A neurite-zippering mechanism, mediated by layer-specific expression of IgCAMs, regulates synaptic laminar specificity in the *C. elegans* nerve ring neuropil

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Abstract

A fundamental design principle of nervous systems is the grouping of neuronal contacts into layers within nerve bundles. The layered arrangement of neurites requires nanoscale precision in their placement within bundles, and this precision, which cannot be exclusively explained by simple tip-directed outgrowth dynamics, underpins synaptic specificity and circuit architecture. Here we implement novel imaging methods and deep learning approaches to document the specific placement of single neurites during the assembly of the *C. elegans* nerve ring. We uncover a zippering mechanism that controls precise placement of neurites onto specific layer subdomains. Nanoscale precision in neurite placement is orchestrated via temporally-regulated expression of specific Ig adhesion molecules, such as SYG-1. Ig adhesion molecules act as instructive signals, defining sublaminar regions and guiding neurite zippering onto target neurons. Our study reveals novel developmental mechanisms that coordinate neurite placement and synaptic specificity within layered brain structures.
Introduction

In brains, neurites are segregated away from the cell bodies into synapse-rich regions termed neuropils: dense structures of nerve cell extensions which commingle to form functional circuits (Maynard, 1962). Neuropils are units of functional organization within brains, and within the precisely arranged architecture of the neuropil, placement of neurites into specific neighborhoods is a major determinant of synaptic specificity and circuit connectivity (Maynard, 1962; Schürmann, 2016; Soiza-Reilly & Commons, 2014; Xu et al., 2020; Zheng et al., 2018). Proper functional connectivity therefore depends on design principles that guide placement of neurites, at nanoscale precision, within neuropils.

A fundamental design principle of neuropil organization is the segregation of neurites into laminae and sublaminae, and this laminar organization principle is conserved and observed in neuropils of both vertebrate and invertebrates (Gabriel et al., 2012; Millard & Pecot, 2018; Sanes & Zipursky, 2010). The laminar structural organization is key to the assembly and functional segregation of specific circuits (Gabriel et al., 2012; Kolodkin & Hiesinger, 2017; Millard & Pecot, 2018; Nevin et al., 2008). For example, in the inner plexiform layer (IPL) of the vertebrate retina, synaptic connections conveying different types of visual information spatially segregate onto distinct sublayers. Placement of neurites with similar functions into specific sublayers restricts synaptic partner choice, driving synaptic specificity (Robles et al., 2013). Therefore, the co-segregation of neurons with similar response properties into specific layers gives rise to topographic maps in which structural principles underpin functional
principles in the precise assembly, and segregation, of distinct circuits (Clandinin & Feldheim, 2009).

Within the layered organization of neuropils, subsets of neurons are capable of specifically projecting onto multiple layers, enabling integration of information across laminar circuits. For example, glycinergic and GABAergic amacrine cells in the IPL of the retina, and the periventricular projection neurons in the zebrafish tectal neuropil, extend single neurites that link multiple sublayers within their respective neuropils (Demb & Singer, 2012; Kolb, 1995; Kunzevitzky et al., 2013; Robles et al., 2011; Strettoi et al., 1992; Taylor & Smith, 2012). Their precise placement, and the distribution of synapses within neurite regions, underlies their functional roles as integrators of information across the modular and layered circuit architecture (Robles et al., 2011; Strettoi et al., 1992; Taylor & Smith, 2012). How the precise placement of these multilayer-spanning neurites is specified during development, particularly in the context of the assembling neuropil, is not understood.

Molecular genetic studies have revealed roles for cell adhesion molecules (CAMs) and guidance factors in neuronal targeting onto specific layers of the brain. For example, studies in both the mouse and fly visual systems have revealed important roles for IgSF proteins, such as the Sidekick, Dscam and Contactin molecules, in targeting neurons to distinct layers or sublaminae (Sanes & Zipursky, 2020; Tan et al., 2015; Yamagata & Sanes, 2008, 2012). These studies indicate that the expression levels, the timing of expression and the co-expression of specific CAMs influence layer-specific targeting of neurites within neuropils (Petrovic & Hummel, 2008a; Poskanzer et al., 2003; Schwabe et al., 2014). These observations also reveal that our current
mechanistic frameworks of axon guidance and cell-cell interactions are insufficient to provide a conceptual understanding of how nanoscale organization within neuropils emerges during development. In particular, how single cells make simultaneous and coordinated use of these molecular cues to achieve precise placement, resulting in synaptic specificity within layered neuropils, is poorly understood.

The *C. elegans* nerve ring is a crowded neuropil tens of micrometers thick in which neuronal processes have to discriminate between targets to assemble functional circuits that underpin specific behaviors (Ware et al., 1975; White et al., 1986). We recently demonstrated that the *C. elegans* nerve ring neuropil is a layered structure, with layers (called “strata”) that functionally segregate sensory information and motor outputs (Moyle et al., 2020). A subset of highly interconnected neurons, which form part of a group called ‘rich-club neurons’, serve as hubs that link functionally distinct strata, analogous to amacrine cells in the inner plexiform layer of the vertebrate retina (Marc et al., 2014; Moyle et al., 2020; Towlson et al., 2013). The precise placement of these rich-club neurons along distinct nerve ring strata, and the specific segregation of their synaptic inputs and outputs between these strata, are important for supporting the structure and function of the nematode brain (Gray et al., 2005; Moyle et al., 2020; Wakabayashi et al., 2004).

To determine how neuronal processes are precisely placed onto specific layers during development, and the implications of this placement for synaptic connectivity, we examined the AIB interneurons, a pair of “rich-club neurons” that integrate sensory and motor information across nerve ring strata (Chalasani et al., 2007; Kang & Avery, 2009; Moyle et al., 2020; Sabrin et al., 2019; Towlson et al., 2013). Each AIB neuron projects
a single neurite, and segments of that single neurite are placed along distinct and specific layers in the *C. elegans* nerve ring. The nanoscale precision of AIB neurite placement informs the specificity of AIB synaptic sites in the dense neuropil structure of the nerve ring.

We labeled the AIB neurons for visualization *in vivo* and implemented novel imaging methods and deep learning approaches to yield high-resolution images of AIB during embryonic development. We discovered that placement of the AIB neurite depends on coordinated zippering mechanisms that align segments of the AIB neurite onto specific sublayers. Through forward and reverse genetic screens we uncovered molecular factors important for placement of the AIB neurite. We identified roles for the IgCAM syg-1 in zippering a segment of the AIB neurite onto a layer boundary. We determined that syg-1 expression is layer specific, and temporally controlled to coincide with AIB neurite outgrowth and zippering onto the correct sublayer. Expression of SYG-1 is sufficient to specify AIB neurite placement, and ectopic expression of just the SYG-1 ectodomain results in segments of the AIB neurite being incorrectly positioned at the ectopic syg-1-expressing layers. Our findings uncover a novel zippering mechanism which acts *in vivo* to place neurites along specific layers of the nerve ring neuropil. This mechanism is based on a temporally-coordinated expression of IgCAMs at specific layer subdomains. The developmental processes uncovered in this *in vivo* study might represent conserved mechanisms that enable placement of neurites, and *en passant* synaptic specificity, in layered neuropil structures.
Results

The AIB neurites are placed along distinct strata in the *C. elegans* nerve ring neuropil

To characterize the precise placement and synaptic distribution of the AIB neurites in the context of the nerve ring neuropil strata, we examined available connectome datasets generated across the larval developmental stages of *C. elegans* (White et al., 1986; Witvliet et al., 2020). Synapses in AIB, like in most nerve ring neurons, are formed *en passant*, or along the length of its single neurite. AIB displays polarity in the distribution of its synaptic specializations: postsynaptic specializations are enriched in the proximal neurite (near the cell body), while presynaptic specializations are enriched in the distal neurite (Fig. 1a-c). Examination of the connectomes revealed that the synaptic polarity of AIB is stereotyped, established by the L1 stage and preserved throughout development (Witvliet et al., 2020) (https://nemanode.org/).

Segments of the AIB single neurite reside along distinct strata of the nerve ring (Supplementary Fig. 1a). The AIB proximal neurite, which is enriched in postsynaptic specializations, forms contacts with neighboring amphid sensory neurons in a sub-bundle coincident with S3/S4 sublaminar region, hereafter termed ‘AIB proximal neighborhood’. The AIB distal neurite, which is enriched in presynaptic specializations, contacts neighboring motor interneurons in a sub-bundle coincident with S2/S3 sublaminar region, hereafter termed ‘AIB distal neighborhood’. In the distal neighborhood, the highest number of AIB contacts are formed with the neurites of motor neuron RIM (Supplementary Fig. 1b,c), a major fasciculation partner and postsynaptic
partner of AIB, located at the S2/S3 sublaminar region (Supplementary Fig. 1b). In the proximal neighborhood, AIB receives synaptic inputs from sensory neurons in the S3/4 sublaminar region (Supplementary Fig. 1b,d). Therefore, placement of the AIB neurite onto these distinct neighborhoods links sensory information (from the S3/4 sublaminar region) to motor neuron outputs (in the S2/S3 sublaminar region) in the nerve ring neuropil. This design principle of the AIB neurite is present in early larval stages and preserved up to adulthood (Supplementary Fig. 1b) (Witvliet et al., 2020).

The AIB neurite segments reach the proximal and distal neighborhoods via a shift of the neurite along the anteroposterior axis, precisely at the dorsal mid-line of the nerve ring (herein referred to as the “chiasm”, as the shift results in a cross-over of the two AIB neurites; indicated by arrowheads in Fig. 1a-g, Supplementary Fig. 1e-j). Remarkably, the shift of the AIB neurite is precisely the width of the S3 strata, enabling it to bridge the S3/4 and S2/3 sublaminae with the positioning of its neurite segments (Supplementary Fig. 1). Network analyses on the contacts of AIBL and AIBR with other neurons across the available connectomes revealed similarity in number and distribution of AIB contacts across development (pairwise cosine similarity index S, ranges from 0.62-0.97, S>0.5 representing a positive correlation between datasets, Supplementary Fig. 2a-c also see Methods). This is indicative of a developmental program that establishes contact profiles for AIB during embryogenesis, and is allosterically maintained during post-embryonic growth (Fan et al., 2020). We also found that AIBL and AIBR have high betweenness centrality (a standard property for rich-club neurons (Towlson et al., 2013)) in an L1 (first larval stage after embryogenesis) and an adult connectome dataset (Witvliet et al., 2020), suggesting that AIB functions as a rich
club neuron at early as well as later postembryonic developmental stages (Supplementary Fig. 2d). Moreover, examination of AIB in the connectome of the nematode *Pristionchus pacificus*, which is separated from *C. elegans* by 100 million years of evolutionary time, revealed similar design principles in morphology and placement of the AIB neurite (Hong et al., 2019). Therefore AIB morphology, position and polarity are conserved features of the architecture of the nematode brain, established during embryonic development and are uniquely designed to integrate and relay information across functional modules of the nerve ring. The developmental programs that govern these design principles, enabling nanoscale precision in the placement of the AIB neurite onto specific neighborhoods and the segregation of synaptic specializations, are not understood.

