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GluD2 Endows Parallel Fiber–Purkinje Cell Synapses with a High Regenerative Capacity

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Although injured axons usually do not regenerate in the adult CNS, parallel fibers (PFs) regenerate synaptic connections onto cerebellar Purkinje cells (PCs). In this study, we investigated the role of GluD2 in this regenerative process after PF transection using GluD2-knockout (KO) mice. All dendritic spines on distal dendrites were innervated by PFs in sham-operated wild-type controls, whereas one-third were devoid of innervation in GluD2-KO mice. In both genotypes, a steep drop in the number of PF synapses occurred with a reciprocal surge in the number of free spines on postlesion day 1, when the PF territory aberrantly expanded toward the proximal dendrites. In wild-type mice, the territory and number of PF synapses were nearly fully restored to normal on postlesion day 7, although PF density remained low. Moreover, presynaptic and postsynaptic elements were markedly enlarged, and the PF terminal-to-PC spine contact ratio increased from 1:1 to 1:2 at most synapses. On postlesion day 30, the size and contact ratio of PF synapses returned to sham-operated control values and PF density recovered through the sprouting and elongation of PF collaterals. However, GluD2-KO mice showed neither a hypertrophic response nor territorial restoration 7 d postlesion, nor the recovery of PF axons or synapses on postlesion day 30. This suggests that PF wiring regenerates initially by inducing hypertrophic responses in surviving synaptic elements (hypertrophic phase), followed by collateral formation by PF axons and retraction of PF synapses (remodeling phase). Without GluD2, no transition to these regenerative phases occurs.

Key words: cerebellum; climbing fiber; GluD2; parallel fiber; Purkinje cell; synapse regeneration

Introduction

Dendrites of Purkinje cells (PCs) are arborized in the parasagittal plane of the cerebellum, and consist of two compartments, distal and proximal, that differ in the origin of their inputs as well as in the mode of innervation. Spiny branchlets in the distal compartment are innervated by $10^5–10^6$ parallel fibers (PFs), T-shaped granule cell axons running in the transverse plane, but each PF usually forms only one synapse onto a given PC (Palay and Chan-Palay, 1974; Napper and Harvey, 1988; Kurihara et al., 1997). In comparison, a single climbing fiber (CF) originating from the inferior olivary nucleus forms hundreds of synapses onto a single PC by twisting around the dendritic shafts in the proximal compartment (Crepel et al., 1981; Rossi et al., 1993; Ichikawa et al., 2002). The segregated innervation territories of the two orthog-
oral afferents and the mono-innervation by single CFs are the two distinguishing features of excitatory synaptic wiring on PCs, and are established in the first 3–4 postnatal weeks in rodents (Altman, 1972; Watanabe and Kano, 2011). These salient features are thought to be the neural basis for information processing and integration underlying cerebellar motor function and learning (Ito, 2012).

Various molecular mechanisms regulate the formation and maturation of PC circuits (Yuzaki, 2009; Watanabe and Kano, 2011; Hashimoto and Kano, 2013). Of these, the glutamate receptor GluD2 (GluRΔ2), which is exclusively concentrated on the postsynaptic membrane at PF–PC synapses (Takayama et al., 1995; Landsend et al., 1997), plays a key role in synapse formation. Ablation of the Grid2 gene (which encodes GluD2) causes a severe reduction of PF–PC synapses and results in numerous spines lacking innervation, i.e., free spines (Guastavino et al., 1990; Kashiwabuchi et al., 1995; Kurihara et al., 1997; Lalouette et al., 2001; Takeuchi et al., 2005). This input-selective synaptic strengthening is mediated by a tripartite molecular complex comprised of GluD2, Cbln1, and neurexin, in which Cbln1 released from PFs interacts with GluD2 on PC spines and neurexin on PF terminals (Matsuda et al., 2010; Uemura et al., 2010). This molecular interaction is also important for suppressing aberrant extension of the CF territory; without it, CFs invade distal dendrites to take over free spines on innervating and neighboring PCs (Ichikawa et al., 2002; Hirai et al., 2005). Furthermore, this GluD2–Cbln1–neurexin system is essential for maintaining the PC circuitry in adulthood (Takeuchi et al., 2005; Ito-Ishida et al., 2008; Torashima et al., 2009; Miyazaki et al., 2010).

