A synthetic synaptic organizer protein restores glutamatergic neuronal circuits


INTRODUCTION: Synapses are fundamental structural and functional units within neural circuits, where they define the connectivity between neurons and provide avenues for communication. At a molecular level, synapses are highly dynamic and their remodeling is essential for all aspects of brain physiology. However, errors in this process can happen and often lead to an imbalance of excitatory and inhibitory signaling. This is thought to be a major cause of neuropsychiatric or neurological disorders, including autism spectrum disorders, epilepsy, schizophrenia, and Alzheimer's disease. Thus, molecular tools to control the number and/or function of synapses would be highly desirable. Physiologically, synapse formation is driven by synaptic organizer proteins. Among these, extracellular scaffolding proteins (ESPs) such as cerebellin-1 (Cbln1) and neuronal pentraxin-1 (NP1) are distinctive in that they could rapidly induce synapse differentiation by binding pre- and/or postsynaptic cell surface proteins at the synaptic cleft. We hypothesized that synthetic molecules that would combine structural features of Cbln1 and NP1 could be used to efficiently reverse the loss of excitatory synapses and promote the structural and functional recovery of damaged neuronal circuits in animal models of neurological disease.

RATIONALE: NP1 recruits postsynaptic AMPA-subtype ionotropic glutamate receptors (AMPARs), responsible for excitatory neurotransmission, through its pentraxin domain. However, NP1 does not seem to induce presynaptic specializations in vivo. By contrast, Cbln1 promotes presynaptic differentiation by interacting with the cell adhesion molecule neurexin (Nrx) through its N-terminal multimerization domain, but cannot bind AMPARs. Guided by structural information, we developed a hexameric synthetic soluble ESP, termed CPTX, which includes the multimerization domain of Cbln1 and the pentraxin domain of NP1. We hypothesized that CPTX should induce Nrx–CPTX–AMPAR transsynaptic molecular bridges, and thus accumulate and align presynaptic vesicle release machinery and postsynaptic neurotransmitter receptors.

RESULTS: Recombinant CPTX selectively bound presynaptic Nrx containing the spliced sequence 4[Nrx(+4)] with nanomolar affinity and most AMPAR subtypes with micromolar affinity. When administered to cerebellar granule cells and hippocampal neurons in vitro, CPTX acted as a bidirectional synapse organizer and induced excitatory pre- and postsynaptic sites. In vivo, CPTX increased the number of functional excitatory synapses and improved gait performance upon injection into the cerebellum of the ataxic Cbln1-null and GluD2-null mice. Furthermore, when injected into the hippocampus of 5xFAD mice, a model of familial Alzheimer's disease, CPTX restored dendritic spine numbers, excitatory synaptic transmission, and long-term potentiation and improved hippocampus-dependent learning. Finally, in mouse models of spinal cord injury, single injections of CPTX into the damaged tissue were sufficient to reorganize excitatory circuits and restore locomotion for more than 7 to 8 weeks.

CONCLUSION: We developed a synthetic, structure-guided, synaptic organizer termed CPTX, which induced functional and structural excitatory synapses in the cerebellar, hippocampal, and spinal cord neuronal circuits in vivo. Molecular components involved in excitatory synapses are considerably different among neuronal circuits. Rationally designed ESPs targeting distinct pre- and postsynaptic molecules may be useful to modulate neural circuit connectivity. This approach may inspire the development of a variety of innovative molecular tools for basic neuroscience as well as the treatment of neurological disorders.
A synthetic synaptic organizer protein restores glutamatergic neuronal circuits

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Neuronal synapses undergo structural and functional changes throughout life, which are essential for nervous system physiology. However, these changes may also perturb the excitatory–inhibitory neurotransmission balance and trigger neuropsychiatric and neurological disorders. Molecular tools to restore this balance are highly desirable. Here, we designed and characterized CPTX, a synthetic synaptic organizer combining structural elements from cerebellin-1 and neuronal pentraxin-1. CPTX can interact with presynaptic neurexins and postsynaptic AMPA-type ionotropic glutamate receptors and induced the formation of excitatory synapses both in vitro and in vivo. CPTX restored synaptic functions, motor coordination, spatial and contextual memories, and locomotion in mouse models for cerebellar ataxia, Alzheimer’s disease, and spinal cord injury, respectively. Thus, CPTX represents a prototype for structure-guided biologics that can efficiently repair or remodel neuronal circuits.

A broad range of neuropsychiatric and neurological disorders, including autism spectrum disorders, epilepsy, schizophrenia, and Alzheimer’s disease (AD), are thought to be caused by an imbalance between excitatory and inhibitory (E/I) synaptic functions (6–7). During organism development, but also throughout life, the formation and remodeling of complex yet precise neuronal circuits rely on specific synaptic organizing proteins (6, 7). These include cell adhesion molecules, such as neurexins (Nrxs), neureligins (8, 9), and receptor protein tyrosine phosphatases (10), as well as secreted proteins, such as fibroblast growth factors, semaphorins, Wnt, and extra-cellular scaffolding proteins (ESPs) (7, 11, 12). ESPs directly connect pre- and postsynaptic membrane proteins to form molecular bridges that span the synaptic cleft and mediate bidirectional signaling. For example, cerebellin-1 (Cbln1) is released from parallel fibers (PFs; axons of cerebellar granule cells) and contributes to the synapse-spanning tripartite complex Nrx–Cbln1–GluD2 (the ionotropic glutamate receptor family member delta-2) (Fig. 1A, left panel). Cbln1 can simultaneously bind Nrx isoforms containing the 30-residue “spliced sequence-4” (SS4) insert [Nrx(+4)] expressed at PF terminals and the amino-terminal domains (ATDs) of GluD2 on Purkinje cells (PCs) (13–15). Notably, a single injection of recombinant Cbln1 into the cerebellum can restore ~75% of PF–PC synapses and normal motor coordination within 1 day in adult Cbln1-null mice in vivo (16). Cerebells (Cbln1–4) are expressed in nearly all brain regions with distinct patterns and developmental dynamics (17, 18). Cbln1 promotes the formation or maintenance of excitatory synapses in the nucleus accumbens (19), whereas Cbln2 is necessary for excitatory synapses in the interpeduncular nucleus (20) and the hippocampus (21, 22). Cbln4 mediates the formation of inhibitory synapses between pyramidal neurons and cortical interneurons (23) by interacting postsynaptically with GluD1 (24), a receptor closely related to GluD2 (25). Other ESPs, such as the C1q-related CtgI, promote excitatory synaptic formation and maintenance in the cerebellum by binding to the Glu protein–coupled receptor BA13 (26, 27). CtgI3 also mediates synapse formation and maintenance in the medial prefrontal cortex (28). CtgII2 and CtgIII3 are produced by mossy fibers and recruit kainate-subtype ionotropic glutamate receptors (KARs) by directly binding to the ATDs of GluK2/4 KAR subunits in CA3 hippocampal neurons (29, 30). Molecular components of excitatory synapses differ considerably among neuronal circuits (31, 32), and other ESPs likely remain to be discovered. Nevertheless, the Cbln1–Cbln4 and CtgII1–CtgII3 examples prompted us to hypothesize that synthetic molecules with defined pre- and postsynaptic binding specificities could be designed and employed to reverse the loss of synapses and promote the structural and functional recovery of damaged neuronal circuits.

Design of a synthetic ESP-type synaptic organizer

Neuronal pentraxins (NPs) are a family of oligomeric secreted (NP1 and NP2) or membrane-attached (NPR) proteins that induce clustering of postsynaptic AMPA-type ionotropic glutamate receptors (AMPARs) through direct interactions between pentraxin (PTX) domains and the AMPAR ATDs (33–36). Astrocytosecreted glypican 4 indirectly leads to AMPAR recruitment by enhancing the release of NP1 from presynaptic axons (37). However, unlike Cbln1 (13, 16) and Cbln4 (23, 24), NPs do not seem to induce presynaptic specializations in vivo (38, 39). (Fig. 1A, middle panel). Thus, we hypothesized that we could exploit the modular architecture of NP1 and Cbln1 to develop a synthetic hexameric ESP, termed CPTX, which would include the N-terminal cysteine-rich region (CRR) of Cbln1 that binds Nrx(+4) and the PTX domain of NP1 (Fig. 1A, right panel). CPTX should have the potential to induce the formation of excitatory synapses by organizing Nrx nanoclusters (40) and to form a trans-synaptic complex by recruiting postsynaptic AMPARs (41).

To define the correct domain boundaries for CPTX assembly, we first solved the crystal structure of the PTX domain of NP1 (NP1PTX) at 1.45 Å resolution (Fig. 1B and table S1). Reminiscent of short pentraxins (42), such as the C-reactive protein and serum amyloid P component, NP1PTX formed a two-layered β sheet with a flattened jellyroll topology containing two Ca2+ ions (Fig. 1B and fig. S1). However, unlike short pentraxins, NP1PTX is monomeric in the crystal as well as in solution, as confirmed by multi-angle light scattering (MALS; Fig. 1C). Surface plasmon resonance (SPR) assays, performed to identify the minimal domain requirements for NP1–GlulA interactions, revealed that NP1PTX bound the ATDs of GluA1, GluA3, and GluA4 AMPARs with affinities in the high micromolar range (Fig. 1D). A recent analysis of the Nrx–Cbln1–GluD2 transsynaptic complex showed a similarly weak interaction between the globular domain of Cbln1 and the GluD2 ATD; however, avidity effects arising from the oligomeric nature of Cbln1 (hexamer) and GluD2 (tetramer) increase the apparent affinity between the full-length partners considerably (15). To mimic the structural organization of Cbln1 in CPTX, we first...
added a triple-coil–forming mutant GCN4 peptide to the N terminus of NP1PTX, resulting in the NP1PTX-3Cl construct, and confirmed its trimeric stoichiometry (Fig. 1C). The Cbln1 CRR (I5) was then added to the NP1PTX-3Cl N terminus, leading to the hexameric CPTX molecule (Fig. 1C and fig. S1).

