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Enriched Expression of GluD1 in Higher Brain Regions and Its Involvement in Parallel Fiber–Interneuron Synapse Formation in the Cerebellum

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Of the two members of the δ subfamily of ionotropic glutamate receptors, GluD2 is exclusively expressed at parallel fiber–Purkinje cell (PF–PC) synapses in the cerebellum and regulates their structural and functional connectivity. However, little is known to date regarding cellular and synaptic expression of GluD1 and its role in synaptic circuit formation. In the present study, we investigated this issue by producing specific and sensitive histochemical probes for GluD1 and analyzing cerebellar synaptic circuits in GluD1-knock-out mice. GluD1 was widely expressed in the adult mouse brain, with high levels in higher brain regions, including the cerebral cortex, striatum, limbic regions (hippocampus, nucleus accumbens, lateral septum, bed nucleus stria terminalis, lateral habenula, and central nucleus of the amygdala), and cerebellar cortex. In the cerebellar cortex, GluD1 mRNA was expressed at the highest level in molecular layer interneurons and its immunoreactivity was concentrated at PF synapses on interneuron somata. In GluD1-knock-out mice, the density of PF synapses on interneuron somata was significantly reduced and the size and number of interneurons were significantly diminished. Therefore, GluD1 is common to GluD2 in expression at PF synapses, but distinct from GluD2 in neuronal expression in the cerebellar cortex; that is, GluD1 in interneurons and GluD2 in PCs. Furthermore, GluD1 regulates the connectivity of PF–interneuron synapses and promotes the differentiation and/or survival of molecular layer interneurons. These results suggest that GluD1 works in concert with GluD2 for the construction of cerebellar synaptic wiring through distinct neuronal and synaptic expressions and also their shared synapse-connecting function.

Key words: Cerebellum; GluD1; GluRδ1; glutamate receptor; mouse; synapse

Introduction

The δ subfamily of ionotropic glutamate receptors shares 17–28% amino acid identity with other subfamilies and consists of GluD1 (GluRδ1) and GluD2 (GluRδ2) (Yamazaki et al., 1992; Lomeli et al., 1993). Despite well conserved membrane topology and amino acid residues critical to ligand binding and Ca²⁺ permeability, the δ subfamily do not function as conventional glutamate-gated ion channels (Hirai et al., 2005; Kakegawa et al., 2007a, 2007b; Ady et al., 2014). Essential roles of GluD2 in synaptic development and plasticity have been established through intensive analyses on GluD2-defective mice (Yuzaki, 2009; Watanabe and Kano, 2011).

GluD2 is expressed exclusively at parallel fiber (PF) synapses on cerebellar Purkinje cells (PCs) (Takayama et al., 1996; Landsend et al., 1997) and weakly in PF synapses on interneuron dendrites (Yamasaki et al., 2011). GluD2 plays a key role in the formation and maintenance of PF–PC synapses (Guastavino et al., 1990; Kashiwabuchi et al., 1995; Kurihara et al., 1997; Lalouette et al., 2001; Takeuchi et al., 2005); this is mediated through selective binding of the N-terminal domain of GluD2 with presynaptic neurexs via granule cell-derived Cbln1 (Matsuda et al., 2010; Uemura et al., 2010). GluD2 is also essential for induction of cerebellar long-term depression at PF–PC synapses (Kashiwabuchi et al., 1995), through interaction of the C-terminus with megakaryocyte protein tyrosine phosphatase (Kohda et al., 2013), and through binding of D-serine to the ligand-binding site (Hirai et al., 2003; Kakegawa et al., 2011; Kohda et al., 2013). Mutation of the GluD2 gene in mice and humans (GRID2 and GRID2, respectively) causes cerebellar ataxia (Kashiwabuchi et al., 1995; Hills et al., 2013; Utine et al., 2013).
In comparison, GluD1 is transcribed weakly in various brain regions (Lomeli et al., 1993). The functional significance of GluD1 is inferred from human genetic studies reporting that the GRID1 is a strong candidate gene for schizophrenia, bipolar disorder, major depressive disorder, and autism spectrum disorder (Fallin et al., 2005; Guo et al., 2007; Glossner et al., 2009; Smith et al., 2009; Treutlein et al., 2009; Cooper et al., 2011; Greenwood et al., 2011; Edwards et al., 2012 Griswold et al., 2012). GluD1 knock-out (KO) mice also exhibit deficits in emotional and social behaviors and in learning and memory (Yadav et al., 2012, 2013). Like GluD2, GluD1 binds to neurexins via the Cnb family and their interaction induces synaptogenesis in vitro (Matsuda et al., 2010, Yasumura et al., 2012; Ryu et al., 2012). These findings suggest that GluD1 is involved in higher brain functions through controlling synaptic connectivity and plasticity. However, fundamental information on cellular and synaptic expression of GluD1 and its functional role, which is essential to link the genetic level to the behavioral and disorder level, is missing to date.

In the present study, we report that GluD1 is widespread in adult mouse brains, with abundant expression in the cerebral cortex, striatum, limbic regions, and cerebellar cortex. In the cerebral cortex, GluD1 is enriched at PF synapses on molecular layer interneurons and GluD1-KO mice manifested significant losses of interneurons and PF-interneuron synapses.

### Materials and Methods

**Animals.** In the present study, we used adult C57BL/6 mice, GluD1-KO mice (Gao et al., 2007), and GluD2-KO mice (Yamasaki et al., 2011) at 2–3 months of age. In each experiment, we used three male mice for qualitative and quantitative analyses. All experiments were performed according to the guidelines laid down by the animal welfare committees of Hokkaido University, Keio University, and Niigata University.

**Antibodies.** Primary antibodies raised against the following molecules were used: microtubule-associated protein 2 (MAP2), vesicular glutamate transporter (VGluT) 1, VGluT2, parvalbumin, and bassoon. In the present study, we produced GluD1 and GluD2 antibodies. For expression of glutathione S-transferase fusion proteins, we subcloned cDNA fragments encoding the corresponding C-terminal regions of mouse GluD1 (895–932 aa residues) and GluD2 (897–934 aa residues) into the BamHI/EcoRI site of pGEX4T-2 plasmid (GE Healthcare). Immunization and affinity purification were performed as described previously (Watanabe et al., 1998). Information on the antigen, host species, source, reference, and specificity is summarized in Table 1.

