#### **RESEARCH ARTICLE**



# βIII spectrin controls the planarity of Purkinje cell dendrites by modulating perpendicular axon-dendrite interactions

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#### ABSTRACT

The mechanism underlying the geometrical patterning of axon and dendrite wiring remains elusive, despite its crucial importance in the formation of functional neural circuits. The cerebellar Purkinje cell (PC) arborizes a typical planar dendrite, which forms an orthogonal network with granule cell (GC) axons. By using electrospun nanofiber substrates, we reproduce the perpendicular contacts between PC dendrites and GC axons in culture. In the model system, PC dendrites show a preference to grow perpendicularly to aligned GC axons, which presumably contribute to the planar dendrite arborization in vivo. We show that BIII spectrin, a causal protein for spinocerebellar ataxia type 5, is required for the biased growth of dendrites. BIII spectrin deficiency causes actin mislocalization and excessive microtubule invasion in dendritic protrusions, resulting in abnormally oriented branch formation. Furthermore, diseaseassociated mutations affect the ability of BIII spectrin to control dendrite orientation. These data indicate that ßIII spectrin organizes the mouse dendritic cytoskeleton and thereby regulates the oriented growth of dendrites with respect to the afferent axons.

KEY WORDS: Dendrite development, Mouse, Neuron, Purkinje, SCA5, Spectrin, Spinocerebellar ataxia

#### INTRODUCTION

Accurate information processing through the neuronal network requires proper development of dendrite patterns, which determine the receptive field, afferent types and the number of synapses they form (London and Häusser, 2005). During development, neurons arborize their dendrites by a set of branch dynamics, including extension, branching and retraction, which are controlled by either cell-intrinsic programs or extrinsic cues from the surrounding tissue (Dong et al., 2015; Jan and Jan, 2010; Valnegri et al., 2015).

Cerebellar Purkinje cells (PCs) develop highly branched dendrites in a single parasagittal plane. The dendrites are innervated by parallel fiber axons of cerebellar granule cells (GCs), which run perpendicularly across the aligned PC dendrites along the coronal axis of the cerebellum. The planar dendrites with a space-filling and non-overlapping arrangement are a distinctive feature of PCs, which is thought to be advantageous for efficient network formation. For instance, planar dendrites connected with perpendicularly oriented axonal bundles contribute to maximization of possible synaptic

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connections with minimal redundancy (Cuntz, 2012; Wen and Chklovskii, 2008). Indeed, each PC dendrite contacts more than 100,000 parallel fibers in rodents (Napper and Harvey, 1988). PCs arborize their dendritic branches via dynamic remodeling during postnatal development (Fujishima et al., 2018; Joo et al., 2014; Kaneko et al., 2011; Kapfhammer, 2004; Sotelo and Dusart, 2009; Takeo et al., 2015; Tanaka, 2009). We have previously identified the basic rules of dendrite formation in PCs grown on coverslips (Fujishima et al., 2012). Typical fan-shaped branching structures were constructed by constant extension and dichotomous terminal branching. Furthermore, we have shown that retraction and stalling induced by dendro-dendritic contact (self-avoidance) play pivotal roles in the non-overlapping and space-filling distribution of branches (Fujishima et al., 2012; Kawabata Galbraith et al., 2018). Indeed, it has been shown that loss of cell-surface molecules such as clustered protocadherins and slit/robo, which recognize the contact between dendritic branches, affects the non-overlapping arrangement of PC dendrites (Gibson et al., 2014; Ing-Esteves et al., 2018; Kuwako and Okano, 2018; Lefebvre et al., 2012; Toyoda et al., 2014). Although these basic rules of dendritic growth explain the dendritic configuration, including the non-overlapping pattern on a twodimensional (2D) substrate, they cannot illustrate how dendrites acquire the planar arrangement in three-dimensional tissue.

Unlike other planar dendrites, such as those of Drosophila dendritic arborization neurons that extend along the structural boundary (Han et al., 2012; Kim et al., 2012), PC dendrites grow in a plane with no obvious scaffolds in the cerebellum. Several lines of evidence suggest that the growth of PC dendrites is oriented in a direction perpendicular to the parallel fibers, which are formed earlier than the extension of PC dendrites (Altman and Anderson, 1972; Crepel et al., 1980; Nagata et al., 2006). The perpendicular growth seems to be involved in the flat dendrite arborization (Altman and Anderson, 1972; Crepel et al., 1980; Nagata et al., 2006; Sotelo and Dusart, 2009). Pioneering studies by Altman demonstrated that disruption of parallel fiber arrays via X-ray irradiation of the developing cerebellum led to the disruption and realignment of PC dendrites in a direction perpendicular to the misoriented parallel fibers (Altman, 1973). However, the molecular machinery underlying the dendrite growth perpendicular to the axonal arrays is largely unknown (Gao et al., 2011; Kim et al., 2014).

Spectrins are structural molecules that form a tetrameric complex with two  $\alpha$  and two  $\beta$  subunits. Spectrin tetramers interact with actin filaments beneath the plasma membrane. Recent studies using super-resolution microscopy have revealed that spectrin, actin and their associated proteins form membrane periodic skeletal (MPS) structures lining the circumference of the axons and dendrites (D'Este et al., 2015; Han et al., 2017; Vassilopoulos et al., 2019; Xu et al., 2013). In the MPS structures, ring-like structures of actin are connected by spectrin tetramers with a periodicity of ~190 nm. Alternatively, spectrin and associated proteins can form a 2D polygonal lattice structure in the soma or dendrites, reminiscent of

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those found in erythrocytes (Han et al., 2017). These structures are considered to have multiple functions, such as the maintenance of membrane stiffness and/or elasticity (Krieg et al., 2017) and acting as diffusion barriers to membrane molecules (Albrecht et al., 2016).

 $\beta$ III spectrin, one of the spectrin  $\beta$  subunits, is highly expressed in the PC dendrites and soma. BIII spectrin (Sptbn2) has been identified as a gene responsible for spinocerebellar ataxia type 5 (SCA5), which manifests as a progressive dysfunction of motor coordination (Ikeda et al., 2006). PCs in SCA5 patients and BIII spectrin knockout animals display aberrant dendrite morphologies owing to abnormal arbor development and maintenance (Gao et al., 2011; Ikeda et al., 2006). Notably, PCs in the knockout mouse exhibit a defect in flat dendrite arborization (Gao et al., 2011), implicating BIII spectrin in the regulation of the orientation of dendrite growth. The loss of  $\beta$ III spectrin leads to the mislocalization and/or diminished level of neurotransmitter receptors and transporters, which might cause the progressive neurodegeneration via excitotoxicity (Armbrust et al., 2014; Perkins et al., 2010; Stankewich et al., 2010). However, the mechanisms by which BIII spectrin controls dendrite development remain elusive.

Here, we show that  $\beta$ III spectrin is indispensable for the directional PC dendrite arborization perpendicular to parallel fibers, which is a prerequisite for the flat dendrite formation. We recapitulated a 2D model of the orthogonal interaction between the dendrite and axonal bundles by using nanofibers as artificial scaffolds to reproduce parallel fiber arrays *in vitro*. We demonstrated that  $\beta$ III spectrin-deficient PCs failed to form dendrites perpendicular to axonal bundles. Super-resolution observations confirmed the repeated lattice-like  $\beta$ III spectrin structures in PC dendrites. Loss of  $\beta$ III spectrin caused abnormal cytoskeletal dynamics and misoriented dendrite growth. Our data provide insights into the role of  $\beta$ III spectrin in controlling the perpendicular connectivity of dendrites and axons to form planar dendrites.

#### RESULTS

# $\beta$ III spectrin is required for a cell-autonomous mechanism of planar dendrite arborization in PCs

Earlier studies have demonstrated that the planarity of PC dendrites is disturbed in loss-of-function mutants of BIII spectrin (Gao et al., 2011). We first confirmed the cell-autonomous role of  $\beta$ III spectrin in PC dendrite formation by short hairpin RNA (shRNA) knockdown in the wild-type background in vivo. We used a plasmid encoding an shRNA targeting BIII spectrin that efficiently knocks down βIII spectrin expression in PCs in a dissociation culture (Fig. S1). Dendrites of BIII spectrin-knockdown cells exhibited the characteristic tapering shape seen in knockout cells (Gao et al., 2011). Besides, there was no notable difference in dendritic shape between control (ctr) and knockdown cells. We delivered the plasmid sparsely into PC precursors via in utero electroporation at embryonic day (E) 11.5. Consistent with previous observations using BIII spectrin knockout mice, we observed almost no differences in the total length of dendrites between control and knockdown cells at postnatal day (P) 14 (Fig. 1A-C). Compared with the normal planar dendrites in control, knockdown PCs exhibited misoriented dendrites that extended away from the main dendritic plane (Fig. 1A,B), similar to those in knockout animals (Gao et al., 2011). Quantitative analysis revealed that the number of dendritic branches extruded away from the main plane was significantly increased in knockdown PCs (Fig. 1A,B,D; Fig. S2; see Materials and Methods), suggesting that a cell-autonomous function of BIII spectrin is necessary for the planar dendrite formation.