We next examined whether CPTX was equipped with the intended dual binding capacity in vitro. SPR assays showed that CPTX and NP1PTX bound the ATD of GluA4 with comparable apparent affinities (dissociation constant $K_D = 4.5 \pm 0.5 \text{ mM}$ and $11.0 \pm 0.9 \text{ mM}$, respectively; Fig. 1D and fig. S2). Like NP1PTX, CPTX also bound to the other GluA ATDs with a GluA3 > GluA1 > GluA2 preference. However, these interactions were weaker, and accurate $K_D$ values could not be determined reliably (Fig. 1D and fig. S2). In addition, CPTX bound specifically to the Nrx1β(+4) ectodomain ($K_D = 4.9 \pm 0.9 \text{ nM}$), but not to Nrx1β(−4) (Fig. 1E), thus retaining the strict isoform recognition specificity of Cbln1.

Cell-based binding assays also showed that CPTX bound specifically to human embryonic kidney (HEK) cells displaying GluA1–GluA4 ATDs on their surface, whereas Cbln1 only bound...
those displaying the GluD2 ATD (fig. S3A). By contrast, both Cbln1 and CPTX bound to HEK293 cells expressing full-length Nrx1α, Nrx1β, Nrx2β, and Nrx3β only when the SS4 insert was present (fig. S3B). The ectodomain of Nrx1β(+4) specifically bound to HEK293 cells expressing GluA4 ATD under CPTX application (fig. S3C). Thus, CPTX can simultaneously bind GluAs and neurexin isoforms containing the SS4 insert.

CPTX induces excitatory pre- and postsynaptic sites in vitro

We examined whether CPTX could serve as a synaptic organizer with the designed GluA versus GluD specificity in neuronal cell cultures. Application of CPTX to Cbln1-null cerebellar granule cells induced punctate accumulation of endogenous synaptophysin, a synaptic vesicle marker, in axons that contact HEK293 cells displaying GluA ATDs on their surface, but not in those that contact cells displaying the GluD2 ATD (Fig. 2A and fig. S4A). Similarly, CPTX induced accumulation of synaptophysin (fig. S4B) or vesicular glutamate transporter 1 (VGLuT1) (fig. S4C) in wild-type hippocampal neurons that contacted HEK293 cells displaying GluA ATDs. By contrast, NPI could bind to the surface of HEK293 cells displaying the Glu4 ATD but failed to accumulate synaptophysin (fig. S4D).

Conversely, application of Cbln1 induced presynaptic synaptophysin accumulation in axons that contacted co-cultured HEK293 cells expressing GluD2, but not GluA ATDs (Fig. 2A and fig. S4, A and B). To examine the direct effect of CPTX, we applied beads coated with CPTX to hippocampal neurons in culture. Cbln1- or CPTX-coated beads accumulate pre-synaptic sites positive for endogenous synaptophysin, Nrx, and VGLuT1 (fig. S4B and fig. S4C), indicating that CPTX can directly induce presynapses in vitro.

We further asked whether CPTX could induce postsynaptic sites in vitro. Coexpression of Nrx1β(+4) and CPTX, but not Cbln1, on the
surface of HEK293 cells facilitated accumulation of endogenous GluA1–3 in dendrites of contacted hippocampal neurons (fig. S6A). Similarly, beads coated with CPTX, but not Cbln1, accumulated endogenous GluA1–GluA3 in dendrites of most hippocampal neurons (Fig. 2B and fig. S6B) and GluA4 in parvalbumin-positive (PV+) interneurons (fig. S6C). Immunocytochemical analyses revealed that soluble CPTX added to the culture medium accumulated between Homer- and Bassoon-positive puncta (fig. S7A) and at puncta double immunopositive for GluA1–4 and Nrx or VGluT1 (Fig. 2C and fig. S7B). CPTX also increased Nrx and GluA intensities in GluA/Nrx- and GluA/VGluT1–double immunopositive areas, respectively. Thus, CPTX serves as a bidirectional synaptic organizer by binding presynaptic Nrx(+4) and postsynaptic AMPARs in vitro.

Rescue of synapse formation and motor coordination by CPTX in cerebellar ataxia mice
To test the impact of CPTX in vivo, we injected it into the cerebellum (lobules VI and VII) of adult Cbln1-null mice. Immunohistochemical analyses 1 day after the injection showed that CPTX was preferentially confined to the injected molecular layer (fig. S8A) and colocalized with VGluT1 (a PF terminal marker), but not VGluT2 (a climbing fiber marker) or VGAT (an inhibitory input marker). The scatter plots show the 2D pixel intensity histograms for CPTX (green) and each synapse marker (magenta). Pearson’s correlation coefficient R values are indicated. Scale bar, 2 μm. (C) Representative electron microscopic images show free dendritic spines (f, blue) and contacted spines (c, red) innervated by PFs (green) in the GluD2-null cerebellum 3 days after injection of Cbln1 or CPTX. Scale bar, 500 nm. The fractions of contacted PC spines are quantified in the lower graph. ***P < 0.001, n = 10 to 20 sections from one or two mice, χ² test. (D) CPTX restores functional PF–EPSCs in GluD2-null mice 3 days after injection. Representative traces are shown. The middle graph shows averaged input-output relationships of PF–EPSCs for each treatment. **P < 0.01, *P < 0.05; n.s., not significant; n = 30 cells each, two-way ANOVA followed by Scheffe post-hoc test. The right graph shows the paired-pulse ratio of the second to first PF–EPSC amplitudes. *P < 0.05, n = 30 cells each. Kruskal-Wallis test followed by Scheffe post-hoc test. (E) CPTX improves the gait of GluD2-null mice 3 days after the injection. Representative footprints before and after CPTX injection are shown near the top. The lower graphs show the quantification of gait parameters. Each line represents an individual score before and after injection with either Cbln1 or CPTX. ***P < 0.001, **P < 0.01, *P < 0.05, n = 5 or 6 mice, Student’s paired t test. Scale bar, 50 mm. The error bars in (C) to (E) represent the mean ± SEM.

Fig. 3. CPTX restores PF-PC synapses and motor coordination in GluD2-null mice.
(A) HIS-tagged CPTX injected into the GluD2-null cerebellum localizes at PF–PC synapses. Mock, vehicle. Scale bar, 500 μm. (B) HIS immunoreactivity in the molecular layer of the injected area (A) colocalized with VGluT1 (a PF marker), but not VGluT2 (a climbing fiber marker) or VGAT (an inhibitory input marker). The scatter plots show the 2D pixel intensity histograms for CPTX (green) and each synapse marker (magenta). Pearson’s correlation coefficient R values are indicated. Scale bar, 2 μm. (C) Representative electron microscopic images show free dendritic spines (f, blue) and contacted spines (c, red) innervated by PFs (green) in the GluD2-null cerebellum 3 days after injection of Cbln1 or CPTX. Scale bar, 500 nm. The fractions of contacted PC spines are quantified in the lower graph. ***P < 0.001, n = 10 to 20 sections from one or two mice, χ² test. (D) CPTX restores functional PF–EPSCs in GluD2-null mice 3 days after injection. Representative traces are shown. The middle graph shows averaged input-output relationships of PF–EPSCs for each treatment. **P < 0.01, *P < 0.05; n.s., not significant; n = 30 cells each, two-way ANOVA followed by Scheffe post-hoc test. The right graph shows the paired-pulse ratio of the second to first PF–EPSC amplitudes. *P < 0.05, n = 30 cells each. Kruskal-Wallis test followed by Scheffe post-hoc test. (E) CPTX improves the gait of GluD2-null mice 3 days after injection. Representative footprints before and after CPTX injection are shown near the top. The lower graphs show the quantification of gait parameters. Each line represents an individual score before and after injection with either Cbln1 or CPTX. ***P < 0.001, **P < 0.01, *P < 0.05, n = 5 or 6 mice, Student’s paired t test. Scale bar, 50 mm. The error bars in (C) to (E) represent the mean ± SEM.
Fig. 4. CPTX restores spines and LTP in the hippocampus of an Alzheimer’s disease model. (A) CPTX restores dendritic spine density in 5xFAD mice. Representative Golgi-Cox staining of apical dendrites of CA1 pyramidal neurons in the dorsal hippocampus. Red marks indicate counted spines. Mock, vehicle (HBS buffer) controls. Scale bar, 5 μm. The graph shows the averaged spine density for seven or eight secondary dendrites for each animal. The bars represent the mean ± SEM. **P < 0.01, n = 4 or 5 mice, one-way ANOVA followed by Holm-Sidak post-hoc test. (B) CPTX restores Schaffer collateral (SC)–evoked field excitatory postsynaptic potentials (fEPSPs) in 5xFAD mice. The graph shows averaged input-output relationships of SC–fEPSPs for each treatment. Mock, vehicle (HBS buffer) controls. The bars represent the mean ± SEM. ***P < 0.001, **P < 0.01, *P < 0.05, n = 6 to 9 slices, repeated two-way ANOVA. (C) CPTX does not affect the presynaptic release probability. The graph shows the paired-pulse ratio of the second to first SC-fEPSP amplitudes at 50-ms intervals. The bars represent the mean ± SEM. n.s., not significant, n = 8 or 9 slices, Student’s t test. (D) CPTX increases the frequency and the amplitude of mEPSCs, but not mIPSCs, in the 5xFAD hippocampus. Representative traces are shown near the top. **P < 0.01, *P < 0.05, n = 9–15 cells, Student’s t test. (E) CPTX restores LTP in 5xFAD mice. Application of theta-burst stimulation to SC three times induced a robust LTP at SC-CA1 synapses in the mock-treated WT, but not in the mock-treated 5xFAD hippocampus. Representative SC-fEPSP traces are shown. The bars represent the mean ± SEM. ***P < 0.001, **P < 0.01, *P < 0.05, n = 8 or 9 slices, one-way ANOVA on ranks followed by Holm-Sidak post-hoc test for the mean values from the last 15 min of the recording.

