis-regulatory sequences (Figure 3G). To determine whether lin-28 acts upstream of let-7 to regulate the timing of PVD dendrite arborization, we first compared the endogenous LIN-41 protein level in PVD neurons between wild type and lin-28(n719lf) mutants. The mNG reporter gene was knocked in the endogenous lin-41 locus to generate a mNG::lin-41 fusion gene using the CRISPR/Cas9 technology. We found that lin-28(n719lf) mutations significantly reduced LIN-41 protein levels in PVD neurons compared to wild type at the early L3 stage (Figures 3H and 3I), suggesting that lin-28 inhibits the let-7-lin-41 circuit in PVD neurons (Figure 7E). We
performed laser dendritomy on PVD primary dendrites in lin-28(n719lf) mutants and observed significantly reduced dendrite growth ability in lin-28(n719lf) mutants compared to wild-type animals (Figure 3J), a phenotype that is opposite to the let-7(n2853rf) mutant phenotype of enhanced dendrite growth ability (Figure 3F). Furthermore, lin-41 overexpression in PVD neurons suppressed the lin-28(n719lf) mutant phenotype of reduced dendrite growth ability (Figure 3J). Together, these results support that lin-28 inhibits the let-7-lin-41 circuit to regulate the timing of PVD dendrite arborization.

Recent studies in sexually dimorphic nervous system differentiation and male tail tip morphogenesis revealed that the lep-5 lncRNA brings together LIN-28 and the LEP-2 Makorin to promote ubiquitination and degradation of LIN-28. To determine whether lep-5 plays a role in regulating PVD dendrite growth ability, we first analyzed the endogenous LIN-41 protein level in PVD neurons in wild-type animals versus lep-5(ny28lf) mutants. We found there is no difference of LIN-41 protein levels in PVD neurons between wild type and lep-5(ny28lf) mutants (Figures S2A and S2B). In addition, lep-5(ny28lf) mutants displayed similar dendrite growth ability to wild-type animals (Figure S2C). Thus, lep-5 is unlikely to regulate the timing of PVD dendrite arborization by modulating the lin-28-let-7-lin-41 circuit.

**The lin-4 to let-7 promoter replacement delays dendrite arborization**

To further support our statement that lin-4 and let-7 microRNAs set start and end times for PVD dendrite arborization, we manipulated the timing of their expression by swapping their promoters with each other. We found that delayed expression of lin-4
microRNA expressed from the let-7 promoter from the L3 stage onward) postponed dendrite arborization in PVD neurons (Figure S3). The transgene that expressed the lin-4 microRNA from the let-7 promoter (Plet-7::lin-4) led to retarded growth in quaternary dendrites (the final order of PVD dendrites) in a lin-4 loss-of-function mutant (Figures S3A and S3B). However, the growth of quaternary dendrites in these transgenic animals (lin-4(e912lf); Ex[Plet-7::lin-4]) was able to catch up later in the adult stage (Figure S3C). To further strengthen this conclusion, we utilized the CRISPR/Cas9 technology to replace the endogenous lin-4 promoter at the lin-4 locus with a late-onset let-7 promoter (Figure 4A’). The repair templates that have been developed recently were used to facilitate the identification of the CRISPR recombinants\textsuperscript{39,40} lin-4(xr70) and lin-4(xr71) in which the endogenous lin-4 promoter has been replaced by the let-7 promoter. We used stem-loop RT-PCR to globally survey temporal expression of mature lin-4 microRNA during animal development. Expression of the lin-4 microRNA in the lin-4 to let-7 promoter replacement CRISPR allele was indeed delayed compared to its expression in wild-type animals (Figure 4A). In these CRISPR lines, lin-4(xr70) and lin-4(xr71), we observed retarded growth of the quaternary dendrites at the early L4 stage (Figure 4D compared to 4B of WT; Figure 4F), which was able to catch up later in the adult stage (Figure 4E compared to 4C of WT; Figures 4G-4I). These results demonstrate that, by manipulating the timing of lin-4 expression through the lin-4 to let-7 promoter replacement, we can delay dendrite arborization.