Axons injured in the adult mammalian CNS of higher species do not usually regenerate (Dusart et al., 2005; Maier and Schwab, 2006). But intriguingly, PF synapses on PCs rapidly regenerate after surgical transection of PFs. Chen and Hillman (1982) were the first to show that, in the adult rat cerebellum, degenerating PF terminals and free spines almost completely disappear within 5 d after surgical transection, and that PF synapses are restored by takeover of preexisting PC spines and new synapse formation. However, the molecular mechanisms underlying this high regenerative ability remain unknown. In the present study, we investigated whether GluD2 plays a role in the regeneration of PF synapses using electron microscopic reconstruction of dendritic innervation and of the territories of PF-transected PCs. We show in wild-type mice that the regeneration of PF–PC synapses proceeds in three distinct phases: the degenerative, hypertrophic, and remodeling phases. In GluD2-knock-out (KO) mice, lesioned PF wiring remains in the degenerative phase, without shifting to the hypertrophic and remodeling phases of regeneration.

Materials and Methods

Animals. Wild-type (C57BL/6N) and grid2 (GluD2)-KO mice of either sex at 2 months of age were used in the present study. GluD2-KO mice were obtained by mating heterozygous pairs, which had been produced and maintained on a pure C57BL/6N genetic background (Yamasaki et al., 2011). The genotype was determined by PCR, as reported previously (Takeuchi et al., 2005). All experiments were performed according to guidelines laid down and approved by the animal care and welfare committees of Sapporo Medical University and Hokkaido University. Three wild-type and three GluD2-KO mice were analyzed for each qualitative and quantitative analysis at each time point.

PF transection. Under anesthesia with chloral hydrate (350 mg/kg body weight, i.p.), a transfolial incision was made in the cerebellar vermis (lobules 4–7) to produce PF-transected “cerebellar islands.” A pair of incisions, with a 2.0 mm intervening space, were made by inserting a microknife to a depth of 0.6 mm from the pial surface, 30° to the parasagittal plane (Fig. 1A). Then, 1, 7, and 30 d later, mice under chloral hydrate anesthesia were perfused transcardially with 4% paraformaldehyde in 0.1 M phosphate buffer (PB), pH 7.4, for light microscopic immunohistochemical analyses, or with 4% paraformaldehyde and 0.1% glutaraldehyde in 0.1 M PB for electron microscopic reconstruction. After excision from the skull, brains were immersed overnight in the same fixative. All analyses were done for cerebellar islands on lobule 6 in PF-transected mice or the corresponding region in sham-operated mice.
Tracer labeling of PFs. To follow the regeneration of PF synapses, biotinylated dextran amine (BDA; molecular weight, 10,000; Invitrogen) was injected into the vermis (lobule 6) of PF-transected mice 2 d before transcardial perfusion. Briefly, under chloral hydrate anesthesia, a 1-MΩ-resistance glass pipette, filled with 0.5 μl of 10% BDA in PBS, pH 7.4, was inserted into the left side of the cortical islands, and BDA was iontophoretically applied with a 7 μA positive current for 20 min (current parameters: 700 ms on and 1300 ms off; Ichikawa et al., 2011). To visualize BDA-labeled PFs, coronal microslicer sections were immersed overnight in streptavidin–peroxidase conjugate (Nichirei) diluted with PBS containing 0.5% Triton X-100 for light microscopy and visualized with 3′,3′-diaminobenzidine (DAB). Using a light microscope (AX-80, Olympus), the density of PF varicosities was measured from >20 labeled PFs from each mouse and averaged. The p value was calculated using ANOVA with Dunnett’s test (see Fig. 3G).