In vivo visualization of AIB synaptic distribution and placement in the nerve ring neuropil

To examine the developmental programs of AIB, we developed cellular and subcellular fluorescent labels for imaging of AIB, its neighborhoods and its synapses in vivo in embryonic stages and postembryonic larval stages. The synaptic distribution that we observed by fluorescent labeling was consistent with the EM connectomic characterizations: presynaptic proteins RAB-3 and CLA-1 localized exclusively to the distal segment of the neurite, while postsynaptic protein GLR-1 localized primarily to the proximal neurite segment (Fig. 1d,e; Polarity index, PI = 0.92 for RAB-3 and 0.24 for GLR-1, where PI>0.5 represents a distal localization and PI<0.5 represents a proximal localization). These markers also enable visualization of the chiasm that result in the
separation of the two AIB neurite segments onto distinct neighborhoods along the antero-posterior axis of the worm (Fig. 1f-I, Supplementary Fig. 1g,h). We found that the chiasm is stereotyped and similar in length across L4 stage animals, as measured from confocal micrographs (mean length = 2.97±0.22 µm, number of neurons measured, n =22), and electron micrographs (dorsal midline shift length in AIBL and AIBR in electron micrographs of an L4 stage animal, JSH, are 3.01 µm and 3.16 µm respectively).

To visualize the AIB neurite in the context of the proximal and distal neighborhoods, we co-labeled neurons that extensively contact (fasciculate) with AIB in the proximal and distal neighborhoods: motor neuron RIM for the distal neighborhood and sensory neurons AWC and ASE for the proximal neighborhood (Supplementary Fig. 1c,d). Consistent with the EM connectomic studies, these representative neighborhood markers overlap with the expected neighborhoods (Fig. 1f-m). Our findings are consistent with and extend previous studies, demonstrating that the AIB unique morphology emerges early in development, is stereotyped across animals and is conserved throughout evolution (Hong et al., 2019; White et al., 1983). Our examination of AIB in the context of electron micrographs from connectomic studies and in vivo imaging reveal that the AIB neurite is designed, by its position and distribution of synapses, to occupy distinct neighborhoods and integrate information across strata of the nerve ring.
Modular developmental programs underlie precise placement and synaptic polarity of the AIB neurites

To understand the cellular and molecular mechanisms underpinning the precise placement of the AIB neurites in the context of the nerve ring neuropil, and how their placement relates to synaptic polarity, we performed unbiased forward genetic screens and candidate screens in postembryonic larval stage animals. Our screens uncovered mutants with defects in AIB neurite outgrowth and placement: a novel mutant allele of transcription factor daf-16/FOXO (ola337) (Supplementary Table 1), lesions in genes encoding cytoskeletal regulators (unc-33 and zyg-8) and lesions in axon guidance genes (vab-1, unc-6 and sax-3). These mutants result in defects in AIB neurite placement in the context of the neighborhoods by affecting AIB outgrowth and/or nerve ring neighborhood development (Fig. 2; Supplementary Fig. 3), (Christensen et al., 2011; Grossman et al., 2013; Yoshimura et al., 2008; Zallen et al., 1999)). Interestingly, inspection of AIB subcellular distribution of pre- and postsynaptic specializations in these mutant backgrounds revealed that when AIB morphology and the nerve ring neighborhoods are affected, the AIB neurites still exhibit a polarized distribution of pre- and postsynaptic proteins (Fig. 2). For instance, even in cases in which the distal AIB neurite segment was largely absent, the neurite retained the polarized distribution of pre- and postsynaptic proteins, with presynaptic components accumulating specifically in the distal segment, or tip, of the truncated neurite and not in the proximal neurite (Fig. 2e-l). The retainment of synaptic polarity even in extreme cases of neurite outgrowth or neighborhood placement defects suggests modularity between the developmental
processes dictating neurite placement and the polarized distribution of pre- and postsynaptic specializations in the context of the proximal and distal neurite segments.

Our genetic findings suggest that synaptic specificity in AIB requires the correct deployment of two modular developmental programs: polarized placement of pre- and postsynaptic specializations along the neurite, and the placement of the neurite onto the correct strata. To specifically understand the mechanisms that regulate placement of the AIB neurite onto the correct strata, we focused our analyses on mutants with nanoscale defects in the placement of the AIB neurite within specific neighborhoods.

The IgCAM SYG-1 is required for precise placement of the AIB distal neurite in the nerve ring

From our screens we discovered that loss-of-function mutant alleles of the Ig superfamily CAM gene syg-1 result in significant defects in placement of the AIB neurite in the nerve ring. In wild type animals, we observed overlap between the AIB distal neurite, and its distal neighborhood partner, RIM (Fig. 3a-d), consistent with electron microscopy characterizations of AIB and RIM fasciculation (Supplementary Fig. 1b). In contrast, in syg-1 mutants the AIB neurites are frequently detached or separated from the RIM neurites (Fig. 3e-h). 70.7% of syg-1(ky652) animals and 62.5% of syg-1(ok3640) animals (as compared to 0% of wild type animals) show regions of AIB-RIM detachment, i.e., failure of the AIB and RIM neurites to fasciculate along their entire length in the distal neighborhood. Cosmid C54A10, containing the syg-1 genomic region, rescued the distal neurite placement defects in syg-1(ky652) (in 87.5% of animals with the rescuing construct, the AIB and RIM neurites show complete overlap.
as in wild type animals) (Fig. 3i-l). In the syg-1(ky652) animals that exhibit defects in AIB neurite placement with respect to RIM, we calculated the percentage of AIB-RIM neurite contact length that exhibits detachment or separation, and found the average percent detachment to be 21.49±4% of total contact length in syg-1(ky652) mutant animals (Fig. 1m, also see Methods). The AIB chiasm is also significantly reduced in length in syg-1 mutants (Supplementary Fig. 4a), and the distal neurite is not positioned at a uniform distance from the proximal AIB neurite (Fig. 1n), consistent with a placement defect of the AIB distal neurite onto the distal neighborhood. We note that in the syg-1(ky652) mutant animals, the cell body position of AIB, AIB’s neurite length and placement of the AIB proximal neurite segment are unaffected (Supplementary Fig. 4b,c), suggesting that syg-1 specifically regulates the placement of the distal neurite segment in the distal neighborhood. Our findings indicate that correct placement of the AIB proximal neurite and the distal neurite are genetically separable. Importantly, our findings indicate that Ig superfamily cell adhesion molecule SYG-1 is necessary for nanoscale placement of the AIB neurite to the distal nerve ring neighborhood.

SYG-1 and its orthologs (Rst and Kirre in Drosophila and Kirrel1/2/3 in mammals) are multipurpose adhesion molecules which function in a wide variety of developmental contexts, including synapse formation in the C. elegans egg-laying circuit, muscle cell fusion, eye patterning and olfactory axon convergence in Drosophila, and formation of the kidney filtration barrier in mammals (S. Bao & Cagan, 2005; Garg et al., 2007; Kim et al., 2015; Serizawa et al., 2006; Shen & Bargmann, 2003; ztokatli et al., 2012). Given SYG-1’s known role in synaptogenesis in C. elegans, we examined the distribution of synaptic sites along the AIB distal neurite in syg-1 mutants. We noted reduction in RAB-
signal in regions of the AIB distal neurite, but specifically for areas lacking RIM contacts (Supplementary Fig. 4g-p). We hypothesized that SYG-1 could mediate synaptogenesis in AIB, and that synapses might then help place AIB in the distal neighborhood. To test this hypothesis we identified, from our forward and reverse genetic screens, additional lesions resulting in synaptogenesis defects in AIB, including a novel allele of syd-2(ola341) (Supplementary Table 1), and cla-1(ok560) (Barstead et al., 2012; Xuan et al., 2017; Zhen & Jin, 1999). We observed that while syd-2 and cla-1 mutants result in an abnormal distribution of presynaptic specializations in AIB (Supplementary Fig. 4q-t), they do not display phenotypes in AIB neurite placement within the distal neighborhood (Supplementary Fig. 4u). Our findings indicate that molecules that affect synaptogenesis do not necessarily result in fasciulation defects for AIB. Together, our findings suggest that SYG-1 plays a role in mediating neurite placement specifically onto the distal neighborhood via fasciculation.

RIM neurons express SYG-1 and regulate AIB neurite placement

To understand how syg-1 regulates the precise placement of the AIB neurite in its distal neighborhood, we next investigated where the syg-1 gene is expressed. We achieved this by co-expressing an AIB reporter and a transcriptional reporter of syg-1 (Schwarz et al., 2009) and examining its expression in the nerve ring of wild type animals. We observed robust expression of the syg-1 transcriptional reporter in a banded pattern in the nerve ring neuropil, with specific enrichment in the AIB proximal and distal neighborhoods (Fig. 4a-d). By quantifying the ratio of mean intensities of the syg-1 reporter in the proximal and distal neighborhoods, we found that on average, syg-
1 expression levels were 3.37 times (S.E.M. = 0.12) higher in the distal neighborhood relative to the proximal neighborhood (Fig. 4i). We then tested whether RIM, the primary distal neighborhood fasciculaton partner of AIB, expresses syg-1. Indeed, co-localization studies revealed syg-1 reporter expression in RIM, but not in AIB, consistent with previous observations (Schwarz et al., 2009) (Fig. 4a-h). Therefore, neurons in the AIB distal nerve ring neighborhood, including RIM, express SYG-1.