The let-7 to lin-4 promoter replacement precociously inhibits dendrite growth ability
Conversely, premature expression of let-7 (let-7 microRNA expressed from the lin-4 promoter from the L1 stage onward) precociously inhibited dendritic growth ability in PVD neurons (Figures 5A and 5B). We utilized the CRISPR/Cas9 technology to generate the let-7(xr67) CRISPR line in which the endogenous let-7 promoter at the let-7 locus has been replaced with an early-onset lin-4 promoter (Figure 5A’). Stem-loop RT-PCR analysis showed that expression of mature let-7 microRNA in the let-7 to lin-4 promoter replacement CRISPR line was indeed precocious compared to its expression in wild-type animals (Figure 5A). In the let-7(xr67) CRISPR line, PVD dendrite growth ability at the L3 stage was significantly lower than that in wild-type animals at the L3 stage and similar to that in wild-type animals at the young adult stage (Figure 5B). PVD dendrite growth ability in the let-7(xr67) line was further reduced at the young adult stage (Figure 5B). Interestingly, PVD dendrites, both proximal and distal segments to the injured site, in let-7(xr67) animals degenerated 24 hours following laser dendritomy at the D1 A stage (one day into the adult stage). Because of this reason, we were unable to determine the dendrite growth ability at the D1 A stage (Figure 5B). Thus, by manipulating the timing of let-7 expression through the let-7 to lin-4 promoter replacement, we can precociously inhibit dendrite growth ability. Taken together, these findings support a model in which the lin-4-lin-14 regulatory circuit sets the start time whereas the let-7-lin-41 regulatory circuit sets the end time for PVD dendrite arborization (Figure 7E).

*lin-14 and lin-41 antagonistically regulate the DMA-1 receptor level on PVD dendrites*
Two transmembrane ligands, SAX-7 and MNR-1, work together with LECT-2, a secreted ligand, to instruct dendrite arborization in PVD neurons through direct interactions with the dendrite receptor DMA-1\(^{32,41-44}\). SAX-7, MNR-1, LECT-2 and DMA-1 form a multi-protein receptor-ligand signaling complex that directs the growth of stereotyped dendritic branches. One way to control dendrite growth ability during PVD dendrite arborization is through regulation of responsiveness of dendrites to growth signals, which can be accomplished by adjusting the abundance of the DMA-1 receptor on dendrites\(^{45}\). To determine whether the \textit{lin-4-lin-14} and the \textit{lin-28-let-7-lin-41} regulatory circuits control PVD dendrite arborization through the regulation of the DMA-1 receptor on PVD dendrites, we analyzed the level of DMA-1 proteins on specific PVD dendrite branches in \textit{lin-14}(n179rf) and \textit{lin-41}(n2914lf) mutants versus wild-type animals. Using the CRISPR/Cas9 technology to knock in the YFP reporter gene to the 3’ end of the endogenous \textit{dma-1} locus to generate a \textit{dma-1::yfp} fusion gene\(^{46}\), we showed that the endogenous level of DMA-1 proteins based on the fluorescent intensity of YFP was significantly increased on the tertiary dendrites in \textit{lin-14}(n179rf) mutants (Figures 6A-6C), consistent with higher dendrite growth ability, which is opposite to lower dendrite growth ability displayed by \textit{lin-4}(e912lf) mutants (Figure 6D). In contrast, the endogenous level of DMA-1 proteins was significantly reduced on the quaternary dendrites in \textit{lin-41}(n2914lf) mutants (Figures 7A-7C), consistent with lower dendrite growth ability. This C-terminal tagging seems to partially affect the function of DMA-1 receptors as the number of tertiary and quaternary dendrites is reduced compared to wild-type animals. To further examine the relationship between the \textit{lin-4-lin-14} regulatory circuit and \textit{dma-1}, we analyzed the dendrite growth ability in \textit{dma-1}(xr50lf); \textit{lin-14}(n179rf) double mutants. The \textit{dma-}
1(xr50lf) mutation, which significantly reduced tertiary dendrite outgrowth, suppressed the excessive tertiary dendrite outgrowth caused by the lin-14(n179rf) mutation (Figure 6E). These results suggest that the lin-4-lin-14 regulatory circuit promotes dendrite outgrowth through up-regulation of the DMA-1 receptor level on dendrites (Figure 7E). To further investigate the relationship between the lin-28-let-7-lin-41 regulatory circuit and dma-1, we performed laser dendritomy on PVD neurons in lin-41(ma104rf) mutants with or without dma-1 overexpression in PVD neurons. The dma-1(xr50lf) mutation significantly reduced dendrite growth ability (Figure 7D). In contrast, dma-1 overexpression in PVD neurons, which significantly enhanced dendrite growth ability, suppressed the reduced dendrite growth ability caused by the lin-41(ma104rf) mutation (Figure 7D). Thus, the lin-28-let-7-lin-41 regulatory circuit inhibits dendrite growth ability through down-regulation of the DMA-1 receptor level on dendrites (Figure 7E). lin-41 has been reported to negatively regulate the lin-29 zinc finger transcription factor in certain cellular contexts in C. elegans15,21,23,47. Although lin-29 is expressed in PVD neurons strongly from the L4 stage onward (Figures S4A and S4B), the lin-29(n333lf) mutation did not affect dendrite growth ability (Figure S4C). Thus, lin-29 is unlikely to act between lin-41 and dma-1 to regulate the timing of PVD dendrite arborization. Together, these results support a model in which the lin-4-lin-14 and the lin-28-let-7-lin-41 regulatory circuits control the timing of PVD dendrite arborization through antagonistic regulation of the DMA-1 receptor level on PVD dendrites (Figure 7E).
Discussion