# Recapitulation of 2D perpendicular contact between PC dendrites and GC axons using artificial nanofibers *in vitro*

Next, we examined the mechanism underlying the effect of  $\beta$ III spectrin deficiency on dendrite planarity. Considering that flat dendrites are closely aligned with neighboring PCs (Fig. 2A), it seems unlikely that gradients of guidance molecules instruct such dendritic configurations. It has previously been suggested that PC dendrites grow perpendicularly to GC axons (Nagata et al., 2006). If so, dendrites will extend in parasagittal planes perpendicular to the bundle of parallel fibers (GC axons) that run along the coronal axis (Fig. 2B, left). We thus speculated that  $\beta$ III spectrin might be involved in dendrite growth perpendicular to GC axons.

To test this hypothesis, we first confirmed whether normal PC dendrites preferentially grow perpendicularly to GC axons. We established a simplified model in which parallel GC axons are recapitulated in 2D spaces (Fig. 2B, right). We used electrospun polycaprolactone nanofibers, which have been used to navigate axonal extensions in culture (Hyysalo et al., 2017). We plated dissociated cells from P0 cerebella on aligned or randomly oriented nanofibers. GCs were labeled with green fluorescent protein (GFP) and identified by the expression of the GC marker Pax6 (Engelkamp et al., 1999). As expected, GC axons ran parallel to each other on aligned nanofibers (arrowheads in Fig. 2C), whereas they extended randomly on non-oriented nanofibers (arrowheads in Fig. 2E).

We previously reported that PCs in dissociated cultures initiate dendrites at ~8 days in vitro (DIV) and extend branches radially until at least 14 DIV (Fujishima et al., 2012). Likewise, PC dendrites on the randomly oriented nanofibers were observed to grow radially, with no directional preference (Fig. 2F). By contrast, the distribution of PC dendrites on aligned nanofibers was highly biased toward the direction perpendicular to fibers, supporting the notion that PC dendrites grow perpendicularly to GC axons (Fig. 2D). Indeed, quantitative analysis revealed that the majority of dendritic segments oriented perpendicular to aligned fibers (polar histogram: dendrite orientation in Fig. 2D; see Materials and Methods; Fig. S3). The perpendicular growth was specific to PC dendrites, because PC axons grew parallel to GC axons (Fig. 2D). Furthermore, GC dendrites grew in a random orientation on aligned fibers (Fig. 2C). These results suggest that the perpendicular interaction is not ubiquitous but is specific to certain synaptic partners, including PC dendrites and GC axons.

To verify that PC dendrites are navigated by GC axons but not by nanofibers, we confirmed the interactions between PC dendrites and GC axons without nanofibers (Fig. S4A). We prepared microexplants of the external granule layer populated with differentiating GCs from the P2 cerebellum (Kawaji et al., 2004; Nagata and Nakatsuji, 1990). Isolated PCs were co-cultured at 1 DIV on the GC axons that extended radially from the explant. We confirmed that PCs developed dendrites perpendicular to the radially extended GC axons at 10 DIV (Fig. S4A), consistent with a previous report (Nagata et al., 2006). Surface rendering showed close apposition of the dendrites and axons (Fig. S4B). These results suggest that direct PC dendrite-GC axon interaction determines the dendrite orientation.

To exclude the possibility that the nanofibers act as scaffolds for PC dendrite growth, we prepared PC cultures with GCs of different densities on nanofibers (Fig. S5). Although PCs established perpendicular dendrites in high- and medium-density cultures, they could not grow dendrites on free nanofibers devoid of GC axons in a low-density culture, suggesting that nanofibers do not serve as growth scaffolds (Fig. S5A,B). Taken together, we conclude that PC dendrites grow perpendicular to GC axons.

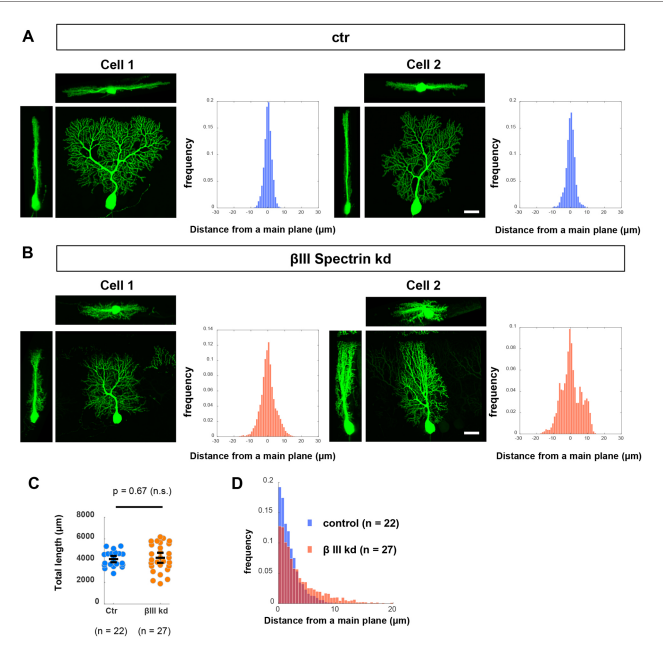
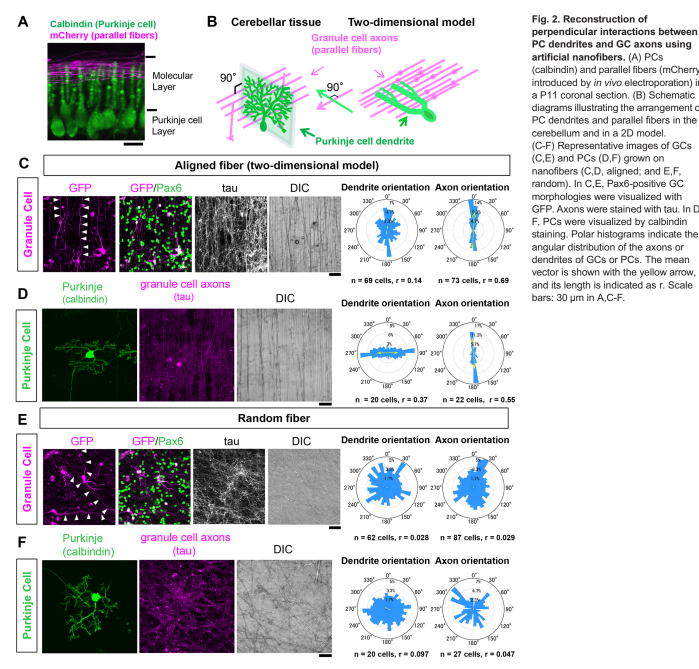


Fig. 1. A cell-autonomous role of  $\beta$ III spectrin in the planar dendrite formation *in vivo*. (A,B) Morphology of PCs expressing the GFP/shRNA-control (A) or GFP/shRNA- $\beta$ III spectrin plasmid (B) in P14 sagittal sections. Top, front (sagittal) and side (coronal) views were taken from the three-dimensionally reconstructed images. Histograms show the proportions of dendritic segments locating at the indicated distances from the main dendritic plane. (C) Quantification of total dendritic length at P14. Statistical analysis: Welch's *t*-test. (D) Proportions of dendritic segments locating at the indicated distances from the main dendritic plane. *P*=2.7×10<sup>-46</sup>, two-sample Kolmogorov–Smirnov test. Scale bars: 20 µm in A,B.

# Knockdown of $\beta III$ spectrin interferes with directed dendrite growth perpendicular to GC axons

To examine whether  $\beta$ III spectrin is required for the perpendicular interaction between PC dendrites and GC axons, we introduced shRNA plasmids to PCs and plated them on nanofibers. In contrast to the dendrites perpendicular to GC axons observed in control cells, a significant proportion of the branches in  $\beta$ III spectrin-knockdown dendrites extended in different orientations (Fig. 3A). Quantitative morphometry revealed that the total length and number of branches were increased in knockdown cells, suggesting excess misoriented branch formation (Fig. 3B,C). This is inconsistent with the observations *in vivo*, where the total length was not affected by the loss of  $\beta$ III spectrin (Fig. 1). We attribute this discrepancy to a homeostatic mechanism of dendritic growth, which adjusts the size of PC dendrites to match the amount of available neurotrophic factors from afferent axons *in vivo* (Joo et al., 2014). Co-expression of shRNAresistant  $\beta$ III spectrin reversed the changes in dendrite orientation, total length and number of branches, negating the off-target effects (Fig. 3A-C,  $\beta$ III kd+rescue). Furthermore, the CRISPR/Cas9-based  $\beta$ III spectrin knockout dendrites exhibited similar misoriented phenotypes (Fig. S6). These results indicate that  $\beta$ III spectrin is required for directed dendrite formation perpendicular to GC axons.