As syg-1 mutant animals exhibit defects in AIB neurite placement in the distal neighborhood, and as SYG-1 is expressed in RIM, we hypothesized that the RIM neurons might regulate placement of the AIB distal neurite onto the distal neighborhood. Via promoter screening and a systematic lineage tracing and cell identification pipeline (Z. Bao et al., 2006; Boyle et al., 2006; Duncan et al., 2019; Murray et al., 2006; Santella et al., 2014), we identified two promoters expressed in the RIM neurons prior to embryonic growth and placement of the AIB distal neurite (which occurs approximately 520-570 m.p.f. – see next section and Fig. 5). The promoters identified were inx-19p (expressed in RIM starting ~370 m.p.f (minutes post fertilization)) and tdc-1p (expressed in RIM starting ~445 m.p.f) (see Methods, Supplementary Fig. 5), http://promoters.wormguides.org). We then used these early promoters to drive an in-vivo split caspase ablation system (Chelur & Chalfie, 2007) to successfully eliminate the RIM neurons during embryogenesis (Supplementary Fig. 5d-g). Consistent with our hypothesis, we observed that ablation of RIM results in defects in placement of the AIB distal neurite to the distal neighborhood. These defects in RIM-ablated animals phenocopy the defects seen for the syg-1 mutants. 80.6% and 51.5% of animals had defects in AIB distal neurite placement in RIM-ablated animals (using two strategies-
see Supplementary figure 5 and Methods), compared to 3.1% in a wild type population; Fig. 4j-q). These observations demonstrate that SYG-1-expressing RIM neurons are necessary for the precise placement of the AIB neurite in the distal neighborhood.

**The AIB neurite is positioned in the distal neighborhood during embryonic development via a novel zippering mechanism**

To understand how the RIM neurons contribute to placement of the AIB distal neurite segment, we examined the placement of the AIB and RIM neurites during embryogenesis. Imaging AIB and RIM in embryos required integration of subtractive labeling strategies for sparse labeling and *in vivo* tracking of the AIB neurites (Supplementary Fig. 6; Armenti et al., 2014a). It also required isotropic imaging via the use of dual-view light-sheet microscopy (diSPIM) (Kumar et al., 2014; Wu et al., 2013) and the development of a triple-view confocal microscope combined with a deep-learning framework for enhanced resolution ((Weigert et al., 2018; Wu et al., 2016), see details in Methods). Our high resolution and continuous imaging approaches in embryos revealed that the stereotypic placement of the AIB neurite involves *(a)* an initial phase of tip-directed outgrowth, and *(b)* relocation of part of the growing neurite to the distal neighborhood by a ‘zippering’ mechanism, as further described below.

*(a) Initial tip-directed outgrowth:* We observed that the cell bodies of the bilaterally symmetric AIB neurons display outgrowth of a neurite that enters the nerve ring ~ 400 m.p.f. The two AIB neurites then circumnavigate the nerve ring at opposite sides of the neuropil—with AIBL growing dorsally on the left side of the nerve ring, and AIBR also projecting dorsally, but at the right side of the ring (Fig. 5a,b). They grow along the
proximal neighborhood, fasciculating with the axons of amphid sensory neurons (from which they later receive synaptic inputs) (Supplementary Fig. 6d). The simultaneous outgrowth of the AIBL and AIBR neurons results in their neurites meeting at the dorsal midline of the nerve ring at approximately 460 m.p.f. The AIB outgrowth continues past their meeting point at the dorsal midline, and each AIB grows along each other, still along the proximal neighborhood, at 480 m.p.f. (Fig. 5c).

(b) Relocation to the distal neighborhood: We observed that around 500 m.p.f., the segment of each AIB neurite that has grown past the midline (i.e., segment that will constitute the future distal neurite) starts separating from its lateral counterpart (the other AIB), starting from the growing tip (Fig. 5d-f, Supplementary Fig. 6e-g). The exit of the tip from the proximal neighborhood takes place as it extends straight, instead of following the ventral turn of the nerve ring, likely due to a loss of adhesion to proximal neighborhood neurons (Supplementary Fig. 6h,i). The growing tip then encounters the distal neighborhood (marked by RIM) at approximately 520 m.p.f. (Fig. 5h). Following this, the entire shaft of the future distal neurite relocates to the distal neighborhood, starting from the tip, and progressively ‘zippering’ towards the midline (Fig. 5i,j). At 510 minutes, 8% of the relocating distal neurite overlaps with RIM and the distal neighborhood. This overlap increases to 60% at around 535 mins and 96% at around 560 mins, demonstrating progressive overlap over time, analogous to the fastening of a zipper (see Methods). The separation or ‘unzippering’ from the proximal neighborhood and ‘zippering’ onto the distal neighborhood leads to repositioning of the entire distal neurite of AIB to a new neighborhood (Fig. 5j-n).
To investigate if placement of the AIB distal neurite by zippering has implications for synaptic protein distribution, we imaged presynaptic protein RAB-3 localization in the AIB distal neurite during the time when it is repositioned by zippering (520-570 m). We discovered that presynaptic proteins populate the neurite starting from the tip, progressively towards the dorsal midline in the same spatial pattern as AIB-RIM zippering-mediated contact (Supplementary Fig. 7). This suggests that the zippering mechanism might influence the spatiotemporal pattern of synaptic protein distribution in AIB upon contact with postsynaptic RIM. These findings are consistent with the observation in syg-1(ky652) that partial contact between AIB and RIM results in altered distribution of synaptic proteins to the sites of RIM contact.

Altogether, we discovered that the AIB neurite is positioned by a combination of tip-directed growth in the proximal neighborhood, circumferential exit of the distal neurite from the proximal neighborhood and its subsequent relocation and zippering to the distal neighborhood. The phenomenon of zippering of neurite shafts as a mechanism of fasciculation had been previously described in the context of dense primary neuron cultures (Barry et al., 2010; Šmit et al., 2017; Voyiadjis et al., 2011) but whether this occurs in developmental contexts in vivo is unknown. We demonstrate that the distal segment of the AIB neurite is placed in the correct neighborhood via a zippering mechanism onto a specific sublayer of the nerve ring.
Increase of local expression of SYG-1 at the distal neighborhood correlates with zippering of the distal AIB neurite onto its neighborhood

To understand how SYG-1 coordinates the developmental sequence of events that place the AIB neurite in specific neighborhoods, we examined the spatiotemporal dynamics of expression of the syg-1 transcriptional reporter during embryogenesis. By long-term imaging using light sheet microscopy (diSPIM) we discovered that the syg-1 promoter exhibits dynamic changes in spatial expression within nerve ring sublayers during the different events that occur to sequentially and precisely place the AIB neurite.

Similar to postembryonic animals, in embryos prior to hatching (~780 m.p.f.), the syg-1 reporter is expressed in both proximal and distal AIB neighborhoods, with enrichment in the distal neighborhood (Fig. 6a-d). To investigate how this expression pattern emerges during development, we imaged the dynamics of expression of the syg-1 promoter throughout the developmental period during which the AIB neurites are placed in the neighborhoods (400-570 m.p.f.) (Fig. 5a-j). We observed:

1. Prior to 470 m.p.f., expression of the syg-1 reporter in the nerve ring is primarily restricted to a single band corresponding to the AIB proximal neighborhood (Fig. 6e,i,i'). This coincides with periods of outgrowth by the AIBs in the proximal neighborhood.

2. Over the next two hours of embryogenesis (470-590 m.p.f.), a pair of neurons (one on each side of the nerve ring) robustly expressing syg-1 grow into, and are placed in, the distal neighborhood (Fig. 6f,j,j'). Through colocalization studies we identified these pair of neurons to correspond to RIMs (Supplementary Fig. 8a-c). As RIMs grow into the distal neighborhood, we also observe the onset of syg-1 expression in other distal neighborhood neurons (Fig. 6f-h). Therefore, during this period (470-590 m.p.f.) syg-1
expression levels increase in the distal neighborhood due to (a) ingrowth of the syg-1-expressing RIM neurons into the neighborhood and (b) onset of syg-1 expression in other neurons in the neighborhood. This increase in syg-1 expression in the distal neighborhood is accompanied by a progressive decrease in syg-1 levels in the proximal neighborhood (Fig. 6j-l). Mean intensities of the syg-1 reporter in the distal neighborhood relative to the proximal neighborhood increases by 2-2.4 times (between 470-600 m.p.f; Supplementary Fig. 8d,e).

These observations indicate that dynamic changes in relative levels of syg-1 between specific nerve ring neighborhoods might underlie the specific placement of the single AIB neurite across these neighborhoods. Altogether we provide evidence of layered expression of an IgCAM in two specific nerve ring neighborhoods, observe that the expression dynamically switches between these neighborhoods and describe how these dynamic changes correlate with the nanoscale precise placement of a single neurite (AIB) across these neighborhoods.

Ectopic syg-1 expression is sufficient to alter placement of the AIB distal neurite

To investigate if syg-1 is instructive for the placement of the AIB distal neurite in the nerve ring, we ectopically increased syg-1 levels of expression in the proximal neighborhood. Through promoter screening and lineage-based cell identification, we determined that during embryogenesis, promoters of the nphp-4 and mgl-1b genes are primarily expressed in neurons of the proximal neighborhood (http://promoters.wormguides.org) (Z. Bao et al., 2006; Boyle et al., 2006; Duncan et
Many of the neurons in which these promoters are expressed are fasciculation partners of AIB in the proximal neighborhood. We used these promoters to drive ectopic expression of a syg-1 cDNA in the proximal neighborhood in the syg-1(ky652) mutant background. Unlike in wild type and syg-1 mutants (Fig. 7a-f, Supplementary Fig. 9a,b), in the animals with ectopic syg-1 expression, we observed a gain-of-function phenotype in which the AIB distal neurite remains partially positioned in the proximal neighborhood through postembryonic larval stages (Fig. 7g-j, Supplementary Fig. 9c). Therefore ectopic expression of syg-1 in neurons of the proximal neighborhood is sufficient to misposition the AIB distal neurite segment. When we ectopically expressed a syg-1 cDNA lacking the intracellular domains, we observed similar gain-of-function effects, indicating that SYG-1’s extracellular Ig domains are sufficient in this context (Supplementary Fig. 9d). These observations suggest that SYG-1 is instructive in the placement of the AIB distal neurite into specific neighborhoods, and that its spatiotemporally regulated expression in banded patterns in the nerve ring is important for placement of specific neurites onto distinct neuropil layers.

