Adult Drosophila sensory neurons specify dendritic territories independently of dendritic contacts through the Wnt5–Drl signaling pathway

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Sensory neurons with common functions are often nonrandomly arranged and form dendritic territories in stereotypic spatial patterns throughout the nervous system, yet molecular mechanisms of how neurons specify dendritic territories remain largely unknown. In Drosophila larvae, dendrites of class IV sensory (C4da) neurons completely but nonredundantly cover the whole epidermis, and the boundaries of these tiled dendritic fields are specified through repulsive interactions between homotypic dendrites. Here we report that, unlike the larval C4da neurons, adult C4da neurons rely on both dendritic repulsive interactions and external positional cues to delimit the boundaries of their dendritic fields. We identify Wnt5 derived from sternites, the ventral-most part of the adult abdominal epidermis, as the critical determinant for the ventral boundaries. Further genetic data indicate that Wnt5 promotes dendrite termination on the periphery of sternites through the Ryk receptor family kinase Derailed (Drl) and the Rho GTPase guanine nucleotide exchange factor Trio in C4da neurons. Our findings thus uncover the dendritic contact-independent mechanism that is required for dendritic boundary specification and suggest that combinatorial actions of the dendritic contact-dependent and -independent mechanisms may ensure appropriate dendritic territories of a given neuron.

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Precise patterning of the dendritic fields is essential for the correct wiring of neuronal circuitry. In many cases, after dendrites cover their territory, growth beyond the boundaries of their territory is curtailed in order to prevent any overlap of the receptive fields of neighboring neurons and the consequent compromise of neuronal circuit properties. Indeed, neuronal diseases characterized by the formation of enlarged dendritic fields result in severe mental retardation (Purpura 1975; Kaufmann and Moser 2000). Notwithstanding recent progress in our knowledge of molecular mechanisms that promote dendritic elaboration (Jan and Jan 2010; Masland 2012), deletion of RGCs causes surrounding neurons to reorient their dendrites toward the depleted area (Perry and Linden 1982), suggesting that mutual repulsion between like dendrites mediates establishment of the RGCs’ dendritic boundaries. However, it has also been suggested that certain types of RGCs can define their unique territories independently of the dendrite–dendrite contacts. For example, in Brn3b−/− and Math5−/− retinas in which only ~10% RGCs are formed, the dendritic arbors of at least two classes of surviving neurons are indistinguishable from normal in shape and size (Lin et al. 2004), raising a possibility that, in addition to the dendritic contact-dependent mechanisms, dendritic contact-independent...
mechanisms likely function to specify dendritic territories even in a two-dimensional (2D) space.

The *Drosophila* dendrite arborization (da) sensory neurons provide a suitable system for the systematic analysis of dendritic territory development (Parrish et al. 2007; Jan and Jan 2010; Emoto 2012). Similar to mammalian RGCs, dendritic arborizations of larval class IV da (C4da) neurons are confined to the 2D space between the epidermis and muscles, and the dendrites cover the whole epidermis in a complete but nonredundant manner. Several lines of evidence demonstrate that dendritic boundaries in larval C4da neurons are defined by homotypic repulsion between neighboring dendrites. First, C4da dendrites display typical avoidance behaviors when two dendritic branches meet at dendritic boundaries (Grueber et al. 2003; Emoto et al. 2004, 2006). Second, duplication of C4da neurons resulted in a partitioning of the receptive field (Grueber et al. 2003). Last, embryonic ablation of C4da neurons caused the remaining C4da neurons to expand their dendritic fields to occupy the territory of the ablated neurons (Grueber et al. 2003). Genetic studies in C4da neurons have revealed that the contact-dependent dendrite repulsions are mediated by multiple molecules, including cell surface proteins (Gao et al. 2000; Hughes et al. 2007; Matthews et al. 2007; Soba et al. 2007) and intracellular signaling molecules (Emoto et al. 2004, 2006; Koike-Kumagai et al. 2009; Matsubara et al. 2011).

During metamorphosis, the larval dendritic arbors are completely replaced with adult-specific processes as a result of extensive pruning and subsequent regeneration of dendritic trees (Kuo et al. 2005; Shimono et al. 2009; Yasunaga et al. 2010; Kanamori et al. 2015). In this study, we investigated how adult C4da neurons rebuild the dendritic territory and found that, unlike the larval dendrites, adult C4da dendrites specify a dendritic boundary independently of dendritic contacts. Our genetic screen revealed that Wnt5 derived from sternites, the ventral-most region of the adult abdomen, is required for specification of the ventral boundaries of C4da dendrites. We further show that Wnt5 promotes dendrite termination on the periphery of sternites through the Ryk receptor family kinase Derailed (Drl) and the Rho guanine nucleotide exchange factor (GEF) Trio in C4da neurons. Our findings thus reveal a novel role of the Wnt5–Drl signaling pathway in the contact-independent dendritic boundary specification and raise a possibility that multiple distinct mechanisms function in parallel to define the dendritic territories of a given neuron.

**Results**

**Dendritic field organization of adult C4da neurons**

During metamorphosis, all three C4da neurons degrade their larval dendrites, and two of them, the dorsal ddaC neuron and the ventro–lateral v’ada neuron, subsequently reconstruct adult-specific dendrites, while the ventral vdaB neuron undergoes apoptosis (Fig. 1A; Kuo et al. 2005; Shimono et al. 2009; Yasunaga et al. 2010). To examine how the remaining two adult C4da neurons recon-
mCD8GFP [Kuo et al. 2005; Yasunaga et al. 2010; Kanamori et al. 2013]. We focused on v’ada neurons that innervate the ventro–lateral epidermis (Shimono et al. 2009; Yasunaga et al. 2010), as the dorsal ddaC dendrites are difficult to image due to tanning of the cuticle. Similar to the larval C4da neurons, adult v’ada neurons positioned their lateral dendritic terminals around the segmental boundaries, and two neighboring dendrites appeared to cover completely the segmental boundaries [Fig. 1B1, B2]. To further examine the relative positions of the neighboring dendrites in detail, we stochastically labeled v’ada neurons with EGFP and mCitrine by using the Flybow system [Hadjieconomous et al. 2011]. This stochastic labeling experiment clearly showed that adult C4da neurons in neighboring segments cover the segment boundaries with dendrites in a complete but nonredundant manner [Supplemental Fig. S1]. In contrast, unlike the larval dendrites, adult v’ada dendrites did not extend to the ventral midline. Instead, growth of adult v’ada dendrites terminated ∼20 µm away from the ventral midline (larva, −33.8 µm ± 3.8 µm, n = 32; adult, 17.1 µm ± 4.8 µm, n = 32) [Fig. 1B1–D]. As a result, the dendritic fields of the two adjacent v’ada neurons were separated by ∼40 µm, and that space was not covered by C4da dendrites. These data suggest that adult v’ada dendrites might use different mechanisms to specify lateral and ventral boundaries.