PVD neurons lose dendrite growth ability as they age, but it is not known why. One theory is that the developmental decline in dendrite growth ability provides a switch in intrinsic states from long-range exploratory growth to short-range dendritic targeting. In this report, we show that two microRNA regulatory circuits are used in timing dendritic arborization in postmitotic PVD neurons, restricting dendrite growth within a defined time frame. The precisely timed lin-4-lin-14 regulatory circuit sets off initial dendrite outgrowth until the L4 stage, at which time point the precisely timed lin-28-let-7-lin-41 regulatory circuit decelerates dendrite growth as terminal dendrite branches are reaching final targets (Figure 7E). These two regulatory circuits control the timing of PVD dendrite arborization through opposed regulation of the DMA-1 receptor level on PVD dendrites (Figure 7E).

The neuronal timers that restrain dendrite arborization within a specific time window during neural circuit formation are poorly understood and remain a mystery. In this study, we uncover two microRNA regulatory circuits that temporally control the choreography of dendrite arborization in PVD neurons. Our study illustrates how neuronal timers regulate the intrinsic potential of dendrite growth. From this point forward, our goal is to describe the required timing mechanisms in dendritic arborization at a resolution that ultimately will allow us to reconstitute the process. Further dissection of genetic networks that regulate the timing of lin-4 and let-7 expression, and identification of downstream targets of the LIN-14 transcription factor and the LIN-41 tripartite motif protein that regulate the DMA-1 receptor level will be indispensable to achieving this goal.
Here, we show that the *lin-4-lin-14* regulatory circuit initiates PVD dendrite arborization through up-regulation of the DMA-1 receptor level. However, the *dma-1* gene is unlikely a direct target of the LIN-14 transcription factor as the *lin-14* mutation not only enhanced the expression of the endogenous *dma-1* gene from its own promoter (Figure 6), but also enhanced the expression of the *dma-1* transgene from a heterologous *ser-2* promoter (data not shown). We report that the *let-7-lin-41* regulatory circuit slows down PVD dendrite arborization as it comes to an end through down-regulation of the DMA-1 receptor level. The regulation of *dma-1* expression by *lin-41* appears to be dependent on regulatory sequences outside of the *dma-1* coding region since the expression of the endogenous *dma-1* gene from the *dma-1* locus (Figure 7), but not the expression of the *dma-1* transgene from a heterologous *ser-2* promoter and a control *unc-54* 3’ UTR is reduced in *lin-41(lf)* mutants (data not shown). It remains to be seen whether *lin-41* functions as RNA-binding proteins to post-transcriptionally regulate the *dma-1* expression that is dependent on either 5’ UTR or 3’ UTR of the *dma-1* gene. Although *lin-29*, a known direct target of LIN-41 in certain cellular contexts, is expressed in PVD neurons (Figures S4A and S4B), no enhancement in dendrite growth ability by the *lin-29(n333lf)* mutation was observed (Figure S4C), which suggests that *lin-29* is unlikely to play a role in regulating the timing of PVD dendrite arborization. Thus, further studies are required to understand the mechanisms by which LIN-14 and LIN-41 regulate the DMA-1 receptor level on PVD dendrites.