or the vesicular γ-aminobutyric acid (GABA) transporter (VGAT, a marker for inhibitory inputs) (fig. S8B). Thus, CPTX was mainly located along PC dendrites where PFs that specifically express Nr3c1 (43) make synapses. Electron microscopy analysis revealed that single injections of CPTX partially restored PF–PC synapses in Cbln1-null mice (fig. S8C). The effect was weaker than for Cbln1 (13, 16), probably because there is a large excess of GluD2 relative to GluAs at PF–PC synapses (44). Whole-cell patch-clamp recordings in acute slice preparations showed that CPTX, but not NPs1PTX-3Cl, increased the amplitudes of PF-evoked excitatory postsynaptic currents (EPSCs) (fig. S8D). Thus, CPTX most likely regulates synaptic functions by bridging pre- and postsynaptic sites. Furthermore, CPTX was as effective as Cbln1 in restoring normal gait patterns (figs. S8E and S9A and movies S1 and S2). We also observed recovery of motor coordination as measured by the rotor-rod test in the ICR mouse genetic background (fig. S9B), but not in the C57BL/6 one where the loss of the skilled motor coordination phenotype was too severe. Similar to Cbln1 (16), the effect of CPTX on the rotor-rod performance recovery was maximal at 3 days after injection but decayed afterward (fig. S9C), suggesting that continued presence of CPTX is necessary to maintain PF–PC synapses in the longer term. Thus, CPTX can induce functional PF–PC synapses that contribute to the restoration of the motor phenotypes in adult Cbln1-null mice.

We next investigated whether CPTX could serve as a synaptic organizer in the absence of GluD2 in vivo. As observed in Cbln1-null mice, injected CPTX was localized at PF–PC synapses in GluD2-null mice (Fig. 3, A and B). Although Cbln1 and GluD2 form a transsynaptic complex, GluD2-null mice are reported to show more severe motor phenotypes with less reduced PF–PC synapses (~40 to 60% of wild-type) than Cbln1-null mice (~20 to 30%) (13, 16). Whereas Cbln1 was completely ineffective in GluD2-null mice in restoring morphological (Fig. 3C) and functional (Fig. 3D) PF–PC synapses, CPTX partially restored these phenotypes. CPTX, but not Cbln1, reset the increased paired-pulse ratio (PPR) of PF-EPSCs to wild-type values (15) (Fig. 3D). Furthermore, unlike Cbln1, CPTX improved the gait of GluD2-null mice (Fig. 3E and movie S3). Thus, CPTX can induce PF–PC synapses that contribute to cerebellar functions independent of GluD2 in vivo.

CPTX induces synaptic AMPAR accumulation in the wild-type hippocampus

To test if CPTX could act in brain regions outside the cerebellum, we injected CPTX into the dorsal hippocampus of Thy1–green fluorescent protein (GFP) and wild-type mice. Injected CPTX diffused throughout the entire CA1 regions of dorsal hippocampi and was detected between Bassoon-positive presynaptic terminals and GFP-positive postsynaptic sites of the CA1 pyramidal neurons in Thy1-GFP mice at 3 days after injection (fig. S10A). Super-resolution microscopy revealed that CPTX was localized between VGluT1 and AMPAR puncta in the CA1 stratum radiatum at 1 day after injection (fig. S10, B and C). By contrast, CPTX
did not show any colocalization with inhibitory synaptic markers (fig. S11). Immunohistochemistry of CPTX and each AMPAR subunit (fig. S12A) revealed that CPTX shifted the intensity histogram of GluA1 to higher values, while causing redistribution of GluA2/3 signals among puncta (fig. S12, B to D). As a result, CPTX increased the size of high-intensity (>mean + 3 SD) puncta of GluA1 and GluA2/3 (fig. S12, E and F). By contrast, although CPTX also shifted the intensity histogram of GluA4 to higher values, the intensity (fig. S17) than in the stratum radiatum of middle-aged 5xFAD mice (Fig. 4E). Because CPTX did not increase GluN1 immunoreactivities, it is plausible that newly formed spines (fig. 4A) and redistribution of PSD95 and AMPARs (fig. S18) failed to increase SC-fEPSPs (fig. S19). A result consistent with the lack of effect of NP1 in inducing synapses in vitro (fig. S4D). CPTX did not affect the PPR of SC-fEPSPs, probably because the PPR was normal in 5xFAD hippocampus (Fig. 4C), a phenotype different from GluD2-null cerebellum (Fig. 3D). Moreover, whole-cell patch-clamp recordings from CA1 pyramidal neurons revealed that the frequency and amplitude of mEPSCs were increased in 5xFAD mice injected with CPTX (Fig. 4D). By contrast, miniature inhibitory synaptic currents (mIPSCs) were unaffected (Fig. 4D). Thus, CPTX specifically improved excitatory transmission between principal cells without changing GABAergic transmission in 5xFAD mice.

Impaired long-term potentiation at SC-CA1 synapses (SC-LTP) is believed to underlie certain cognitive abnormalities in 5xFAD mice (47). CPTX injection rescued impaired SC-LTP in middle-aged 5xFAD mice (Fig. 4E). Because CPTX did not increase GluN1 immunoreactivities, it is plausible that newly formed spines (Fig. 4A) and redistribution of PSD95 and AMPARs (fig. S18) may contribute to restoration of LTP. To evaluate spatial memory in a paradigm with minimal stress exposure, we placed three groups of mice in a labyrinth with food pellets as a reward and measured the total distance that mice traveled to reach the goal during the encoding and retrieval sessions (Fig. 5A). Although there was no difference in the travel distance between each group during the encoding session, 5xFAD mice traveled longer distances than wild-type mice during the retrieval session. CPTX injection 3 days before the learning test decreased the distance that 5xFAD mice traveled in the retrieval session, suggesting an improvement of spatial learning after the CPTX treatment. After reversal learning, when mice had to find a reward in a new position, the performance of 5xFAD mice was impaired in the mock- but not in the CPTX-injected group, as compared to wild-type mice (Fig. 5A). The modest effect of CPTX in this test may be caused by the insufficient spread of CPTX because spatial memory requires localization, together with synaptophysin, GluA1–3, and GluN1, an essential N-methyl-D-aspartate (NMDA) receptor subunit in the hippocampus (fig. S18A). CPTX caused the redistribution of PSD95 as well as GluA1–3 signal intensities without affecting synaptophysin or GluN1 signals (fig. S18, B and C). CPTX also increased the proportion of large puncta positive for PSD95 and GluA1–3, but not GluN1 or synaptophysin (fig. S18D). Extracellular field recordings confirmed that excitatory postsynaptic potentials (fEPSPs) induced by SC stimulation were reduced in acute slices prepared from 5xFAD mice (Fig. 4B), as reported earlier (47). CPTX restored the amplitude of SC-fEPSPs, which became even larger than in the wild-type hippocampus (Fig. 4B). By contrast, NP1PTX-3Cl failed to increase SC-fEPSPs (fig. S19), a result consistent with the lack of effect of NP1 in inducing synapses in vitro (fig. S4D).

Fig. 5. CPTX restores hippocampus-dependent behaviors in an Alzheimer’s disease model. (A) CPTX improves spatial memory in 5xFAD mice. CPTX or Mock (vehicle) was injected in 5xFAD mice on day 0. On day 3, mice were placed at the start point (S) of a 3D-printed maze and the pellet was placed at the reward point (R). Two hours after the initial encoding (E) session, mice were returned to the same start point to examine memory integrity in the retrieval (R) session. On the next day, the position of reward was changed, and reversal learning (relearning) was evaluated. Scale bar, 30 cm. The averaged total distances that mice traveled to reach the goal during the encoding and the retrieval sessions are shown in the lower graph. Log10 scaling of the y axis facilitates comparison of distances before and after training. The bars represent the mean ± SEM. **P < 0.01, *P < 0.05, #P < 0.1, n = 8 to 11 mice, two-way repeated measures ANOVA followed by Fisher’s post-hoc least significant difference (LSD) test. (B) CPTX restores context discrimination in 5xFAD mice. CPTX or Mock (vehicle) was injected in 5xFAD mice on day 0. Electrical shock was applied to wild-type and 5xFAD mice in context A and context B on day 5. Freezing time was measured in context A and context B on day 6. The lower graph shows the mean (± SEM) freezing time of each group. ***P < 0.001, n = 8 to 11 mice, Student’s paired t test.
integration of information from head direction cells, grid cells, and place cells with landmarks (context). To more directly assess the function of the CA1 region, we next evaluated contextual fear conditioning by exposing the same three groups of mice to an electrical shock in context A and measuring the freezing time in the conditioned context A and a neutral context B 1 day later. 5xFAD mice failed to discriminate context A from B, i.e., showed generalized rather than context-specific learned fear responses. CPTX injection 5 days before fear conditioning improved context discrimination in 5xFAD mice to be even better than in mock-treated wild-type controls (Fig. 5B), indicating that CPTX could rescue the context generalization in 5xFAD mice. Thus, CPTX injection rescues the impaired basal excitatory transmission, SC-LTP, and hippocampus-dependent learning in middle-aged 5xFAD mice.