Ventral boundaries of adult v’ada dendrites are specified independently of dendrite contacts

To test whether neighboring neurons affect the dendritic boundary specification of adult v’ada neurons, we ablated v’ada neurons in the early pupal stage by using the FLP-out system that induces stochastic expression of the toxic ricin A chain in C4da neurons [Smith et al. 1996]. Figure 2B shows an example of a v’ada neuron at the abdominal segment 4 (A4) whose neighboring neurons were ablated. In this situation, both the width and the field size of the v’ada dendrites were significantly expanded compared with those of control v’ada neurons [field size: control, 21.9 × 10⁴ µm² ± 1.1 × 10⁴ µm², n = 6; ablation, 26.9 × 10⁴ µm² ± 1.1 × 10⁴ µm², n = 9; width: control, 273 µm ± 9 µm, n = 10; ablation, 352 µm ± 14 µm, n = 13] (Fig. 2A–Figure 2. Neighboring neurons are required for proper specification of the lateral boundaries but are dispensable for the ventral boundaries in v’ada neurons. (A, B) Lateral views of adult v’ada dendrites in the A3–A5 segments of an adult ventral abdomen. (A,A’) Dendrites of three v’ada neurons cover the body wall completely but redundantly. (B,B’) Ablation of neighboring v’ada neurons leads to expansion of dendritic fields of the remaining neuron. Sternites, the ventral-most epithelial region in the adult abdominal segments, are labeled in yellow. Bar, 100 µm. (C, D) Quantification of the dendritic field area [C] and the field width [D] of v’ada neurons in control and cell-ablated abdomens. Error bars indicate standard error of the mean. In C, P = 0.01. In D, P < 0.001. (E, F) Ventral views of adult v’ada dendrites in A3–A5 segments. (E,E’) Ventral boundaries of v’ada neurons are established on the periphery of sternites. (F,F’) The ventral boundaries are unaffected by ablation of v’ada neurons in the contralateral hemisegments. Magnified views of the blue boxed regions are shown in E” and F”. Sternites are labeled in yellow. Bars: E, 100 µm; E’, 50 µm. (G,H) Quantification of the distance from the ventral midline to the branch terminals [G] and the total dendritic length within single sternites [H] in control and cell-ablated abdomens. In G, P = 0.486. In F, P = 0.669. [*] P < 0.05; [**] P < 0.01, unpaired Student’s t-test.
D). These data suggest that the lateral boundaries of v'ada dendrites are specified at least in part by repulsive interactions between neighboring dendrites. Consistent with these ablation data, time-lapse imaging of dendritic terminals at the segmental boundary revealed the typical avoidance behaviors of dendritic terminals when they met at the segmental boundaries [Supplemental Movie S1]. In contrast to the lateral boundaries, ablation of contralateral v'ada neurons had no obvious effect on the relative position of ventral boundaries of v'ada neurons (distance: control, 29.9 µm ± 6.7 µm, n = 16; ablation, 22.8 µm ± 7.5 µm, n = 15; length: control, 349 µm ± 45 µm, n = 16; ablation, 322 µm ± 43 µm, n = 15) [Fig. 2E–H], suggesting that the ventral boundaries of v'ada neurons can be specified independently of neighboring dendrites. In both the control and the contralateral ablated cases, the ventral boundaries were coincident on the peripheral of sternites, the ventral-most epithelial region in the adult abdominal segments, which contains arrays of mechanosensory bristles. These data together suggest that dendritic territories of adult v'ada neurons are specified by two distinct mechanisms: Ventral boundaries are specified independently of dendritic contacts, whereas the lateral boundaries are established through repulsive interactions between homotypic dendrites.

**Sternites are required to specify the ventral boundaries of adult v'ada dendrites**

Adult abdominal segments are subdivided along the dorso–ventral axis into a dorsal tergite, ventro–lateral pleura, and a ventral sternite [Kopp et al. 1999]. Given that v'ada neurons positioned the ventral boundaries on the periphery of sternites, we reasoned that sternites might play a role in the establishment of the ventral boundaries. Previous studies reported that the Wingless–Frizzled signaling pathway promotes sternite identity in the adult abdomen and that suppressing the Wingless–Frizzled signaling in the pupal abdomen results in the transformation of sternites into pleura [Shirras and Couso 1996; Kopp et al. 1999]. To test the role for sternites in the dendritic boundary specification, we induced somatic clones mutant for both *frizzled* (*fz*) and *frizzled 2* (*fz2*), which are known as the major Wingless receptors in the adult abdomen [Shirras and Couso 1996; Kopp et al. 1999], and successfully generated hemisegments in which sternites were largely transformed into pleura (Fig. 4A–B; Supplemental Table S1). Among the 37 genes tested, Wnt5 derived from sternites restricted the ventral boundaries of v'ada dendrites (also called *Dwnt5*) mutants showed robust defects in ventral boundary formation. Although wild-type dendrites formed their ventral boundaries on the periphery of sternites, dendritic territories of Wnt5 neurons expanded significantly toward the ventral midline. Importantly, in both the control and transformed cases, the ventral boundaries of v'ada dendrites are established on the periphery of sternites. Arrowheads indicate the ventral midline. Bar, 100 µm.

To uncover the potential factors derived from sternites, we carried out a genetic screen, focusing on genes that encode secreted molecules and their receptors [Supplemental Table S1]. Among the 37 genes tested, Wnt5 also showed robust defects in ventral boundary formation. Although wild-type dendrites formed their ventral boundaries on the periphery of sternites, dendritic territories of Wnt5 neurons expanded significantly toward the ventral midline, and dendrites of two v'ada neurons seemed to cover the whole ventral epidermis (distance: wild-type, 20.1 µm ± 4.7 µm, n = 36; Wnt5, −13.0 µm ± 6.6 µm, n = 24; length: wild-type, 778 µm ± 63 µm, n = 18; Wnt5, 2188 µm ± 107 µm, n = 12) [Fig. 4A–D]. Interestingly, two v'ada dendrites seemed to tile a sternite with an obvious boundary at the ventral midline [Fig. 4B], suggesting that repulsive interactions...
between two v’ada dendrites likely specify the boundary. Unlike mutants for the Wingless–Frizzled signaling pathway [Fig. 3], sternites were correctly formed at the ventral-most position in Wnt5 mutants, suggesting that Wnt5 is dispensable for sternite development. Indeed, no significant difference was detectable in the number of bristles and the average size of the sternites (Supplemental Fig. S2G,H). In addition, neighboring Wnt5 dendrites seemed to cover the whole segmental boundary, which is indistinguishable from that in wild-type dendrites (Supplemental Fig. S2A–D). Further quantitative analysis indicated that the lengths and number of dendritic branches outside of sternites were unaffected in Wnt5 mutants [Supplemental Fig. S2E,F]. Simultaneous observation of C4da dendrites and the extracellular matrix (ECM) suggest that the dendrite–ECM interaction was also unaffected in Wnt5 mutants [Supplemental Fig. S2I,J]. These data suggest that Wnt5 plays a specific role in the establishment of the ventral boundary of v’ada dendrites rather than general roles in dendrite growth and branching.