In the *let-7(xr67)* CRISPR line, the *let-7* microRNA expressed from the *lin-4* promoter does not block PVD dendrite growth in early development. This could be due to many
different reasons. For example, the *Plin-4::let-7* genomic configuration may not be able to support expression and processing of the primary *let-7* microRNA as efficiently as the *Plin-4::lin-4* genomic configuration supports expression and processing of the primary *lin-4* microRNA. Alternatively, between the *lin-4-lin-14* and the *let-7-lin-41* regulatory circuits, the *lin-4-lin-14* regulatory circuit could impact the DMA-1 receptor level to a larger extent than the *let-7-lin-41* regulatory circuit.

It remains to be seen whether the timing mechanisms of dendrite arborization by *lin-4* and *let-7* microRNA regulatory circuits can be extended beyond PVD neurons, especially knowing that these two timing microRNAs are expressed broadly in many neurons in *C. elegans* (Figures 1A-1F). Since *let-7* and *lin-4* microRNAs are evolutionarily conserved, it is possible that these microRNA regulatory circuits control the timing of dendrite arborization in other organisms as well. This possibility is supported by the findings that *let-7* is expressed in the nervous system in various species, including *C. elegans, Drosophila, mouse, and human* and that *miR-125*, the *lin-4* homolog, functions in the *Drosophila* and mouse nervous system. Future studies in other systems could determine whether temporal control of dendrite arborization by the two microRNA regulatory circuits is also evolutionarily conserved. Although little is known about the timing of dendrite arborization in vertebrates, research has suggested that failed stabilization of dendritic branches in adults could lead to psychiatric and neurodegenerative disorders. Our studies thus provide important mechanistic insights into the timing of dendrite arborization, a critical step in forming functional neuronal circuits.
Experimental Procedures

Strains and plasmids

*C. elegans* strains were cultured using standard methods\(^53\). All strains were grown at 20°C, except for experiments that involved the *lin-14(n179ts)* allele, which was grown at 25°C. Standard protocol was used for the plasmid constructions. Strains and plasmids used in this study are listed in Tables S1 and S2.

Transgenic animals

Germline transformation of *C. elegans* was performed using standard techniques\(^54\). For example, the *Plin-41::GFP* transgene was injected at 50 ng/µl along with the coinjection marker *Podr-1::RFP* at 50 ng/µl. Transgenic lines were maintained by following the *Podr-1::RFP* fluorescence.

CRISPR/Cas9 genome editing

We generated the *lin-4(xr70), lin-4(xr71), let-7(xr67),* and *let-7(xr68)* CRISPR lines using the CRISPR/Cas9 genome editing technology. CRISPR recombinants were identified using a self-excising drug selection cassette as previously described\(^39,40\). The strategies are illustrated in the diagrams in Figures 4A’ and 5A’.