CPTX restores synapses and locomotion in spinal cord injury models

Reorganization and the E/I balance of spared intraspinal networks have recently been shown to contribute to functional recovery after spinal cord injury (SCI) (48, 49). To test whether CPTX could restore glutamatergic circuits in the spinal cord, we performed dorsal hemisections at the thoracic vertebra 10 (T10) in mice and injected CPTX rostral to the injured region (Fig. 6A). Spinal cords were dissected 2 to 5 days after injections, and coronal serial sections were immunostained for VGluT2 and GluA4 upon Mock (vehicle) or CPTX injection. Scale bar, 0.5 mm. (B) Representative orthogonal images obtained by Airyscan super-resolution microscopy indicating the localization of CPTX in proximity of VGluT2- and GluA4-immunopositive puncta. Scale bar, 0.5 μm. (C) Representative immunohistochemical staining images of coronal sections stained for VGluT2 (blue), GluA4 (green), and HIS (CPTX; magenta) from mock- or CPTX-treated spinal cords. Scale bar, 5 μm. (D) Quantification of the fraction of GluA4+/VGluT2+–double-positive puncta. The bars represent the mean ± SEM. *P < 0.05, n = 16 slices from eight mice, Student’s t test. (E to G) Time-course analyses of locomotion [Basso mouse scale (BMS) score] in SCI mice after injections. Mock (vehicle), chondroitinase ABC (ChABC), Cbln1, or CPTX were injected into the spinal cord immediately (E) or 1 week after (F) hemisection, or immediately after contusion by impactors (G) (70 kdyn impact force). For the sham controls, the spinal cord was surgically exposed without imposing injury or injection. Mice that showed a BMS score of 1.5 at 1 week after hemisection were selected for (F). **P < 0.01, *P < 0.05, n = 9 mice for each treatment, repeated two-way ANOVA with post-hoc Bonferroni-Dunn test (comparing the various treatments with Mock for each time point).
CPTX induced GluA4+ excitatory synapses within regions spared by the injury (fig. S20, A and B).

Finally, we assessed the recovery of locomotion after spinal cord hemisections. For comparison, we separately injected chondroitinase ABC (ChABC), an enzyme that promotes robust nerve regeneration (52), as well as Cbln1 to the lesioned spinal cord. During the acute stage, single injections of CPTX as well as ChABC, but not Cbln1, significantly restored the Basso mouse scale (BMS) score (53) (Fig. 6E and movie S5) and other locomotive parameters examined until 6 weeks after the injury (fig. S21A and movie S6). ChABC delivered 1 week after the hemisection was less effective (Fig. 6F), as reported previously (54). By contrast, when CPTX was delivered 1 week after injury, it was most effective (Fig. 6F, fig. S21B, and movies S7 and S8). CPTX also efficiently restored locomotion in a contusion SCI model, which is considered closer to pathophysiology that occurs commonly in humans (Fig. 6G and fig. S21C). In this paradigm, a combination of CPTX and ChABC treatment showed the most prominent recovery of locomotion, especially in the early stages after contusion. Immunoblot analyses revealed that by 7 days after injection, CPTX became undetectable in the spinal cord as well as in the hippocampus and the cerebellum (fig. S22). Thus, CPTX may directly enhance excitatory connectivity, which may be further stabilized by endogenous synaptic organizers, to promote sustained functional recovery in spinal cord injury models. Mechanistically, this process appears to be distinct from the one induced by ChABC treatment.

Discussion

Here, we developed a synthetic, structure-guided, synaptic organizer termed CPTX that can induce functional and structural synapses in the cerebellar, hippocampal, and spinal cord neuronal circuits in vivo. CPTX induced accumulation of AMPARs, as revealed by increased GluA1 intensity in all puncta and redistribution of GluA2/3 signals in the hippocampus 1 day after injection. CPTX application to the wild-type hippocampus led to increased mEPSC frequency in ~4 hours, and then both the frequency and the amplitude of mEPSCs ~3 days later. Thus, we speculate that CPTX first accumulates preexisting surface AMPARs at immature synapses with loose perisynaptic extracellular matrix (55), leading to redistribution of AMPARs and the increased mEPSC frequency. Considering that GluA1-containing AMPARs traffic to the surface in an activity-dependent manner (56), CPTX may stabilize newly inserted GluA1-containing AMPARs in mature synapses, leading to increased GluA1 surface amounts and mEPSC amplitudes. NP1 or NP1PTX-3Cl, which could bind to AMPARs, failed to induce excitatory synapses in the hippocampus. Moreover, NP1PTX-3Cl could not restore PF-EPSCs in Cbln1-null mice. Thus, we propose that CPTX induces functional synapses by connecting presynaptic neuromins with postsynaptic AMPARs and facilitating transsynaptic alignment of AMPARs with the synaptic vesicle release sites. Such “molecular bridges” may initiate presynaptic maturation by inducing Nrx nanoclusters (40) and may eventually integrate within transsynaptic nanocolumns at excitatory synapses (41). Because molecular components are considerably different among neuronal synapse types and circuits, further studies are warranted to test this hypothesis and clarify the detailed molecular mechanisms by which CPTX restores functional and structural excitatory synapses under pathological conditions.

Potential adverse effects of CPTX could include inhibition of synapses where CPTX binds GluAs but does not have a presynaptic Nrx(+4) partner. However, injected CPTX did not replace endogenous NPs that target the ATD of GluA4. CPTX could also cause artificial hyperconnectivity, but we did not observe obvious abnormal behaviors, such as seizures or hyperactivities, in mice injected with CPTX into the hippocampus, the spinal cord or the cerebellum. Most likely this is because CPTX selectively strengthens synapses that contain Nrx(+4) at presynaptic sites. However, constitutive expression of Nrx(+4) in genetically modified mice reduces mEPSC frequency and decreases LTP (57). Thus, the dose and the affinity of CPTX-like molecules should be optimized for future therapeutic use.

The catalog of ESPs continues to grow. For example, LGI1, an ESP produced by the hippocampal neurons, is particularly interesting because it indirectly recruits AMPARs by binding to pre- and postsynaptic ADAM23 and ADAM22, respectively (57, 58). Owing to their modular structure, we anticipate that the toolkit of synthetic ESPs can be expanded to include a variety of pre- and postsynaptic specificities, with a range of affinities and enhanced resistance to proteolytic degradation in vivo. Such ESPs could be used to restore or modify synaptic connectivity in various neuronal circuits. The structure-guided design of ESPs that target specific GABA_A receptor subtypes (59), as well as specific AMPAR or KAR subunits, will be useful to restore E/I balance in specific neuronal circuits affected by autism spectrum disorders and schizophrenia (1, 5, 60), epilepsy (3), and AD (2). Notably, although the effect of CPTX was transient in Cbln1-null ataxia mice, a single injection of CPTX greatly enhanced locomotion for at least 7 to 8 weeks in SCI models, long after injected CPTX was degraded. In the cerebellum, PF-PC synapses rely heavily on Cbln1 and GluD2 as synaptic organizers. Thus, once the injected CPTX is degraded, cerebellar synapses are rapidly lost and ataxic phenotypes return quickly. By contrast, in 5xFAD mice and SCI models, the new synapses formed upon CPTX injection might survive longer because other (endogenous) synaptic organizers could be recruited by increased neuronal activities. Thus, synthetic ESPs built upon the principles described here may become powerful tools to investigate and potentially cure disorders associated with impaired neuronal connectivity.

Materials and methods

Mice

Wild-type (Charles River Laboratories and Japan SLC) and Cbln1-null mice were used for primary culture. In experiments assessing cerebellar functions, we used male and female Cbln1-null mice with the C57BL/6J or ICR genetic backgrounds and GluD2-null mice (13) with the C57BL/6J genetic background. They were housed on a conventional 12 hours light/dark cycle (light on at 8:30 a.m.) and their behavior was tested in the light phase of the cycle. Mice expressing enhanced green fluorescent protein (EGFP) under the control of a modified Thy1 promoter region (The Jackson Laboratory, cat. no. 007788) were used for immunohistochemistry. In experiments assessing hippocampal functions, we used male C57BL/6J, PV-Cre/Ai9 double transgenic mice (The Jackson Laboratory, cat. no. 00720 and 007909), 5xFAD mice (The Jackson Laboratory, cat. no. 006554) which carry five familial AD mutations, and their wild-type littermates with C57BL/6J genetic background (Charles River Laboratories). At least one week before starting the experiments, mice were transferred from the main animal facility of DZNE (Magdeburg, Germany) to a small vivarium, where they were housed individually with food and water ad libitum before food restriction diet (FRD) was given, on a reversed 12 hours light/dark cycle (light on at 9:00 p.m.). All behavioral experiments assessing hippocampal functions were performed in the late afternoons during the dark phase of the cycle when mice were active, under constant temperature (22° ± 1°C) and humidity (55 ± 5%). For the spinal cord injury (SCI) model, wild-type mice with the ICR genetic background (Japan SLC) were used. They were housed on a conventional 12 hours light/dark cycle. Locomotor functions were assessed at a fixed time in the evening. All treatments and behavioral procedures were conducted in accordance with ethical animal research standards defined by German and Japanese law and approved by the Ethical Committee on Animal Health and Care of the State of Saxony-Anhalt, Germany, with license numbers 42502-2-1343 DZ/NE and 42502-2-1322 DZ/NE, by the Animal Resource Committee of Keio University (no. 09050) and by the Institutional Animal Care and Use Committee of Aichi Medical University (no. 1559).