Immunohistochemistry. For immunofluorescence, sections (50 μm thick) were made on a microslicer (MAN52, Campden) and were prepared in the parasagittal and coronal planes. Sections were incubated overnight at room temperature in a mixture of rabbit and guinea pig antiserum to GluD2 (1:10,000; Nakagawa et al., 1998) and guinea pig vesicular glutamate transporter type 2 (VGluT2) antibody (1.0 μg/ml; Miyazaki et al., 2003) for parasagittal sections (see Fig. 4A–H) or rabbit antiserum to GluD2 and guinea pig vesicular glutamate transporter type 1 (VGluT1) antibody (1.0 μg/ml; Miyazaki et al., 2003) for coronal sections (Fig. 1B,C). Subsequently, sections were incubated in a mixture of AlexaFluor 594-labeled anti-guinea pig IgG and AlexaFluor 488-labeled anti-rabbit IgG for 2 h (1:200; Invitrogen). Coronal images were obtained on a fluorescence microscope (BX-9000, Keyence), and parasagittal images were captured using a confocal scanning laser microscope (Radiance 2100, Zeiss) equipped with an Ar/Kr laser and a Plan-Apochromat (40×, 1.29 NA, oil-immersion) objective lens (Zeiss). To avoid cross talk between fluorophores, fluorescent signals were acquired sequentially using the 488 and 568 nm excitation laser lines. Five images (512 × 512 pixels; pinhole size, 1.6 mm) at different levels along the z-axis were compiled into single images. The heights of CFs and the molecular layer were measured using 10 images from each mouse, and averaged. Statistical differences between sham-operated control and PF-transected mice were assessed using ANOVA with Dunnett’s test, whereas those between wild-type and GluD2-KO mice were assessed using Student’s t test (see Fig. 4F–K).

For immunoelectron microscopic labeling of CFs, parasagittal microslicer sections were incubated overnight with guinea pig VGluT2 antibody (1.0 μg/ml) and peroxidase-labeled guinea pig IgG antibody (1:200, Jackson Immunoresearch), and visualized with DAB. To label GABAergic terminals, sections were incubated overnight with rabbit VIAAT antibody (1.0 μg/ml), and subjected to silver-enhanced pre-embedding immunogold microscopy using 1.4 nm gold particle-conjugated rabbit IgG antibody (1:100; Nanoprobes) and the HQ silver kit (Nanoprobes).

Serial electron microscopy and 3D reconstruction. To measure the density, size, and contact ratio of PFs, 10 sets of five serial electron microscopic images were taken from superficial portions of the molecular layer (0–40 μm depth from the pia mater) for each of three PF-transected or sham-operated mice. We measured the short diameter of PF terminals and the density of PF axons. The p value was calculated using ANOVA with Dunnett’s test (Fig. 2I–L).

To reconstruct 3D images of PC dendrites, we prepared sets of 1000–1400 ultrathin sections (100 nm thick) in the horizontal plane. A ribbon of serial sections, each consisting of at least 20–30 sections, was mounted on a single-slot copper grid (1 × 2 mm) supported with a formvar membrane. Electron micrographs were taken from every second section at an original magnification of 15,000×. Using electron microscopic images, the path length (L) of curved or leaning dendrites was calculated using the long (D1) and short (D2) diameters of dendritic profiles and using the thickness (t) and number (n) of sections: L = D1/D2 × t × n. From the reconstructed images, we obtained the mean number and density of synapses and free spines, as well as the composition of three dendritic segments innervated by CFs only, CFs plus PFs, and PFs only. The p value was calculated using ANOVA with Dunnett’s test (see Fig. 8).

Results

Surgical transection of PFs

A pair of transfolial incisions were made in the cerebellar vermis of adult wild-type (C57BL/6N) and GluD2-KO mice (Fig. 1A) to investigate the role of the receptor in the degeneration and regeneration of PF–PC synapses in PF-deafferented cerebellar islands. The corresponding cerebellar regions in sham-operated mice were used as experimental controls for each genotype. Transection of PFs was ascertained by the discontinuity at incision sites by immunofluorescence labeling for VGluT1, a marker of PF terminals (Fig. 1B,C), and by anterograde tracer labeling of BDA injected into the cerebellar islands (Fig. 1D,E). Long transverse beams of BDA-labeled PFs extended bilaterally from the injection sites in sham-operated controls (Fig. 1D), whereas in transected mice, the PF beams did not extend beyond the incision sites, even by postlesion day 30 (Fig. 1E). These findings demonstrate the successful transection of PFs and the lack of regeneration beyond the lesion site, at least until postlesion day 30.

Degeneration and regeneration in cerebellar islands

Changes in PF axons, terminals, and synapses were examined in cerebellar islands on postlesion days 1, 7, and 30 by electron microscopy (Fig. 2). In this analysis, we prepared 10 sets of parasagittal serial sections from each mouse, and photos were taken from superficial portions of the molecular layer. Using serial images, we identified PF axons (red) and terminals (orange) running orthogonal to the section plane, and distinguished PC spines forming PF synapses (blue) from those lacking synaptic contact, ie, free spines (green).