Stem-loop reverse transcription-PCR

We quantified the mature microRNA level by modifying the microRNA assay developed previously\(^55\). Equal amounts of the RNA preparation from staged wild type, *lin-4(xr71)*, and *let-7(xr67)* animals were used for RT-PCR amplification of mature microRNAs and of
*pmp-3* transcripts. Reverse transcription reactions contained purified total RNA, 50 nM stem-loop RT primer, 1X RT first strand buffer, 0.25 mM each of dNTPs, 10 mM MgCl₂, 0.1 M DTT, 200 U SuperScript III reverse transcriptase and 40 U RNase inhibitor. The mixture of RNA template, dNTPs and RT primer was incubated for 5 min at 65°C. The mixture was then placed on ice for at least 1 min before adding RT buffer, DTT, MgCl₂, SuperScript III and RNase inhibitor. The reaction was incubated for 50 min at 50°C before heat inactivation at 85°C for 5 min. PCR was conducted using 0.25 µl RT products as template in 20 µl PCR for 17 cycles.

RT_primer_lin-4   TCAACTGGTGTCGTTGGAGCTCGGCAATTCACTTTCTGGTGTCGTTGGAGCTCGGCAATTCACTTT
lin-4_F          CGGCGGTCCCTGAGACCTCAA
RT_primer_let-7   CTCAACTGGTGTCGTTGGAGCTCGGCAATTCACTTTCTGGTGTCGTTGGAGAACTATAC
let-7_F          CGGCGGTGAGGTTAGTGGTTGT
Universal reverse primer  CTGGTGTCGTTGGAGGCGAATTCC

*pmp-3*  TGGCCGGATGATGTTGTCGC
*pmp-3*  ACGAACAATGCGCCAAGGCCAGC

**Laser dendritomy**

Animals were mounted on 2% agarose pads and anesthetized with 5 mM sodium azide, the lowest possible concentration to keep adult animals immobilized. Laser dendritomy was performed on PVD primary dendrites using either a cavity-dumped Ti:sapphire laser oscillator (Cascade Laser, KMLabs Inc.), which generates laser pulses ~100 fs in duration.
and 200 kHz in repetition rate, or a MicroPoint Laser Ablation System (Andor/Oxford Instruments), which generates 337 nm laser pulses 2-6 ns in duration. The laser pulses were tightly-focused onto targeted primary dendrites using a Plan Apo VC 100x, 1.4 NA oil-immersion objective on a Nikon ECLIPSE Ti microscope. Successful laser dendritomy was confirmed by visualizing the targeted area immediately after surgery. Worms were recovered within 10 minutes of sodium azide treatment and placed on fresh plates with bacterial food.

**Quantification of dendrite lengths**

PVD neurons in recovered worms were imaged 24 hours after dendritomy and the dendrite regrowth was quantified. 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**

Animals were mounted on 2% agarose pads and anesthetized with 20 mM sodium azide. Fluorescence microscopy was performed using either a Plan-Apochromat 60x, 1.4 NA objective on a Zeiss Axio Imager M2 microscope with a Hamamatsu ORCA-Flash4.0 LT+ camera, or a Plan Apo VC 60x, 1.4 NA objective on a Nikon ECLIPSE Ti microscope with a Hamamatsu ORCA-ER camera. The morphology of neuronal cell bodies and dendrites was based on high-magnification Z-stacks. Images of DMA-1-YFP were captured in live
animals using a Plan-Apochromat 40x, 1.3 NA objective on a Zeiss LSM 880 confocal microscope.

**Statistical analysis**

Average data of dendrite number, dendrite length, promoter reporter expression intensity, and relative DMA-1::YFP intensity are presented as means ± SEM. Data of % PVD neurons with excessive or limited dendrites are presented as proportions ± SEP. Statistical analyses were carried out by Student’s t-tests or two-proportion Z-tests using GraphPad Prism 7.0 or the Primer of Biostatistics software. The statistical test used for each panel is described in the figure legend. Sample sizes for experiments are shown in the respective panels. In all figures, n represents number of animals examined.
Author Contributions

N.S. conceived, designed, performed, analyzed experiments, made constructs, and drafted the article. H.C. conceived, designed, performed, analyzed experiments, and drafted the article. Y.Z. conceived, designed, performed, analyzed experiments, and made constructs. M.S. made the promoter replacement CRISPR lines, performed, and analyzed experiments. W.Z. made the \textit{dma-1(wy996)} [\textit{dma-1::YFP}] CRISPR line and contributed unpublished essential data and reagents. K.S. conceived experiments, contributed unpublished essential data and reagents, and drafted the article. C.C. conceived, designed, analyzed and interpreted data, and drafted the article.