To investigate temporal and spatial expression patterns of Wnt5 in the pupal/adult abdomen, we generated a novel Wnt5 reporter line in which EGFP is inserted into the first exon of the Wnt5 locus [Supplemental Fig. S3]. We first examined EGFP expression in the pupal optic lobes, since the anti-Wnt5 antibody detected high levels of Wnt5 in the medulla of pupal optic lobes [Srahna et al. 2006]. Similarly, we found that the Wnt5-EGFP reporter was highly expressed in medulla neurons, and the expression patterns were consistent with the anti-Wnt5 staining, confirming that the Wnt5 reporter mimics expression patterns of endogenous Wnt5 [Supplemental Fig. S3]. Using this Wnt5 reporter, we next examined Wnt5 expression in the pupal/adult abdomen. The larval abdomen is reorganized during metamorphosis, and sternites are initially induced ∼60 h after pupal formation (APF) in the ventral-most region of the pupal abdomen (Kopp et al. 1999). Consistent with sternites functioning as the source for Wnt5, the Wnt5-EGFP reporter signal became visible weakly in the ventral-most abdominal regions ∼60 h APF, and the expression levels were gradually increased as the pupa developed to the adult [Fig. 4E]. By the 1-d adult stage, EGFP expression was eventually concentrated to the peripheral region of sternites [Fig. 4E]. Remarkably, double staining of Wnt5-EGFP and v’ada dendrites revealed that Wnt5 expression is consistent with the ventral boundaries of v’ada dendrites [Fig. 4F1,F2]. Taken together, these data suggest that Wnt5 is a sternite-derived factor that specifies the ventral boundaries of adult v’ada dendrites.

To further investigate the role of Wnt5 in dendrite boundary specification, we assessed whether ectopic Wnt5 might affect dendritic boundaries in larval v’ada...
neurons. Unlike the adult peripheral tissue, no obvious Wnt5-GFP expression was detectable in larval epithelial cells, whereas larval v’ada neurons express Drl receptor (Supplemental Fig. S4D,E). We ectopically expressed Wnt5 in the larval epithelial cells around which C4da dendrites establish lateral boundaries and found that v’ada dendrites were significantly excluded from the region where ectopic Wnt5 was highly expressed, whereas no obvious growth defect was observed in the region without ectopic Wnt5 (Supplemental Fig. S4A–C). These data suggest that Wnt5 is necessary and sufficient to specify C4da dendrite boundaries.

**Wnt5 promotes dendrite termination on the ventral boundary**

How might Wnt5 specify the ventral boundary of v’ada dendrites on the appropriate position? One possibility is that dendrites initially innervate into sternites, followed by specific retraction or pruning of the dendritic branches that innervate into sternites, since dendrite pruning/retraction is often observed in developing mammalian sensory circuits to refine dendritic fields (Wong and Ghosh 2002; Emoto 2011). An alternative scenario is that Wnt5 might prevent dendritic branches from innervating sternites. To distinguish between these possibilities, we first measured the ventral extent of v’ada dendrites during the first 5 d of adult development in wild-type and Wnt5 mutants. At eclosion, no significant difference was observed in both the distance from dendritic terminals to the ventral midline (Fig. 5A) and the dendrite length within single sternites (Fig. 5B) in wild-type controls and Wnt5 mutants. Over the next 24 h, wild-type neurons appeared to reduce dendrite growth, whereas Wnt5 dendrites grew continuously at least during next several days (Fig. 5A,B). Thus, v’ada dendrites appear to avoid sternites, as they initially arrive at the periphery of sternites without going through a noticeable period of invasion followed by retraction/pruning.

To further examine this model, we imaged single v’ada dendrites of live wild-type controls and Wnt5 mutants for 48 h, starting at the time of eclosion. At the 0-d adult stage, many of the terminal branches reached the periphery of sternites in both the wild-type control and Wnt5 mutants (Fig. 5C). Over the next 24 h, Wnt5 dendrites grew into sternites, while wild-type dendrites persisted at the periphery of sternites (Fig. 5C). Quantification of the branch dynamics revealed that dendrite extension was significantly reduced in the wild type over the 48 h, whereas branch extension was much less affected in the Wnt5 mutants compared with wild type (median: 36 µm for 0–1 d and 19 µm for 1–2 d in wild type; 48 µm for 0–1 d and 38 µm for 1–2 d in Wnt5) (Fig. 5D). In contrast,
no significant difference was detected in the dendrite retraction between the wild-type controls and Wnt5 mutants during the 48 h (median: 18 µm for 0–1 d and 17 µm for 1–2 d in wild type; 20 µm for 0–1 d and 15 µm for 1–2 d in Wnt5) (Fig. 5D). These observations are consistent with the scenario that Wnt5 dendrites are able to grow and retract normally but are specifically defective in termination on the periphery of sternites (Fig. 5E).

**Drl and Drl-2 receptor kinases in v'ada neurons mediate Wnt5-dependent boundary determination in a redundant manner**

Wnt5 typically acts through cell surface receptors on target cells (Fradkin et al. 1995, 2010; Yoshikawa et al. 2003; Kikuchi et al. 2007). To identify the functional Wnt5 receptors required for dendrite boundary specification in v'ada neurons, we examined nine potential Wnt5 receptors and found that mutants for the Ryk receptor family kinase Drl-2 showed a weak but significant defect in the boundary specification of v'ada dendrites (distance: wild type, 31.9 µm ± 5.2 µm, n = 28; Drl-2, 8.8 µm ± 5.1 µm, n = 26; length: wild type, 1000 µm ± 106 µm, n = 14; Drl-2, 1176 µm ± 93 µm, n = 13) (Fig. 6A,B,E,F; Supplemental Table S1). Mutants for Drl, the other member of the Drosophila Ryk receptor kinase family, also showed weak boundary defects (distance: drl, 29.6 µm ± 6.1 µm, n = 28; length: drl, 1000 µm ± 106 µm, n = 14) (Fig. 6A,B,E,F). We thus next examined drl Drl-2 double mutants and found robust boundary defects, which were comparable with those in Wnt5 mutants (distance: drl Drl-2, −13.0 µm ± 4.9 µm, n = 26; length: drl Drl-2, 1847 µm ± 85 µm, n =...
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13) (Fig. 6D–F). Similar to Wnt5 mutants, drl Drl-2 double mutants showed no significant defects in the lateral boundary formation [Supplemental Fig. S5A–H]. These results suggest that Drl and Drl-2 function together to establish the ventral boundaries of v’ada dendrites.