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**Plasmids**

For surface plasmon resonance (SPR) experiments, cDNAs encoding the extracellular human glutamate receptor D2 amino-terminal domain (ATD; UniProt ID O43424; GluD2 ATD: Asp24-Gly440), the mouse glutamate receptor A1 ATD (UniProt ID P23818; GluA1 ATD: Ala9-Thr42), the mouse glutamate receptor A2 ATD (UniProt ID P23819; GluA2 ATD: Val22-Thr49), the mouse glutamate receptor A3 ATD (UniProt ID Q9Z2W9; GluA3 ATD: Gly23-Thr42), the mouse glutamate receptor A4 ATD (UniProt ID Q9Z2W8; GluA4 ATD: Gly21-Thr42), and full-length mouse NPR (UniProt ID Q62443) was cloned into the pCAGGS vector as described previously (sequences: 5′-CAGCATATTGGGCTGGCCTTTCTGTA-3′) to the C terminus of the cysteine-rich region (CRR) (64) of HEK293 cells. Briefly, for large-scale transient transfections, the pHLsec expression plasmids encoding the proteins of interest (NP1ptx, Nrx1±SS4, Nrx1±SS4, GluA1ATD, GluA2ATD, GluA3ATD, GluA4ATD, and GluD2ATD) were transfected into 3-liter adherent HEK293T (ATCC cat. no. CRL-1753) expression cell cultures. The supernatants were collected 5 days after transfection and filtered (0.22 μm). For the lentiviral infections, the pHR-CMV-TetO2 transfection plasmid (Addgene cat. no. 113887) encoding the protein of interest (CPTX), the psPAX2 packaging plasmid and the pMD2.G envelope plasmid were cotransfected into the HEK293T Lenti-X producer cell line (Takara Bio cat. no. 632180). The supernatant containing the viral particles was harvested after 3 days, filtered (0.45 μm) and directly used for the infection of HEK293T expression cells. The infected cells were grown and expanded to 3-L adherent cell cultures. The supernatants were collected 5 days after infection and filtered (0.22 μm). All supernatants were concentrated to ~1.0 g/liter using a 30,000 MWCO Amicon Ultra-15 Centrifugal Filter Centrifugation System (Millipore) and directly used for the immunoblotting analyses to check the selectivity of anti-NPI and anti-NPR antibodies (fig. S15, A and B). Mouse NPR (without the signal sequence: ΔMet1-Ala22), NP2 (without signal sequence: ΔMet1-Ala14) and NPR (without the transmembrane domain, ΔMet1-Ile23) were cloned into pCAGGS downstream of the Igs secretion signal sequence, followed by a 2xHA tag.

**Production of recombinant proteins**

For large-scale production, proteins were expressed using established procedures for transient transfection (61) or lentiviral transduction (64) of HEK293 cells. Briefly, for large-scale transient transfections, the pHLsec expression plasmids encoding the proteins of interest (NP1ptx, Nrx1±SS4, Nrx1±SS4, GluA1ATD, GluA2ATD, GluA3ATD, GluA4ATD, and GluD2ATD) were transfected into 3-L adherent HEK293T (ATCC cat. no. CRL-1753) expression cell cultures. The supernatants were collected 5 days after transfection and filtered (0.22 μm). For the lentiviral infections, the pHR-CMV-TetO2 transfection plasmid (Addgene cat. no. 113887) encoding the protein of interest (CPTX), the psPAX2 packaging plasmid and the pMD2.G envelope plasmid were cotransfected into the HEK293T Lenti-X producer cell line (Takara Bio cat. no. 632180). The supernatant containing the viral particles was harvested after 3 days, filtered (0.45 μm) and directly used for the infection of HEK293T expression cells. The infected cells were grown and expanded to 3-L adherent cell cultures. The supernatants were collected 5 days after infection and filtered (0.22 μm). All supernatants were concentrated to ~1.0 g/liter using a 30,000 MWCO Amicon Ultra-15 Centrifugal Filter Centrifugation System (Millipore) and directly used for the immunoblotting analyses to check the selectivity of anti-NPI and anti-NPR antibodies (fig. S15, A and B). Mouse NPR (without the signal sequence: ΔMet1-Ala22), NP2 (without signal sequence: ΔMet1-Ala14) and NPR (without the transmembrane domain, ΔMet1-Ile23) were cloned into pCAGGS downstream of the Igs secretion signal sequence, followed by a 2xHA tag.

**Protein crystalization**

Crystallization trials, using 100 nl protein solution plus 100 nl reservoir solution in sitting drop vapor diffusion format, were set up in 96-well Greiner plates using a Cartesian Technologies robot (65). Purified human NP1ptx (Pro224-Ile431), produced from HEK293T cells and concentrated to 6.5 g/liter, crystallized in 15% (v/v) glycerol, 25.5% (w/v) polyethylene glycol (PEG) 8000, 0.17M ammonium sulfate and 0.085 M sodium cacodylate (pH 6.5). Crystals were cryoprotected using a reservoir solution containing 30% (v/v) ethylene glycol.

**Crystallographic data collection and structure determination**

Diffraction data for NP1ptx were collected at Diamond Light Source (DLS) beamline I03 to a nominal resolution of 1.45 Å in space group (SG) C2. All data were indexed, integrated, and scaled using the automated XIA2 expert system (66), using the Labelit, POINTLESS and AIMLESS (68, 69) and XDS (70) software. The structure of NP1ptx was solved by molecular replacement using Phaser (71) and using the human SAP (Serum Amyloid P Component; PDB code ISAC) crystal structure (42) as a search model. Crystallographic data collection and refinement statistics are presented in table S1.

**Structure refinement and model analysis**

Maximum-likelihood refinement of NP1ptx was initially performed with Refmac5 using “jelly body” restraints (72), and finally with phenix.refine (73) using automated X-ray and atomic displacement parameter (ADP) weight optimization applied throughout. Automated model building was performed using ARP/WARP (74) and further manual model building was performed using Coot (75). Structure validation was performed using the MolProbity routines within the PHENIX software suite (73, 76). Molecular representations were made using PyMOL (77).

**Multi-angle light scattering (MALS)**

Protein samples concentrated to ~1.0 g/liter were injected into an HPLC-driven SEC column (Superdex 200 10/30 column, GE Healthcare) equilibrated with HBS-C buffer. The SEC column was coupled to an online UV detector (Shimadzu), an 18-angle light scattering detector (DAWN HELOS; Wyatt Technology), and a refractive index detector (Optilab T-rEX; Wyatt Technology). CPTX contains an N-linked sugar on the Cbln1 N-terminal cysteine-rich region (CRR), and molecular mass determination was performed using an adapted RI increment value (dn/dc standard value: 0.185 ml/g) to account for the glycosylation state. NP1ptx and NP1ptx±SS4 are not glycosylated hence the standard dn/dc value was used for molecular mass determination. Data
analysis was carried out using the ASTRA V software (Wyatt Technology).

**Surface plasmon resonance (SPR)**

cDNA for the immobilized proteins (CPTX, NP1PTX, Cbn1) was cloned into the pHSec-AviTag3 vector (47), resulting in proteins carrying a C-terminal biotin ligase (BirA) recognition sequence (Avitag). Constructs were cotransfected with pDisplay-BirA-ER (Addgene cat. no. 20856; coding for an ER-resident biotin ligase) (78) for in vivo biotinylation in HEK293T cells in small-scale six-well plates in a 3:1 pHSec-pDisplay stoichiometric ratio. A concentration of 100 μM D-biotin was maintained in the expression medium to ensure biotinylation of the Avitag. After 48 hours of expression, conditioned medium was collected and dialyzed against 10 mM Tris (pH 7.4), 150 mM sodium chloride, 3 mM calcium chloride and 0.005% (v/v) Tween-20 (TBS-CT). SPR experiments were performed on a Biacore T200 (GE Healthcare) operated at a data collection frequency of 10 Hz. Streptavidin (Sigma-Aldrich cat. no. S4762) was chemically coupled via amine coupling chemistry onto CM5 chips to a response unit (RU) level of 5000 RU. Then, biotinylated proteins were captured to the desired RU level. In each instance, for every two analyte binding cycles, a buffer injection was performed, allowing for double referencing of the binding responses (79).