**SYG-1 is required for layer-specific placement of rich-club neuron AVE**

We next examined if syg-1 is also important for placement of other neurites in the layers in which it is highly expressed. We focused on the rich-club AVE neurons, the neurites of which are also placed in two distinct neighborhoods, different from the AIB neighborhoods and separated by a posterior-anterior chiasm at the dorsal midline. (Fig. 8a-c)(Moyle et al., 2020; Sabrin et al., 2019; Towlson et al., 2013; White et al., 1986).
Reconstructions from electron micrographs reveal that the AVE neurons have a morphology similar to AIB, however its neurite is more anteriorly placed (by one stratum) with respect to AIB (Moyle et al., 2020). Therefore the proximal neurite of AVE occupies the S2/S3 neighborhood occupied by the AIB distal neurite. Since syg-1 expression is enriched in this “AIB distal/AVE proximal” neighborhood, we tested, by examining AVE neurite placement relative to the RIM neurons, if placement of the AVE neurite in this neighborhood is also affected in syg-1(ky652) mutants.

Although AVE and RIM are not synaptic partners, they form non-synaptic contacts with each other along their neurites, as indicated by EM connectome data (White et al., 1986; Witvliet et al., 2020). When we fluorescently labeled RIM and AVE in wildtype animals, we observed that the proximal AVE neurite runs along the RIM neurite, consistent with the EM studies (Fig. 8d,e,e’). By contrast, in syg-1 mutants the AVE proximal neurite is frequently detached from RIM (seen in 50% of syg-1(ky652) mutants versus 9.1% in wild type (Fig. 8f,g,g’,h). The dorsal midline shift of AVE is also affected in syg-1 mutant animals (mean length = 2.73 µm in syg-1(ky652) and 3.99 µm in wild type animals; Fig. 8i). Together with the AIB studies, these observations are consistent with syg-1 expression in a S2/S3 sublayer resulting in nanoscale placement of both the AIB and AVE neurites into that neighborhood.

In the previous sections we demonstrated that RIM has robust syg-1 expression during nerve ring development and throughout postembryonic stages. So we next investigated if RIM acts as a guidepost, positioning AVE onto the S2/S3 neighborhood to facilitate formation of AVE synapses onto correct partners. We observed that caspase-mediated ablation of the RIM neurons results in partial defects in the
placement of the AVE proximal neurite (37.03% of RIM-ablated animals had defects in AVE neurite placement, compared to 6.37% in wild type animals) (Fig. 8j,k). Our observations suggest that layer-specific expression of syg-1 regulates nanoscale precision in placement of the AVE neurites in the AVE proximal neighborhood. Together, our studies uncover spatiotemporal dynamics of expression of an adhesion molecule between nerve ring neighborhoods, and its regulation of a novel zipper ing mechanism for positioning neurites onto these specific neuropil strata.

Discussion

Nanoscale placement of rich-club interneuron AIB results in the wiring of modular circuits across distinct layers of the neuropil. Neuropil layers are a conserved organizational principle present in evolutionarily diverse brain regions, ranging from the C. elegans nerve ring to the Drosophila lamina and the inner plexiform layer of the mammalian retina (Moyle et al., 2020; Robles et al., 2011; Tan et al., 2015). Precise placement of neuronal processes and their synapses, particularly for neurons spanning across multiple layers, underpin brain topographic maps, and their functional and structural principles (Clandinin & Feldheim, 2009). We find that in the layered nematode nerve ring, “rich-club” interneuron AIB facilitates network connectivity between the modular S2, S3 and S4 layers, and that its precise placement and synaptic distribution in the context of the neuropil is tightly linked to its functional properties as a rich-club informational hub across layers. The conserved and stereotyped design principles observed for AIB, both by EM analyses and in vivo, are reminiscent of those seen for cellular motifs in the inner plexiform layer of the vertebrate retina. For example, All
amacrine cells, which distribute their neurites and synapses across distinct, but specific sublaminae, receive inputs from rod bipolar axon terminals (in lower sublamina b) and produce outputs onto ganglion cell dendrites (in sublamina a) (Kolb, 1995; Strettoi et al., 1992). Our forward genetic screens in AIB reveal that the developmental processes dictating neurite placement, and those instructing polarized distribution of pre- and postsynaptic specializations across laminae, are genetically separable. Therefore in AIB, the correct emergence of conserved design principles requires the modular deployment of two developmental programs: polarized placement of pre- and postsynaptic specializations along the neurite, and the nanoscale placement of the neurite onto the correct neighborhood of the neuropil.

A novel zippering mechanism governs precise AIB placement onto specific neighborhoods in the neuropil. The correct development of the neuropil structure requires that 181 neurites, which form synapses en passant or along the length of their axon, are precisely placed onto neighborhoods within layers. Our current mechanistic frameworks of neurodevelopment are insufficient to explain how single cells make simultaneous and coordinated use of similar molecular cues to achieve precise placement onto specific neighborhoods within the layered neuropil. By combining light sheet microscopy, a new triple-view confocal microscope and machine learning approaches, we are able to resolve these developmental events for the AIB neurons of *C. elegans*, and uncover an in vivo mechanism that places neurites onto layers via zippering. Zippering has been observed and characterized in tissue culture cells, and has been hypothesized to be a mechanism that could instruct neurites switching tracts within bundles or fascicles (Honig et al., 1998; Šmít et al., 2017). Evidence for the in
existence of this mechanism, or its importance in development, has been lacking.

We now demonstrate that zippering occurs in vivo in embryos, and that it is important for placing neurites within specific neuropil neighborhoods during development.

Spatiotemporally-regulated expression of IgCAM protein SYG-1 in specific nerve ring neighborhoods guide placement of AIB neurites. Our genetic studies, embryonic expression studies and imaging analyses reveal that expression of this IgCAM in specific nerve ring neighborhoods is dynamic, coincides with AIB developmental decisions, and is necessary, non-autonomously, for AIB neurite placement onto specific layers. Our findings are consistent with studies in the Drosophila medulla, which determined that expression of DIP family of IgCAMs are restricted to specific layers or layer boundaries, regulated during pupal development and probably necessary for matching neuronal pairs within medullar layers (Tan et al., 2015). Moreover, spatiotemporal dynamics of adhesion molecule expression encodes layer specificity in the lamina of the Drosophila visual system, e.g., it was found that expression of the adhesion molecule Cadherin-N renders neurons competent to layer-specific targeting in a temporally segregated manner (Petrovic & Hummel, 2008). Our observations extend these findings, now demonstrating that the spatiotemporal regulation of SYG-1 in specific C. elegans nerve ring neighborhoods is tightly coordinated with the sequential cellular decisions of AIB during the placement of its neurite segments onto specific layers within the neuropil.

SYG-1 expression is sufficient to instruct misplacement of AIB onto ectopic neighborhoods. Expression of the syg-1 gene, first at the S3/S4 nerve ring neighborhood and later at the S2/S3 neighborhood coincides with the AIB outgrowth
dynamics through these neighborhoods. Mis-expression of SYG-1 is sufficient to result in the retention of segments of the AIB neurite onto the incorrect layer, indicative of a role for SYG-1 in instructing zippering and placement of AIB onto the proper neuropil layer. The Ig-CAM SYG-1 is conserved across evolution, and its orthologs (Rst and Kirre in *Drosophila* and Kirrel1/2/3 in mammals) play important roles as adhesion molecules in varying developmental contexts (S. Bao & Cagan, 2005; Garg et al., 2007; Kim et al., 2015; Serizawa et al., 2006; Shen & Bargmann, 2003; ztokatli et al., 2012).

We note that while in the *C. elegans* egg-laying circuit, SYG-1 is known to interact with its partner (SYG-2) to regulate synapse formation, in the context of the nerve ring, these molecules appear to primarily mediate fasciculation. A loss-of-function mutant allele of syg-2 exhibits similar defects in AIB distal neurite placement (Supplementary Fig. 4d-f).

A loss of function mutant in syg-2 has been previously shown to result in defasciculation defects of the HSNL axon (Shen et al., 2004), and while not characterized in this study, we hypothesize that similar mechanisms might underpin syg-2 roles at the nerve ring.

Importantly, our findings suggest that SYG-1 role in neurite placement in nerve ring neighborhoods is analogous to the role for mammalian orthologs Kirrel 2 and Kirrel3, which are involved in axon sorting in the olfactory system, possibly via regulation of axon fasciculation.

Zippering mechanisms via affinity-mediated adhesion might help instruct neighborhood coherence while preserving ‘fluid’, or transient interactions among neurites within neuropil structures. Our analyses of contact profiles for individual neurons in the nerve ring neuropil reveal that most interactions between neuropil neurites are brief, resulting in a ‘tangled’ structure with variable contact profiles across
connectomes (Moyle et al., 2020). Yet, neuropils, including the C. elegans nerve ring, retain structural design features that underlie functional relationships among neurons, and circuit specificity. How does the system solve this tension of flexibility and stereotypicity? We speculate that dynamic expression of adhesion molecules, such as SYG-1, and zippering mechanisms, might help preserve the tissue organization in the tangled context of neuropils by creating affinity relationships of relative strengths. These relationships could result in the separation of neuropil clusters, similar to observations in developing embryos in which cell types expressing compatible cell adhesion molecules were observed to phase separate onto clusters based on affinity relationships (Foty & Steinberg, 2005, 2013; Steinberg, 1962). Our observations of the expression of SYG-1 simultaneously coordinating, not only AIB development, but also laminar placement of rich club neuron AVE, suggest a layer-specific molecular signature that influences cosegregation of neighboring neurites via affinity profiles. We hypothesize that in neuropils, these layer-specific molecular signatures mark specific domains that enable placement of neurites via flexible, differential cell-adhesion mediated zippering mechanisms.

Methods

C. elegans strains

C. elegans strains were raised at 20°C using OP50 Escherichia coli seeded on NGM plates. N2 Bristol is the wild-type reference strain used. We received the NC1750
strain from the Caenorhabditis Genetics Center (CGC). Transgenic strains used in this study are available upon request.

**Molecular biology and Transgenic lines**

We used Gibson Assembly (New England Biolabs) or the Gateway system (Invitrogen) to make plasmids used for generating transgenic *C. elegans* strains. Detailed cloning information or plasmid maps will be provided upon request. Transgenic strains were generated via microinjection with the construct of interest at 5-100 ng/μL by standard techniques (Mello & Fire, 1995). Co-injection markers *unc-122p*: GFP or *unc-122p*: RFP were used.

We generated the *syg-1* transcriptional reporter (Fig. 4a-h, Fig.6, Supplementary Fig. 8) by fusing membrane-targeted PH:GFP to a 3.5 kb *syg-1* promoter region as described (Schwarz et al., 2009).