**Immunoblot.** To confirm the specificity of GluD1 and GluD2 antibodies, human embryonic kidney 293 (HEK293) cells (CRL-1573; American Type Culture Collection) were transfected with 10 μg of pCAGGS vector (provided by Dr. J. Miyazaki, Tohoku University, Sendai, Japan) carrying human influenza hemagglutinin (HA)-tagged GluD1 or GluD2 cDNA using the calcium phosphate method. HA CDNA was added immediately downstream of the signal sequence of GluD1 or GluD2. HEK293 cells 36–48 h after transfection or mouse brain regions were solubilized in TNE buffer (50 mM NaF, 1% NP-40, 20 mM EDTA, 1 μM peptatin A, 2 μg/ml leupeptin, 10 μg/ml aprotinin, and 50 mM Tris- HCl, pH 8.0) with 0.1% SDS. The soluble fractions were analyzed by immunoblotting with 1 μg/ml of GluD1 and GluD2 antibodies or with HA antibody (mouse, 1:1000; Covance Research Products).

The cerebellum, hippocampus, and cerebral cortex were excised and homogenized in homogenate buffer (0.32 M sucrose, 5 mM EDTA, 5 mM HEPES-NaOH, pH 7.4. complete protease inhibitor mixture tablet; Roche) and centrifuged at 1000 x g for 10 min to obtain the S1 (homogenate) fraction. Synaptosomal and postsynaptic density (PSD) fractions were prepared according to the method used by Carlin et al. (1980). Protein samples of each fraction were submitted to SDS-PAGE for immunoblotting as described previously ( Abe et al., 2004). Signal intensities of immunoreacted bands were determined using an Ez-Capture MG (ATTO) and their statistical differences were evaluated using a Student’s unpaired t test with Welch’s correction.

**Immunocytochemistry.** In immunocytochemical experiments, transfected HEK293 cells were fixed with 4% paraformaldehyde for 10 min at room temperature and incubated with a blocking solution (2% goat serum, 2% BSA, and 0.4% Triton X-100 in PBS) for 1 h. Cells were then incubated with mouse HA antibody (1:1000) and with GluD1 or GluD2 antibody (1 μg/ml) for 1 h. To visualize the bound primary antibody, the cells were incubated with Alexa Fluor 546- or Alexa Fluor 488-conjugated secondary antibodies (1:2000) to mouse, guinea pig, or rabbit IgG (Invitrogen).

**Fixation and sections.** Under deep pentobarbital anesthesia (100 mg/kg body weight, i.p.), mice were fixed by transcardial perfusion with 4% paraformaldehyde (PFA) in 0.1 M phosphate buffer (PB; pH 7.2) for immunofluorescence (except for GluD1) and chromogenic in situ hybridization; 2% PFA/2% glutaraldehyde in 0.1 M sodium cacodylate buffer, pH 7.2, for conventional electron microscopy; and 1% PFA/0.1% glutaraldehyde in 0.1 M PB for postembedding immunogold electron microscopy.

For immunofluorescence, except for GluD1 and GluD2, 50-μm-thick sections were prepared on a microslicer (VT1000S; Leica Microsystems) and subjected to free-floating incubation. Perfused brains for chromogenic in situ hybridization were further postfixed for 3 d at room temperature, according to the method of Hioki et al. (2010), and cryoprotected with 30% sucrose in 0.1 M PB; 30 μm cryosections were then prepared on a cryostat (CM1900; Leica Microsystems). For fluorescence in situ hybridization (FISH) and immunofluorescence for GluD1, brains were freshly obtained under deep pentobarbital anesthesia and immediately frozen in powdered dry ice for preparation of fresh frozen sections (4 μm). Before incubation for FISH in situ hybridization, fresh frozen sections were air-dried and fixed by dipping in 4% PFA in 0.1 x PB for 15 min. For GluD2 immunofluorescence, paraffin sections (4 μm; SM1000R; Leica Microsystems) mounted on silane-coated glass slides were subjected, before immunohistochemical reaction, to pepsin pretreatment for antigen exposure, as described previously (Watanabe et al., 1998).

For conventional electron microscopy, the straight portion of lobule 4/5 was excised, postfixed with 1% osmium tetroxide for 15 min, stained in block with 2% uranyl acetate for 30 min, dehydrated, and embedded in Epon812. For postembedding immunogold electron microscopy, microslicer sections (400 μm) were cyroprotected with 30% glycerol in PB and frozen rapidly with liquid propane in the EM CPC unit (Leica Microsystems). Frozen sections were immersed in 0.5% uranyl acetate in methanol at –90°C in the AFS freeze-substitution unit (Leica Microsystems), infiltrated at –45°C with Locryml HM-20 resin (Chemische Werke Lowi), and polymerized with ultraviolet light. Ultrathin sections were made using an Ultracut ultramicrotome (Leica Microsystems).