**Antibodies**

The origin, dilution, company and catalog number are as follows: Anti-calbindin (Goat, 1:500, Frontier Institute, Af1000), anti-Parvalbumin (Goat, 1:500, Frontier Institute, Af460), anti-Homer 1 (Guinea pig, 1:1000, Synaptic Systems, 160-004) and anti-Bassoon (Rabbit, 1:500, Synaptic Systems, 141-003 or Mouse, 1:500, Enzo Life Sciences, SAP7F407). Anti-NP1 and NPR antibodies were newly raised by guinea pigs and rabbits immunized with peptides Glu185-Lys227 (NP1) or Ser226-Glu263 (NPR). For the simultaneous detection of various GluAs (GluA1-3 or GluA1-4), the mixture of the primary antibodies described above was used. Secondary antibodies conjugated with DyLight 405, Alexas 488, 546, 647 and Cy3 (Invitrogen or Jackson ImmunoResearch Laboratories) against the respective primary antibody were used in the dilution 1:1000 for immunocytochemistry or 1:200 for immunohistochemistry. Isospecific secondary antibodies against mouse IgG1 or IgG2a were used for the simultaneous staining of two types of mouse primary antibodies.

**Immuno blotting**

For fig. S15, A and B, the hippocampus and conditioned medium were solubilized in lamine buffer (2% SDS, 80 mM Tris-HCl (pH 6.8), 10% glycerol, 0.00625% Coomassie blue G 250) and the proteins were reduced by boiling for 3 min using 2% 2-mercaptoethanol. For fig. S22A, brain tissue from the hippocampus, cerebellum (vermis) and spinal cord were solubilized by 190 to 250 μl 1% SDS buffer, sonicated, and quantified for protein concentration. Samples were diluted in lamine buffer and reduced by boiling for 20 min using 2% 2-mercaptoethanol. Samples were subjected to gradient gel SDS-PAGE (FujiFilm Wako Chemicals) and blotted onto the membrane (Immobilon-P PVDF membranes, Merck Millipore). Membranes were blocked with TS-tween (0.1% Tween-20, 50 mM Tris-HCl pH7.6, 150 mM NaCl) containing 5% skimmed milk (CP-Mejji) and subjected to the primary antibody for 2 hours and HRP-conjugated secondary antibody (GE Healthcare) for 30 min. Chemiluminescence was generated by the ImmunoStar (Fujifilm Wako Chemicals) or Immobilon (Merck Millipore) parking NS21 supplement (80), 50 U/ml Penicillin, 50 mg/ml Streptomycin (Invitrogen), 2 mM l-Glutamine and 2% FBS (HyClone) in 5% CO2 at 37°C. One to 2 hours after the initial incubation, the culture medium was changed to fresh medium without FBS. For induction of presynapses by HEK293 cells (fig. S4), neurons were co-cultured with 2 × 107 HEK293 cells expressing GFP or Myc-tagged GluAl-4 or GluD2 ATDs at day-in-vitro (DIV) 4, 5 or 7 for 1 hour; followed by treatment with vehicle or recombinant Cbn1-HIS or CPTX-HIS (23.6 to 30 nM, calculated for the hexamers), or with conditioned medium containing NP1-HA or CPTX-HIS secreted from HEK293 cells transfected with those plasmids for 2 days. For induction of AMPARs by HEK293 cells (fig. S6), neurons were co-cultured with HEK293 cells expressing Nrx3xFLAG and GFP, Cbn1-HIS or CPTX-HIS at DIV9 for 4 hours. For the in vitro tripartite binding assay (fig. S3C), cells were further incubated with the conditioned medium containing NrxI(4-8)+hFc for 4 hours. Cells were fixed with 4% PFA/PBS for 15 min and washed with PBS three times. After blocking with 3% BSA/PBS (BSA: Sigma-Aldrich) for 30 min without permeabilization, cells were further incubated with primary antibodies against the ligand tag (HIS, HA and/or hFc) in 3% BSA/PBS for 2 hours at room temperature (RT) or 24 hours at 4°C. Following three washes with PBS, cells were permeabilized with PBS containing 0.1% Triton-X 100 (Sigma-Aldrich) and 3% BSA. Cells were stained with primary antibodies against the receptor tag (FLAG or Myc) for 2 hours at RT or 24 hours at 4°C, followed by washing with PBS and incubation with the respective secondary antibodies for 30 min. After washing with PBS, coverslips were mounted on a glass slide with Fluormount-G. Samples in which HEK293 cells expressed GFP were not stained by primary antibodies against GFP but treated with the same primary antibodies against the receptor tag (FLAG or Myc) and the fluorescence of GFP was detected by microscopy.

**In vitro synapse formation assays**

**Hippocampal culture**

Preparation from embryonic day 16-17 (E16-17) wild-type ICR mice was performed as described (13) with some modifications. Neurons (2 × 107) were plated on 12-mm cover glasses coated with PLL and maintained in Neurobasal cell culture medium (Invitrogen) containing N2 supplement (80), 50 U/ml Penicillin, 50 mg/ml Streptomycin (Invitrogen), 2 mM l-Glutamine and 2% FBS (HyClone) in 5% CO2 at 37°C. One to 2 hours after the initial incubation, the culture medium was changed to fresh medium without FBS. For induction of presynapses by HEK293 cells (fig. S4), neurons were co-cultured with 2 × 107 HEK293 cells expressing GFP or Myc-tagged GluAl-4 or GluD2 ATDs at day-in-vitro (DIV) 4, 5 or 7 for 1 hour; followed by treatment with vehicle or recombinant Cbn1-HIS or CPTX-HIS (23.6 to 30 nM, calculated for the hexamers), or with conditioned medium containing NP1-HA or CPTX-HIS secreted from HEK293 cells transfected with those plasmids for 2 days. For induction of AMPARs by HEK293 cells (fig. S6A), neurons were co-cultured with HEK293 cells expressing Nrx3xFLAG and GFP, Cbn1-HIS or CPTX-HIS at DIV9 for 2 days. For synapse induction by beads, neurons were treated with streptavidin Dynabeads M-280 (Thermo Fisher

**In vivo binding assays**

HEK293 tsA201 cells (a gift from R. Horn, Thomas Jefferson University, Philadelphia, USA) were cultured in high glucose DMEM (Sigma-Aldrich) containing 10% FBS (HyClone), 50 U/ml Penicillin, 50 mg/ml Streptomycin (Invitrogen), 2 mM l-Glutamine in 10% CO2 at 37°C. Cells were transfected with pDisplay encoding myc-tagged GluAl-4 ATD or with pCAGGS encoding GFP or FLAG-tagged Nrx, using Lipofectamin 2000 (Invitrogen). On the following day, transfected cells were detached by phosphate-buffered saline (PBS) containing 5 mM EDTA and seeded on 12-mm coverslips coated with poly-L-lysine (PLL; Sigma-Aldrich) at 2 × 105 cells/well. One hour after seeding, cells were treated with Mock (vehicle; HBS buffer), recombinant Cbn1-HIS or CPTX-HIS (23.6 nM, calculated for the hexamers) for 4 hours to overnight. For the in vitro tripartite binding assay (fig. S3C), cells were further incubated with the conditioned medium containing NrxI(4-8)+hFc for 4 hours. Cells were fixed with 4% PFA/PBS for 15 min and washed with PBS three times. After blocking with 3% BSA/PBS (BSA: Sigma-Aldrich) for 30 min without permeabilization, cells were further incubated with primary antibodies against the ligand tag (HIS, HA and/or hFc) in 3% BSA/PBS for 2 hours at room temperature (RT) or 24 hours at 4°C. Following three washes with PBS, cells were permeabilized with PBS containing 0.1% Triton-X 100 (Sigma-Aldrich) and 3% BSA. Cells were stained with primary antibodies against the receptor tag (FLAG or Myc) for 2 hours at RT or 24 hours at 4°C, followed by washing with PBS and incubation with the respective secondary antibodies for 30 min. After washing with PBS, coverslips were mounted on a glass slide with Fluormount-G. Samples in which HEK293 cells expressed GFP were not stained by primary antibodies against GFP but treated with the same primary antibodies against the receptor tag (FLAG or Myc) and the fluorescence of GFP was detected by microscopy.

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were fixed with 4% PFA/PBS for 5 min at RT. Following the incubations, the cells were washed with PBS and fixed with chilled methanol for 8 min on ice, followed by washing with PBS. Cells were permeabilized and blocked with PBS containing 0.1% Triton-X 100, 3% BSA and stained with primary antibodies diluted in blocking buffer for 2 hours at RT or 24 hours at 4°C, followed by washing with PBS and incubation with secondary antibodies diluted in blocking buffer for 30 min. Following washing with PBS, coverslips were mounted on a glass slide with Fluoromount-G (SouthernBiotech).

**Cerebellar culture**

Cerebellar cultures were prepared from postnatal day 0 to 3 (P0 to P3) Chln-null mouse (on the ICR genetic background) as described (J3) with minor modifications. Neurons (2 × 10⁵) were plated on 12 mm cover glasses coated with PLL and were maintained in DMEM/F12 (Sigma-Aldrich) containing 100 μM putrescine, 30 mM sodium selenite, 0.5 μg/ml triiodothyronine, 0.25 mg/ml BSA, 3.9 mM glutamate and N3 supplement (100 μg/ml apo-transferrin, 10 μg/ml insulin and 20 mM progesterone) and 10% FBS in 5% CO₂ at 37°C. One to 2 hours after incubation, the culture medium was changed to fresh medium without FBS. Co-culturing with HEK293 cells (at DIV 4 or 5 for 2 days), fixation, permeabilization and staining were performed as described above for the hippocampal culture.