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Declaration of Interests
The authors declare no competing interests.
References


Figure legends

Figure 1. Expression of lin-4/lin-14 and let-7/lin-41 regulatory circuits in PVD neurons. (A-C) Overlapping expression of Plin-4::GFP and Plin-14::mCherry reporters in neurons of many regions, such as the head ganglia, ventral nerve cord, tail, and mid-body region at the third larval (L3) stage. (D-F) Overlapping expression of Plet-7::GFP and Plin-41::mCherry reporters in many neurons, in the head ganglia, ventral nerve cord, tail, and mid-body region at the L4 stage. PVD neuron expression was also indicated. Scale bar, 20 µm. (G) The timing and steps of PVD dendrite arborization. Schematic drawings of PVD dendrite arbors as they are seen at five different stages of development, late larval stage 2 (L2) to late L4. All views showing the left-side cells of each PVD pair; anterior is to the left. A single axon (grey color) emerges ventralward from the cell body before traveling anteriorly along the ventral nerve cord. The dendrite processes emerging from the cell body elaborate into highly organized dendrite arbors. (H) Both Plin-4::GFP and Plin-14::GFP reporters are expressed in PVD neurons. (I) Detection of both Plet-7::GFP and Plin-41::GFP expression in PVD neurons. Scale bar, 10 µm. (J) Expression of the Plin-4::GFP reporter in PVD neurons was assessed at five different stages of development in wild-type animals. (K) Expression of the Plet-7::GFP reporter in PVD neurons was assessed at four different stages of development in wild-type animals. Each line represents data from a single animal followed over time. Bars represent the average expression intensity of either the Plin-4::GFP or the Plet-7::GFP reporter measured at each time point. Error bars, SEM. ***p < 0.001 by a Student’s t-test.
Figure 2. Initiation of dendritic arborization is affected by mutations in the lin-4-lin-14 regulatory circuit. (A-D) Representative images showing extent of dendrite arborization in wild type, lin-4(e912lf), lin-14(n355gf), and lin-14(n179rf) mutants. Arrowheads point to contacts between neighboring tertiary dendrites. Scale bar, 20 µm. (E) Percentages of PVD neurons at the young adult stage with limited dendrite outgrowth in wild type, lin-4(e912lf), and lin-14(n355gf) mutants. Error bars, SEP. ***p < 0.001, relative to wild type, by a two-proportion Z-test. (F) Percentages of PVD neurons at the mid L3 stage based on the highest order dendrite branch observed in wild type, lin-4(e912lf), lin-14(n179rf), and lin-4(e912lf); lin-14(n179rf) mutants. (G) Quantification of the number of tertiary branches per 250 µm in the anterior direction from the PVD cell body at the young adult stage in wild type, lin-14(n179rf) mutants, lin-14(n179rf) mutants carrying the Pser-2::lin-14 transgene, animals carrying the Pser-2::lin-14 transgene, and lin-14(n355gf) mutants. Tertiary dendrites were divided into two groups: non-overlapped tertiary is defined as those with normal self-avoidance and overlapped tertiary as those with self-avoidance defects. Error bars, SEM. ***p < 0.001 by a Student’s t-test.