**SNP mapping and Whole-Genome Sequencing**

We isolated mutant allele *daf-16(ola337)* and *syd-2(ola341)* from a visual forward genetic screen in an integrated wild type transgenic strain (*olals67*) with AIB labeled with cytoplasmic mCherry and AIB presynaptic sites labeled with GFP: RAB-3. Ethyl methanesulfonate (EMS) mutagenesis was performed and animals were screened for defects in placement of the AIB neurite, and defects in AIB presynaptic site distribution. We screened F2 progeny on a Leica DM 5000 B compound microscope with an HCX PL APO 63x/1.40–0.60 oil objective.
The novel lesions ola337 and ola341 were out-crossed six times to wild type (N2) animals and still retained a defect in AIB distal neurite placement and AIB presynaptic protein distribution, respectively. We then used single-nucleotide polymorphism (SNP) mapping as described (Davis et al., 2005) to map the ola337 and ola341 lesions.

We then performed whole-genome sequencing on ola337 and ola341 (along with 4 other mutants from the screen) at the Yale Center for Genome Analysis (YCGA), as previously described (Sarin et al., 2008). We analyzed the results using the Galaxy platform, with http://usegalaxy.org/cloudmap (Minevich et al., 2012) using the EMS variant density mapping workflow. Lesion information from mutants is listed in Supplementary Table 1.

Confocal imaging of C. elegans larvae and image processing

We used an UltraView VoX spinning disc confocal microscope with a 60x CFI Plan Apo VC, NA 1.4, oil objective on a Nikon Ti-E stand (PerkinElmer) with a Hamamatsu C9100–50 camera. We imaged the following fluorescently tagged fusion proteins, eGFP, GFP, PH:GFP (membrane-tethered), RFP, mTagBFP1 and mCherry at 405, 488 or 561 nm excitation wavelength. We anesthetized C. elegans at room temperature in 10mM levamisole (Sigma) under glass coverslips and mounted them on glass slides for imaging.

We used the Volocity image acquisition software (Improvision by Perkin Elmer) and processed our images using Fiji(Schindelin et al., 2012). Image processing included maximum intensity projection, 3D projection, rotation, cropping, brightness/contrast, line segment straightening, and pseudo coloring. All quantifications from confocal images
were conducted on maximal projections of the raw data. The simple neurite tracer Fiji plugin and a MATLAB code were used to validate estimation of minimum perpendicular distance between neurites. Pseudocoloring of AIBL and AIBR was performed in Fiji. Pixels corresponding to the neurite of either AIBL/R were identified and the rest of the pixels in the image were cleared. This was done for both neurons of the pair and the resulting images were merged.

**Embryo labeling, imaging and image processing**

For labeling of neurites and subcellular structures in embryos, we used membrane tethered PH:GFP. A subtractive labeling strategy was employed for AIB embryo labeling (Supplementary Fig. 6a). Briefly, we generated a strain containing unc-42p::ZF1::PH::GFP and lim-4p::SL2::ZIF-1, which degraded GFP in the sublateral neurons, leaving GFP expression only in the AIB and/or ASH neurons. (Armenti et al., 2014b). Onset of twitching was used as a reference to time developmental events. Embryonic twitching is stereotyped and starts at 430 minutes post fertilization (m.p.f) for our imaging conditions.

Embryonic imaging was performed via dual-view inverted light sheet microscopy (diSPIM) (Kumar et al., 2014; Wu et al., 2013) and a combined triple-view line scanning confocal/DL for denoising, as described below.

**Triple-view line-scanning confocal/DL**

We developed a triple-view microscope that can sequentially capture three specimen views, each acquired using line-scanning confocal microscopy. Multiview
registration and deconvolution can be used to fuse the 3 views (Wu et al., 2013), improving spatial resolution. Much of the hardware for this system is similar to the previously published triple-view system (Wu et al., 2016), i.e., we used two 0.8 NA water immersion objectives for the top views and a 1.2 NA water immersion lens placed beneath the coverslip for the bottom view. To increase acquisition speed and reduce photobleaching, we applied a deep-learning framework (Weigert et al., 2018) to predict the triple-view result when only using data acquired from the bottom view. The training datasets were established from 50 embryos (anesthetized with 0.3% sodium azide) in the post-twitching stage, in which the ground truth data were the deconvolved triple view confocal images, and the input data were the raw single view confocal images resulting in improved resolution (270nm X 250 nm X 335nm).

**Cell lineaging**

Cell lineaging was performed using StarryNite/AceTree (Z. Bao et al., 2006; Boyle et al., 2006; Murray et al., 2006). Light sheet microscopy and lineaging were integrated to uncover cell identities in pre-twitching embryos (Duncan et al., 2019). Lineaging information for promoters is available at [http://promoters.wormguides.org](http://promoters.wormguides.org). Our integrated imaging and lineaging approaches enabled us to identify a promoter region of *inx-19* which is expressed in the RIM neurons prior to RIM neurite outgrowth (~370 m.p.f.) and in additional neurons in later embryonic stages. The *inx-19p* was one of the promoters used for embryonic ablation of RIM (described in the next section).

In addition our integrated imaging and lineaging approach also enabled us to identify two promoters with expression primarily in the AIB proximal neighborhoods (*nphp-4p*...
and mgl-1bp). 4/4 neuron classes that were identified to have nphp4p expression, are in the AIB proximal neighborhood (ADL/R, ASGL/R, ASHL/R, ASJL/R) and 2/3 neuron classes that were identified to have mgl-1bp expression are in the AIB proximal neighborhood (AIAL/R, ADFR) (http://promoters.wormguides.org).

Caspase-mediated ablation of RIM neurons

The RIM neurons were ablated using a split-caspase ablation system (Chelur & Chalfie, 2007). We generated one set of transgenic strains with co-expression of the p12 or p17 subunit of human Caspase-3, both expressed under inx-19p (termed ablation strategy 1), and another set of ablation strains with co-expression of the p12 subunit expressed under inx-19p and p17 under tdc-1p (termed ablation strategy 2) (Supplementary Fig. 5). L3 larvae from the RIM-ablated populations were imaged on the spinning-disk confocal microscope (described earlier).

Rendering of neurites and contacts in the EM datasets

From available EM datasets (C. Brittin et al., 2018; Cook et al., 2019; White et al., 1986; Witvliet et al., 2020) we rendered the segmentations of neuron boundaries in 2D using TrakEM2 in Fiji. TrakEM2 segmentations were volumetrically rendered by using the 3D viewer plugin in Fiji (ImageJ2; downloaded from https://imagej.net/Fiji#Downloads) and saved as object files (.obj), or by using the 3d viewer in CytoSHOW (Duncan et al., 2019), an open source image analysis software. CytoSHOW can be downloaded from http://www.cytoshow.org/ as described.
To generate 3D mappings of inter-neurite membrane contact, the entire collection of 76,046 segmented neuron membrane boundaries from the JSH TEM datasets (C. Brittin et al., 2018; C. A. Brittin et al., 2020; White et al., 1986) were imported from TrakEM2 format into CytoSHOW as 2D cell-name-labelled and uniquely color-coded regions of interest (ROIs). To test for membrane juxtaposition, we dilated each individual cell-specific ROI by 9 pixels (40.5 nm) and tested for overlap with neighboring undilated ROIs from the same EM slice. A collection of 289,012 regions of test-overlap were recorded as new ROIs, each bearing the color code of the dilated test ROI and labeled with both cell-names from the pair of test-overlapped ROIs. These "contact patch" ROIs were then grouped by cell-pair-name and rendered via a marching cubes algorithm to yield 3D isosurfaces saved in .obj files. Each of the 8852 rendered .obj files represents all patches of close adjacency between a given pair of neurons, color-coded and labeled by cell-pair name. Selected .obj files were co-displayed in a CytoSHOW3D viewer window to produce views presented in Figures.

**Cosine similarity analysis for comparing AIB contacts across connectomes**

We performed cosine similarity analysis (Han et al., 2012) on AIB contacts in available connectome datasets (C. Brittin et al., 2018; Cook et al., 2019; Witvliet et al., 2020). For each available adjacency dataset (C. Brittin et al., 2018; Moyle et al., 2020; Witvliet et al., 2020), we extracted vectors comprising of number of AIB contacts with neurons common to all the datasets. We then performed cosine similarity analysis on these vectors using the formula:
\[
\frac{\sum_{i=1}^{n} A_i B_i}{\sqrt{\sum_{i=1}^{n} A_i^2 \sum_{i=1}^{n} B_i^2}}
\]

where \(A\) and \(B\) are the two vectors under consideration with the symbol “\(i\)” denoting the \(i\)-th entry of each vector. The similarity values were plotted as a heat map for AIBL and AIBR using Prism. For the datasets L1_0hr, L1_5hr, L1_8hr, L2_23hr, L3_27hr, L4_JSH and Adult_N2U, the neuron-neuron contacts in the EM sections corresponding to the nerve ring were used.

### Betweenness centrality analysis

We analyzed betweenness centrality for two of the available connectomes of different developmental stages (L1 and adult) (Witvliet et al., 2020). By treating individual components (mostly neurons) of a connectome as the vertices of a graph, we use the following definition of Betweenness Centrality for a vertex \(v\),

\[
BC(v) = \sum_{s,t:s \neq t \neq v} \frac{\lambda_{st}(v)}{\lambda_{st}}.
\]

Here \(\lambda_{st}(v)\) denotes the number of shortest paths between the vertices \(s\) and \(t\), that include vertex \(v\), whereas \(\lambda_{st}\) denotes the total number of shortest paths between the vertices \(s\) and \(t\). We finally divide \(BC(v)\) by \((N-1)(N-2)/2\) to normalize it to lie between 0 and 1. For our implementation we use the Brain Connectivity Toolbox (Rubinov & Sporns, 2010) of MATLAB2020, in particular, the function “betweenness_bin.m” in which we input the binary connection matrix corresponding to the L1 and adult connectomes (Witvliet et al., 2020). We made a Prism box plot (10 to
90 percentile) of betweenness centrality values of all components in each of the two connectomes and highlighted the betweenness centrality values for AIBL and AIBR.

Image representation, quantification and statistical analysis

Representation of AIB and AVE from confocal images – Since we observed that the proximal and distal neurites of AIBL and AIBR completely align and overlap (Supplementary Fig. 1k-m) in confocal image stacks where the worms are oriented on their side, for representation purposes we have used the upper 50% of z-slices in confocal image stacks to make maximum intensity projections. This shows the proximal neurite of AIBL in the context of the distal of AIBR (which has the same anterior-posterior position as the distal neurite of AIBL) (Supplementary Fig. 2k-m), or vice versa. We used the same procedure for AVEL and AVER.