To examine whether Drl receptors could function cell-autonomously in v’ada neurons to specify the ventral boundary, we performed rescue experiments by expressing drl or Drl-2 in drl Drl-2 double-mutant neurons. Expression of either drl or Drl-2 driven by ppp-Gal4 fully rescued the boundary defects in drl Drl-2 double-mutant neurons (distance: drl Drl-2, –8.6 µm ± 5.4 µm, n = 16; +UAS-drl, 14.7 µm ± 11.3 µm, n = 14; +UAS-Drl-2, 8.4 µm ± 9.1 µm, n = 12; length: drl Drl-2, 1828 µm ± 98 µm, n = 8; +UAS-drl, 1187 µm ± 111 µm, n = 7; +UAS-Drl-2, 1064 µm ± 166 µm, n = 6) [Fig. 6G,H, Supplemental Fig. S6], indicating that Drl and Drl-2 act cell-autonomously and redundantly in v’ada neurons to specify the ventral dendritic boundaries. Consistent with this notion, RNAi knockdown of drl and Drl-2 in C4da neurons caused boundary defects similar to those in drl Drl-2 mutants [Supplemental Fig. S5J].

Finally, we examined expression patterns of Drl receptors by using the Gal4 enhancer trap in drl that has been shown to mimic the expression pattern of drl to drive expression of GFP [Moreau-Fauvarque et al. 1998]. In adult peripheral tissues, drl expression was specifically observed in C4da neurons but not in abdominal muscle fibres and epithelial cells, including sternites [Fig. 6I1–I3], further confirming that Drl receptors function in adult C4da neurons to specify the ventral boundaries. We further examined intracellular distribution of Drl by expressing Drl::GFP in C4da neurons. Drl::GFP seemed to be localized evenly in dendritic arbors as well as axons and the soma [Supplemental Fig. S7].

Rho GEF Trio and Rho1 act with the Wnt5–Drl signaling pathway to specify dendrite boundaries

In many contexts, the Ryk receptor family kinases act through their intracellular domains [Fradkin et al. 2010]. Consistently, we found that Drl without the intracellular domain failed to rescue the boundary defects in drl Drl-2 double-mutant neurons (length: 1727 µm ± 151 µm, n = 8) [Fig. 6H], indicating that the intracellular domains of Drl receptors are required for their functions in dendritic boundary specification. Previous studies suggest that Wnt5 signals at least in part through the canonical Wnt signaling pathway [Shimizu et al. 2011]. The Src family kinases are also suggested to associate and function with Drl receptors in axon guidance [Wouda et al. 2008]. We therefore examined whether attenuation of the canonical Wnt signaling pathway or Src kinase signaling could affect ventral boundary formation and found no significant defects [Supplemental Table S2].

The Rho GTPase GEFs and GAPs [GTPase-activating proteins] function downstream from many receptors to regulate neurite dynamics by controlling the actin cytoskeletons [Luo 2000; Dickson 2001; Bashaw and Klein 2010]. In the Drosophila genome, 21 potential Rho GAPs and 20 Rho GEFs are present [Bernards 2003]. To examine whether Rho GEFs or GAPs might be involved in the dendritic boundary specification, we screened all potential Rho GAPS and GEFs by using RNAi knockdown in neurons and found significant boundary defects in adult v’ada dendrites following knockdown of the Rho GEF trio [Supplemental Table S3]. To further confirm this, we conducted MARCM [mosaic analysis with a repressive cell marker] analysis to generate trio homozygous mutant clones in a heterozygous background [Lee and Luo 1999]. We found that trio mutant clones showed significant boundary defects in v’ada dendrites (distance: control, 40.0 µm ± 5.1 µm, n = 13; Trio, –7.3 µm ± 12.1 µm, n = 13; length: control, 447 µm ± 37 µm, n = 13; Trio, 620 µm ± 92 µm, n = 13) [Fig. 7A,B,D]. These boundary defects in trio MARCM clones were fully rescued by expression of the trio gene in v’ada neurons (distance: 41.1 ± 8.6 µm, n = 13; length: 460 ± 53 µm, n = 13) [Fig. 7C–E], indicating that Trio functions autonomously in v’ada neurons.

To investigate whether Trio and Drl receptors might function in the same genetic pathway to regulate ventral boundary formation, we assayed for genetic interactions between trio and the Wnt5–Drl signaling genes and found that transheterozygous combinations of mutations in trio together with Wnt5 caused significant boundary defects, whereas heterozygosity for each of the genes had no detectable effect on boundary formation [Supplemental Fig. S9]. Similarly, transheterozygous combinations of trio and drl Drl-2 caused significant boundary defects [Supplemental Fig. S9]. Thus, Trio and the Wnt5–Drl signaling genes interact genetically to specify dendritic boundaries in v’ada neurons.

Trio could function as a GEF for both Rho1 and Rac1 in a context-dependent manner [Bateman and Van Vactor 2001; Schmidt and Debant 2014]. To distinguish between these possibilities, we overexpressed the dominant-negative version of Rho1 or Rac1 specifically in the pupal/adult C4da neurons by using the Flip-out technique [Kanamori et al. 2013]. We found that neuronal expression of the dominant-negative Rho1 caused significant boundary defects similar to those seen in Wnt5 mutants, whereas no obvious boundary defect was apparent following expression of the dominant-negative Rac1 [Supplemental Fig. S10], suggesting that Trio specifies the dendrite boundary through Rho1 activation.

To further examine whether Wnt5–Drl might signal through the Trio–Rho1 pathway to specify the dendritic boundary in v’ada neurons, we next investigated whether overexpression of Trio or a dominant active version of Rho1 in v’ada neurons could rescue the Wnt5 loss-of-function phenotypes. We found that overexpression of Trio in v’ada neurons significantly rescued the dendritic boundary defects in Wnt5 mutants (distance: Wnt5, –3.8 µm ± 4.2 µm, n = 32; Wnt5 UAS-Trio, 8.0 µm ± 4.3 µm, n = 36; length: Wnt5, 1846 µm ± 115 µm, n = 16; Wnt5 UAS-Trio, 1434 µm ± 125 µm, n = 18) [Fig. 7F–K]. Similarly, overexpression of the constitutively active Rho1 in Wnt5 mutant v’ada neurons completely rescued the boundary defects (distance: Wnt5 UAS-Rho1-CA, 26.6 µm ± 7.6 µm, n = 16; length: Wnt5 UAS-Rho1-CA, 1333 µm ± 251 µm, n = 8)
Importantly, overexpression of Trio or the dominant active Rho1 in wild-type v’ada neurons caused no significant defects in the ventral boundary specification of v’ada dendrites (Fig. 7J), suggesting that Trio and Rho1 activities do not simply arrest dendritic growth and/or branching in adult v’ada neurons; rather, Trio and Rho1 specifically promote dendrite termination on the periphery of sternites downstream from the Wnt5–Drl signaling.

Taken together, our findings indicate that Wnt5–Drl signaling promotes dendrite termination at least in part through the Trio–Rho1 pathway in adult v’ada neurons.