Spectrin molecules form tetrameric complexes composed of  $\alpha$  and  $\beta$  subunits, and the loss of  $\alpha$  subunits destabilizes  $\beta$  subunits in embryonic tissue (Stankewich et al., 2011). Among  $\alpha$ -spectrin subtypes,  $\alpha$ II spectrin is most abundant in non-erythrocytic cells (Cianci et al., 1999; Winkelmann and Forget, 1993). Indeed,  $\alpha$ II spectrin was strongly expressed in PC somata and dendrites, similar



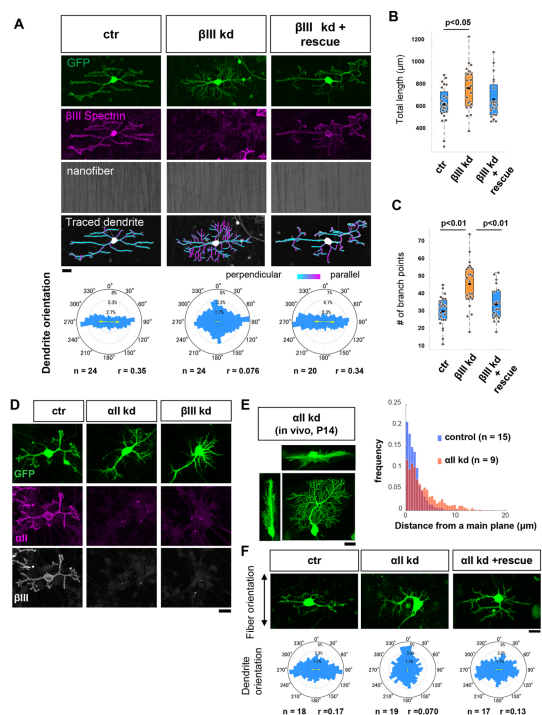
(calbindin) and parallel fibers (mCherry, introduced by in vivo electroporation) in a P11 coronal section. (B) Schematic diagrams illustrating the arrangement of PC dendrites and parallel fibers in the cerebellum and in a 2D model. (C-F) Representative images of GCs (C,E) and PCs (D,F) grown on nanofibers (C,D, aligned; and E,F, random). In C,E, Pax6-positive GC morphologies were visualized with GFP. Axons were stained with tau. In D. F, PCs were visualized by calbindin staining. Polar histograms indicate the angular distribution of the axons or dendrites of GCs or PCs. The mean vector is shown with the yellow arrow, and its length is indicated as r. Scale

to  $\beta$ III spectrin (Fig. 3D). We found that the knockdown of either  $\alpha$ II or BIII spectrin dramatically reduced both proteins in PCs, suggesting the interdependency of the molecules in stabilization of the spectrin complex (Fig. 3D). Accordingly, all spectrin knockdown disrupted planar arborization in vivo (Fig. 3E) in addition to biased dendrite organization on aligned nanofibers (Fig. 3F). Thus, these results indicate that the  $\alpha$ II/ $\beta$ III spectrin complex regulates PC dendrite orientation.

#### Loss of **BIII** spectrin causes aberrant branch formation

We have previously demonstrated that the non-overlapping arrangement of PC dendrites is achieved by contact-dependent branch retraction (Fujishima et al., 2012). Thus, the perpendicular dendrites on nanofibers might be attributed either to biased outgrowth (extension and branch formation) in the perpendicular direction or to the biased retraction of misoriented branches. To

distinguish between these possibilities, we performed time-lapse observations of growing dendrites on aligned nanofibers. We traced the outgrowth of PC dendrites ( $\sim 0.7 \,\mu$ m/h) every 3 h for several days from 8 DIV, when dendrite formation was initiated in the culture (Fujishima et al., 2012). Control PCs continuously grew their dendrites by extension and branch formation perpendicular to the GC axon orientation (Fig. 4A,B). Next, we measured the orientation of retracted branches. Dendritic retractions were triggered by collisions of growing dendritic tips, consistent with our previous report (Fujishima et al., 2012). Of all dendritic retractions observed in control PCs, ~86% were induced after obvious dendritic collisions. In addition,  $\sim$ 73% of branches were eliminated after a collision (Fig. 4D). Disorienting branches were eliminated with high probability because they had more chance of contact with other branches, but perpendicular branches were also retracted when they collided with other growing branches (Fig. 4C).

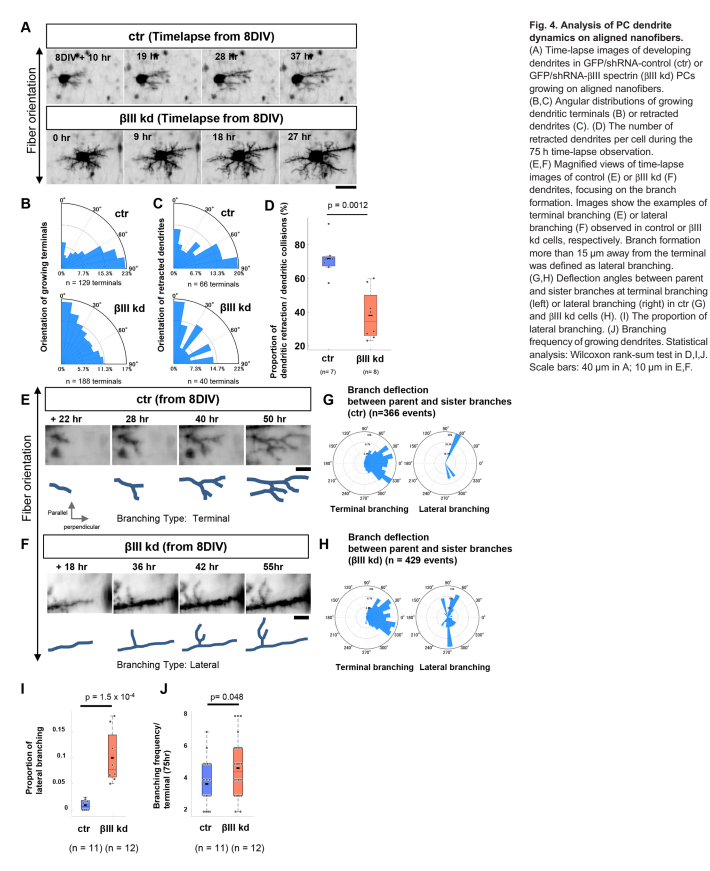


n = 18 r =0.17

Fig. 3. Disruption of perpendicular dendrite growth via the knockdown of ßIII spectrin. (A) Morphology of PCs transfected with GFP/shRNA control (ctr), GFP/shRNA-ßIII spectrin (ßIII kd) and GFP/shRNA-ßIII spectrin with shRNA-resistant ßIII spectrin (ßIII kd+rescue) grown on aligned fibers (12 DIV). Traced dendrites are pseudocolored based on the angles between the dendritic segments and fibers. Polar histograms indicate angular distributions of dendritic segments. P=6×10<sup>-16</sup> for ctr versus βIII kd, P>0.5 for ctr versus rescue, and P=6×10<sup>-16</sup> for βIII kd versus rescue (Watson–Wheeler test with Bonferroni correction). (B,C) Quantification of total dendritic length (B) and number of branches (C). Statistical analysis: ANOVA followed by the Tukey-Kramer post hoc test. (D) all and bill spectrin signals in PCs transfected with GFP/shRNA-control (ctr), GFP/shRNA-all spectrin (all kd) and GFP/shRNA-bill (bill kd). (E) Morphology of PCs transfected with GFP/shRNA-all spectrin (all kd) in P14 sagittal section. The histogram shows the proportions of dendritic segments locating at the indicated distance from the main dendritic plane. P=2.2×10<sup>-39</sup>, two-sample Kolmogorov–Smirnov test. (F) Top panels, morphology of PCs transfected with GFP/ shRNA-control (ctr), GFP/shRNA-all spectrin (all kd) and GFP/shRNA-all spectrin plus shRNA-resistant all spectrin (all kd+rescue) grown on aligned nanofibers. Bottom panels, angular distribution of dendritic segments. Scale bars: 20 µm in A,D-F.

Therefore, we concluded that the perpendicular dendrite orientation in control PCs was likely to be attributable to biased dendritic growth, although the elimination of disoriented branches might contribute to some extent.