Figure 3. Developmental decline in dendrite growth ability is affected by mutations in the lin-28-let-7-lin-41 regulatory circuit. (A-D) Representative images showing extent of dendrite regrowth 24 hours after dendritomy of the primary dendrite at either the L3 or the young adult stage in wild type (A, B) and let-7(n2853rf) mutants (C, D). PVD dendrites were visualized using the xrls37[Pf49H12.4::GFP] marker. Dorsal is up; anterior is to the left. Red arrows indicate lesion sites. Scale bar, 20 µm. (E) Average PVD dendrite length regrown in wild type and let-7(n2853rf) mutants 24 hours following dendritomy at
different stages. Asterisks indicate cases in which later stage animals differ from L3-stage animals at ***p < 0.001 by a Student’s t-test. D1 A indicates one day into the adult stage. Error bars indicate SEM. (F) Average dendrite length regrown in wild type, let-7(n2853rf), lin-41(n2914lf), and lin-41(n2914lf); let-7(n2853rf) mutants. PVD primary dendrites were severed by laser surgery at the young adult stage. Dendrite lengths were measured 24 hours after dendritomy as the actual contour length between the injury site and dendrite termini by tracing dendrites through a 3-dimensional image stack. Asterisks indicate cases in which let-7 or lin-41 mutants differ from wild type at ***p < 0.001 by a Student’s t-test. NS, not significant. (G) Representative images of the expression of LIN-28 proteins in PVD neurons in wild type at the early L3 stage. A lin-28::GFP fosmid-based reporter is expressed in PVD neurons. The Pser-2::mCherry reporter was used to label PVD neurons. Scale bar, 5 µm. (H) Representative images of the expression of endogenous LIN-41 proteins in PVD neurons in wild type and lin-28(n719lf) mutants at the early L3 stage. The Pser-2::mCherry reporter was used to label PVD neurons. Scale bar, 5 µm. (I) Quantification of endogenous LIN-41 proteins based on the mNG::LIN-41 fluorescent intensity in the PVD cell body in wild type and lin-28(n719lf) mutants. (J) Average dendrite length regrown in wild type, lin-28(n719lf) mutants, and lin-28(n719lf) mutants overexpressing lin-41 in PVD neurons. Error bars, SEM. ***p < 0.001 by a Student’s t-test.

Figure 4. Delayed dendrite arborization by the lin-4 to let-7 promoter replacement. Expression of the lin-4 microRNA by a late-onset let-7 promoter postponed growth of the quaternary dendrites. (A) Stem-loop RT-PCR analysis of RNA isolated from populations
of staged animals revealed late onset expression of the lin-4 microRNA in the lin-4(xr71) CRISPR line contrast to early onset expression in wild-type animals. (A’) Strategies of promoter replacements by the CRISPR/Cas9 technology. (B-E) Representative images showing extent of quaternary dendrite arborization in wild type (B, C) and the lin-4(xr71) CRISPR line (D, E), in which the endogenous lin-4 promoter has been replaced by the let-7 promoter. Images were taken at the early L4 (B, D) and the young adult (C, E) stages. Dorsal is up; anterior is to the left. Scale bar, 20 µm. (F-I) Quantification of the number of quaternary branches per 250 µm in the anterior direction from the PVD cell body at the early L4 stage (F) or various time points at the young adult stage (G-I) in wild type, lin-4(xr70) CRISPR line, and lin-4(xr71) CRISPR line. Each dot represents data from a single animal. ***p < 0.001 by a Student’s t-test.

Figure 5. Precocious decline in dendrite growth ability by the let-7 to lin-4 promoter replacement. Expression of the let-7 microRNA by an early-onset lin-4 promoter precociously reduced dendrite growth ability in PVD neurons. (A) Stem-loop RT-PCR analysis of RNA isolated from populations of staged animals revealed early onset expression of the let-7 microRNA in the let-7(xr67) CRISPR line contrast to late onset expression in wild-type animals. (A’) Strategies of promoter replacements by the CRISPR/Cas9 technology. (B) Average PVD dendrite length regrown in wild type and the let-7(xr67) CRISPR line 24 hours following dendritomy of the primary dendrite at different stages. Asterisks indicate cases in which later stage animals differ from L3-stage animals or a comparison between wild type and let-7(xr67) is significantly different at ***p < 0.001.
by a Student’s $t$-test. D1 A indicates one day into the adult stage. ND represents “not determined”. Error bars indicate SEM.