Figure 7. Trio and Rho1 promote dendrite termination downstream from Drl receptors. (A–C) Ventral views of v’ada dendrites in wild-type control (A) and trioE4.1 (B) and trioE4.1 with a UAS-trio transgene (C) MARCM clones. Arrowheads indicate the ventral midline. (D,E) Quantification of the distance from the ventral midline to the dendritic branch terminals [D] and the total dendritic length within single sternites [E]. Note that we quantified the dendritic length of single neurons. Error bars indicate standard error of the mean. The numbers below each bar indicate n values. In D, control versus trioE4.1, P = 0.002; trioE4.1 versus trio rescue, P = 0.002, one-way ANOVA followed by Tukey’s HSD test. In E, P = 0.124, one-way ANOVA. (F–I) Ventral views of v’ada dendrites in wild-type control (F), Wnt5 mutants (G), Wnt5 mutants with a UAS-Trio transgene [H], and Wnt5 mutants with a UAS-Rho1-CA transgene [I]. (J,K) Quantification of the distance from the ventral midline to dendritic branch terminals [J] and the total dendritic length within single sternites [K]. The numbers below each bar indicate n values. In J, control versus UAS-trio versus UAS-Rho1-CA, P = 0.115, one-way ANOVA; Wnt5 versus Wnt5 UAS-trio, P = 0.081; Wnt5 versus Wnt5 UAS-Rho1-CA, P = 0.001, one-way ANOVA followed by Dunnett’s test. In K, control versus UAS-trio, P = 0.228; control versus UAS-Rho1-CA, P = 0.005, one-way ANOVA followed by Dunnett’s test; Wnt5 versus Wnt5 UAS-trio, P = 0.031; Wnt5 versus Wnt5 UAS-Rho1-CA, P = 0.039, one-way ANOVA followed by Dunnett’s test. (*) P < 0.05; (**) P < 0.01.

Discussion

Dendritic territories of sensory neurons in the same functional class are often organized in stereotyped patterns. Previous studies have established that dendritic boundaries of the larval C4da sensory neurons in Drosophila are predominantly established through mutual repulsion between neighboring dendrites (Grueber et al. 2003; Emoto et al. 2004). In this study, we showed that, unlike the larval neurons, adult v’ada neurons establish the ventral boundaries independently of neighboring neurons (Figs. 1, 2). Furthermore, our genetic ablation data, in which transformation of sternites to pleura caused the ventral boundaries to extend ventrally (Fig. 3), indicated that sternite-derived factors are required for the ventral boundary specification (Fig. 5). In contrast to the ventral boundary, v’ada neurons required adjacent neurons to define the lateral boundaries, like the larval v’ada neurons (Figs. 1, 2). Thus, v’ada dendrites establish the lateral boundaries through the dendritic contact-dependent mechanism, whereas the ventral boundaries are specified by the contact-independent mechanism.
The cellular mechanisms underlying contact-dependent and -independent boundary determination seem to be distinct. Our time-lapse observations demonstrated that dendritic terminals of adult v’ada neurons remained dynamic to avoid each other at the lateral boundaries even after they established the boundaries [Supplemental Movie S1]. This is consistent with the previous report that dendrite boundaries of larval C4da neurons are defined by continuous avoidance behaviors of terminal branches [Emoto et al. 2004]. In contrast, in the ventral boundaries, adult v’ada neurons exhibited reduced dendrite dynamics after reaching the boundary on the periphery of sternites and eventually arrested dendrite growth [Fig. 5]. Therefore, the contact-independent boundary specification appears to be mediated by dendrite termination, whereas the contact-dependent mechanism is likely mediated by homotypic repulsion. These two mechanisms appear to act independently, since no obvious defects were observed in lateral boundary formation in v’ada dendrites mutant for Wnt5 and drl despite the fact that the ventral boundaries were significantly impaired [Figs. 4, 6].

Recent studies have identified multiple repulsive signaling pathways that function in dendrite patterning of vertebrate and invertebrate neurons, including semaphorin/plexin [Polleux et al. 2000; Matsuoka et al. 2012; Sun et al. 2013], Dscams [Matthews et al. 2007; Soba et al. 2007; Fuerst et al. 2009], protocadherins [Lefebvre et al. 2012], Ntrin/DCC/UNC-5 [Smith et al. 2012], and Slit/Robo [Furrer et al. 2007; Gibson et al. 2014]. We thus examined potential roles of the repulsive factors in the ventral boundary specification of v’ada dendrites by using genetic mutants or RNAi knockdown, but, to our surprise, none of them showed obvious defects in v’ada dendrite boundary specification [Supplemental Table S1]. Instead, using an unbiased genetic screen, we identified Wnt5, a secreted member of the Wnt family proteins, as the critical factor derived from sternites [Fig. 4]. We also identified the Ryk receptor family kinases Drl and Drl-2 as the receptors for Wnt5 in v’ada neurons (Fig. 6). These data have demonstrated that Wnt5–Drl signaling plays an essential role in the ventral boundary specification of v’ada dendrites.

How does Wnt5–Drl signaling specify the dendritic boundary at the stereotyped position? In the mouse spinal cord, Wnt5a is expressed by the cells surrounding corticospinal tract axons in a gradient along the anterior–posterior axis, and this Wnt gradient repels corticospinal axons down the spinal cord [Liu et al. 2005]. In the present study, we found that Wnt5 expression in the ventral epithelium is dramatically changed during the pupal/adult stage [Fig. 4E]. Wnt5 expression was initially visible in epithelial cells around the entire sternite regions at ∼60 h APF, but Wnt5 production was gradually restricted to the peripheral region of sternites during the adult stages. In contrast, Drl receptors seemed to be expressed in v’ada neurons throughout the larval-to-adult development at the same level [Fig. 6I]. Thus, the spatial and temporal control of Wnt5 expression likely defines the appropriate positioning of the ventral boundary. In addition, our studies using a novel Wnt5 reporter indicated that the ventral dendritic

boundaries of v’ada neurons are coincident with the places where Wnt5 is produced [Fig. 4F]. Therefore, in spite of its diffusive nature, Wnt5 seems to exert its influence on v’ada dendrites in the vicinity of the protein source. Recent studies indicate that Wnt5a contains multiple lipid modifications, including palmitoylation, and that the lipidation of Wnt5a protein is critical for its functions [Kikuchi et al. 2007]. It is thus possible that Wnt5 proteins secreted from sternites might be anchored to cell membranes or nearby in the ECM to produce a sharp boundary for v’ada dendrites at the precise regions. Collectively, it appears likely that the spatiotemporally regulated expression as well as the short-range action enable Wnt5 to define a sharp dendritic boundary at the appropriate timing and position.