In BIII spectrin-knockdown PCs, the fraction of parallel-growing dendritic terminals was markedly increased (Fig. 4A,B). Similar to control PCs, most of the dendritic retractions (~90%) in knockdown cells occurred after the dendritic collisions. However, the frequency



of dendrite retraction after a collision was reduced to 39%, suggesting that contact-dependent dendritic retraction is dysregulated in knockdown cells (Fig. 4D). However, the orientation of retracted dendrites was strongly biased parallel to axons, refuting that the

increase in misoriented dendrites in knockdown cells was caused by the suppression of retraction of errant arbors. Therefore, we focused on how  $\beta$ III spectrin deficiency leads to the dysregulation of growth orientation. DEVELOPMENT

Our previous studies have shown that PCs form dendritic branches primarily via the bifurcation of growing terminals (terminal branching), whereas they rarely extend collaterals from the shaft (lateral branching) (Fujishima et al., 2012). Accordingly, control dendrites on aligned fibers mainly displayed terminal branching (Fig. 4E,I). By contrast, lateral branching was increased by more than 10-fold in  $\beta$ III spectrin-knockdown dendrites (Fig. 4F,I), although the branching frequency was only slightly altered (Fig. 4J). The deflection angle between the bifurcated terminal branches ranged within approximately  $\pm 30^{\circ}$ , whereas that of lateral branching was greater than  $60^{\circ}$  in both the control and knockdown dendrites (Fig. 4G,H). These results suggest that  $\beta$ III spectrin regulates the perpendicular dendrite growth by inhibiting lateral branching. Thus, the frequent lateral branching might contribute to the increase in misoriented dendrites in the  $\beta$ III spectrin-deficient cells.

# Lateral branch formation in $\beta III$ spectrin-knockdown dendrites

We next observed how dendritic planarity was affected in BIII spectrin-knockdown cells in vivo. Control dendrites transfected with GFP aligned in a parasagittal plane in the molecular layer parallel to the neighboring PC dendrites (Fig. 5A,B). These PCs rarely exhibited dendritic branches growing in lateral (coronal) directions. Neighboring dendrites were separated by gaps of  $\sim$ 1-3 µm, showing minimal crossing with adjacent branches. Similar to control PCs, the main dendritic arbors of knockdown PCs were mostly parallel to neighboring dendrites in coronal sections (Fig. 5A,B, BIII kd), although some dendrites bent or tilted into incorrect planes (white arrows in Fig. 5A). Notably, knockdown dendrites exhibited an increased number of laterally oriented branches growing into the territories of the neighboring PC dendrites (yellow arrows in Fig. 5A,C). These misoriented branches often turned and extended parasagittally into the gaps between PC dendrites (yellow arrowhead in Fig. 5A). These misoriented lateral branches are likely to contribute to the disruption of planar dendrites in BIII spectrin-deficient PCs.

It has been demonstrated that the growing PC dendrites are covered with numerous dendritic protrusions, including dendritic filopodia and immature spines (Kawabata Galbraith et al., 2018; Shimada et al., 1998). Given that dendritic protrusions are known to serve as branch precursors in some neurons, we next analyzed the protrusions in PCs with or without BIII spectrin expression. Control dendrites presented numerous protrusions emanating from the shaft, with a mean length of  $1.48\pm0.04 \,\mu\text{m}$  (mean $\pm$ s.e.m., n=231) (Fig. 5D,E). By contrast, BIII spectrin-knockdown dendrites exhibited significantly longer protrusions (2.21±0.17  $\mu$ m, *n*=102) at a lower density, in agreement with previous studies (Fig. 5D-F) (Efimova et al., 2017; Gao et al., 2011). Notably, some protrusions in βIII spectrin-knockdown cells were abnormally elongated ( $>5 \mu m$ ) in lateral directions away from the main sagittal plane of the dendritic shaft (arrows in Fig. 5D,G). These long lateral protrusions seemed to serve as precursors of the ectopic lateral branches in BIII spectrin-deficient cells.

# Abnormal formation of dendritic protrusions in βIII spectrin-knockdown dendrites

To analyze the implications of  $\beta$ III spectrin in the formation of dendritic protrusions and branches, we observed the dendritic structures in neurons grown on aligned nanofibers. Control PCs bore highly dense protrusions that covered the lateral surface of the dendritic shaft, similar to those observed *in vivo* (Fig. 6A). These protrusions expressed glutamate receptor  $\delta$ 2 (GluD2), which functions as a synaptic glue by binding with presynaptic

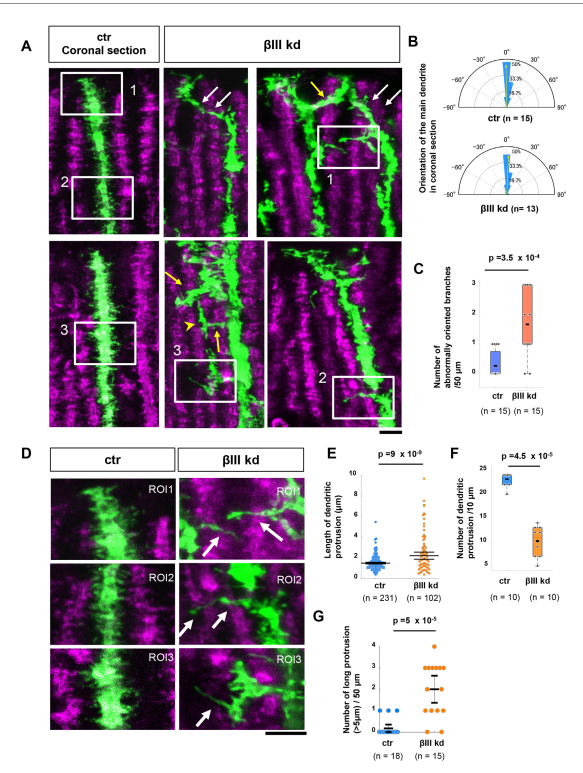
neurexin/Cbln1 in GC axonal terminals (Matsuda et al., 2010; Uemura et al., 2010), suggesting that these protrusions are dendritic spines or spine precursors (Fig. 6A).

Compared with the control PCs, the distal dendrites of BIII spectrin-knockdown cells were significantly thinner (ctr, 1.80± 0.15  $\mu$ m, *n*=19;  $\beta$ III kd, 0.90 $\pm$ 0.06  $\mu$ m, *n*=22, mean $\pm$ s.e.m.,  $P=7\times10^{-6}$ , Student's *t*-test) (Fig. 6B). In contrast to the dendritic protrusions in control cells, which presented a relatively constant length of 1-3 µm, those in knockdown cells presented various lengths at a lower density, with an average length that was significantly longer than that in control cells (Fig. 6C,D). We found some abnormally elongated protrusions of nearly 10 µm in knockdown dendrites that extended parallel to the orientation of GC axons. The extremely long protrusions exhibited multiple GluD2 puncta that were irregularly arranged along their lengths [Fig. 6B, region of interest (ROI)2]. The orientation of the protrusions in the control and knockdown dendrites was mostly parallel to GC axons (Fig. 6E). Other short protrusions often appeared wider at their bases, reminiscent of the shaft synapses with deficient neck formation observed in BIII spectrin-deficient hippocampal neurons (Fig. 6B, ROI1) (Efimova et al., 2017). We also observed ectopic GluD2 puncta on the dendritic shaft. These results suggest that βIII spectrin is required for shaping dendritic protrusions and that its loss results in the conversion of those protrusions to either spineless shaft synapses or abnormally oriented branches extending along the GC axons (Fig. 6F).

We next analyzed the subcellular localization of  $\beta$ III spectrin in growing dendrites in PCs cultured on coverslips. Consistent with previous reports (Efimova et al., 2017; Gao et al., 2011),  $\beta$ III spectrin was strongly localized on the surface of the dendritic shaft and the protrusion base, whereas it was excluded from the protrusion tips (Fig. 6G,H). Actin showed an inverse gradient along the protrusions such that it was densely localized at the tip and sharply declined in the base of the protrusion and the shaft (Fig. 6H,I). By contrast, in  $\beta$ III spectrin-knockdown PCs, actin was more widely distributed along the entire length of both long and thin (Fig. 6J, type 1) and short and stubby (Fig. 6J, type 2) protrusions and was often dispersed in the shaft of distal dendrites (Fig. 6J, type 1; Fig. 6K). These results imply that  $\beta$ III spectrin might be involved in the formation of the structural boundary between the dendritic shaft and protrusions that confines actin filaments within protrusions.