**In situ hybridization.** Mouse cDNA fragments of GluD1 (nucleotides 63–3092 bp; GenBank accession number, NM_008166), VGlut1 (301–1680, BC054462), and 67 kDa glutamic acid decarboxylase (GAD67, 1035–2015; NM_008077) were subcloned into the pBluescript II plasmid vector. Digoxigenin (DIG)- or fluorescein-labeled cRNA probes were

### References

Table 1. Primary antibodies used in the present study

<table>
<thead>
<tr>
<th>Molecule</th>
<th>Sequence (NCBI #)</th>
<th>Host</th>
<th>Specificity</th>
<th>Reference/source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bassoon</td>
<td></td>
<td>Ms</td>
<td>Enzo Life Sciences (SARTI407)</td>
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<tr>
<td>GluD1</td>
<td>895–932aa (NM_008166)</td>
<td>Rabbit, GP</td>
<td>KO</td>
<td>Present study</td>
</tr>
<tr>
<td>GluD2</td>
<td>897–934aa (NM_008167)</td>
<td>Rabbit, GP</td>
<td>KO</td>
<td>Present study</td>
</tr>
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<td>MAP2</td>
<td>927–1104 (NM008632)</td>
<td>Goat</td>
<td>Go</td>
<td>*Miyazaki et al. (2006)</td>
</tr>
<tr>
<td>HA</td>
<td>11–11 (clone 16E2)</td>
<td>Mouse</td>
<td>Molecule Sequence (NCBI #)</td>
<td></td>
</tr>
<tr>
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<td>1–100aa (NM_01645)</td>
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<td>GP</td>
<td>IB</td>
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<tr>
<td>VGluT1</td>
<td>531–560aa (BC054462)</td>
<td>Guinea pig</td>
<td>IB</td>
<td>Miyazaki et al., 2003</td>
</tr>
<tr>
<td>VGluT2</td>
<td>559–582aa (BC03875)</td>
<td>Guinea pig</td>
<td>GP</td>
<td>Miyazaki et al., 2003</td>
</tr>
</tbody>
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Go, Goat polyclonal antibody; GP, guinea pig polyclonal antibody; IB, immunoblot with brain homogenates; KO, lack of immunohistochemical or immunoblot labeling in knockout mice; Ms, mouse monoclonal antibody; Rab, rabbit polyclonal antibody.

*The specificity of the MAP2 antibody is supported by selective labeling of somatodendritic neuronal elements (see the reference).
transcribed in vitro for chromogenic and FISH (Yamasaki et al., 2010). Fragmentation of riboprobes by alkaline digestion was omitted to increase the sensitivity and specificity. For immunohistochemical detection of DIG and fluorescein, sections were blocked with DIG blocking solution [TNT buffer containing 1% blocking reagent (Roche Diagnostics) and 4% normal sheep serum] for 30 min and 0.5% TSA blocking reagent (PerkinElmer) in TNT buffer for 30 min. Sections were then incubated with either alkaline-phosphatase-conjugated sheep anti-DIG (1:500, 1.5 h; Roche Diagnostics) for chromogenic detection or peroxidase-conjugated anti-DIG (1:1000, 1 h; Roche Diagnostics) or anti-fluorescein antibody (1:1500, 1 h; Invitrogen) for fluorogenic detection. After two washes in TNT buffer for 15 min each, chromogenic detection was performed using nitro-blue tetrazolium and 3-bromo-4-chloro-3′-indolylphosphate (1:50; Roche Diagnostics) in detection buffer (0.1 M Tris-HCl, pH 9.5, 0.1 M NaCl, and 50 mM MgCl₂) for 12 h. For double FISH, the first detection was performed with peroxidase-conjugated anti-fluorescein antibody, followed by incubation with the FITC-TSA Plus amplification kit (PerkinElmer). After inactivation of residual peroxidase activity by dipping sections in 1% H₂O₂ for 30 min, the second detection was performed by incubating sections in DIG-labeled cRNA probe, followed by peroxidase-conjugated anti-DIG antibody and the Cy3-TSA plus amplification kit (PerkinElmer). Sections were counterstained with TOTO-3 (1:50 in PBS, 20 min; Invitrogen). Images of chromogenic in situ hybridization were taken with a light microscope (BZ-9000; Keyence) and PlanApo (4×/0.20 and 10×/0.45) objective lenses (Nikon), and images of FISH were captured using confocal laser-scanning microscope as described in the section called Immunofluorescence.

Expression levels of GluD1 mRNA were semiquantitatively evaluated from FISH images taken using the gain level settings. The separate color components were converted to grayscale and the gray level (arbitrary units) and area were measured using MetaMorph software (Molecular Devices) from individual somata of GAD67 mRNA-positive neurons or from regions containing multiple GAD67 mRNA-negative/TOTO-3-labeled neurons (i.e., pyramidal cells in the hippocampus and granule cells in the cerebellum). In the latter, the gray level of a given region was divided by the number of TOTO-3-labeled neurons to calculate the mean relative intensity in each neuron type.

Immunofluorescence. All immunohistochemical incubations were performed at room temperature. Sections were incubated with 10% normal donkey serum for 20 min, a mixture of primary antibodies overnight (1:1000 dilution for bassoon and 1 μg/ml for others), and a mixture of Alexa Fluor 488-, Cy3-, or Alexa Fluor 647-labeled species-specific secondary antibodies for 2 h at a dilution of 1:200 (Invitrogen; Jackson ImmunoResearch). Im-