In the nervous system, Wnt5–Drl/Ryk signaling acts as both attractive and repulsive cues for axons and dendrites presumably through distinct downstream targets, including Src family kinases and CaMKII [Wouda et al. 2008; Hutchins et al. 2011; Wu et al. 2014]. Our time-lapse observations revealed that adult v’ada neurons terminate dendrite growth specifically on the periphery of sternites, where Wnt5 is locally produced, without obvious attractive or repulsive responses to the Wnt5 source [Figs. 4, 5]. Thus, Wnt5–Drl signaling might limit dendrite growth on the periphery of sternites rather than function as attractive/repulsive cues for v’ada dendrites. Consistent with this scenario, neither Src kinases nor CaMKII was required for the dendrite boundary specification [Supplemental Table S2]. Instead, we identified the Rho GTPase GEF Trio and Rho1 as the novel downstream factors that mediate Wnt5–Drl signaling in dendrite termination. This idea is supported by the following lines of evidence. First, reduction of Trio or Rho1 activity in v’ada neurons caused dendrite boundary defects similar to those observed in Wnt5 mutants [Fig. 7; Supplemental Fig. S10]. Second, Wnt5 and drl mutations showed a strong genetic interaction with trio mutations in the dendrite boundary specification of adult v’ada neurons [Supplemental Fig. S9]. Third, overexpression of Trio or the constitutively active Rho1 in Wnt5 mutant neurons partially rescued the boundary defects of v’ada dendrites [Fig. 7]. This model is consistent with previous reports that loss-of-function mutations in Rho1 and Trio cause Drosophila mushroom body neurons to overshoot their dendritic territories, whereas constitutively active Rho1 expression results in a reduction of dendritic fields [Awasaki et al. 2000; Lee et al. 2000]. It is also possible that Trio–Rho1 might function in parallel to the Wnt5–Drl signaling in C4da neurons. How Wnt5 activates Trio–Rho1 signaling remains to be elucidated. Given that the intracellular domain of Drl receptor kinases is indispensable for dendrite boundary specification [Fig. 6G,H], Trio could be activated through the intracellular domain of Drl receptors, including the tyrosine kinase domain. Interestingly, recent reports indicate that Trio is phosphorylated at multiple tyrosine residues and that the tyrosine phosphorylations appear to be essential for its Rho GEF activity [DeGeer et al. 2013; Sonoshita et al. 2014]. It is thus of interest to test whether Trio could be phosphorylated in response to Wnt5 signals in v’ada neurons.
Developing neurons in the mammalian brain often extend their dendritic arbors into a three-dimensional (3D) space and thus have to establish distinct dendritic boundaries within different environments. Therefore, although mutual repulsion is the most effective way to specify dendritic boundaries when dendrites are confined to the 2D space, multiple systems, including dendritic contact-dependent and -independent mechanisms, are probably required to specify distinct dendritic boundaries in the 3D space. Given that we showed the critical role of Wnt5–Ryk signaling in the contact-independent mechanism for specifying dendritic boundaries in v’ada neurons, it is of great interest to investigate whether the Wnt5–Ryk signaling might play a significant role in establishing the dendritic territories in the 3D space, including the mammalian cortex, as well as in the 2D space, such as retinas. It is worth noting that both Wnt5a and Ryk receptor kinases appear to be expressed in the specific layers of the cortex, the olfactory bulb, and the retinas of the developing mouse brain [Allen Brain Atlas, http://www.brain-map.org].

Materials and methods

Drosophila genetics

Fly stocks and crosses were maintained on standard medium at 25°C unless otherwise stated. The following strains were used: ppk-GAL4, ppk-GAL4 (Kanamori et al. 2013), UAS-Wnt5, UAS-drl, UAS-draICD, UAS-Drl-2, UAS-trio, Wnt5D7 (null allele), DrlE124 [null allele], and trioE4 [strong hypomorphic allele]. UAS-Rac1-N17, UAS-Rho1-N19, UAS-Rho1-V14, dra-GAL4, ppk-CD4tdTomato, and UAS-mCD8GFP were obtained from the Bloomington Drosophila Stock Center and Kyoto Drosophila Genetic Resource Center.

To express the ricin proteins in v’ada neurons, we crossed hsFLP122, UFwTRA19/TM6B flies to yw, ppk-Gal4, UAS-mCD8GFP flies. The offspring between third instar larval stage and 24 h APF were heat-shocked for 15–30 min at 38°C and dissected at the 15- to 20-d adult stage.

To generate sterile clones, we crossed ppk-Gal4, UAS-mCD8GFP, hsFLP122, FRT 2A/TM6B flies to yw, f2E351 f2E351 FRT 2A/TM2 flies.

For the rescue experiments, we crossed ppk-Gal4, UAS-mCD8GFP, hsFLP122, FRT 2A/TM6B flies to yw, f2E351 f2E351 FRT 2A/TM2 flies.

Generation of transgenic flies

We generated the Wnt5-EGFP construct from CH322-141N12 [BACpac resources] in attB-P[acman]-ApR. CH322-141N12 contains the entire Wnt5 locus plus 5 kb of upstream sequence. We introduced an EGFP-stop/Kmr cassette after the ATG start codon of Wnt5 using the recombinating technique as described [Venken et al. 2006]. Potential recombinants were selected on chloramphenicol and kanamycin, and the correct targeting of the cassette was verified with PCR. The construct was then introduced into the VK00027 landing site on the third chromosome using ϕ-C31 integrase [BestGene, Inc.].

Immunohistochemistry

Immunostaining of Drosophila abdominal cuticles was performed as described [Yasunaga et al. 2010]. Briefly, abdominal cuticles were dissected and fixed in 4% parafomaldehyde in PBS for 30–60 min at room temperature followed by blocking for 60 min in PBS containing 0.3% Triton X-100 and 5% normal goat serum. Tissues were stained with mouse anti-GFP (1:1000), 3F6, Wako), rabbit anti-GFP (1:500, polyclonal, Medical and Biological Laboratories), and/or rat anti-CD8a (1:100, Caltag). Images were taken on a Leica TCS SP8 confocal microscope (Leica) and adjusted for brightness and contrast with Adobe Photoshop [Adobe Systems].

Quantitative analysis

For visualization of dendrites, we labeled v’ada neurons with UAS-mCD8GFP under the control of ppk-GAL4 and imaged GFP fluorescence in living animals by mounting them in silicon oil [Shin-Etsu] except for Figures 4F and 7C, where immunostaining or the ppk-CD4::tdTomato reporter was used. Maximum projections of Z-stacks were used in all cases. The dendrite length was measured by using the ImageJ plug-in Neuron [National Institutes of Health]. For the quantitative analyses, we focused on the v’ada dendrites in segments A3 and A4, since these neurons exhibit similar and consistent dendrite branch lengths and branch points. For quantification of the total branch length and the branch points, we measured the total branch length of the ventral half of the dendritic fields because the dorsal half partially overlaps with the dendritic fields of the other ppk-positive neurons in the same segment. Dendritic fields were defined by a polygon that connected the distal-most dendritic tips [Grueber et al. 2003]. For quantification of the total length of dendritic branches within single sternites, sternites were visualized by taking advantage of their autofluorescence in the green channel. The distance from the ventral midline to the ventral boundaries of C4da dendrites was measured as the distance between the midline and the dendritic terminals located most ventrally.