**Quantification of signal intensities of in vitro assays**

Fluorescence images of cultured samples were captured using a CCD camera attached to a conventional fluorescence microscope (BX63, Olympus) or a confocal laser scanning microscope (FV1000, Olympus) using a 40× objective for the detection of HEK293 cells or using a 60× objective with 5× digital zoom for the detection of beads. The same parameters, such as exposure time and excitation power, were used for the sample set to be compared, except for GFP signals. Analyses were performed semi-automatically by a customized macro using Image J (Fiji). Background subtraction using the rolling ball method (radius = 20 to 100 pixels (px)) was applied to images to reduce the background noise. Regions of interest (ROIs) of beads or HEK293 cells expressing receptors were determined by the signal intensity and area size fulfilling the threshold (Otsu or Triangle) and particle definition. For quantification of signals on HEK293 cells, the signal intensity within the ROI was determined by subtracting the mean signal intensity outside the ROI. For quantification of signals on beads and to minimize non-specific signals associated with the cross-reactivity of beads with antibodies, the signal intensity was determined by subtracting the mean intensity of beads located in the field where no cells were present (Fig. 2B and figs. S5, A and B, and S6, B and C). For fig. S6C, ROIs of beads were manually classified into two groups, those contacting and non-contacting with PV⁺ interneurons, which were identified by PV and GluA4 immunoreactivities. The signal intensity was calculated by subtracting the intensity on non-contacting beads from that on contacting beads.

**Immunocytochemistry and immunohistochemistry**

**Hippocampal culture**

Dissociated hippocampal neurons were prepared from E17 embryos, infected by AAV CAG-GFP at DIV12 to express GFP (Fig. 2C and fig. S7B) and cultured for 21 DIV. They were incubated overnight (between DIV20 and 21) with 7 nM CPTX added directly to the culture medium. Neurons were either fixed with 4% PFA, permeabilized with 0.1% Triton-X 100 in PBS for 10 min, washed three times and blocked (0.1% Glycine + 0.1% Tween-20 + 10% normal goat serum in PBS for fig. S7A) for 45 min, or fixed with 4% PFA, fixed with chilled methanol, permeabilized and blocked (2% Goat serum 2% BSA + 0.1% Triton-X 100 in PBS for Fig. 2C and fig. S7B) for 30 min at RT. Cells were incubated 1 hour or overnight with primary antibodies diluted in the blocking buffer and washed with PBS three times. Cells were incubated with respective secondary antibodies for 1 hour at RT. Coverslips were washed with PBS, incubated with or without DAPI (Invitrogen, cat. no. D1306) for 10 min and mounted using Fluoromount aqueous mounting media (Sigma Aldrich, cat. no. F4680) or Fluoromount-G (SouthernBiotech). For the acquisition of the images in fig. S7A, a complete Z stack spanning primary hippocampal cultures was acquired on a Leica TCS SP8 3X microscope equipped with a 405 nm diode laser and a white light laser (WLL) for excitation. The format for all images was set to 1024 × 1024 pixels with an optical zoom of 2× resulting in a pixel size of 119 nm in the x-y plane. Images were taken at 400 lines/s and line averaging of 4. For figs. S10B, S11, S12, and S17, 1 day after the unilateral hippocampal injection of CPTX into WT mice, and for fig. S18, 3 days after the unilateral hippocampal injection of CPTX into 5xFAD mice (11 to 12 months old), the brain was freshly frozen with liquid nitrogen after decapitation under deep isoflurane anesthesia. Sections (10 to 20 μm thick) were cut by cryostat (Leica), mounted on glass slides or coverslips and postfixed with 4% PFA in sodium phosphate buffer (PB; pH 7.2) or 3% glyoxal for 15 min to 2 hours at RT. Following washing with PBS + 0.1% Triton X-100, sections were treated with 10% donkey serum for 30 min at RT, incubated with a mixture of primary antibodies overnight and with a mixture of respective secondary antibodies for 2 hours. Finally, the sections were attached.
to the glass slides and mounted with Fluormount-G, VECTASHIELD (Vector Laboratories) or ProLong gold (Invitrogen). Fluorescent images were captured using confocal microscopy (SD-OSR, Olympus or LSM880, ZEISS) using the same parameters as objectives, zoom, laser power, exposure time, gain and offset for the comparison of samples. Super-resolution images were obtained by structural illumination with SD-OSR (Objective: 100×/1.49 oil with 3× zoom lens, pixel size: 40 nm/px, configuration: XY < 120 nm, Z < 300 nm) or by Airyscan with LSM880 (63×/1.40 oil with 3× digital zoom, 35 nm/px, XY < 120 nm, Z < 350 nm). Images obtained by Airyscan were 3D-reconstructed by Imaris (Bitplane).

For the quantification of pre- and postsynaptic molecule clustering (figs. S12, S17, and S18), images were obtained using a 100× objective around the area of the injected side of the hippocampus (stratum radiatum or stratum lacunosum moleculare) and the corresponding area of the contralateral uninjected or control-injected side. Injected areas were confirmed by the HIS immunoreactivity. The dynamic range of signals was first determined by maximal and minimal signal intensities in pair images of the injected and the contralateral area and then each signal intensity was normalized to this value (max-min). Normalized intensity histograms were calculated by dividing the normalized signal intensity into 40 bins on a logarithmic scale. Thus, the signal intensity in the nth bin equals $10^{\alpha x (max-min) / 40}$. To analyze the bright pixels, the mean intensity + 3 SD of the contralateral area was used as the threshold. Selected pixels were analyzed as particles using the “Analyze Particles” command within Image J without any constraint on size or circularity, leading to the calculation of particle size, number, area fraction and intensity.

**Cerebellar sections**

One day after the cerebellar injection of vehicle or CPTX into Cbln1-null mice or GluD2-null mice (3 or 7 months old), they were transcardially perfused with 4% PFA in 0.1 M sodium phosphate buffer (PB; pH 7.2) for 10 min under deep pentobarbital anesthesia, followed by post-fixation of dissected brain samples with 4% PFA in PB for 2 hours at 4°C. Alternatively, the brain was freshly frozen with liquid nitrogen after decapitation under deep isoflurane anesthesia. Free-floating sections (50 μm thickness) from 4% PFA fixed brains were prepared with a microslicer (DKT-1000, Dosaka EM), followed by treatment with 1 mg/ml pepsin (Dako) in PBS containing 0.1% Triton X-100 and 0.2 N HCl for 10 min at 37°C, washing of the sections with PBS containing 0.1% Triton X-100 and blocking with 10% donkey serum for 30 min at RT. Sections of freshly frozen brains were prepared by cryostat (Leica) at 20 μm thickness, mounted on glass slides and fixed with 3% glyoxal for 15 min, followed by blocking with 10% donkey serum for 30 min at RT and permeabilization with PBS containing 0.1% Triton X-100 (wash buffer). Sections were incubated with a mixture of primary antibodies diluted in wash buffer overnight, washed three times, incubated with a mixture of respective secondary antibodies in wash buffer and again washed three times. Finally, the sections were mounted with VECTASHIELD (Vector Laboratories). Fluorescent images were captured using a confocal laser scanning microscope (FV1000, Olympus) using the same parameters such as objectives, zoom, laser power, gain and offset for the comparison of samples. Colocalization of signals was analyzed by the plugin “Coloc 2” in Image J (87).

**Cortical sections**

One-month-old mice, into which miR vectors were transfected by in utero electroporation, were transcardially perfused with 3% glyoxal for 10 min under deep pentobarbital anesthesia, followed by post-fixation of the dissected brain with 3% glyoxal for 2 hours at 4°C. Free-floating sections (100 μm thickness) were prepared with a microslicer (DKT-1000, Dosaka EM). Following washing of the sections with PBS containing 0.1% Triton X-100 (wash buffer), sections were subsequently treated with 10% donkey serum for 30 min at RT, incubated with a mixture of primary antibodies diluted in wash buffer overnight, washed three times, incubated with a mixture of respective secondary antibodies in wash buffer and again washed three times. Finally, the sections were attached to glass slides and mounted with VECTASHIELD (Vector Laboratories). Z-stacked fluorescent images of NPY and NRP were obtained throughout the brain slice in the transfected cortical area at every 2 μm using a confocal microscope (SD-OSR, Olympus) using 20× objectives. Soma area regions of interest (ROIs) in layers 2/3 with abundant expression of NPY/NRP were found in a large population of neurons and were manually selected in an unbiased manner according to the GFP signal. Signal intensities of NPY/NRP were quantified using Image J. Background signals were subtracted using the rolling ball method (radius = 50 px) and the signals were normalized to the surrounding signals enlarged to 5 px because the absolute signal intensity was strongly affected by the z-position of the ROI in the brain section at 100 μm thickness.