In sham-operated controls, most PF terminals in wild-type mice formed synaptic contacts with single PC spines (ie, 1:1 contact), and the postsynaptic density (PSD) was large and well thickened beneath the postsynaptic membrane (Fig. 2A, arrowheads). In contrast, approximately one-third of PC spines were free spines in sham-operated GluD2-KO mouse controls (Fig. 2E,F), a hallmark of synaptic defects in this mutant, and were associated with rudimentary PSD-like condensation, as reported previously (Guastavino et al., 1990; Kurihara et al., 1997; Ichikawa et al., 2002; Takeuchi et al., 2005). On postlesion day 1, transfolial transection caused a marked reduction of PF axons and terminals, with a concomitant emergence or increase of free spines in wild-type and GluD2-KO mice, respectively (Fig. 2B,F). This was reflected in a significant drop in the profile density of PF axons (per μm² of area) in the molecular layer (Fig. 2I) and in the percentage of PC spines having synaptic contact with PF terminals (Fig. 2F), compared with sham-operated controls (n = 3; p < 0.05, ANOVA with Dunnett’s test). Thereafter, genotypic differences became more apparent. In wild-type mice, free spines virtually disappeared (Fig. 2C,F), and PF terminals and PC spines were markedly enlarged on postlesion day 30 (Fig. 2C,K). Furthermore, the PF terminal-to-PC spine contact ratio was increased to 1:2 at most PF–PC synapses (Fig. 2C,L). On postlesion day 30, the profile density of PF axons was increased to 75.2 ± 13.0% of the control level (Fig. 2I), and the size of PF terminals and the PF terminal-to-PC spine contact ratio returned to control levels (Fig. 2D,K,L). In contrast, in GluD2-KO mice, no such hypertrophic response was observed on postlesion day 7 (Fig. 2G,K,L). In addition, >90% of PC spines were free spines (Fig. 2F–H,J), and the profile density of PF axons remained low, at 55.0 ± 3.2% of control level, on postlesion day 30 (Fig. 2I). Thus, regeneration of PF axons and synapses was severely impaired in GluD2-KO mice compared with wild-type animals.
Sprouting, elongation, and varicosity formation in PF collaterals

Light microscopic examination of BDA-labeled PFs on postlesion day 7 confirmed transient hypertrophic responses, i.e., thickening of PF axons and enlargement of PF terminals, in wild-type mice (Fig. 3A–C), but not in GluD2-KO mice (Fig. 3D–F). We also found on postlesion day 30 that PFs in wild-type mice often projected long collateral branches bearing a number of varicosities (Fig. 3C), whereas those in GluD2-KO mice gave off a few short branches bearing only one or a few varicosities (Fig. 3F). No significant differences were noted in the density of PF varicosities (per millimeter of BDA-labeled PFs) between sham-operated controls and PF-transected mice on postlesion day 7 or 30 in any genotype (Fig. 3G). Given the lack of PF regeneration beyond the transection site, these findings suggest that the increase in PF density on postlesion day 30 in wild-type mice is achieved mainly by sprouting and elongation of PF collaterals within the cerebellar islands.

Changes in PC dendrites and CF reach

Next, we examined changes in molecular layer thickness and CF reach after PF transection by double-immunofluorescence labeling for calbindin (green) and VGluT2 (red), markers for PCs and CF terminals, respectively. Confocal microscopic images were taken from parasagittal sections prepared from cerebellar islands and the corresponding regions of sham-operated controls (Fig. 4A–H). In sham-operated controls, the thickness of the molecular layer, i.e., the mean vertical height to the tips of calbindin-positive PC dendrites, was significantly lower in GluD2-KO mice compared with wild-type mice (Fig. 4I; n = 3; p < 0.01, Student’s t test), consistent with our previous report (Kurihara et al., 1997). Nevertheless, the CF reach, i.e., the mean vertical height to the tips of VGluT2-labeled CF terminals, was similar in GluD2-KO and wild-type mice (Fig. 4J), suggesting invasion of the CFs to the distal dendritic compartment in GluD2-KO mice (Fig. 4F). As a result, the CF reach relative to molecular layer thickness was sig-
nificantly elevated in sham-operated GluD2-KO controls compared with wild-type mice (Fig. 4K; \( p < 0.01 \), Student’s \( t \) test).