Statistical analysis

All statistical analysis was performed with Systat 13 (Hulinks, Inc.). For two-sample comparisons, unpaired Student’s t-tests were applied. For comparisons among more than two groups, one-way ANOVA tests were used and followed by Dunnett’s and Tukey’s tests, as indicated in the figure legends.

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References

Awasaki T, Saito M, Sone M, Suzuki E, Sakai R, Ito K, Hama C. 2000. The Drosophila Trio plays an essential role in patterning of axons by regulating their directional extension. *Neuron* 26: 119–131.

Bashaw GJ, Klein R. 2010. Signaling from axon guidance receptors. *Cold Spring Harb Perspect Biol* 2: a001941.

Bateman J, Van Vactor D. 2001. The Trio family of guanine-nucleotide-exchange factors: regulators of axon guidance. *J Cell Sci* 114: 1973–1980.

Bernards A. 2003. GAPs galore! A survey of putative Ras superfamily GTase activating proteins in man and *Drosophila*. *Biochim Biophys Acta* 1603: 47–82.

DeGeer J, Boudeau J, Schmidt S, Bedford F, Lamarche-Vane N, Debant A. 2013. Tyrosine phosphorylation of the Rho guanine nucleotide exchange factor Trio regulates netrin-1/DCC-mediated cortical axon growth. *Mol Cell Biol* 33: 739–751.

Dickson BJ. 2001. Rho GTPases in growth cone guidance. *Curr Opin Neurobiol* 11: 103–110.

Emoto K. 2011. Dendrite remodeling in development and disease. *Dev Growth Differ* 53: 277–286.

Emoto K. 2012. Signaling mechanisms that coordinate the development and maintenance of dendritic fields. *Curr Opin Neurobiol* 22: 805–811.

Emoto K, He Y, Ye B, Grueber WB, Adler PN, Jan LY, Jan YN. 2004. Control of dendritic branching and tiling by the Tricornered-kinase/Furry signaling pathway in Drosophila sensory neurons. *Cell* 119: 245–256.

Emoto K, Parrish IZ, Jan LY, Jan YN. 2006. The tumour suppressor Hippo acts with the NDR kinases in dendritic tiling and maintenance. *Nature* 443: 210–213.

Fradkin LG, Noordermeer JN, Nasu R. 1995. The Drosophila Wnt protein Dwnt-3 is a secreted glycoprotein localized on the axon tracts of the embryonic CNS. *Dev Biol* 168: 202–213.

Fradkin LG, Dura JM, Noordermeer JN. 2010. Ryks: new partners for Wnts on the developing and regenerating nervous system. *Trends Neurosci* 33: 84–92.

Fuest PG, Bruce F, Tian M, Wei W, Elstrott J, Feller MB, Erskine L, Singer JH, Burgess RW. 2009. DSCAM and DSCAML1 function in self-avoidance in multiple cell types in the developing mouse retina. *Neuron* 64: 484–497.

Furrer MP, Vassenkova I, Kamiyama D, Rosado Y, Chiba A. 2007. Slit and Robo control the development of dendrites in Drosophila CNS. *Development* 134: 3795–3804.

Gao FB, Kohwi M, Brennan JE, Jan LY, Jan YN. 2000. Control of dendritic field formation in *Drosophila*: the roles of flamingo and competition between homologous neurons. *Neuron* 28: 91–101.

Gibson DA, Tymansky S, Yuan RC, Leung HC, Lefebvre JL, Sanes JR, Chedotal A, Ma L. 2014. Dendrite self-avoidance requires autonomous Slit-Robo signaling in cerebellar Purkinje cells. *Neuron* 81: 1040–1056.

Grueber WB, Ye B, Moore A, Jan LY, Jan YN. 2003. Dendrites of distinct classes of Drosophila sensory neurons show different capacities for homotypic repulsion. *Curr Biol* 13: 616–626.

Hadjieconomous D, Rotkopf S, Alexandre C, Bell DM, Dickson BJ, Salecker I. 2011. Flybow: genetic multicolor cell labeling for neural circuit analysis in Drosophila melanogaster. *Nat Methods* 8: 260–266.

Hughes ME, Bortnick R, Tsubouchi A, Baumr P, Kondo M, Uemura T, Schmucker D. 2007. Homophilic Dscam interactions control complex dendrite morphogenesis. *Neuron* 54: 417–427. 

Hutchins BI, Li L, Kalil K. 2011. Wnt/calcium signaling mediates axon growth and guidance in the developing corpus callosum. *Dev Neurobiol* 71: 269–283.

Jan YN, Jan LY. 2010. Branching out: mechanisms of dendritic arborization. *Nat Rev Neurosci* 11: 316–328.

Kanamori T, Kanai M, Dairyo Y, Yasunaga K, Morikawa R, Emoto K. 2013. Compartmentalized calcium transients trigger dendrite pruning in *Drosophila* sensory neurons. *Science* 340: 1475–1478.

Kanamori T, Yoshino J, Yasunaga K, Dairyo Y, Emoto K. 2015. Local endocytosis triggers dendritic thinning and branching in *Drosophila* sensory neurons. *Nat Commun* 6: 6515.

Kaufmann WE, Moser HW. 2000. Dendritic anomalies in disorders associated with mental retardation. *Cereb Cortex* 10: 981–991.

Kikuchi A, Yamamoto H, Kishida S. 2007. Multiplicity of the interactions of Wnt proteins and their receptors. *Cell Signal* 19: 659–671.

Koike-Kumagai M, Yasunaga K, Morikawa R, Kanamori T, Emoto K. 2009. The target of rapamycin complex 2 controls dendritic tiling of *Drosophila* sensory neurons through the Tricornered kinase signaling pathway. *EMBO J* 28: 3879–3892.

Kopp A, Blackman RK, Duncan I. 1999. Wingless, Decapentaplegic and EGF Receptor signaling pathways interact to specify dorso-ventral pattern in the adult abdomen of *Drosophila*. *Development* 126: 3495–3507.

Kuo CT, Jan LY, Jan YN. 2005. Dendrite-specific remodeling of *Drosophila* sensory neurons requires matrix metalloproteases, ubiquitin–proteasome, and exdysone signaling. *Proc Natl Acad Sci* 102: 15230–15235.

Lee T, Luo L. 1999. Mosaic analysis with repressible cell marker for studies of gene function in neuronal morphogenesis. *Neuron* 22: 451–461.

Lee T, Winter C, Marticke SS, Lee A, Luo L. 2000. Essential roles of *Drosophila* RhoA in the regulation of neuroblast proliferation and dendritic but not axonal morphogenesis. *Neuron* 25: 307–316.

Lefebvre JL, Kostadinov D, Chen WY, Maniatis T, Sanes JR. 2012. Protocadherins mediate dendritic self-avoidance in the mammalian nervous system. *Nature* 488: 517–521.