Figure 1. In situ hybridization for GluD1 mRNA in the adult mouse brain. A, B, H, Chromogenic in situ hybridization for GluD1 in the whole brain (A), hippocampus (B), and cerebellum (H). Hybridization with a sense probe yielded no significant signal (A, inset). C, D, K, Fluorescent in situ hybridization for GluD1 (red) and VGlut1 (green) mRNAs in the hippocampus (C, D) and cerebellar cortex (K). Arrowheads and double arrowheads in K indicate molecular layer interneurons or Golgi cells, respectively. E, F, I, J, FISH for GluD1 (red) and GAD67 (green) mRNAs in the hippocampus (E, F) and cerebellar cortex (I, J). Arrows in F and J indicate interneurons that express GAD67 mRNA. Arrowheads and double arrowheads in I indicate molecular layer interneurons or Golgi cells, respectively. Asterisks indicate PC somata. G, I, Histograms showing the mean relative fluorescence intensity of GluD1 mRNA in hippocampal (G) and cerebellar (I) neurons. The intensity in each neuron type was normalized to the intensity in CA1 pyramidal cells in the hippocampus (n = 133 for CA1 pyramidal cells, n = 43 for CA2 pyramidal cells, n = 48 for CA3 pyramidal cells, n = 65 for dentate gyrus granule cells, and n = 53 for hippocampal interneurons) and to the intensity in molecular layer interneurons in the cerebellum (n = 53 for molecular layer interneurons, n = 49 for Golgi cells, n = 57 for PCs, and n = 243 for granule cells). CA1–3, CA1–3 regions of the Ammon’s horn; Ch, cerebellum; Cs, cortex; DIG, digoxigenin; GL, granular layer; GCL, granule cell layer; Hi, hippocampus; LM, stratum lacunaeum-molecularae; Mb, midbrain; ML, molecular layer; MO, medulla oblongata; Mo, molecular layer; OB, olfactory bulb; Or, striatum orien.; PCL, Purkinje cell layer; Py, pyramidal cell layer; Ra, stratum radiatum; Sc, striatum; Th, thalamus. Scale bars: A, 1 mm; B, C, E, H, 500 μm; D, F, I, J, K, 20 μm.
...saline, pH 7.4 (TTBS) for 10 min, blocking solution containing 2% normal goat serum (Nichirei) in TTBS for 10 min, GluD1 or GluD2 antibody (20 μg/ml for each) diluted in 2% normal goat serum in TTBS overnight, and colloidal gold-conjugated (10 nm) anti-guinea pig IgG (1:100; British BioCell International) in blocking solution for 2 h. For double labeling against VGluT2 and GluD1 or GluD2, sections were incubated in blocking solution containing 2% normal goat serum, followed by VGluT2 antibody (20 μg/ml) diluted in 2% normal goat serum in TTBS overnight and then colloidal gold-conjugated (15 nm) anti-rabbit IgG in blocking solution for 2 h. After extensive washing in distilled water, sections were incubated in blocking solution containing 2% rabbit serum (Nichirei) in TTBS for 10 min, GluD1 or GluD2 antibody diluted with 2% normal rabbit serum in TTBS overnight, and colloidal gold-conjugated (10 nm) anti-guinea pig IgG in blocking solution for 2 h. After extensive washing in distilled water, sections were fixed with 2% OsO4 for 15 min and then stained with 5% uranyl acetate/40% EtOH for 90 s and Reynold’s lead citrate solution for 60 s. Photographs were taken from the molecular layer with the H-7100 electron microscope. For quantitative analysis, postsynaptic-membrane-associated immunogold particles, defined as those <20 nm apart from the cell membrane, were counted on scanned electron micrographs and analyzed using MetaMorph software. Statistical significance was evaluated using the Mann–Whitney U test. The vertical distribution of GluD1 was examined by sampling synaptic profiles with presynaptic and postsynaptic membranes that were cut perpendicularly to the plane of the synaptic cleft and measuring the distance from the midline of the synaptic cleft to the center of the immunogold particles.

Results

GluD1 mRNA expression in the brain

We first examined regional and cellular expression of GluD1 mRNA in the adult mouse brain using in situ hybridization (Fig. 1). Use of a long antisense riboprobe and a long fixation protocol greatly enhanced the intensity of specific signals for GluD1 mRNA. Chromogenic in situ hybridization revealed higher levels in the olfactory bulb, hippocampus, cerebral cortex, striatum, and cerebellar cortex (Fig. 1A). In the hippocampus, GluD1 mRNA was intense in the pyramidal cell layer of the Ammon’s horn, especially the CA2 subregion, and in the granule cell layer of the dentate gyrus (Fig. 1B). In the cerebellar cortex, cells expressing GluD1 mRNA were scattered in the molecular and granular layers (Fig. 1H). Specificity was ascertained by the lack of hybridizing signals using a sense riboprobe (Fig. 1A, inset).

Neurochemical properties of GluD1 mRNA-expressing neurons were examined in the hippocampus and cerebellar cortex by double-labeling FISH. In the hippocampus, GluD1 mRNA (Fig. 1C–F, red) was detected in pyramidal and granule cells expressing VGluT1 mRNA (Fig. 1CD, green), and in GABAergic interneurons expressing GAD67 mRNA (Fig. 1E,F, green, arrows). The mean fluorescence intensity for GluD1 mRNA was calculated for each type of neuron and normalized to that in CA1 pyramidal cells (see Materials and Methods). The mean relative intensity...
was highest in CA2 pyramidal cells and lowest in hippocampal interneurons (Fig. 1G). In the cerebellar cortex, GluD1 mRNA (Fig. 1I–K, red) was detected in GABAergic interneurons expressing GAD67 mRNA (Fig. 1F,J, green), including basket and stellate cells in the molecular layer and Golgi cells in the granular layer (Fig. 1I,J). Signals for GluD1 mRNA were very low in PCs (Fig. 1J, asterisks), and undetectable in granule cells expressing VGluT1 mRNA (Fig. 1K). The intensity of fluorescent signals was measured and normalized to that in molecular layer interneurons. The mean relative intensity varied greatly from the highest level in molecular layer interneurons to the lowest or background level in granule cells (Fig. 1L). Therefore, GluD1 is widely expressed in the adult mouse brain, but its transcription levels are variable according to neuron type.

Production of specific GluD antibodies
In the present study, we produced GluD1 and GluD2 antibodies against the corresponding C-terminal regions with low sequence homology. Immunoblotting experiments showed that rabbit and guinea pig GluD1 antibodies selectively recognized multiple protein bands at 110–130 kDa in HEK293 cells transfected with HA-tagged GluD1 cDNA, but not GluD2 cDNA (Fig. 2A). Likewise, guinea pig GluD2 antibody selectively recognized protein bands at similar molecular masses in HEK293 cells transfected with HA-tagged GluD1 cDNA, but not GluD2 cDNA (Fig. 2A). These GluD1 and GluD2 antibodies yielded selective immunocytochemical labeling for HEK293 cells transfected with HA-tagged GluD1 or GluD2 cDNA, respectively (Fig. 2B).