After PF transection, the molecular layer thickness was reduced significantly and progressively in both genotypes (Fig. 4Ij), and the reduction in CF reach was mild in both genotypes (Fig. 4Jf). Accordingly, the CF reach relative to the molecular layer thickness remained significantly high in GluD2-KO mice (92–97%) compared with wild-type mice (80–86%) on each postlesion day examined (Fig. 4K; \( n = 3; p < 0.05 \) for each day, Student’s \( t \) test). Of note, the distal dendrites were severely atrophied in GluD2-KO mice on postlesion days 7 and 30, as reflected by an increase in calbindin-negative space between calbindin-positive dendritic shafts, and atrophied dendrites were associated with more VGluT2-labeled CF terminals (Fig. 4G–H).

Reconstruction of dendritic innervation

To clarify changes in synaptic composition and location, we reconstructed PC dendrites by serial electron microscopy using horizontal sections cut parallel to the pial surface. From each genotype at each time point, we selected three PCs that followed a single dendritic tract from the apical pole of PC somata up to the dendritic tips ascending to the pia mater. We identified CF synapses by the presence of dark immunoperoxidase precipitates for VGluT2 (Fig. 5A) and PF synapses by innervation by VGluT2-negative transverse axons (Fig. 5B). All of the identified CF and PF terminals formed asymmetrical synapses on dendritic spines (Fig. 5A, B). After PF transection, unusual synaptic elements were frequently encountered, including degenerating PF synapses with high electron density (Fig. 5C, right), symmetrical synapses on dendritic spines, and free spines lacking presynaptic contact (Fig. 5C, left and middle). Symmetrical synapses on PC spines were atypical GABAergic synapses by molecular layer interneurons, as their presynaptic terminals were labeled for VIAAT (Fig. 5D).

Three kinds of spine-type synapses and free spines were reconstructed along dendritic tracts in wild-type and GluD2-KO PCs (Fig. 5E–L). PF synapses (red dots) and free spines (pink squares) are plotted to the right of each dendrite, whereas CF synapses (yellow circles) and spine-type GABAergic synapses (green) are to the left. We also categorized PC dendrites into three domains according to the location of PF and CF synapses (Fig. 5E–L): PC dendrite-I (PCD-I), innervated by CFs alone (blue columns); PCD-II, innervated by both CFs and PFs (cyan); and PCD-III, innervated by PFs alone (green). Because free spines result from failed synaptic contact with PFs in GluD2-KO mice (Ichikawa et al., 2002; Takeuchi et al., 2005) and also emerge after surgical transection of PFs (Chen and Hillman, 1982; the present study), we assumed that free spines represented PF synapses in categorization of dendritic domains. Short segments from the distal and proximal dendrites were 3D-reconstructed and are presented in Figures 6 and 7, respectively.

Changes in PF and CF innervation onto PCs

Using the reconstructed data, we analyzed differences in PC innervation, including differences in territories, and measured the path length of each dendritic domain in GluD2-KO and wild-type mice. Consistent with our previous study (Ichikawa et al., 2002), the PCD-II domain was abnormally elongated with reciprocal shortening of the PCD-I and PCD-III domains in sham-operated GluD2-KO control mice compared with wild-type mice (Fig. 8A). Moreover, reflecting progressive thinning of the molecular layer (Fig. 4l), the total path length of traced PC dendrites was significantly shorter on postlesion days 7 and 30 compared with sham-operated GluD2-KO and wild-type controls (Fig. 8A; \( n = 3; p < 0.05 \) for each day, ANOVA with Dunnett’s test).

In wild-type mice, the mean path length of the PF territory, ie, the summed length of the PCD-I and PCD-III domains, was significantly longer on postlesion day 1 (Fig. 8B; \( n = 3; p < 0.01 \), ANOVA with Dunnett’s test), when the mean number of PF synapses in the PF territory showed significant reductions (Fig. 8C; \( n = 3; p < 0.01 \), ANOVA with Dunnett’s test). These changes were transient and returned to control levels on postlesion days 7 and 30 (Fig. 8B, C). GluD2-KO mice showed similar changes on postlesion day 1, ie, elongation of the PF territory and a reduction in the number of PF synapses (Fig. 8B, C). However, the number of PF synapses showed no recovery on postlesion day 7 or 30 (Fig. 8C; \( p < 0.01 \) at each time point, ANOVA with Dunnett’s test). Thus, the PF territory expands through the massive degeneration of PF synapses in both strains of mice, but whereas these changes are transient in wild-type mice, they persist in the GluD2-KO.