#### Membrane periodic skeletal structure formed by $\beta III$ spectrin

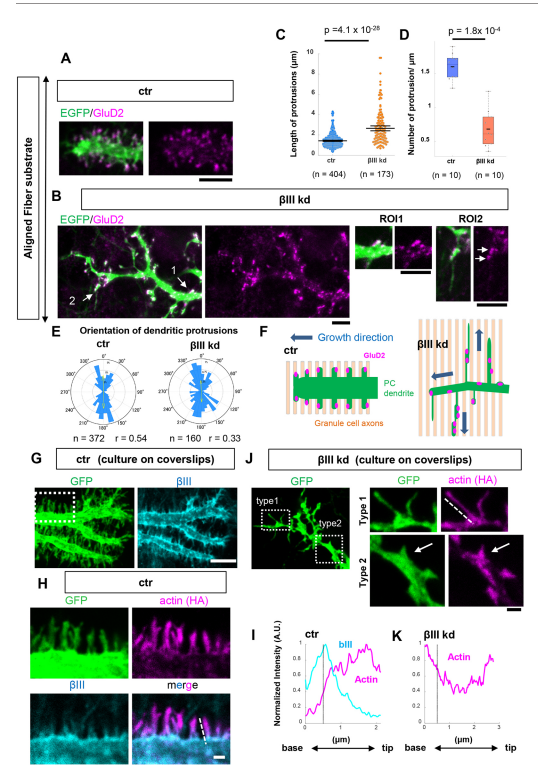
Super-resolution microscopy has revealed that the spectrin-actin complex forms MPS structures, which might function as a barrier to diffusion for membrane proteins (Albrecht et al., 2016; Leite et al., 2016). Using stimulated emission depletion microscopy (STED), we confirmed the existence of MPS-like repeated structures composed of BIII spectrin in the shaft of developing PC dendrites (Fig. S7A,C, ROI1). The average interval of the repeated structures was  $186\pm5 \,\mu\text{m}$  (Fig. S7D), consistent with previous studies (Xu et al., 2013). Actin rings were not clearly observed in the structures, probably owing to very low actin signals in the shafts compared with the dendritic filopodia (data not shown; Fig. 6H). Repeated BIII spectrin signals were also observed along the narrow corridor of the protrusion neck (Fig. S7B,C, ROI2 and ROI3) with an interval of  $187\pm5 \,\mu m$  (Fig. S7D). The repeated structures were not continuous but were often interrupted by irregularly arranged subsets, consistent with previous observations demonstrating the lower propensity for repeated structure formation in dendrites (Fig. S7E,F) (D'Este et al., 2015; Han et al., 2017). Thus, BIII spectrin formed random meshwork or repeated structures in dendritic shafts and filopodial bases.



**Fig. 5.** βIII spectrin knockdown induces the extension of laterally oriented branches. (A) Growing dendritic arbors in PCs transfected with GFP/shRNA-control (ctr) or GFP/shRNA-βIII spectrin (βIII kd) in the vermis region in P14 coronal slices. βIII spectrin signals are shown in magenta. (B) Angular distribution of the main dendritic arbor. (C) The number of laterally oriented branches extruded from the main dendrites. (D) Images are magnified views of the insets in A. (E) Length of dendritic protrusions. (F) Number of dendritic protrusions per 10 µm of the dendritic segment. (G) The number of abnormally long (more than 5 µm) dendritic protrusions per 50 µm of the dendritic segment. Statistical analyses: Wilcoxon rank-sum test in C; Student's *t*-test in E-G. Scale bars: 5 µm in A,D.

# $\beta III$ spectrin suppresses microtubule entry into dendritic protrusions

It has previously been demonstrated using young hippocampal neurons that the transition from filopodia to neurites is triggered by microtubule invasion into the filopodia after local actin remodeling (Dent et al., 2007; Flynn et al., 2012). To test whether  $\beta$ III spectrin contributes to blocking microtubule entry into dendritic protrusions, we monitored microtubule polymerization in dendrites by transfecting end-binding protein 3 fused with enhanced green fluorescent protein (EB3-EGFP), a plus-end marker of dynamic microtubules. We analyzed the growing



# Fig. 6. Abnormal extension of dendritic protrusions is induced by the loss of $\beta$ III spectrin.

(A,B) Representative images of the growing dendritic terminals of PCs transfected with GFP/shRNA-control (A) or GFP/shRNA-βIII spectrin (B). Cells were cultured on aligned nanofibers and stained with postsynaptic GluD2 (magenta) at 10 DIV. ROI1 and ROI2 are magnified views of the regions indicated by arrows 1 and 2 in the left image. (C) Length of dendritic protrusions. (D) Number of dendritic protrusions per micrometer. Statistical analysis: Wilcoxon rank-sum tests. (E) Orientation of dendritic protrusions. (F) Schematic explanation of growing dendrites (green) in control or BIII kd PCs on aligned GC axons (light orange). (G) Localization of βIII spectrin (cyan) in the growing dendrite in a control PC grown on a coverslip. (H) Magnified views of actin and BIII spectrin signals in the dendritic protrusions in the boxed region of G. HA-actin was transfected to visualize actin signals. (I) Intensity profiles of βIII spectrin and actin along the dotted line in H. The black dotted line in the graph indicates the boundary between the protrusion and shaft. (J) Left: representative image of the growing dendrites of GFP/BIII spectrin kd PCs grown on coverslips. Right: magnified views of actin signals in the dendritic protrusions (type 1, thin and long; type 2, short and stubby) in βIII kd cells. (K) Intensity profiles of actin along the dotted line in J. Scale bars: 5 µm in A,B,G,J (left panel); 1 µm in H,J (right panels).

dendritic terminals (within 10  $\mu$ m from the terminal) and more proximal dendrites (more than 10  $\mu$ m away from the terminal) separately to examine the regional difference in EB3 dynamics.

In control PCs,  $48\pm5\%$  (mean±s.e.m., 226 protrusions from 18 dendrites) of dendritic protrusions around the growing terminal were targeted by EB3-EGFP within 150 s of observation (Fig. 7A,C). By contrast, a significantly lower proportion of protrusions were invaded by EB3 in proximal dendrites (5±2%, 149 protrusions from 11 dendrites), in line with the notion that microtubule entry into filopodia triggers neurite extension at dendritic tips.

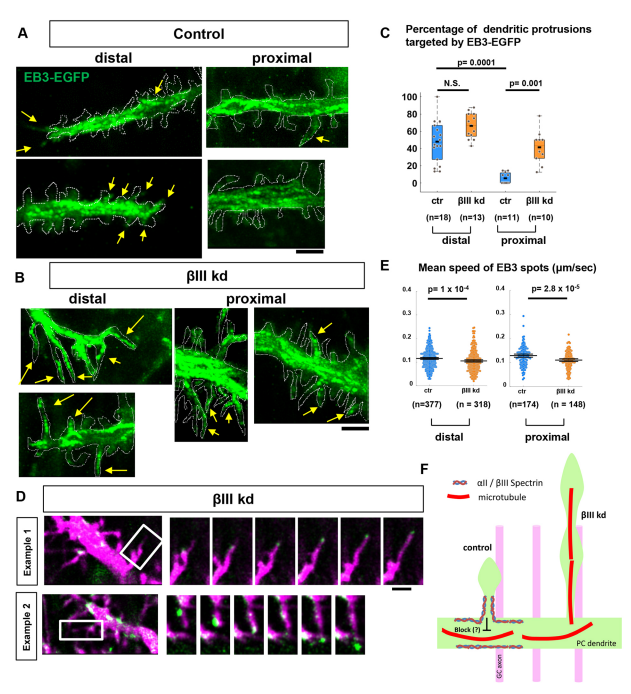
We found that  $\beta$ III spectrin knockdown significantly increased the proportion of EB3-targeted protrusions in proximal regions (41±6%, 91 protrusions from 10 dendrites) (Fig. 7B,C), whereas only a slight increase was observed in the distal area (67±4%, 102 protrusions from 13 dendrites). These data support the idea that  $\beta$ III spectrin interferes with microtubule invasion into protrusions in proximal dendrites. We often observed that the dendritic protrusions targeted by EB3 puncta extended aberrantly (Fig. 7D). Motility of those protrusions was immediately suppressed by treatment with the microtubule depolymerizer colchicine (Fig. S8), supporting that

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**Fig. 7. The invasion of EB3 into dendritic protrusions is facilitated by the loss of βIII spectrin.** (A,B) Time-lapse imaging of EB3-EGFP in the dendrites of PCs transfected with mCherry/shRNA-control (A) or mCherry/shRNA-βIII spectrin (B). Images are the maximum projections of EB3-EGFP spots during 150 s of observation. Distal (left) and proximal (right) dendritic regions were recorded every 3 s. Dotted lines indicate the contour of dendrites (mCherry). EB3-EGFP signals in dendritic protrusions are indicated by arrows. (C) The proportion of filopodia targeted by EB3-EGFP within 150 s of observation. (D) Example of filopodial extension associated with EB3-EGFP signals at the proximal dendrite region in βIII kd cells. Right panels show time-series images (12 s intervals) of boxed regions in the left images. (E) Speed of EB3 spots traveling within PC dendrites. (F) Schematic hypothesis of the function of βIII spectrin in the regulation of microtubule dynamics. Statistical analyses: Steel–Dwass test in C; Wilcoxon rank-sum test in E. Scale bars: 3 µm in A,B; 2 µm in D.

microtubule entry drives aberrant elongation of protrusions in knockdown cells. The speed of the EB3 movement was instead slightly downregulated in knockdown cells, refuting that excessive microtubule entry in knockdown cells was caused by increased microtubule polymerization activity (Fig. 7E). These results suggest that  $\beta$ III spectrin controls directed dendritic arborization by suppressing microtubule invasion and ectopic branch formation from proximal dendritic protrusions (Fig. 7F).