The specificity was further tested by immunoblot using homogenates prepared from the hippocampus, cerebral cortex, and cerebellum of wild-type, GluD1-KO, and GluD2-KO mice (Fig. 2C). In each brain region, rabbit and guinea pig GluD1 antibodies recognized protein bands in wild-type mice, but not GluD1-KO mice. Likewise, guinea pig GluD2 antibody recognized specific protein bands in all three brain regions of wild-type mice, but not in GluD2-KO mice. Of note, low levels of GluD2 were observed in the hippocampus and cerebral cortex in addition to the cerebellum. Therefore, these GluD antibodies are sensitive and specific to respective GluD.

GluD1 distribution in the brain
In conventional immunofluorescence using 4% PFA-perfused brain sections, the intensity of GluD1 immunolabeling was low (data not shown). When using fresh-frozen sections, GluD1 staining was markedly intensified all over the brains of wild-type mice (Fig. 3A). GluD1 immunoreactivity was distributed widely, with higher levels in the hippocampus, cerebral cortex, striatum, and cerebellar cortex (Fig. 3A). GluD1 was generally low in the brainstem. These protein expression patterns are consistent with the mRNA expression patterns (Fig. 1A). The absence of immunostaining in GluD1-KO mouse brains (Fig. 3A, inset) verified the specificity of GluD1 immunohistochemistry, including the lack of cross reactivity to GluD2.

Using coronal brain sections, regional distribution of GluD1 was investigated in detail. Laminar distribution was remarkable in the hippocampus (Fig. 3C). GluD1 was enriched in the stratum lacunosum-moleculare of the Ammon’s horn and in the molecular layer of dentate gyrus (Fig. 3C). AC, anterior commissure; CA1–3, CA1–3 regions of the Ammon’s horn; DG, dentate gyrus; Or, granule cell layer; LV, lateral ventricle; Mb, midbrain; MO, medulla oblongata; OB, olfactory bulb; Or, stratum oriens; Py, pyramidal cell layer; Ra, stratum radiatum; Th, thalamus. Scale bars: A, 1 mm; B, C, 500 μm.

Selective expression of GluD1 at PF–interneuron synapses
GluD2 has been shown to accumulate in the cerebellar molecular layer, where it is high at PF synapses on PC spines and
weak at PF synapses on interneuron dendrites (Takayama et al., 1996; Landsend et al., 1997; Yamasaki et al., 2011). In the present study, we selected this region for detailed analysis of synaptic localization of GluD1.

Immunofluorescence for GluD1 exhibited preferential interneuron labeling in the molecular layer, showing a basal-to-superficial gradient of immunofluorescence intensity (Fig. 4A, arrows). This staining was not detected in GluD1-KO mice (Fig. 4A, inset), indicating the specificity of the interneuron labeling. At high magnifications, GluD1 formed basket-like clustering around somata of molecular layer interneurons. To determine whether perisomatic clusters of GluD1 were associated with particular synapses, we applied triple immunofluorescence for GluD1 (Fig. 4B,C, red), presynaptic active zone protein bassoon (Fig. 4B,C, green), and PF terminal marker VGluT1 (Fig. 4B,C, blue). In both the superficial (Fig. 4B) and basal (Fig. 4C) molecular layer, where stellate and basket cells reside, respectively (Palay and Chan-Palay, 1974), perisomatic clusters of GluD1 were apposed to bassoon-positive/VGluT1-positive PF terminals, with GluD1 being always located inside these terminals (Fig. 4B,C, arrowheads). PF synapse-associated clustering was also true for GluD1-positive puncta in the neuropil because they were distributed around MAP2-positive dendrites of molecular layer interneurons (Fig. 4D, green) and apposed to VGluT1-positive PF terminals (Fig. 4E, blue). These results suggest that GluD1 accumulates at PF synapses on somata and dendrites of molecular layer interneurons.

Reciprocal distribution of GluD1 and GluD2 among PF synapses

To confirm GluD1 expression at PF-interneuron synapses and to compare it quantitatively with expression of GluD2, postembedding immunogold electron microscopy was applied to the molecular layer. Immunogold particles for GluD1 often fell on the postsynaptic membrane at PF-InS synapse, which were defined as asymmetrical synapses between small round terminals and interneuron somata (Fig. 5A). Compared with PF-InS synapses, immunogold labeling was less frequent at PF synapses on interneuron dendrites (PF-InD synapse), which were defined as asymmetrical synapses on aspiny dendrites (Fig. 5B). The postsynaptic membrane at the corresponding synapses in GluD1-KO mice was rarely labeled (Fig. 5A,B, insets). The density of immunogold labeling per 1 μm² of the postsynaptic membrane was significantly higher at PF-InS and PF-InD synapses in wild-type mice than the background level, as determined from the corresponding synapses in GluD1-KO mice (p < 0.001, 0.01, or 0.05, respectively, Mann–Whitney U test; Fig. 5E). Moreover, the density of immunogold labeling was apparently higher at PF-InS synapses than at PF-InD synapses in both the basal and superficial molecular layer of wild-type mice, although statistical significance was noted in the basal molecular layer only (p < 0.001; Fig. 5E).

In contrast, synaptic labeling for GluD2 was overwhelming at PF–PC synapses (p < 0.001; Fig. 6C,E), and was also detected at a much lower level at PF–InD synapses (p < 0.001; Fig. 6B,E). However, no significant labeling was observed at PF–InS synapses (Fig. 6A,E) or CF–PC synapses (Fig. 6D,E). Therefore,
GluD1 and GluD2 are selective to PF synapses and are expressed reciprocally among the three kinds of PF synapses.

**Opposing changes in PF–InS synapse density in GluD1-KO and GluK2-KO mice**

GluD2 strengthens the connectivity of PF–PC synapses, and the ablation of the Grid2 gene in mice results in reduced PF–PC synapse number per PC to half the level found in wild-type mice (Kashiwabuchi et al., 1995; Kurihara et al., 1997). To address whether GluD1 plays a similar role, we analyzed PF–InS synapses in wild-type, GluD1-KO, and GluD2-KO mice by triple immunofluorescence for bassoon (green), VGluT1 (red), and parvalbumin (blue; Fig. 7A–C). Although parvalbumin is expressed in both interneurons and PCs, they are readily distinguished by the size and location of somata (Fig. 8A–C). In wild-type mice, parvalbumin-positive somata of molecular layer interneurons were frequently contacted by bassoon-positive/VGluT1-positive PF terminals (Fig. 7A, arrowheads). Such PF terminals attaching to interneuron somata were apparently reduced in GluD1-KO mice (Fig. 7B) but increased in GluD2-KO mice (Fig. 7C).