Quantification of penetrance of mutant phenotypes – The percentage of mutant or ablation animals exhibiting a normal AIB distal neurite trajectory was determined by visualizing and scoring under the Leica compound microscope described. Animals in which distal neurites of both AIBL or AIBR were placed at a uniform distance from the proximal neurites were scored as having normal AIB distal neurite trajectory. The same scoring protocol was followed for determining percentage of animals having normal AVE distal neurite trajectory.

Quantification of minimum perpendicular distance between neurites – Minimum perpendicular distances between neurites were measured by manually creating a straight line selection (on Fiji) between the neurites (perpendicular to one of the
neurites) in the region where the gap between them is estimated to be the smallest. The measurements were done on maximum intensity projections of raw confocal image stacks where the worms are oriented on their side (z-stacks acquired along left right axis of the worm, producing a lateral view of the neurons).

**Quantification of percent detachment between neurites** – The percent detachment for defasciculated neurites (AIB and RIM or AVE and RIM) is calculated by the formula

\[
\text{\% detachment} = \frac{\text{detached length } (L_d)}{\text{total length } (L_t)} \times 100
\]

Also shown in Fig. 3m. \(L_d\) is calculated by making a freehand line selection along the detached region of the RIM neurite and measuring its length and \(L_t\) is calculated by making a freehand selection along the RIM neurite for the entire length over which it contacts AIB or AVE, and measuring the length of the selection. All the measurements were performed on maximum intensity projections of confocal image stacks where the worms are oriented on their side (z-stacks acquired along left right axis of the worm, producing a lateral view of the neurons).

**Quantification of relative (distal) enrichment of syg-1 reporter expression in neighborhoods** – Relative (distal) enrichment of syg-1 reporter expression is calculated using the formula (also shown in Fig. 4i),

\[
\text{Relative enrichment (syg-1p)} = \frac{\text{mean distal neighborhood intensity } (I_d)}{\text{mean proximal neighborhood intensity } (I_p)}
\]

These measurements were done in transgenic animals co-expressing the AIB reporter and the syg-1 transcriptional reporter. For calculation of \(I_p\), a freehand line selection was made (using Fiji) along the band of syg-1 expression along the AIB proximal neurite (as visualized with the AIB marker) and mean intensity along the selection is calculated. Same was done for calculation of \(I_d\) except along the AIB distal neurite. The ratios of \(I_d\)
and $l_p$ were plotted as relative (distal) enrichment values. These values were calculated from maximum intensity projections of confocal image stacks where the worms are oriented on their side (z-stacks acquired along left right axis of the worm, producing a lateral view of the neurons). $l_d/l_p$ was calculated from the side of the animal closer to the objective, consistently across images to avoid differences due to depth artifacts.

**Quantification of the dorsal midline shift (chiasm) length of AIB and AVE** – The dorsal midline shift (chiasm) lengths of AIB and AVE were calculated by making 3D maximum intensity projections of confocal z-stacks and orienting the neuron pair to a dorsal-ventral view. A straight line selection is made along the posterior-anterior shift of each neuron and its length measured using Fiji.

**Quantification of distal neurite length of AIB** – The length of the distal neurite of AIB was measured by drawing a freehand line along the region of the neurite past the proximal neurite in maximum intensity projections of confocal image stacks where the worms are oriented on their side (z-stacks acquired along left right axis of the worm, producing a lateral view of the neurons).

**Quantification of the relative position of the AIB neurite during embryogenesis** – The distances of the unzippering and zippering forks from the dorsal midline in Fig. 5l and 5m are calculated from deconvolved maximum intensity projections of diSPIM images where the neurons are oriented in an axial view. These distances are lengths along the AIB neurite from the unzippering/zippering forks to the dorsal midline.

The distance of the zippering fork from the midline is subtracted from the total length of the neurite at each timepoint to obtain the length of overlap between AIB and
RIM. The fraction of the length of AIB-RIM overlap to the total AIB neurite length multiplied by 100, yields a percentage overlap value at every timepoint. The reported values of percent overlap are averages across the three independent embryo datasets used for the Fig. 5m plot. These measurements are performed with CytoSHOW. We performed the same steps as with the confocal images to pseudocolor the neurites for representation.

Quantification of the angle of exit of the developing AIB distal neurite and ventral turn of nerve ring in embryos – The angle of exit of the developing AIB distal neurite is measured as the angle between straight line tangents drawn along the separating distal segment of AIBL and the proximal neurite of AIBR and vice versa. These measurements are performed on deconvolved maximum intensity projections of diSPIM images where the neurons are oriented in an axial view. The angle of ventral turn of the nerve ring is measured as the angle between straight line tangents drawn along segments of the nerve ring on either side of the ventral bend of the nerve ring (see Supplementary Fig. 6h,i). These measurements are performed with CytoSHOW.

Imaging and representation of synaptic protein RAB-3 in AIB in embryos – Time-lapse imaging of presynaptic protein RAB-3 in AIB in embryos was performed using diSPIM. To visualize the distribution of RAB-3 along the neurite we straightened the distal neurite of each AIB neuron from maximum intensity projections where the AIB neurons are oriented in the axial view.
Statistical analyses

Statistical analyses were conducted with PRISM 7 software. For each case, the chosen statistical test is described in the figure legend and “n” values are reported. Briefly, for continuous data, comparisons between two groups were determined by the Student’s t test. Error bars were reported as standard errors of the mean (SEM). For categorical data, groups were compared with Fisher’s exact test. The p values for significant differences are reported in the figure legend.

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**Figure Legends**

**Figure 1. Design principles of AIB neurite placement and synaptic positions enable it to link two neighborhoods in the nerve ring**

(a) Schematic of an adult/larval *C. elegans* showing an AIB neuron (cyan) and its proximal (orange) and distal (magenta) neighborhoods. The AIB neurite has a proximal neurite segment (orange arrow), a posterior-anterior shift (black arrowhead, herein also referred to as chiasm) at the dorsal midline (dashed straight line) and a distal neurite segment (magenta arrow; on the other side of the worm, AIB cyan color muted in the distal segment to reflect the position behind the pharynx, which is in gray). The region occupied by the nerve ring neuropil is depicted in light brown. Note that this schematic only shows one neuron of the AIB pair (AIBR and AIBL, for right and left, respectively, see Supplementary Fig. 1e-j for both AIBs). Cell body is marked with an asterisk.

(b,c) Volumetric reconstruction of AIBR from the JSH electron microscopy connectome dataset (White et al., 1986) in lateral (b) and axial (c) views. Postsynaptic (red) and presynaptic (yellow) regions of the neurite, based on connectivity maps of AIBR, are indicated. Note that the postsynaptic and presynaptic regions coincide with the proximal
and distal segment of the neurites, in (a). Arrowhead points to the chiasm (formed by
the posterior-anterior shift at the dorsal midline). Scale bar = 1 µm

(d,e) Representative confocal image showing lateral (d) and axial (e) view of an AIB
neuron with postsynaptic sites (red, labeled by GLR-1:GFP) and presynaptic sites
(yellow, labeled by mCh:RAB-3). Cell-specific expression of GLR-1 and RAB-3 was
achieved by using a bashed *inx-1* promoter (http://promoters.wormguides.org) (Altun et
al., 2008). Arrowhead points to the chiasm. Cell body is marked with an asterisk. Scale
bar = 10 µm, also applies to (e).

(f,g) Schematic of AIB (corresponding to the dashed box region in (a) and also images
in (b-e)), showing lateral (f) and axial (g) views of an AIB neuron (cyan) in the context of
the proximal (orange) and distal (magenta) neighborhoods in the nerve ring (light
brown). Dashed line represents the dorsal midline.

(h,i) Representative confocal image from a wild type L4 animal showing an AIB neuron
labeled with cytoplasmic mCherry (cyan) in lateral (h) and axial (i) views. Scale bar = 10
µm, also applies to (i-m).

(j,k) Representative confocal image from a wild type L4 animal showing an AIB
interneuron labeled with cytoplasmic mCherry (cyan); and RIM motor neuron labeled
with cytoplasmic GFP (magenta) in lateral (j) and axial (k) views. RIM-specific labeling
was achieved by a *cex-1* promoter (Piggott et al., 2011)
(http://promoters.wormguides.org). Note the colocalization of the AIB distal neurite with
the distal neighborhood marker RIM. The orange and magenta arrows indicate the
positions of the proximal and distal neighborhoods along the anterior-posterior axis.
Arrowhead indicates the chiasm.
As (j,k), but with AIB (cyan) and AWC and ASE sensory neurons (orange). AWC and ASE labeling was achieved by a ceh-36 promoter (Lanjuin et al., 2003; Walton et al., 2015) (http://promoters.wormguides.org). Note the colocalization of the AIB proximal neurite with the proximal neighborhood markers AWC and ASE.

Figure 2. Polarized distribution of synaptic proteins in the AIB neurite, and AIB neurite neighborhood placement in the neuropil, are genetically separable

(a-d) Representative confocal image (a-c) and schematic (d) of AIB expressing cytoplasmic mCherry (a) and GFP:RAB-3 (b) in wild type L4 animals for simultaneous visualization of neurite morphology and distribution of presynaptic sites. Merged image (c). Cell body is marked with an asterisk. Scale bar = 10 µm, applies to panels (a-k).

(e-l) As (a-d) but in the genetic background of a novel daf-16 mutant allele, ola337 ((e-h; see also Supplementary Table 1) and in unc-33(e204) mutants (i-l). Note GFP:RAB-3 polarized distribution to the distal parts of the neurite even in animals displaying severe axon truncation, or displaying morphology and placement defects of the AIB neurite in the context of the neuropil neighborhoods (also see Supplementary Fig. 3).

Figure 3. SYG-1 is required for precise placement of the AIB distal neurite to the distal, RIM-containing neighborhood

(a-d) Representative confocal images of AIB (a) and RIM (b) neurons in a wild type animal (L4 stage), and the merged image (c). The dashed box represents the region of
contact between the AIB and RIM neurites, magnified in (d). RIM is the main
postsynaptic partner for AIB (White et al., 1986) and co-localizes extensively with AIB
distal neurite (Arrow in (d) and Supplementary Fig. 1b,c). Scale bar = 10 µm in (a)
applies to (e-g) and (i-k). Scale bar = 1 µm in (d) applies to (h) and (l). Cell bodies are
marked with an asterisk.