#### **Mutations causing SCA5**

Mutations in  $\beta$ III spectrin are known to cause SCA5. We wondered whether the disease mutations affect biased dendrite growth perpendicular to GC axons. We focused on three mutations identified in earlier studies (Ikeda et al., 2006): L253P (a leucine to proline substitution in the calponin homology domain found in a German family); E532-M544 del (an in-frame 13 amino acid deletion in the third spectrin repeat found in an American family); and L629-R634 delinsW (a five amino acid deletion with tryptophan insertion in the third spectrin repeat found in a French family) (Fig. 8A). The amino acid sequences related to these mutations are conserved among humans and mice. Thus, we generated mouse  $\beta$ III spectrin mutant constructs harboring the corresponding mutations. These constructs were designed to be shRNA resistant and were tagged with a myc epitope at their N-termini for imaging.

Dissociated PCs on aligned nanofibers were knocked down for endogenous  $\beta$ III spectrin and concomitantly transfected with wildtype  $\beta$ III spectrin or disease-related mutants. Wild-type  $\beta$ III spectrin was distributed in the somatodendritic area up to the most distal dendritic regions. In sharp contrast, the L253P form was localized in intracellular vesicular structures in the somatic area, whereas almost no signal was observed in dendrites (Fig. 8B). On the contrary, mutants with deletions in the third spectrin repeat (E532-M544 del and L629-R634 delinsW) localized to the dendritic plasma membrane to a lesser extent than the wild type. Quantitative image analysis revealed the differential localization of mutants in PC dendrites (Fig. 8C).

PCs expressing the shRNA-resistant wild-type molecule exhibited normal perpendicular dendrites. By contrast, all disease mutants were defective in the regulation of perpendicular dendrite formation. L253P and E532-M544 del were completely incompetent in perpendicular guidance, whereas the L629-R634 delinsW, which showed modest mislocalization, retained weak but significant guidance activity (Fig. 8D). Given that the disease mutants examined are associated with dominant forms of SCA5, we next asked whether the disease mutants act dominantly in the wild-type background without the knockdown plasmid. Although the effects were weak, these disease mutants affected the dendrite orientation (Fig. S9), supporting that these mutants mimic disease phenotypes.

To confirm the effect of the L253P and E532-M544 del mutations in the planar dendrite arborization *in vivo*, we delivered βIII spectrinknockdown plasmid and shRNA resistant βIII spectrin wild type, L253P or E532-M544 del to immature PCs by *in utero* electroporation. Consistent with *in vitro* observation, L253P and E532-M544 del exhibited abnormal localization in vesicular structures in the soma and proximal dendritic surface, respectively, in contrast to the wild type spreading over the entire dendritic surface (Fig. 8E). PCs expressing the wild type showed planar dendrites, whereas cells expressing either L253P or E532-M544 del displayed disorganized dendrites growing away from the main dendritic plane (Fig. 8E,F). These results suggest that these disease mutations of βIII spectrin disrupt dendritic configuration in PCs.

#### DISCUSSION

In this study, we established a simplified 2D model of axon-dendrite topology using aligned nanofibers and confirmed that PC dendrites grew preferentially in the direction perpendicular to the bundles of afferent parallel fiber axons. Directional arborization is likely to be a prerequisite for the planar dendrite formation *in vivo*. Moreover, we revealed that biased dendrite arborization was affected by the loss of  $\alpha$ II/ $\beta$ III spectrins. In control PCs, dendritic branches were formed mainly by terminal bifurcation, with only a few collateral branches emerging from proximal dendrites (Fujishima et al., 2012). By contrast, lateral branching events were significantly increased in  $\beta$ III spectrin-knockdown cells (Fig. 4G).

Dendritic protrusions in differentiating neurons serve as either branch precursors or immature spines (Heiman and Shaham, 2010; Yuste and Bonhoeffer, 2004). Dendrites of developing PCs in culture exhibited numerous lateral protrusions expressing the postsynaptic protein GluD2, suggesting that these lateral protrusions at the proximal dendrites are immature spine precursors. Notably,  $\beta$ III spectrin-knockdown dendrites abnormally extended some proximal protrusions to a length indistinguishable from that of dendritic branches. These elongated protrusions bore multiple GluD2 puncta along their length, which interact with neurexins/Cbln1 expressed in GC axons. Thus, the loss of  $\beta$ III spectrin seems to alter the fate of proximal dendritic protrusions from immature spines to misoriented lateral branches extending along the GC axons away from the main parasagittal plane.

#### Microtubule dynamics and dendritic branching

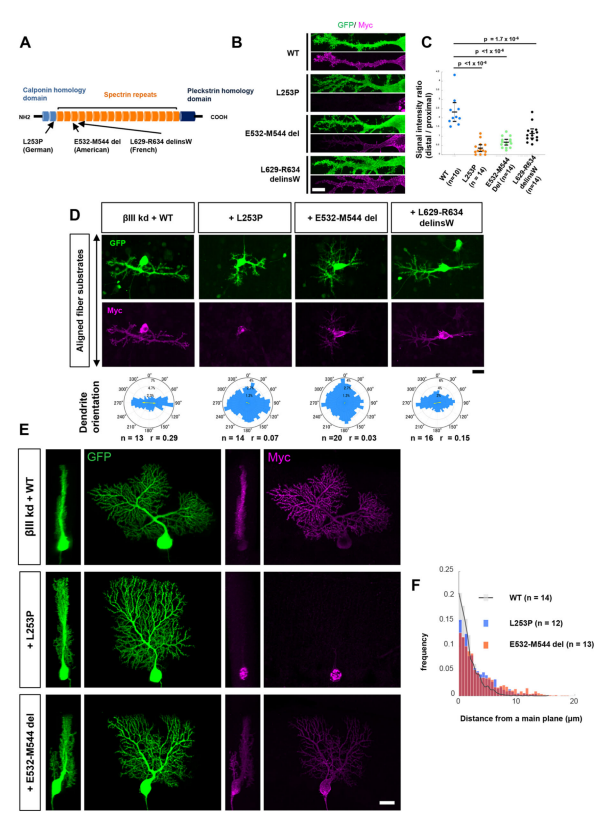
Branch formation requires microtubule extension into the precursor protrusions or filopodia, which is regulated by dynamic interplay with actin (Burnette et al., 2007; Flynn et al., 2012; Hu et al., 2012). It has been demonstrated that bundled actin in dendritic filopodia guides microtubules into filopodial protrusions from dendritic shafts, while the cortical actin meshwork in the shaft region confines microtubules and suppresses the interaction with bundled actin filaments in filopodia (Dent et al., 2007).

βIII spectrin was enriched in the thin neck regions of dendritic protrusions and the dendritic shaft and formed regular or irregular membrane skeletons (Fig. 6H; Fig. S7). We assume that βIII spectrin forms the membrane skeletons in coordination with cortical actin that functions as a molecular fence dividing the dendritic shaft and protrusions, confining microtubule dynamics along dendrites. Consistently, it has been shown that  $\beta$ III spectrin regulates constricted neck formation in dendritic protrusions (Efimova et al., 2017). Thus, in βIII spectrin-deficient PCs, dendritic protrusions were often misshapen, with actin signals expanded to the protrusion base and dendritic shaft, in contrast to the confined localization in the tip of protrusions in normal cells (Fig. 6G,J). Mislocalized actin filaments and, possibly, actin-microtubule crosslinkers, such as Navigator1, Drebrin1 and p140Cap (Geraldo et al., 2008; Jaworski et al., 2009; van Haren et al., 2014), might interact with dynamic microtubules in the shaft and enhance their entry into the protrusion. Alternatively, it is also possible that some spectrin-binding proteins (4.1 proteins, ankyrins), which also interact with microtubules (Chen et al., 2020; Leterrier et al., 2011; Ruiz-Saenz et al., 2013), might contribute to the formation of a molecular fence at the protrusion base to block microtubule entry.

Although  $\beta$ III spectrin distributes throughout the entire dendritic length, abnormal microtubule entry to protrusions was seen only in proximal dendrites in  $\beta$ III spectrin-knockdown cells. We speculate that spectrin structures, including MPS structures, are gradually stabilized along the proximal to distal axis. In normal PCs, proximal dendritic shafts would already be covered by stable spectrin structures that impede microtubule invasion into protrusions, whereas the growing end of dendrites with immature spectrin structures allow microtubule entry that enhances dendrite extension. In  $\beta$ III spectrin-knockdown cells, inhibition of microtubule entry into the proximal dendritic protrusions would be disrupted and thus induce abnormal lateral branching. Further studies are required to elucidate this hypothesis.