To quantify this difference, the number of PF–InS synapses on somatic profiles of molecular layer interneurons was counted by conventional electron microscopy (Fig. 7D–G). The mean number of PF–InS synapses per 1 μm of somatic membrane was reduced by 47% in GluD1-KO mice (Fig. 7E,G) and increased by 60% in GluD2-KO mice (Fig. 7F,G) compared with wild-type mice (Fig. 7D,G). A Kruskal–Wallis test revealed a significant effect of group (p < 0.001). A post hoc test using Mann–Whitney U test with Bonferroni correction showed significant differences between wild-type and GluD1-KO mice (p < 0.001) or GluD2-KO mice (p < 0.001) (Fig. 7G). Therefore, the density of PF–InS synapses is altered in opposite directions in GluD1-KO and GluD2-KO mice compared with wild-type mice.

**Figure 5.** Postembedding immunogold electron microscopy for GluD1 in the cerebellar molecular layer. A–C, Single-labeling postembedding immunogold for GluD1 (ϕ = 10 nm) at a PF–InS synapse (A), a PF–InD synapse (B), and a PF–PC synapse (C). Double-labeling postembedding immunogold for GluD1 (ϕ = 10 nm) and VGluT2 (ϕ = 15 nm) at a CF–PC synapse. Insets in A–D indicate negative labeling at the corresponding synapses in GluD1-KO mice. Arrows and arrowheads indicate immunogold labeling or the edge of the PSD, respectively. E, Histogram showing the mean density of immunogold particles for GluD1 (per 1 μm of the postsynaptic membrane). The density was 4.53 ± 0.72 (n = 71, basal) at PF–InS synapses, 1.75 ± 0.33 (n = 69, basal) at PF–InD synapses, 3.96 ± 1.33 (n = 26, superficial) at PF–InS synapses, 1.13 ± 0.42 (n = 41, superficial) at PF–InD synapses, 0.56 ± 0.17 at PF–PC synapses (n = 89), and 0.10 ± 0.10 (n = 34) at CF–PC synapses of wild-type mice. In GluD1-KO mice, the density was 0.35 ± 0.25 (n = 36, basal) at PF–InS synapses, 0.42 ± 0.42 (n = 36, basal) at PF–InD synapses, 0.34 ± 0.34 (n = 15; superficial) at PF–InS synapses, 0.18 ± 0.18 (n = 20; superficial) at PF–InD synapses, 0.23 ± 0.23 at PF–PC synapses (n = 22), and 0.19 ± 0.19 (n = 20) at CF–PC synapses. Bars on each column represent SEM. ***p < 0.001; **p < 0.01; *p < 0.05; N.S. not significant; Mann–Whitney U test. F, Histogram showing the vertical distribution of GluD1 epitope at PF–InS synapses sampled from the basal molecular layer. The distance was measured from the midline of the synaptic cleft to the center of immunogold particles (n = 114 particles from two C57BL/6 mice). On the x-axis, − and + represent presynaptic or postsynaptic side, respectively, from the midline of the synaptic cleft. Sp, Spine. Scale bars, 200 nm.
Opposing changes in interneuron density in GluD1-KO and GluD2-KO mice

In addition, we noticed alterations in the number of parvalbumin-positive interneurons in the molecular layer. Compared with wild-type mice, parvalbumin-positive interneurons appeared to be scarce in GluD1-KO mice but numerous in GluD2-KO mice (Fig. 8A–C). The density of parvalbumin-positive interneurons in the molecular layer was significantly reduced by 35% in GluD1-KO mice and increased by 76% in GluD2-KO mice compared with wild-type mice (Kruskal–Wallis test, \(p < 0.001\); Mann–Whitney U test with Bonferroni correction, \(p < 0.001\) for each; Fig. 8D). This opposing change did not result merely from downregulation or upregulation of parvalbumin expression because the number of GAD67 mRNA-expressing interneurons was also changed in parallel with parvalbumin-positive interneurons (Fig. 8F–H).

We further noted diminished somatic size of interneurons in GluD1-KO mice (Fig. 8B). Compared with wild-type mice, the mean somatic area of parvalbumin-positive interneurons was significantly reduced by 24% in GluD1-KO mice (Kruskal–Wallis test, \(p < 0.001\); post hoc test using Mann–Whitney U test with Bonferroni correction, \(p < 0.001\); Fig. 8E), whereas no significant difference was found in GluD2-KO mice (Mann–Whitney U test with Bonferroni correction, \(p > 0.05\); Fig. 8E). These results suggest that GluD1 promotes the differentiation and/or survival of molecular layer interneurons, whereas GluD2 counteracts this phenomenon.

Compensatory upregulation of GluD1 in GluD2-KO mice

To understand the opposing changes by the ablation of GluD1 and GluD2, we pursued possible compensatory changes of one GluD expression in mutant mice lacking another GluD. In GluD2-KO mice, GluD1 labeling was clearly enhanced on somata and dendrites of molecular layer interneurons; this enhancement was confirmed in all three pairs of wild-type and GluD2-KO mice (Fig. 9A, B). By measuring the intensity of immunofluorescence signals in the molecular layer, the mean intensity for GluD1 was significantly increased by 54% in GluD2-KO mice compared with wild-type mice (\(p < 0.001\), Mann–Whitney U test; Fig. 9E, left). In contrast, no significant upregulation or downregulation was discerned for GluD2 in the molecular layer of GluD1-KO mice (\(p > 0.05\); Fig. 9C,D, 9E, right).
This immunohistochemical finding was in parallel with immunoblot data using cerebellar samples prepared from wild-type, GluD1-KO, and GluD2-KO cerebella (Fig. 9F). The intensity of GluD1 bands in GluD2-KO cerebella was significantly elevated in the homogenate (by 48% compared with wild-type cerebella; \( p < 0.01 \), Student’s t test with Welch’s correction, \( n = 3 \)), synaptosome fraction (by 64%; \( p < 0.01 \)), and PSD fraction (by 79%; \( p < 0.001 \); Fig. 9G, top). No significant increase or decrease was found for GluD2 bands in any cerebellar samples prepared from GluD1-KO cerebella (\( p > 0.05 \) for each, \( n = 3 \); Fig. 9G, bottom). Therefore, ablation of the Grid2 gene in mice significantly upregulates GluD1 expression in molecular layer interneurons.