The mean path length of the CF territory, ie, the summed length of the PCD-I and PCD-II domains, showed a mild reduction after PF transection in both genotypes (Fig. 8D). The shortening of the CF territory showed a tendency to correlate with the total path length of PC dendrites (Fig. 8A) and molecular layer thickness (Fig. 4I). After PF transection, no significant changes were found in the number of CF synapses in the CF territory in either genotype, although the values were consistently higher in
wild-type mice than in GluD2-KO mice on each postlesion day (Fig. 8E). Thus, the reorganization and restoration of PF innervation of PCs proceeds swiftly and dynamically in a GluD2-dependent manner, whereas changes in CF innervation and territory are subtle, if any, in either strain of mice.

Free spines and atypical GABAergic synapses
Reflecting the steep increase in the number of free spines per dendritic tract (Fig. 8G), the mean path length of dendrites studied with free spines increased strikingly on postlesion day 1 in both strains of mice (Fig. 8F; n = 3; p < 0.01 or 0.05 on postlesion
day 1 in wild-type and GluD2-KO mice, respectively, ANOVA with Dunnett’s test). Most free spines had disappeared by postlesion day 7 and were completely gone by postlesion day 30 in wild-type mice (Fig. 8G). In comparison, in GluD2-KO mice, free spines were numerous along substantial lengths of PC dendrites on postlesion days 7 and 30 (Fig. 8F,G; n = 3; p < 0.01 on each postlesion day, ANOVA with Dunnett’s test).

Atypical GABAergic synapses on PC spines were not found in sham-operated wild-type or GluD2-KO control mice (Fig. 5E,F). On postlesion days 1 and 7, they appeared in wild-type mice but not
in GluD2-KO mice (Fig. 8H). These findings indicate that the rapid disappearance of free spines and the transient appearance of atypical GABAergic synapses are GluD2-dependent processes.

Discussion
Animal models of perturbed cerebellar circuitry have provided important insight into the mechanisms underlying the developmental regulation and plasticity of synaptic wiring (Altman and Anderson, 1972; Strata and Rossi, 1998; Sotelo, 2004; Yuzaki, 2009; Watanabe and Kano, 2011). In the present study, we examined the regeneration of PF wiring by reconstructing dendritic innervation onto PCs after surgical transection of PFs. The findings revealed striking differences between wild-type and GluD2-KO mice following PF transection, highlighting the pivotal role of GluD2 in PF regenerative rewiring.

Comparison with previous studies
Chen and Hillman (1982) documented the rapid regeneration of PF–PC synapses.Collapsed PF ghosts, phagosomes in Bergmann glia and free spines emerge and peak within a day after PF transection. Dual innervation of single spines by degenerating and normal PFs or by two normal PFs also appear in transected cerebellar islands. Because 1:1 contact of presynaptic and postsynaptic structures is typical of normal PF–PC synapses (~90% of synapses; Palay and Chan-Palay, 1974; Kurihara et al., 1997), the emergence of dual innervation is considered to represent rapid regeneration of the synapse. Thereafter, PC spines enlarge, elongate, and branch approximately postlesion day 4, and giant spines in contact with enlarged PF terminals prevail on postlesion days 5–30. Enlarged PF terminals often contact multiple spines. Our present observations in mice are consistent with Chen and Hillman’s (1982) study in rats, except that we did not observe innervation of single spines by dual PF terminals. In the current study, we also quantitatively evaluated changes in innervation along dendritic trees.

The three phases of PF regenerative wiring
Here, we propose three distinct phases following PF lesioning in wild-type mice: the (1) degenerative, (2) hypertrophic, and (3) remodeling phases (Fig. 9).