Lin B, Wang SW, Masland RH. 2004. Retinal ganglion cell type, size, and spacing can be specified independent of homotypic dendritic contacts. *Neuron* 43: 475–485.

Liu Y, Shi J, Lu CC, Wang ZB, Lyuksyutova AI, Song XJ, Zou Y. 2005. Ryk-mediated Wnt repulsion regulates posterior-directed growth of corticospinal tract. *Nat Neurosci* 8: 1151–1159.

Luo L. 2000. Rho GTPases in neuronal morphogenesis. *Nat Rev Neurosci* 1: 173–180.

Masland RH. 2012. The neuronal organization of the retina. *Neuron* 76: 266–280.

Matsubara D, Horiuchi SY, Shimono K, Usui T, Uemura T. 2011. The seven-pass transmembrane cadherin Flamingo controls dendritic self-avoidance via its binding to a LIM domain protein, Espinas, in *Drosophila* sensory neurons. *Genes Dev* 25: 1982–1996.

Matsuzaka RL, Jiang Z, Samuels IS, Nguyen-Ba-Charvet KT, Sun LO, Peachey NS, Chedotal A, Yau KW, Kolodkin AL. 2012. Guidance-cue control of horizontal cell morphology, lamina- tion, and synapse formation in the mammalian outer retina. *J Neurosci* 32: 6859–6868.

Matthews BJ, Kim ME, Flanagan JJ, Hattori D, Clemens JC, Zipursky SL, Grueber WG. 2007. Dendrite self-avoidance is controlled by Dscam. *Cell* 129: 593–604.

Moreau-Fauvarque C, Talliebou E, Boissonneau E, Mesnard J, Dura JM. 1998. The receptor tyrosine kinase gene linotte is...
required for neuronal pathway selection in the *Drosophila* mushroom bodies. *Mech Dev* 78: 47–61.

Morikawa R, Kanamori T, Yasunaga K, Emoto K. 2011. Different levels of the TRIM protein Asap regulate distinct axonal projections of *Drosophila* sensory neurons. *Proc Natl Acad Sci* 108: 19389–19394.

Parrish JZ, Emoto K, Kim MD, Jan YN. 2007. Mechanisms that regulate establishment, maintenance, and remodeling of dendritic fields. *Annu Rev Neurosci* 30: 399–423.

Perry VH, Linden R. 1982. Evidence for dendritic competition in the developing retina. *Nature* 297: 683–685.

Polleux F, Morrow T, Ghosh A. 2000. Semaphorin 3A is a chemoattractant for cortical apical dendrites. *Nature* 404: 567–573.

Puram SV, Bonni A. 2013. Cell-intrinsic driver of dendrite morphogenesis. *Development* 140: 4657–4671.

Purpura DP. 1975. Dendritic differentiation in human cerebral cortex: normal and aberrant developmental patterns. *Adv Neuro* 12: 91–134.

Schmidt S, Debant A. 2014. Function and regulation of the Rho guanine nucleotide exchange factor Trio. *Small GTPases* 5: e29769.

Shimizu K, Sato M, Tabata T. 2011. The Wnt5/planar cell polarity pathway regulates axonal development of the *Drosophila* mushroom body neuron. *J Neurosci* 31: 4944–4954.

Shimono K, Fujimoto A, Tsuyama T, Yamamoto-Kochi M, Sato M, Hattori Y, Sugimura K, Usui T, Kimura K, Uemura T. 2009. Multidendritic sensory neurons in the adult *Drosophila* abdomen: origin, dendritic morphology, and segment- and age-dependent programmed cell death. *Neural Dev* 4: 37.

Shirras AD, Couso JP. 1996. Cell fates in the adult abdomen of *Drosophila* are determined by wingless during pupal development. *Dev Biol* 175: 24–36.

Smith HK, Roberts LL, Allen MJ, Connolly JB, Moffat KG, O’Kane CJ. 1996. Inducible ternary control of transgene expression and cell ablation in *Drosophila*. *Dev Genes Evol* 206: 14–24.

Smith CJ, Watson JD, VanHoven MK, Colon-Ramos DA, Miller DM III. 2012. Netrin (UNC-6) mediates dendritic self-avoidance. *Nat Neurosci* 15: 731–737.

Soba P, Zhu S, Emoto K, Younger S, Yang SJ, Yu HH, Lec T, Jan LY, Jan YN. 2007. *Drosophila* sensory neurons require Dscam for dendrite self-avoidance and proper dendritic organization. *Neuron* 54: 403–416.

Sonoshita M, Itatani Y, Kakizaki F, Sakimura K, Terashima T, Katsuyama Y, Sakai Y, Taketo MM. 2014. Promotion of colorectal cancer invasion and metastasis through activation of Notch–Dab1–Abl–RhoGEF protein TRIO. *Cancer Discov* 5: 198–211.

Srahna M, Leyssen M, Choi CM, Fradkin LG, Noordermeer JN, Hassan BA. 2006. A signaling network for patterning of neuronal connectivity in the *Drosophila* brain. *PLoS Biol* 4: e348.

Sun LO, Jang Z, Rivlin-Etzion M, Hand R, Brady CM, Matsuoka RL, Yau KW, Feller MB, Kolodkin AL. 2013. On and off retinal circuit assembly by divergent molecular mechanisms. *Science* 342: 1241974.

Venken KJ, He Y, Hoskins RA, Bellen HJ. 2006. P[acman]: a BAC transgenic platform for targeted insertion of large DNA fragments in *D. melanogaster*. *Science* 314: 1747–1751.

Wassle H, Boycott BR. 1991. Functional architecture of the mammalian retina. *Physiol Rev* 71: 447–480.

Wong RO, Ghosh A. 2002. Activity-dependent regulation of dendritic growth and patterning. *Nat Rev Neurosci* 3: 803–812.

Wouda RR, Bansraj MR, de Jong AW, Noordermeer JN, Fradkin LG. 2008. Src family kinases are required for WNT5 signaling through the Derailed/Ryk receptor in the *Drosophila* embryonic central nervous system. *Development* 135: 2277–2287.

Wu Y, Helt JC, Wexler E, Petrova IM, Noordermeer JN, Fradkin LG, Hing H. 2014. Wnt5 and dl/rk/ryk gradients pattern the *Drosophila* olfactory map. *J Neurosci* 34: 14961–14972.

Yasunaga K, Kanamori T, Morikawa R, Suzuki E, Emoto K. 2010. Dendrite reshaping of adult *Drosophila* sensory neurons requires matrix metalloproteinase-mediated modification of the basement membranes. *Dev Cell* 18: 621–632.

Yoshikawa S, McKinnon RD, Kokei M, Thomas JB. 2003. Wnt-mediated axon guidance via the Drosophila Derailed receptor. *Nature* 422: 583–588.

Zipursky SL, Grueber WB. 2013. The molecular basis of self-avoidance. *Annu Rev Neurosci* 36: 547–568.