Discussion

Although cellular and synaptic expressions of GluD2 and its regulatory mechanisms for synaptic development and plasticity have been well characterized (Yuzaki, 2009; Watanabe and Kano, 2011), those for GluD1 remain largely elusive. In the present study, we investigated this issue using expression analyses on GluD1 and neuroanatomical analyses on the cerebellar molecular layer of GluD1-KO mice. Here, we have demonstrated that GluD1 is distinct from GluD2 in regional, neuronal, and synaptic expressions in adult mouse brains and that specific types of synapses and neurons expressing GluD1 are significantly lost in GluD1-KO mice.

Enriched expression of GluD1 in higher brain regions

Using in situ hybridization with a radiolabeled oligonucleotide probe, Lomeli et al. (1993) first reported low and diffuse expression of GluD1 mRNA in adult rat brains, with the highest level in the hippocampus and transient elevation at neonatal stages. In the present study, we developed histochemical probes that can detect GluD1 mRNA and immunoreactivity with high sensitivity and specificity. As a result, GluD1 expression was clearly shown to be widespread in the adult mouse brain, with higher levels in the cerebral cortex, striatum, cerebellar cortex, and many regions of the limbic system, including the hippocampus, nucleus accumbens, lateral septum, bed nucleus stria terminalis, lateral habenula, and central nucleus of the amygdala.

The significance of GluD1 in a variety of brain functions has been inferred from animal and human studies. Aberrant

Figure 7. Opposing changes in the density of PF synapses on interneuron somata in GluD1-KO and GluD2-KO mice. A–C, Triple immunofluorescence for VGluT1 (red), bassoon (green), and parvalbumin (blue) in wild-type (A), GluD1-KO (B), and GluD2-KO mice (C). Arrowheads indicate VGluT1(+/+)/bassoon(+/+) puncta around parvalbumin(+/+) interneuron somata. D–F, Electron microscopic images of somata of molecular layer interneurons in wild-type (D), GluD1-KO (E), and GluD2-KO mice (F). Arrows indicate PF–InS synapses formed between interneuron somata (pseudocolored in green) and PF terminals (blue). G, Histogram showing the mean density of asymmetrical synapses on interneuron somata (per 1 \( \mu m \) of somatic membrane): 0.12 ± 0.01 synapses (\( n = 54 \) cells) in wild-type, 0.06 ± 0.01 (\( n = 40 \)) in GluD1-KO, and 0.18 ± 0.01 (\( n = 67 \)) in GluD2-KO mice. Bars on each column represent SEM. *** \( p < 0.001 \), Mann–Whitney U test. Scale bars: A–C, 5 \( \mu m \); D–F, 1 \( \mu m \).
Distinct synaptic targeting of GluD1

We selected the cerebellar molecular layer for the present analysis because this region has been best studied for synaptic expression and functional role of GluD2. In this region, GluD1 was enriched at PF synapses in the following order: PF–InS synapse > PF–InD synapse ≫ PF–PC synapse. This distribution was distinct from that of GluD2, which was expressed in the following order: PF–PC synapse ≫ PF–InD synapse > PF–InS synapse (Takayama et al., 1996; Landsend et al., 1997; Yamasaki et al., 2011; present study). In contrast, CF–PC synapses were at the background level for both GluD1 and GluD2. These expression properties indicate that GluD1 and GluD2 are similar in that they are selective to PF synapses in the cerebellar molecular layer, but their synaptic targeting is reciprocally regulated in target cell- and target site-dependent manners. Perisomatic baskets are formed around GABAergic molecular layer interneurons by excitatory PF terminals (Palay and Chan-Palay, 1974; Zanjani et al., 2012) and GluD1 was most enriched at PF–InS synapses forming this pericellular basket. This molecular-anatomical configuration is in a sharp contrast with that in the cortex and hippocampus, where perisomatic baskets are formed around glutamatergic principal neurons by terminals of inhibitory interneurons (Freund, 2003). This suggests that synaptic targeting of GluD1 is strategically controlled to construct distinct synaptic organization in specific brain regions and neurons.

Because previous studies have highlighted exclusive expression of GluD2 in the cerebellar cortex (Araki et al., 1993; Lomeli et al., 1993; Yamazaki et al., 2011), low but specific detection of GluD2 in the cortex and hippocampus was an unexpected finding in the present study. Recently, biallelic deletions of GRID2 gene have been identified in human families with severe cerebellar atrophy and symptoms (Hills et al., 2013; Utine et al., 2013). It is interesting to note that all of the affected individuals also showed delay in speech and cognitive development, implying the role of GluD2 in the cerebral cortex (Hills et al., 2013). Therefore, it is important in future studies to clarify how synaptic targeting of GluD1 and GluD2 is regulated in extracerebellar regions.