Reflecting massive degeneration, the number of PF–PC synapses dropped steeply, whereas there was a reciprocal increase in
the number of free spines on postlesion day 1. At this time point, the PF territory, which is confined to the distal one-third of dendritic trees in sham-operated adult wild-type controls (Ichikawa et al., 2002, 2016; Fig. 5E), markedly elongated to encompass the proximal dendritic portions. This also occurred, to an even greater extent, for free spines. These observations suggest that free spines are produced not only by detachment of degenerating PF terminals, but also by de novo protrusion from proximal dendrites, and that some of these free spines on proximal dendrites are innervated by PFs. Aberrant interneuron innervation of PC spines in the expanded PF territory also suggests that regenerative responses start as immediately as degeneration does. Nevertheless, the reduction of PF synapse number to less than half that in sham-operated controls indicates that degeneration exceeds regeneration on postlesion day 1. Importantly, all events in this degenerative phase, including aberrant expansion of the PF territory, elongation of the PCD-II domain, massive emergence of free spines, and aberrant spine innervation by interneurons, recapitulate the developmental events in the early postnatal cerebellum (Kurihara et al., 1997; Ichikawa et al., 2016). Therefore, disorganized synaptic circuitry in the degenerative phase mirrors the immature synaptic circuitry during development.

In wild-type mice on postlesion days 7 and 30, most measured parameters of PF wiring returned to control levels, with retraction of the PF territory, restoration of PF synapse number, and disappearance of free spines and spine-type interneuron synapses. Nevertheless, there were significant differences between the two postlesion days. For example, although the size of PF terminals and PC spines and the PF terminal-to-PC spine contact ratio were increased on postlesion day 7, they returned to control levels by postlesion day 30. In addition, the profile density of PF axons remained low on postlesion day 7, but substantially recov-
tered by postlesion day 30, when long PF collaterals with numerous varicosities were observed. These findings suggest that PF synapses regenerate first via hypertrophic responses in surviving PFs (the hypertrophic phase), and second, through an increase in PF density by collateral sprouting and elongation (the remodeling phase). In the latter phase, the shape, size, and connectivity of PF synapses are also restored to a relatively normal compact architecture.

Role of GluD2 in PF synapse regeneration

The most notable difference in GluD2-KO mice was the lack of any sign of PF synapse regeneration. On postlesion day 7, GluD2-KO mice showed no hypertrophic response. Moreover, the PF territory remained elongated, and free spines were further increased in number. This continued until postlesion day 30, when the density of PF axon profiles remained low, suggesting that GluD2 is essential for the regeneration of PF innervation and territory after lesioning.

The connectivity of PF synapses is strengthened and maintained by synaptic adhesion systems comprised of GluD2, Cbln1 and neurexin (Guastavino et al., 1990; Kashiwabuchi et al., 1995; Kurihara et al., 1997; Hirai et al., 2005; Matsuda et al., 2010; Uemura et al., 2010). GluD2-KO and Cbln1-KO mice share many common phenotypes, including the severe loss of PF–PC synapses, mismatched alignment of presynaptic and postsynaptic structures at PF–PC synapses, emergence of free spines, distal invasion of the CF territory, persistent multiple CF innervation of PCs, impaired long-term depression at PF–PC synapses, and ataxia (Kashiwabuchi et al., 1995; Ichikawa et al., 2002; Hirai et al., 2005; Miyazaki et al., 2010). Importantly, cerebellar administration of recombinant Cbln1 alleviated these phenotypic defects in Cbln1-KO mice (Ito-Ishida et al., 2008). Furthermore, administration of recombinant Cbln1 to cultured Cbln1-KO cerebellar neurons induced the formation of PF protrusions, which often form enlarged circular structures encapsulating PC spines, facilitated the accumulation of synaptic vesicles and GluD2, and promoted the maturation of PF–PC synapses (Ito-Ishida et al., 2012). Therefore, rapid regeneration of PF wiring is likely mediated by the interaction of GluD2 with Cbln1 released from surviving and regenerating PFs, as in cerebellar development (Yuzaki, 2009; Watanabe and Kano, 2011). Without GluD2, this interaction is disrupted and PF wiring remains in the degenerative phase.

Expansion of the PF territory in the degenerative phase

Robust expansion of the PF territory occurred in the degenerative phase in both strains of mice, whereas the subsequent territorial retraction occurred only in wild-type mice. Recent studies provide insight into how these territorial changes might be produced. In normal development, PF and CF territories on PC dendrites are highly overlapping until postnatal day 15 in mice, and then segregated through the elimination of PF synapses from proximal dendrites (Ichikawa et al., 2016). This territorial segregation is arrested in mutant mice defective in the type-1 metabotropic glutamate receptor (mGluR1) or the P/Q-type voltage-dependent Ca2+ channel CaV2.1 (Miyazaki et al., 2004, 2012; Ichikawa et al., 2016). In PCs, mGluR1 is readily activated by PF stimulation and induces inositol 1,4,5-trisphosphate-mediated Ca2+ release in local dendrites (Finch and Augustine, 1998; Takechi et al., 1998), whereas CaV2.1 is activated upon strong depolarization by CF