#### SCA5-related mutations affect dendrite growth in PCs

SCA5 is one of the autosomal dominant cerebellar ataxias caused by heterozygous mutation in the  $\beta$ III spectrin gene (Ikeda et al., 2006). Although our experiment might not completely mimic the disease condition, some SCA5-related mutants misexpressed in  $\beta$ III spectrin-knockdown cells failed to substitute for wild-type  $\beta$ III spectrin to control the dendrite orientation (Fig. 8), and the mutants dominantly affected dendritic orientation in the wild-type background (Fig. S9).



**Fig. 8. Functional analysis of the effect of SCA5-related mutations on the regulation of dendrite orientation.** (A) Domain structure of βIII spectrin and familial mutations associated with SCA5. (B) Localization of myc-tagged βIII spectrin wild type (WT), L253P, E532-M544 del or L629-R643 delinsW mutant in PCs (9 DIV). (C) Quantification of the signal intensity ratio (distal versus proximal) of βIII molecules in PC dendrites. Statistical analysis: Kruskal–Wallis test followed by Dunnett's test. (D) Morphologies of PCs transfected with GFP/shRNA-βIII spectrin plus βIII spectrin WT, L253P, E532-M544 del or L629-R643 delinsW grown on aligned nanofibers. Polar histograms indicate the angular distribution of dendritic segments. (E) Morphologies of PCs expressing the GFP/shRNA-βIII plasmid with βIII spectrin WT, L253P or E532-M544 del in P14 sagittal slices. (F) Proportions of dendritic segments locating at the indicated distances from the main plane. Scale bars: 10 µm in B; 20 µm in D,E.

L253P proteins were not delivered to dendritic membranes nor did they replace the function of the wild-type molecule in regulating the oriented dendrite growth. Consistently, previous studies have shown that the L253P mutation affects the trafficking of β-spectrin from the Golgi apparatus (Clarkson et al., 2010). It has also been suggested that the L253P mutation might reduce the plasticity of the actinspectrin network by enhancing the actin-spectrin affinity. This change might affect the proper dendritic localization of BIII spectrin (Avery et al., 2017). By contrast, L629-R634 delinsW had only minor effects on dendritic localization and the oriented arborization of dendrites. Interestingly, E532-M544 del was severely defective in controlling dendrite growth orientation despite its relatively normal dendritic localization except for the distalmost region. It has been proposed that the deletion of E532-M544 possibly affects the triple  $\alpha$ helical structures of the spectrin repeats, which might result in the alteration of overall  $\alpha/\beta$  structures (Ikeda et al., 2006). We assume that the E532-M544 del mutation might affect the stabilization of the spectrin architecture in dendrites and interfere with dendrite growth in the normal direction.

The SCA5 patients and BIII spectrin knockout animals exhibit progressive neurodegeneration that has been attributed to excitotoxicity owing to the mislocalization and the decreased level of glutamate transporters (Gao et al., 2011; Perkins et al., 2010, 2016b). Given that SCA5 is a late-onset cerebellar ataxia, it might be difficult to relate the developmentally disorganized dendrites observed in the present study directly to symptoms of the adult-onset disease. However, it is suggested that the thinner dendritic branches in the knockout PCs might alter signal propagation and contribute to the hyperexcitability (Gao et al., 2011). Abnormal branch growth in the coronal direction might lead to redundant and inefficient connectivity with parallel fibers (Cuntz, 2012). Furthermore, disruption of planarity is thought to affect the compartmentalization of cerebellar circuits. In the mature cerebellar cortex, each PC is innervated by a single climbing fiber axon of an inferior olive neuron. Coronally extended PC dendrites invading the neighboring PC territory might receive abnormal innervation by climbing fibers of neighboring PCs (Gao et al., 2011; Kaneko et al., 2011; Miyazaki and Watanabe, 2011; Perkins et al., 2016a). Developmental defects in dendrite arborization owing to the loss of BIII spectrin might deteriorate the cerebellar function before degeneration.

# Remaining questions in the perpendicular axon-dendrite interactions

We demonstrate that PC dendrites grow perpendicular to parallel fibers, which seemingly contribute to the planar arborization *in vivo*. We also demonstrate that  $\beta$ III spectrin is required for the perpendicular dendrite arborization by suppressing ectopic branch formation. However, considering that main dendritic frameworks still form planar patterns in the  $\beta$ III spectrin-deficient PCs (Fig. 5B),  $\beta$ III spectrin might function as a gatekeeper to maintain the perpendicular and planar dendrites but not as the main determinant regulating the directional dendrite extension.

The perpendicular interactions might serve as a permissive mechanism for the planar PC dendrite arborization in parasagittal planes. However, other mechanisms should also be involved in the spatial organization of dendrites, because perpendicular interactions would not assure arborization in the exact parasagittal plane. For instance, contact-based repulsion might support the flat dendrite formation by regulating the space between neighboring dendritic planes (Fig. 5A) (Fujishima et al., 2012; Ing-Esteves et al., 2018).

Nagata et al. (2006) previously proposed that the perpendicular interaction between PC dendrites and parallel fibers might be

regulated by 'contact guidance', whereby cells recognize anisotropy of substrates to determine their directionality. Previous studies using carcinoma cells have demonstrated that anisotropic substrates affect the orientation of focal adhesions and actin fibers anchored to the adhesions, thereby leading to the directional movement (Ray et al., 2017). We assume that PC dendrites might sense the anisotropy of GC axons and adjust the orientation of adhesions and cytoskeletal components, including spectrin tetramers, thereby affecting the orientation of dendrite growth.

Previous studies using diffusion magnetic resonance imaging have revealed a three-dimensional grid-like organization of axonal bundles in the forebrain, in which two distinct fibers interweave and cross at nearly right angles (Wedeen et al., 2012). Such a grid-like organization might be formed by perpendicular contact between different types of axons. It is of interest to confirm whether perpendicular contact guidance is a general mechanism in neural network formation.

#### **MATERIALS AND METHODS**

#### Mice

Mice were handled in accordance with the guidelines of the Animal Experiment Committee of Kyoto University and were housed in a dedicated pathogen-free environment with a 12 h/12 h light/dark cycle.

#### Plasmids

The pAAV-CAG-GFP (or mCherry)-hH1 vector, including the human H1 promoter, was used to express shRNA to knockdown target gene expression, as previously described (Fukumitsu et al., 2015). The targeting sequences were designed by using the Web-based software siDirect (Naito et al., 2009): control shRNA (5'-GCATCTCCATTAGCGAACATT-3'), ßIII spectrin shRNA (5'-GTCAATGTGCACAACTTTACC-3') and all spectrin shRNA (5'-GTAAAGACCTCACTAATGTCC-3'). To generate resistant mutants of  $\alpha$ II spectrin and  $\beta$ III spectrin that contained three silent mutations within shRNA target sequences, complementary DNA (cDNA) of mouse all spectrin or BIII spectrin was cloned from a mouse brain cDNA library and mutagenized by using a PCR-based method. To generate the L253P, E532-M544 del and L629-R634 delinsW BIII spectrin mutants, PCR-based mutagenesis was performed by using the resistant mutant of BIII spectrin as a template. all spectrin tagged with hemagglutinin (HA) at the N-terminus and BIII spectrin wild-type and mutant sequences tagged with myc at the N-terminus were cloned into the pCAGGS vector. To generate the EB3-EGFP construct, the coding sequence of EB3 was amplified from a mouse brain cDNA library and inserted into the pAAV-CAG-EGFP plasmid. For the CRISPR/Cas9-based knockout of BIII spectrin, the guide RNA sequence was selected by using the Web-based software CRISPRdirect (Naito et al., 2015). The βIII spectrin target sequence (5'-GAGACC-TGTACAGCGACCTG-3') was inserted into pSpCas9(BB)-2A-GFP (PX458) (Addgene plasmid #48138) (Ran et al., 2013).

#### In utero electroporation

The *in utero* electroporation of plasmids was performed as described previously (Nishiyama et al., 2012). Briefly, pregnant mice on day 11.5 of gestation were deeply anesthetized via the intra-abdominal injection of a mixture of medetomidine, midazolam and butorphanol. Plasmid DNA ( $1-5 \mu g/\mu l$ ) was microinjected into the fourth ventricle of the embryos (FemtoJet; Eppendorf). Then, five current pulses (amplitude, 33 V; duration, 30 ms; intervals, 970 ms) were delivered with a forceps-shaped electrode (CUY650P3; NepaGene) connected to an electroporator (CUY21; NepaGene).