(e-h) As (a-d) but in the syg-1(ky652) mutant background. Note the gap between the
AIB distal neurite and the RIM neurites (h), indicating loss of contact between the AIB
and RIM neurites due to defective placement of the AIB distal neurite in the RIM (distal)
neighborhood.

(i-l) As (a-d) and (e-h) but for a syg-1(ky652) mutant animal expressing a rescuing
cosmid (C54A10) containing the genomic region of wild type syg-1.

(m) Schematic and scatter plot of quantifications of the loss of contacts between the AIB
and RIM neurites. The extent of detachment of the AIB distal neurites from RIM, and
hence its deviation from the RIM neighborhood, was quantified using the indicated
formula (see also Methods). The cyan and magenta neurites in the schematic represent
the AIB distal neurite and the RIM neurites, respectively, as in the images. Scatter plot
depicts % detachment values for wild type (n=41), syg-1(ky652) mutant (n=39) and
cosmid rescue animals (n=16). Error bars indicate standard error of the mean (S.E.M.).

****p<0.0001 by unpaired Student’s t-test between WT and syg-1(ky652), and between
syg-1(ky652), and syg-1(ky652) rescued with syg-1 containing cosmid (C54A10)
called syg-1 rescue in graph).

(n) Schematic and scatter plot of quantification of minimum perpendicular distance
between the AIB proximal and distal neurites. Schematic indicates AIB (cyan) and RIM
(magenta) neurons in the context of the nerve ring (light brown) in WT and syg-1(ky652). Double-headed arrows indicate perpendicular distance between the AIB proximal and distal neurites in WT and syg-1(ky652), respectively. Scatter plot depicts minimum perpendicular distance values for wild type (n=19), syg-1(ky652) mutant (n=29) and cosmid rescue animals (n=12) (see Methods). Error bars indicate standard error of the mean (S.E.M.). ****p<0.0001 by unpaired Student’s t-test between WT and syg-1(ky652), and between syg-1(ky652) and syg-1(ky652) rescued with syg-1 containing cosmid (C54A10) (called syg-1 rescue in graph).

**Figure 4. The SYG-1-expressing RIM neurons regulate AIB distal neurite position**

(a-d) Representative confocal image of a wild type L3 animal co-expressing (a) a membrane-targeted syg-1 transcriptional reporter (see Methods) and (b) cytoplasmic mCherry driven by the AIB-specific promoter (inx-1p). (c) is a merge image of (a) and (b). Since the syg-1 reporter is membrane-targeted, it labels cell body outlines and neurites (a,c). The dashed box or inset in (c) represents the region of contact between AIB and neurites expressing the syg-1 reporter, magnified in (d). Note that the syg-1 reporter shows two bands of expression in the nerve ring (a and c) which coincide with the proximal and distal AIB neurites (b) and neighborhoods (orange and magenta arrows). Note also that there is no membrane outline corresponding to the AIB cell body (a, we drew a dashed siluette of the AIB cell body position as determined in (b)), consistent with the syg-1 reporter not being expressed in AIB. Asterisk indicates cell
(e-h) As (a-d), but with mTagBFP1 driven by RIM-specific promoter, *cex-1p*. Note the RIM neurite colocalizes with the anterior band of *syg-1* expression, coincident with the AIB distal neighborhood (magenta arrow). The white arrowhead in (e-g) and semi-transparent magenta outline in (e) indicates colocalization of the RIM cell body with the *syg-1* reporter.

(i) Schematic (left) and scatter plot quantification (right) of the expression pattern of the *syg-1* reporter at the two AIB neighborhoods. The mean intensities of the *syg-1* reporter in the two neighborhoods were calculated, and the ratios of the mean distal neighborhood intensity (*I_d*) to the mean proximal neighborhood intensity (*I_p*) were plotted as the ‘Relative enrichment’, as shown in schematic and explained in Methods.

(j-o) Confocal images showing AIB (labeled with cytoplasmic mCherry; (j,m)) and RIM (labeled with PH:GFP; (k,n)) and merged images (l,o) for wild type animals (j-l) and animals in which RIM was genetically ablated (m-o). RIM ablation was achieved using Strategy 2, explained in Supplementary Fig. 5 and below. Scale bar = 10 μm, also applies to (k-o).

(p) Quantification of the penetrance of the AIB neurite placement defect as the percentage of animals with normal AIB distal neurite trajectory. Strategy 1 and Strategy 2 refer to split caspase ablations (Chelur & Chalfie, 2007) using two different combinations of promoters expressed in RIM. In Strategy 1, expression of both caspase fragments was driven by an *inx-19* gene promoter. In Strategy 2, expression of one caspase fragment was driven by *inx-19p* and the other fragment by a *tdc-1* gene
promoter (Alkema et al., 2005); Supplementary Fig. 5 and http://promoters.wormguides.org). tdc-1p drives expression of caspases in RIM later in embryonic development as compared to inx-19p (data not shown), resulting in later ablations of RIM, and weaker phenotype. ****p<0.0001 by Fisher’s exact test between WT and RIM-ablated populations.

(q) Quantification of the expressivity of the AIB neurite placement defect by measuring the minimum perpendicular distances between the AIB proximal and distal neurites (see Fig. 3n) in WT (n=28) and RIM-ablated populations (n=10 for strategy 1 and n=14 for strategy 2).****p<0.0001 by unpaired Student’s t-test between WT and each of the RIM-ablated populations. Error bars indicate standard error of the mean (S.E.M.).

Figure 5. The AIB neurite repositions onto the distal neighborhood via a zippering mechanism

(a) Schematic of axial view of the AIB neuron pair - AIBL (cyan) and AIBR (yellow) in the context of the nerve ring (light brown) and the pharynx (grey), with distal and proximal neighborhood labeled (see Fig. 1).

(b-f) Time-lapse showing initial placement of AIBL and AIBR in the proximal neighborhood and their subsequent separation from this neighborhood. Images are deconvolved diSPIM maximum intensity projections obtained from developing embryos. Neurons were individually pseudocolored to distinguish them (see Methods). The dashed boxes represent the dorsal half of the nerve ring and are magnified in (b’-f’). (b’-f’’) are schematic diagrams representing the images in (b-f). Dashed vertical lines
represents the dorsal midline. Note in (b,b',b''), the AIBL and AIBR neurites approaching the dorsal midline in the proximal neighborhood, see schematic in (a). In (c,c',c''), AIBL and AIBR have met at the dorsal midline and continue growing along each other, past the midline. The latter part of the neurite, past the midline, becomes the future distal neurite. (d,d',d'') shows the tip of the AIBL future distal neurite moving away from the proximal neighborhood and its counterpart, AIBR. The arrowhead indicates the point of separation of the AIBL distal neurite and the AIBR proximal neurite. (e,e',e'') shows further separation of the two neurites and by (f,f',f''), they have completely separated. The arrowheads in (e,e'e'') and (f,f'f'') also indicate the junction between the separating AIBL distal neurite and the AIBR proximal neurite. Scale bar = 10 \( \mu m \) for (b-f) and 2 \( \mu m \) for (b'-f'). All times are in m.p.f. (minutes post fertilization). A similar sequence of events is visualized at higher spatial resolution in Supplementary Fig. 6 using triple-view line scanning confocal microscopy.

(g) Schematic of one AIB neuron (cyan) in the context of distal neighborhood marker RIM (magenta), the nerve ring (light brown) and the pharynx (grey).

(h-j) Time-lapse showing placement of the AIB neurite (cyan) relative to the distal neighborhood marked by RIM (magenta). As in (b-f), images are deconvolved diSPIM maximum intensity projections and the neurons were pseudocolored. The dashed boxes represent the dorsal half of the nerve ring and are magnified in (h'-j'). Dashed line indicates dorsal midline (where the dorsal shift, or chiasm, in the adult is positioned, see Fig. 1). (h''-j'') are schematic diagrams representing the images in (h-j). Note in (h,h',h''), the tip of the AIB neurite encounters the growing RIM neurite (green arrowhead in (h'); black arrowhead in (h''). In (i,i',i''), the AIB distal neurite has partially
aligned along the RIM neurites. The green arrowhead now indicates point of initial
encounter of the two neurites (same as in (h’)) and the white arrowhead (in i’) indicates
the zippering event bringing the AIB and RIM neurons together in the distal
neighborhood (as black arrowhead in i”). In (j,j’,j”) the two neurites have zippered up to
the dorsal midline as indicated by the white arrowhead (j’) and black arrowhead (j”).
Arrow in (i’) indicates direction of zippering. Scale bar = 10 µm for (h-j) and 2 µm for (h’-
j’). All times are in m.p.f. (minutes post fertilization).

(k) Confocal micrograph of a postembryonic L4 animal in axial view showing the
relationship of AIB-RIM as in (j). The same image as Fig. 1(k) was used as it allows
visualization of one AIB of the pair due to mosaic array expression. Scale bar = 10 µm.
The region in the dashed box represents dorsal part of the nerve ring, magnified in (k’).
Scale bar in (k’) = 2 µm.

(l,m) The white arrowheads in d’ and i’, which correspond to the points at which the
neurites are separating or joining, are defined as the unzipping fork and zippering fork
respectively. The distances of these points, from the dorsal midline, are quantified in (l)
and (m) for different developmental stages in synchronized embryos as indicated in the
timepoints on the x-axis (±5 mins). Error bars represent standard error of the mean
(S.E.M.), n=5 in (l) and n=3 in (m).

(n) Schematic highlights the three steps by which the AIB distal neurite is repositioned
to a new neighborhood – (i) encounter with the new neighborhood; (ii) partial zippering
and (iii) complete zippering onto the distal neighborhood (marked by RIM). The term
“unzipping” is used to refer to the concomitant AIB detachment from the proximal
neighborhood as it is placed to the distal neighborhood via zippering.
Figure 6. Spatiotemporal regulation of syg-1 expression during embryogenesis correlates with AIB placement in the distal neighborhood

(a) Schematic of the axial view of the AIB neurons (cyan) with the proximal (orange arrows and dashed line) and distal (magenta arrow and dashed line) neighborhoods.

(b-d) Deconvolved diSPIM image of a late stage embryo (~1 hr prior to hatching, or 780 m.p.f.) showing colocalization of the syg-1 reporter with the AIB proximal and distal neurites (and neighborhoods). This expression pattern of syg-1 at this late embryonic stage is similar to that observed in postembryonic larvae (Fig. 4a-h). Scale bar = 10 µm, applies in (b-d).