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**Figure 8.** Changes after PF transection in the path length of innervation territories and in the number of PF synapses, CF synapses, free spines, and spine-type GABAergic synapses. A, The path length (μm) of whole single dendritic tracts and of PCD-I (blue), PCD-II (cyan), and PCD-III (green). B, The path length of the PF territory (ie, PCD-I + PCD-II). C, PF synapse number in the PF territory. D, The path length of the CF territory (ie, PCD-I – PCD-II). E, CF synapse number in the CF territory. F, The path length of dendrites studded with free spines. G, Free spine number. H, Spine-type GABAergic synapse number. Left and right bars represent scores in wild-type and GluD2-KO mice, respectively. All scores were measured from three mice at each time point for each analysis, and expressed as the mean ± SD. Differences between sham-operated control and PF-transected mice in each genotype were assessed using ANOVA with Dunnett’s test. **p < 0.01, *p < 0.05 (actual p values are indicated in the parenthesis).
firing and mediates global Ca\(^{2+}\) influx in the whole dendritic tree (Kano et al., 1992; Konnerth et al., 1992; Regehr and Mintz, 1994). Ectopic PF innervation of proximal dendrites also occurs following deprivation of CF input to PCs (Desclin and Escubi, 1974; Sotelo et al., 1975; Rossi et al., 1991; Cesa et al., 2003, 2005) or blockade of electrical activity in the cerebellar cortex (Bravin et al., 1999; Morando et al., 2001; Cesa et al., 2003, 2007). These findings suggest that PF and CF activities inducing local and global Ca\(^{2+}\) dynamics in PC dendrites work in concert to promote PF synapse elimination from proximal dendrites. Therefore, expansion of the PF territory in the degenerative phase might be caused by a lack of mGluR1

![Image of dendritic domains and synapse regulation phases]

Figure 9. Summary schematic illustrating the three distinct phases of regenerative PF wiring in wild-type mice (top) and the arrested regeneration in GluD2-KO mice (bottom). Three dendritic domains are colored in blue for PCD-I, cyan for PCD-II, and green for PCD-III. In wild-type mice, PF axons and synapses massively degenerate after PF transection. In this degenerative phase, numerous free spines emerge and the PF territory expands aberrantly. Then, PF synapses enlarge, and there is an increase in the contact ratio with PC spines (to 1:2) in the hypertrophic phase. In this phase, the number and density of PF synapses nearly recover to normal levels, and free spines nearly completely disappear, but PF density remains low. In the remodeling phase, PF density increases, at least partly through sprouting and elongation of PF collaterals, and PF synapses reacquire a compact structure and a 1:1 contact ratio with PC spines. This suggests that competitive synaptic wiring by PFs and CFs onto PCs is regained in wild-type mice. In contrast, in the GluD2-KO mice, the degenerative phase persists, without a shift to the hypertrophic or remodeling phases of PF–PC regenerative rewiring.
activation resulting from the massive loss of PF synapses in both strains of mice. The subsequent retraction of the PF territory in wild-type mice may be triggered by the restoration of mGlur1 activation following PF synapse regeneration.

High regenerative capacity of PF wiring

Although axotomy of adult central neurons induces sprouting from proximal axon stumps and upregulation of growth-associated genes, these initial responses are transitory and are rapidly followed by growth arrest and the formation of retraction bulbs (Maier and Schwab, 2006). The major obstacle to regeneration is the presence of neurite growth inhibitory factors and molecules in the environment in the adult CNS, several of which are upregulated after injury, and associated with myelin and glial scars (Fawcett et al., 2012). In comparison, PFs are unmyelinated axons, and oligodendrocytes and astrocytes that form myelin and glial scars are lacking in the molecular layer of the cerebellum (Palay and Chan-Palay, 1974). Instead, the Bergmann glia, specialized astrocytes in the molecular layer, thoroughly enwrap the somatodendritic elements of PCs and support their metabolism and function (Yamada and Watanabe, 2002). These unique environments in the cerebellar molecular layer may also help GluD2 work for rapid and effective regeneration of PF–PC wiring. Therefore, our findings shed light on the role of synaptic adhesion systems in the regeneration of synaptic connection in the adult brain.

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