GluD1 strengthens synaptic connectivity

We found that the number of PF–InS synapses was significantly reduced by 47% in GluD1-KO mice, suggesting that GluD1 strengthens the connectivity of PF–InS synapses, as GluD2 does for PF–PC synapses (Kurihara et al., 1997; Takeuchi et al., 2005). It has been demonstrated that the connectivity of PF–PC synapses is mediated by binding of GluD2 on PC spines to neurexin I (Hirai et al., 2005; Ito-Ishida et al., 2008; Matsuda et al., 2010; Uemura et al., 2010). Cbln1 is produced and secreted from granule cells and accumulates in the

Figure 6. Opposing changes in the density of molecular layer interneurons between GluD1-KO and GluD2-KO mice and reduced somatic size in GluD1-KO mice. A–C, Immunofluorescence for parvalbumin in the cerebellar cortex of wild-type (A), GluD1-KO (B), and GluD2-KO mice (C). Asterisks indicate the soma of molecular layer interneurons. D, Histogram showing the mean density of parvalbumin-positive interneurons: 646.0 ± 24.3 cells/mm² (measured area, 1.36 mm²) in wild-type, 423.0 ± 16.6 (1.59) in GluD1-KO, and 1138.1 ± 42.8 (0.93) in GluD2-KO mice. E, Histogram showing the mean somatic area of parvalbumin-positive interneurons: 72.1 ± 1.7 μm² (n = 109 cells) in wild-type, 54.5 ± 1.1 (1.29) in GluD1-KO, and 70.2 ± 1.3 (n = 118) in GluD2-KO mice. Bars on each column represent SEM. ***p < 0.001, Mann–Whitney U test. F–H, Immunofluorescence for parvalbumin (green) combined with FISH for GAD67 mRNA (red) in wild-type (F), GluD1-KO (G), and GluD2-KO mice (H). Arrows indicate molecular layer interneurons. Scale bars, 20 μm.

emotional and social behaviors have been documented in GluD1-KO mice, including higher spontaneous locomotor activity in the open-field test, lower anxiety-like behavior in the elevated plus maze test, depression-like behavior in the forced swim test, robust aggression in the resident-intruder test, and deficits in the social interaction test (Yadav et al., 2012). The mice also manifest enhanced working memory and deficits in the fear conditioning test (Yadav et al., 2013). Human genome-wide association and copy number variation studies identified the GRID1 gene, which codes for GluD1, as a strong candidate gene for schizophrenia, bipolar disorder, major depressive disorder, and autism spectrum disorder (Fallin et al., 2005; Guo et al., 2007; Glessner et al., 2009; Smith et al., 2009; Treutlein et al., 2009; Cooper et al., 2011; Greenwood et al., 2011; Edwards et al., 2012; Griswold et al., 2012). Importantly, neural regions with rich GluD1 expression are the neural centers for cognition, memory and learning, emotion, voluntary movement, and motivation and rewarding. Therefore, detailed expression and phenotypic analyses in these higher brain regions are expected to provide important insights toward understanding the role of GluD1 in neural development and function.
synaptic cleft of PF synapses on PCs and molecular layer interneurons (Miura et al., 2009). Furthermore, Cbln1 also binds to GluD1 and their interaction induces synaptogenesis in vitro experiments (Matsuda et al., 2010, Yasumura et al., 2012; Ryu et al., 2012). These findings suggest that GluD1 strengthens the connectivity of PF–InS synapses by interacting with Cbln1 and neurexins. This possibility needs to be tested in future studies.

GluD1 promotes survival and differentiation of interneurons

Another phenotype in GluD1-KO mice was a significant (35%) loss of molecular layer interneurons. We speculate that this phenotype is secondary to reduced PF–InS synapses for the following reasons.

Neurotrophin signaling, particularly that mediated by brain-derived neurotrophic factor (BDNF) and the BDNF receptor TrkB, is indispensable for development of cerebellar interneurons (Lindholm et al., 1997; Schwartz et al., 1997; Carter et al., 2002). BDNF is highly expressed in granule cells, whereas high-affinity neurotrophin receptors are expressed in various cerebellar neurons, including molecular layer interneurons (Klein et al., 1990; Alvarez-Dolado et al., 1994; Gao et al., 1995; Segal et al., 1995; Benisty et al., 1998). BDNF is required for the survival, differentiation, and synaptogenesis of GABAergic interneurons in vitro (Mertz et al., 2000; Seil, 2003; Spatkovski and Schilling, 2003; Adcock et al., 2004) and ablation of the trkB gene in mice severely impairs synaptic differentiation in GABAergic interneurons (Rico et al., 2002). Therefore, we assume that GluD1-mediated formation and/or maintenance of PF–InS synapses increases both excitatory drive and neurotrophic support to molecular layer interneurons, thereby promoting their differentiation and survival. The lack of compensatory upregulation of GluD2 further supports that these phenotypes in GluD1-KO mice are primarily due to the loss of GluD1.

In our previous study, no significant decrease or increase was observed for the density of PF–InD synapses in GluD2-KO mice (Yamasaki et al., 2011). Therefore, it was unexpected to find significant increase of PF–InS synapses by 60% in GluD2-KO mice here. We assume that this synaptic increase is explained, at least partly, by compensatory upregulation of GluD1 at interneuron somata in GluD2-KO mice. With respect to unaltered density of PD–InD synapses in GluD2-KO mice (Yamasaki et al., 2011), compensatory upregulation at interneuron dendrites might also contribute to the maintenance of PD–InD synapses.

Functional implications

Molecular layer interneurons are the node of a feedforward inhibition from granule cells to PCs. This pathway is driven by PF activities and casts two modes of inhibition: GABAergic inhibition onto the soma and dendrites of PCs and electrical inhibition onto the axon initial segment of PCs (Korn and Axelrad, 1980; Fritschy et al., 2006; Iwakura et al., 2012; Blot and Barbour, 2014). Therefore, impaired PF–interneuron synapse formation in GluD1-KO mice should lead to substantial diminishment of feedforward inhibition to PCs. Conversely, the formation of PF–PC synapses that construct the major neural pathway and the maturation process leading to CF mono-innervation are severely disrupted in GluD2-KO mice (Kashiwabuchi et al., 1995; Kurihara et al., 1997; Ichikawa et al., 2002; Miyazaki et al., 2010). Together, GluD1 and GluD2 work in concert to construct the basic synaptic wiring in the cerebellar cortex through distinct neuronal and synaptic expressions and also through their shared synapse-connecting function.

**References**


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