(e-h) Time-lapse images of syg-1 reporter expression in earlier embryonic stages (450-630 m.p.f.). Images are deconvolved diSPIM maximum intensity projections. The dashed boxes represent the dorsal half of the nerve ring and are magnified in (i-l). (i'-l') are schematic diagrams representing the images in (i-l). In (e,i,i') syg-1 expression is primarily visible in a single band containing amphid neurites, and therefore coincident with the AIB proximal neighborhood. The magenta dashed line and magenta arrows point to the distal neighborhood and the orange arrow, to the proximal neighborhood. (f,j,j') show onset of weak syg-1 expression in a second neuropil band (white arrow in (j)) and ingrowth of syg-1-expressing RIM neurites along this band (white arrowhead, see also Supplementary Fig. 8) onto the distal neighborhood. syg-1 expression increases in the distal neighborhood and decreases in the proximal neighborhood as embryonic development progresses, and coincident with AIB developmental events that
enable its transition from the proximal to the distal neighborhood (j-I), (Fig. 5a-j). Scale bar = 10 µm in (e-h) and 1 µm in (i-l).

Figure 7. Ectopic syg-1 expression is sufficient to redirect placement of the AIB distal neurite to ectopic neighborhoods

(a) Schematic of lateral view of a wild type AIB neuron (cyan) in the context of the proximal (orange) and distal (magenta) neighborhoods, and the nerve ring (light brown). SYG-1 endogenous expression higher in the distal neighborhood represented by yellow arrowhead.

(b-c) Confocal image of a wild type L4 animal with AIB (labeled with cytoplasmic mCherry and pseudocolored in cyan) and the proximal neighborhood neurons AWC and ASE (labeled with cytoplasmic GFP and pseudocolored in orange). The dashed box represents the region of contact between AIB and the proximal neighborhood neurons, magnified in (c). Scale bar = 10 µm in (b) (also applies to (e) and (h)) and 1 µm in (c), (also applied to (f) and (i). Cell body is marked with an asterisk.

(d-f) As (a-c) but in the syg-1(ky652) lof (loss of function) mutant background. Note that the distal neurite is positioned away from the proximal neighborhood, as in wild type, although these animals display defects in fasciculation with the distal neighborhood (see Fig. 3).

(g-i) As (a-c) and (d-f) but with ectopic overexpression of syg-1 cDNA in the proximal neighborhood neurons and in the syg-1(ky652) mutant background. In the schematic (g), expression of SYG-1 in the proximal neighborhood (achieved using nphp-4p, also
see Supplementary Fig. 9c,d) is represented by yellow arrowhead. Note that the AIB distal neurite is now partially positioned in the proximal neighborhood in which syg-1 cDNA was ectopically expressed (h,i).

(j) Schematic (left) and scatter plot quantification (right) of minimum perpendicular distances ($d_{\text{min}}$, indicated by black double-headed arrow) between the AIB distal neurite and proximal neighborhood neurons in WT (n=11), syg-1(ky652) (n=12), and two syg-1(ky652) populations with WT syg-1 cDNA overexpressed in two different sets of proximal neighborhood neurons via the use of *nphp-4p* and *mgl-1bp* (n=12 and 10 respectively). The *nphp-4p* and *mgl-1bp* promoters are expressed primarily in proximal neighborhood neurons during embryogenesis (http://promoters.wormguides.org).

**p<0.01 by unpaired Student’s t-test between syg-1(ky652) and animals with ectopic proximal WT syg-1 expression. Error bars indicate standard error of the mean (S.E.M.).

**Figure 8. SYG-1 dictates layer-specific placement of rich club interneuron AVE**

(a,b) Schematic of the lateral and axial views of command interneuron AVE (green) in the context of its neighborhoods: proximal (magenta) and distal (yellow), with the nerve ring (light brown) and pharynx (grey). Black arrowhead in (a) indicate a posterior-anterior chiasm. The magenta and yellow arrows indicate the positions of the AVE proximal and AVE distal neighborhoods, respectively. Note that while the design principles of AVE are similar to those of rich-club interneuron AIB, their positions in the nerve ring, and the strata they connect, are different (see (c), compare to Supplementary Fig. 1a for AIB).
(c) Volumetric reconstruction of the AVE neuron (green) in the context of the nerve ring strata S2 (purple) and S3 (orange). Note the placement of the AVE proximal neurite along the border of S2 and S3, and the AVE distal neurite at the anterior boundary of S2 (the anterior boundary abuts S1, not shown here). The dashed lines indicate the layer borders. Scale bar = 1 µm.

(d) Confocal image of an L4 animal with AVE and RIM co-labeled. The magenta and yellow arrows indicate the positions of the AVE proximal and AVE distal neighborhoods, respectively. White arrowhead indicates AVE chiasm, corresponding to its anterior shift. Dashed box shows region of contact of the AVE and RIM neurites, magnified in (e). (e’) is a schematic of the image in (e). Note the region of contact between the AVE proximal neurite and the RIM neurites (e,e’), in the context of AVE, in the AVE proximal neighborhood (as compared to RIM position in AIB distal neighborhood). Scale bar corresponds to 10 µm in (d) and 1 µm in (e). Scale bars in (d) and (e) apply to (f) and (g) respectively. Cell bodies are marked with an asterisk.

(f,g,g’) As (d,e,e’) but in syg-1(ky652) mutant background. Note the gap between the AVE proximal neurite and the RIM neurites (f,g,g’) and defect in the dorsal midline shift.

(h) Scatter plot showing quantification of the loss of contacts between the AVE and RIM neurites. The extent of detachment of the AVE proximal neurites from RIM, and hence its deviation from the RIM neighborhood, was quantified using the indicated formula in Fig. 3m (also see Methods). Scatter plot depicts % detachment values for wild type (n=22) and syg-1(ky652) (n=16). Error bars indicate standard error of the mean (S.E.M.). **p<0.01 by unpaired Student’s t-test between WT and syg-1(ky652).
(i) Quantification of length of the posterior-anterior shift, quantified for each AVE neurite, for WT(n=32) and syg-1(ky652) mutants (n=40) and displayed as a scatter plot. Error bars indicate standard error of the mean (S.E.M.). **p<0.001 by unpaired Student’s t-test between WT and syg-1(ky652).

(j) Confocal image of an AVE neuron in L3 animals with RIM neurons ablated using ablation strategy 2 (Supplementary Fig. 5 and Methods). Note altered placement of the AVE proximal neurite with respect to the proximal neighborhood (indicated by the magenta dashed line and the magenta arrow). Scale bar = 10 µm.

(k) The percentage of animals having altered AVE distal neurite trajectory are plotted for WT (n=31) and RIM ablated populations (n=27). *p<0.1 by Fisher’s exact test between WT and the RIM-ablated population.

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Figure 1

AIB

GLR-1(post)
RAB-3(pre)

Post-
synaptic
Pre-
synaptic

Nerve ring
Pharynx

distal proximal

distal(RIM) proximal

LATERAL VIEW

AXIAL VIEW

Post-
pytic

GLR-1(post)
RAB-3(pre)

distal proximal

distal(RIM) proximal

Proximal

(AWC, ASE)
Figure 3

\[ \text{% detachment} = \frac{\text{detached length (}L_d\text{)} \times 100}{\text{total length (}L_t\text{)}} \]

Separation between AIB segments

WT
- uniform separation

syg-1
- non-uniform separation

Minimum dist between AIB segments, \(d_{\text{min}}\) (\(\mu\)m)
**Figure 4**

(a) syg-1p
(b) AIB
(c) merge
(d) syg-1p
(e) syg-1p
(f) RIM
(g) merge
(h) syg-1p
(i) **Relative enrichment** (syg-1p)

\[ \text{Relative enrichment} = \frac{I_d}{I_p} \]

(i) distal proximal

Mean distal intensity = \(I_d\)
Mean proximal intensity = \(I_p\)

(j) AIB
(k) RIM
(l) merge
(m) AIB
(n) RIM
(o) merge

WT

WT

WT

RIM ablation

RIM ablation

RIM ablation

(p) % animals with normal distal neurite trajectory

(q) Minimum dist between AIB segments (\(d_{min}\))
Figure 5

(a) Axial view of the post-embryonic L4 segment showing the AIB and AIBR neuron pairs.

(b) AIBL neuron in an axon, with the AIBR neuron next to it.

(c) AIBL neuron in an axon, with the AIBR neuron next to it.

(d) AIBL neuron in an axon, with the AIBR neuron next to it.

(e) AIBL neuron in an axon, with the AIBR neuron next to it.

(f) AIBL neuron in an axon, with the AIBR neuron next to it.

(g) Axial view of the post-embryonic L4 segment with the AIB and RIM neuron pairs highlighted.

(h) AIBL neuron in an axon, with the AIBR neuron next to it.

(i) AIBL neuron in an axon, with the AIBR neuron next to it.

(j) AIBL neuron in an axon, with the AIBR neuron next to it.

(k) AIBL neuron in an axon, with the AIBR neuron next to it.

(l) Distance of unzippering fork from dorsal midline (μm).

(m) Distance of zippering fork from dorsal midline (μm).

(n) Zippering/zippering direction.

(i) Proximal neurite encounters RIM neighborhood.

(ii) Partially zippered.

(iii) Completely zippered.

→ Unzippering/zippering direction.
Figure 6

(a) AXIAL VIEW

(b) AIB

(c) Psyg-1

(d) 780 mins merge

(e) 457 mins Psyg-1s

(f) 487 mins

(g) 529 mins

(h) 628 mins

(i) i' j' k' l'

(j) j' k' l'

(k) k'

(l) l'

AXIAL VIEW
**Figure 7**

**a** syg-1 expression higher in distal neighborhood neurons

**d** syg-1 lof mutant

lof mutant - no functional SYG-1

**g** syg-1 mutant+ proximal ectopic WT syg-1

ectopic syg-1 expression in proximal neighborhood neurons

**j** distal proximal

Min dist, d_{min} of distal neurite to proximal neighborhood (µm)

**nphp-4p;syg-1b**

**mgl-1bp;syg-1b**

WT syg-1

WT syg-1

WT syg-1

WT syg-1

WT syg-1

WT syg-1
Figure 8

(a) Lateral view

(b) Axial view

(c) EM reconstruction

(d) WT AVE RIM

(e) syg-1 AVE RIM

(f) WT AVE

(g) syg-1 AVE

(h) % detachment

(i) Dorsal midline shift length (μm)

(j) RIM ablation

(k) % animals with normal AVE neurite trajectory