#### In vivo electroporation to label parallel fibers

The *in vivo* electroporation of plasmids was performed as described previously (Umeshima et al., 2007). P8 ICR mice were cryoanesthetized. A small burr hole was made in the skull over the cerebellum with a 27-gauge needle. The plasmid DNA (pAAV-CAG-mCherry) was microinjected through the hole by using a syringe with a 33-gauge needle (Ito). A forceps-shaped electrode

connected to the cathode of an electroporator (CUY21; NepaGene) was placed in the occipital region. A needle used for DNA injection was connected to the anode. Then, six current pulses (amplitude, 70 V; duration, 50 ms; intervals, 150 ms) were delivered. After the wound was sutured, the pups were warmed at 37°C and returned to the home cage.

### Primary cerebellar culture and nucleofection of cerebellar neurons

The primary culture of cerebellar neurons was performed as previously described (Fujishima et al., 2012) with slight modifications. Cerebella from P0 mice were dissected in Hank's balanced salt solution (Gibco) and dissociated using a Neuron Dissociation Kit (FUJIFILM Wako Pure Chemicals). Cells were plated on a 12 mm coverslip coated with poly-D-lysine in Dulbecco's modified Eagle's medium/Ham's F12 (DMEM/F12) supplemented with 10% fetal bovine serum at a density of 1.5 cerebella per coverslip. Following incubation, the media were replaced with maintenance media containing DMEM/F12, 0.1 mg/ml bovine serum albumin, 2.1 mg/ml glucose,  $2\times$  Glutamax, 8  $\mu$ M progesterone, 20  $\mu$ g/ml insulin, 200  $\mu$ g/ml transferrin, 100  $\mu$ M putrescine, 30 nM selenium dioxide, 4  $\mu$ M AraC and 1% penicillin-streptomycin. For cultures on nanofibers, dissociated neurons were plated on aligned or random nanofiber plates (Nanofiber Solutions).

For the transfection of plasmid DNA into PCs, nucleofection was performed as described previously (Kawabata Galbraith et al., 2018). Briefly, dissociated cerebellar cells from between 1.5 and two cerebella were washed twice with OptiMEM and resuspended in 100  $\mu$ l of OptiMEM containing 5-8  $\mu$ g of plasmid DNA. Cells were transferred to a cuvette and nucleofected with a Nepa21 electroporator (NEPA GENE).

To separate the large cell fraction (containing PCs, interneurons and glia) and the small cell fraction (containing GCs), dissociated cells from P0 cerebellar tissue were purified with a step gradient of Percoll (Hatten, 1985). Isolated GCs from P0 mice were plated on the aligned fibers at a density of  $10 \times 10^5$ ,  $7 \times 10^5$  or  $5 \times 10^5$  cells/cm<sup>2</sup>. Then, the large cell fraction including PCs was plated on the culture at a density of  $10 \times 10^4$ ,  $7 \times 10^4$  or  $5 \times 10^4$  cells/cm<sup>2</sup>, respectively.

For the co-culture of cerebellar microexplants and PCs, a microexplant culture from cerebellar tissues was prepared as described previously (Nakatsuji and Nagata, 1989). Briefly, the external granular layer of the cerebellar cortex from P2 mice was dissected into 300-500  $\mu$ m pieces and plated on the poly-D-lysine- and laminin-coated coverslips. One day after plating, the isolated large cell fraction including PCs from P0 mice was added to the culture.

#### Immunofluorescence and image acquisition

For immunocytochemistry, cells cultured on coverslips or nanofibers were fixed for 15 min at room temperature in 4% paraformaldehyde (PFA) in PBS. Cells were washed and permeabilized with PBS containing 0.25% Triton (PBS-0.25 T). The cells were then blocked with a blocking solution [PBS (-0.25 T) with 2% bovine serum albumin] for 30 min at room temperature. The cells were incubated with primary antibodies at 4°C overnight in the blocking solution, washed with PBS and incubated with fluorescently labeled secondary antibodies in blocking solution (for details of the antibodies used, see Table S1).

For immunohistochemistry, the mice were anesthetized with isoflurane and perfused with PBS followed by 4% PFA in phosphate buffer. Their brains were removed and postfixed overnight in 4% PFA/PBS at 4°C. After washing with PBS, the brains were embedded in a 3.5% low-melt agarose in PBS. Sagittal or coronal sections were cut at a thickness of 50-100  $\mu$ m using a vibratome (Dosaka). The sections were permeabilized in PBS with 0.5% Triton (PBS-0.5 T) and blocked with 2% skim milk in PBS (-0.5 T) for 30 min. The sections were incubated with primary antibodies at 4°C overnight in blocking solution, washed with PBS and incubated with fluorescently labeled secondary antibodies in 2% skim milk in PBS (-0.5 T).

Images of the fixed samples were acquired on a laser scanning confocal microscope (Fluoview FV1000; Olympus) equipped with UPLSAPO  $40\times$  dry (NA 0.95),  $60\times$  water-immersion (NA 1.2) and  $100\times$  oil-immersion objectives (NA 1.40). For the imaging of PC dendrites in the cerebellar tissue, serial confocal *z*-stack images were acquired from midsagittal regions

in lobes IV-V with the  $60 \times$  objective at a *z*-step of 0.57 µm. The images were three-dimensionally reconstructed by using Imaris software (Bitplane). For the imaging of PC dendrites in the dissociated culture, serial confocal *z*-stack images were obtained with  $40 \times$  or  $100 \times$  objectives.

#### **Quantification of dendrite morphology**

For the analysis of dendritic flatness, captured confocal *z*-stacked images were binarized and skeletonized in ImageJ ['*Skeletonize* (*2D*/*3D*)' plugin]. To identify the plane with the closest fit to the given dendritic arbor, a principal components analysis was performed in MATLAB software (Fig. S2). To analyze the distance of the dendritic branches from the fitted plane, 3000 points (for Fig. 1A,B) or 100 points (for Fig. 1D; Fig. 3E; Fig. 8E) in the skeletonized dendritic images from each cell were randomly selected, and the distance between each point and the fitted plane was calculated.

For the morphometric analysis of PC dendrites on aligned fibers, *z*-projected images were binarized and skeletonized in the ImageJ plugin or MATLAB software. To analyze the branch angle, dendritic branches were divided into  $3.5 \,\mu\text{m}$  segments, and the angle between each segment and fiber was quantified (Fig. S3).

#### **STED** imaging

We used a Leica TCS SP8 STED with an oil immersion  $100 \times objective lens$  with NA 1.4 (HC-PL-APO  $100 \times /1.4$  OIL, Leica) to analyze the subcellular localization of  $\beta$ III spectrin. The protein was labeled with anti- $\beta$ III spectrin (Santa Cruz, SC-28273, 1:100) and a secondary antibody conjugated with Alexa555 (Thermo Fisher A-31572, 1:1000). The fluorophore was excited with a white laser tuned to 555 nm and depleted with a 660 nm STED laser. A time gate window of 0.35-3.85 ns was used to maximize the STED resolution.

#### Time-lapse imaging, image processing and image analysis

For long-term time-lapse imaging, fluorescently labeled PCs were observed every 1-3 h with an incubation microscope (LCV100; Olympus) equipped with a 20× objective (NA 0.7; Olympus). Serial *z*-stacked images were obtained at *z*-steps of 1  $\mu$ m (1  $\mu$ m×five steps).

For the high-resolution live imaging of dendritic arbors (used for EB3 imaging experiments), for the analysis of EB3 dynamics, we used 10-11 DIV PC dendrites transfected with pAAV-CA-EB3-EGFP and pAAV-CAG-mCherry/hH1-control or hH1-BIII spectrin. Time-lapse images were obtained with a confocal microscope (FV1000, BX61W1; Olympus) equipped with a LUMFI 60× objective (NA 1.10; Olympus) under 5% CO2 supplementation. Images were recorded with 3× digital zoom at an interval of 3 s. To eliminate background cytosolic signals, average subtraction was performed (Schätzle et al., 2016), in which an average projection of all the time-lapse images of EB3-EGFP signals was generated and subtracted from each frame of the time-lapse images. The images were then processed via an unsharp masking procedure in ImageJ software to obtain the enhanced images. For the quantitative analysis, the number of dendritic protrusions in the dendritic segments of 10 µm at the most distal regions or proximal regions (more than 10 µm away from the most distal end) were counted in ImageJ. Of those, dendritic protrusions invaded by the EB3-EGFP were identified manually with the assistance of the ImageJ plugin TrackMate (Tinevez et al., 2017).

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#### **Competing interests**

The authors declare no competing or financial interests.

#### Author contributions

Conceptualization: K.F., M.K.; Methodology: K.F., M.K.; Formal analysis: K.F., J.K., M.Y.; Investigation: K.F., J.K., M.Y.; Writing - original draft: K.F., M.K.; Writing - review & editing: K.F., M.K.

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#### Supplementary information

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