**Figure 6.** *lin-14* negatively regulates DMA-1 protein levels on PVD dendrites. (A, B) Representative images showing DMA-1::YFP fusion proteins on PVD dendrites in wild type (A) and *lin-14(n179rf)* mutants (B). Arrowheads point to tertiary dendrites. Dorsal is up; anterior is to the left. Scale bar, 20 µm. (C) Average fluorescent intensity of DMA-1::YFP fusion proteins on PVD tertiary dendrites in wild type (*dma-1(wy996)*) versus *lin-14(n179rf)* mutants (*dma-1(wy996); lin-14(n179rf)*). Error bars, SEM. ***p < 0.001 by a Student’s $t$-test. (D) Quantification of the number of secondary dendrites per 300 µm in the anterior direction from the PVD cell body at the young adult stage in wild type and *lin-4(e912lf)* mutants. Each dot represents data from a single animal. (E) Percentages of PVD neurons at the young adult stage with excessive or limited tertiary dendrite outgrowth in *lin-14(n179rf)*, *dma-1(xr50lf)*, and *dma-1(xr50lf); lin-14(n179rf)* mutants. ***p < 0.001 by a Student’s $t$-test.

**Figure 7.** *lin-41* positively regulates DMA-1 protein levels on PVD dendrites. (A, B) Representative images showing DMA-1::YFP fusion proteins on PVD dendrites in wild type (A) and *lin-41(n2914lf)* mutants (B). Arrowheads point to quaternary dendrites. (C) Average fluorescent intensity of DMA-1::YFP fusion proteins on PVD quaternary dendrites in wild type (*dma-1(wy996)*) versus *lin-41(n2914lf)* mutants (*dma-1(wy996) lin-41(n2914lf)*). (D) Average PVD dendrite length regrown in wild type, *lin-41(ma104rf)* mutants, *dma-1(xr50lf)* mutants, animals overexpressing *dma-1* in PVD neurons, and *lin-
41(ma104rf) mutants overexpressing dna-1 in PVD neurons 24 hours following dendritomy of the primary dendrite at the young adult stage. Error bars, SEM. *p < 0.05 and ***p < 0.001 by a Student’s t-test. (E) Model of sculpting dendritic arbors by two precisely timed microRNA regulatory circuits. Initially, the lin-4 microRNA down-regulates the lin-14 transcription factor to trigger dendrite arborization. Later, the let-7 microRNA down-regulates the lin-41 tripartite motif protein to slow down dendrite growth in the final stage of dendrite arborization.
% PVD neurons with limited dendrite arborization

$\%$ mid L3-stage animals with the specified dendrite as the highest order dendrite branch

Average number of tertiary dendrites per 250 μm

Figure 2 (Chang)
Figure 3 (Chang)
Figure 4 (Chang)
**Figure 5 (Chang)**

**A**

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**B**

Average dendrite length (μm)

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**A’**

5' homologous arm (1.5Kb) let-7 promoter (2.9Kb) 3' homologous arm (1.5Kb)

sgRNA 1 target

sgRNA 2 target

LoxP LoxP

Self Excising Cassette (SEC)

3xFLAG

Homologous recombination

Heat shock-induced SEC excision

lin-4 promoter

let-7

**Figure 5 (Chang)**
**Figure 6 (Chang)**

**A** dma-1(wy996) [dma-1::yfp]

**B** dma-1(wy996) [dma-1::yfp]; lin-14(n179rf)

**C**

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**D**

Number of secondary dendrites per 300 μm

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**E**

% PVD neurons

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<tr>
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*** indicates significant difference.
**Figure 7 (Chang)**

**A** `dma-1(wy996) [dma-1::yfp]`

**B** `dma-1(wy996) [dma-1::yfp] lin-41(n2914lf)`

**C**

![Graph showing relative DMA-1::YFP intensity on quaternary dendrites](image)

- `n = 23`
- `n = 26`

**D**

![Bar graph showing average dendrite length (μm)](image)

- `n = 20`
- `n = 20`
- `n = 21`
- `n = 27`
- `n = 40`

**E**

- `lin-4`
- `lin-14`
- `dma-1` on 3° dendrites
- `lin-28`
- `let-7`
- `lin-41`
- `dma-1` on 4° dendrites
- `4° dendrite outgrowth`

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