**Spinal cord sections**

Two to 5 days after the injection the mice were transcardially perfused with 3% glyoxal for 10 min under deep pentobarbital anesthesia, followed by post-fixation of the dissected spinal cord with 3% glyoxal overnight at 4°C and by cryoprotection in 30% sucrose/PBS for several days. Spinal cords embedded in Tissue-Tek O.C.T. compound (Sakura Finetek, cat. no. 4583) were horizontally or coronally sectioned by cryostat (Leica) at 20 to 40 μm thickness in 4 to 5 mm around the epicenter of the injury and the slices were mounted on APS-coated glass slides (Matsunami) at 200 to 400 μm intervals. Following washing with PBS + 0.1% Triton X-100, sections were treated with 10% donkey serum for 30 min at RT and incubated with a mixture of primary and respective secondary antibodies overnight. Finally, the sections were attached to glass slides and mounted with Fluormount-G (SouthernBiotech). Fluorescence images of all sections were captured and digitized as a virtual slide (OlyVIA, Olympus) using a conventional fluorescence microscope (BX63, Olympus) with a 4× objective to determine the section at 1.4 to 1.6 mm upstream of the precise epicenter of the injury. Fluorescence images of the section were subsequently captured using a confocal microscope (SD-OSR, Olympus; 63× objective) around the gray matter ventral root using the same parameters such as objectives, laser power, exposure time, gain and offset for the comparison of samples. The super-resolution images were obtained by Airyscan 2 with a confocal laser scanning microscope (LSM980, ZEISS; 63×/1.40 oil objective with 2.5x digital zoom, 35 nm/px, XY < 120 nm, Z < 350 nm). To define the particles of GluA4, VGlut2 and their intersections, the fluorescent images were subjected to background subtraction (50 px), Laplace filtering ((9 x 9), box filtering (5 x 5), auto thresholding (Otsu method), particle extraction (> 5 px), binarization and segmentation by watershed. The defined particles were used as ROI to quantify the size and mean intensity of each channel. The percentage of GluA4+/VGlut2+ -double-positive puncta was calculated as the number of VGlut2 particles accompanied by more than 1 px of defined GluA4 particles in the ROI, divided by the total number VGlut2 particles. All image processing was performed by ImageJ. For fig. S16, D and E, slices of the spinal cord and hippocampus from the same mouse were mounted on glass slides, stained using the same procedures and the images were taken using the same microscope settings.

**Golgi-Cox staining and dendritic spine analysis**

WT and 5xFAD mice (11 to 12 months old) were sacrificed and decapitated 3 days after CPTX or Mock injection. Brains were quickly removed from the skull and washed with distilled water to remove blood from the surface. Golgi-Cox impregnation of neurons was performed using the FD Rapid GolgiStain kit (FD NeuroTechnologies, cat. no. PK401). Dye-impregnated brains were rapidly frozen on methyl butane and dry ice, embedded in Tissue-Tek O.C.T. compound (Sakura Finetek, cat. no. 4583), cryo-sectioned coronally at 100 μm...
thickness and directly mounted on gelatin-coated slides (FD NeuroTechnologies, cat. no. PO010) with the help of solution C provided in the kit. Sections were stained according to the manufacturer’s protocol and mounted using the ROTI Histokit embedding medium (Carl Roth, cat. no. 6638). Secondary apical dendrites from CA1 pyramidal neurons with cell bodies located in the dorsal part of the hippocampus were imaged for spine analysis. Sections were imaged by an experimenter blinded to experimental conditions using a Leica DMRXe microscope (Leica) as Z stacks with 0.25 μm intervals and at a magnification of 100×. 7-8 dendrites of 60 to 80 μm length were imaged for each animal. The analysis was done by an experimenter blinded to experimental conditions using the Neuro lucida software v11 (MBF Bioscience). Dendritic spines were defined as the protrusions greater than 0.4 μm that emerged from the dendrites and were identified going through the focal planes of the Z stacks. To facilitate visual inspection of the Z-minimal projections in Fig. 4A and of the 3D images in Movie S4, the background was removed using the rolling ball method (radius = 30 px) and images were corrected using an unsharp mask filter (radius = 2 px, mask weight = 0.7) in Fiji (87). Then, 3D projections (made using the 3D projection tool in Fiji) were additionally processed using the rolling ball method (radius = 15 px), an unsharp mask filter (radius = 1 px, mask weight = 0.6) and contrasted to remove a remaining blur around dendritic shafts and enhance the visibility of small protrusions.

Electron microscopy
For the in vivo Cbln1 and CPTX administration experiments, Cbln1-null or GluD2-null mice (C57BL/6J genetic background, 3 to 6 months old) were perfused transcardially with 2% PFA + 2% glutaraldehyde in 0.1 M PB (pH 7.2) under deep pentobarbital anesthesia 3 days after the injection into the cerebellum. Parasagittal microslicer sections of the cerebellum (300 μm thickness) were postfixed for 2 hours with 1% OsO4 in 0.1 M PB. After block staining in 1% aqueous uranyl acetate solution and dehydration with graded alcohols, the sections were isolated and placed on an agar block to cut transverse slices (350 μm thick) using a vibrating microtome (VT1200S, Leica) in an ice-cold solution containing 2 mM KCl, 1 mM MgCl2, 2 mM MgSO4, 1.25 mM NaH2PO4, 26 mM NaHCO3, 1 mM CaCl2, 10 mM D-glucose, and 230 mM sucrose for mice younger than 6 months or 2.5 mM KCl, 10 mM MgSO4, 12 mM NaH2PO4, 30 mM NaHCO3, 0.5 mM CaCl2, 20 mM HEPES, 25 mM glucose, 93 mM NaMDG, 5 mM sodium ascorbate, and 3 mM sodium pyruvate for 15- to 18-month-old mice (pH 7.4 with HCl). All solutions were saturated with 95% O2 and 5% CO2 and the osmolality was adjusted to 300 ± 5 mOsm. For in vitro CPTX treatments, slices were incubated in a 3-mL chamber for 4 hours at room temperature in the artificial cerebrospinal fluid (ACSF) solution containing 124 mM NaCl, 2.5 mM KCl, 1.3 mM MgSO4, 1 mM NaH2PO4, 26.2 mM NaHCO3, 2.5 mM CaCl2, and 11 mM D-glucose. CPTX was added at 20 μg/ml to slices in the chamber, whereas control slices were incubated in ACSF without CPTX. Next, the slices were transferred to the recording chamber and were continuously perfused with the same ACSF for mEPSC measurements. Picrotoxin (50 μM, Tocris Bioscience), CGP 55845 (3 μM, Tocris), and tetrodotoxin (Tocris) were added to ACSF to block GABAa, GABab receptor and Na+ channels, respectively, and hence to isolate mEPSCs. Whole-cell patch recordings from CA1 pyramidal neurons were obtained using glass electrodes (4 to 6 MΩ, outer diameter of 1.5 mm, wall thickness of 0.315 mm; Hilgenberg). The glass electrodes were filled with a solution containing 140 mM K-gluconate, 8 mM NaCl, 0.2 mM CaCl2, 10 mM HEPES, 5 mM lidocaine N-ethyl bromide (QX-314 Br), 0.5 mM Na3GTP, and 2 mM MgATP (pH 7.2, 290 ± 3 mOsm). The membrane potential was clamped at −70 mV. For miniature inhibitory post synaptic current (mIPSC) recordings, NBQX (25 μM, AMPAR antagonist, Tocris), D-AP5 (50 μM, NMDAR antagonist, Tocris) and GTP 58546 were added to the ACSF solution. The glass electrodes were filled with a solution containing 120 mM CsCl, 8 mM NaCl, 0.2 mM MgCl2, 10 mM HEPES, 2 mM EGTA, 5 mM QX-314 Br, 0.3 mM Na3GTP, and 2 mM MgATP (pH 7.2, 290 ± 3 mOsm). mEPSC and mIPSC events were analyzed offline using the Mini Analysis software (v6.0.3, Synaptosoft).

ACSF solution containing 120 mM NaCl, 2.5 mM KCl, 1.5 mM MgCl2, 125 mM NaH2PO4, 24 mM NaHCO3, 2 mM CaCl2, and 25 mM D-glucose was used for field excitatory postsynaptic potential (fEPSP) recording. Thin glass electrodes (outer diameter 1.5 mm, wall thickness 0.188 mm, −2 MΩ) filled with ACSF were used for stimulation and recording of fEPSPs. The Shaffer collaterals pathway was stimulated using a glass electrode and theta-burst stimulation (TBS) trains were applied three times to induce LTP (83). The stimulation intensity was determined based on the input-output curve and was set to give fEPSPs with a slope of ~50% of the supramaximal fEPSP. Single stimuli were repeated every 20 s for at least 10 min for baseline recording before and for 60 min after LTP induction. The paired-pulse ratio (PPR) was evaluated at 50-ms intervals under the same conditions. All recordings were obtained at 30 ± 1°C using an EPC-10 amplifier (HEKA Elektronik). The recordings were filtered at 1 to 3 kHz and digitized at 10-20 kHz.

Cerebellum
Parasagittal cerebellar slices (200 μm thick) were prepared from 4- to 6-week-old GluD2-null or Cbln1-null mice at 3 days after vehicle, Cbln1, NPtX-3Cl or CPTX injection, as described previously (84). Whole-cell patch-clamp recordings were made from PCs using a 60× water-immersion objective attached to an upright microscope (BX51WI, Olympus) at RT. The resistance of the patch pipettes was 1.5–3 MΩ when filled with an intracellular solution composed of 150 mM Cs-glucocan, 10 mM HEPES, 4 mM MgCl2, 4 mM Na2ATP, 1 mM Na3GTP, 0.4 mM EGTA, and 5 mM QX-314 Br (pH 7.3, 298 mOsm/kg) for measurements of the input-output relationship and paired-pulse ratio of PF-EPSC amplitudes, and of 65 mM Cs-methanesulfonate, 65 mM K-glucocan, 20 mM HEPES, 10 mM KCl, 1 mM MgCl2, 4 mM Na2ATP, 1 mM Na3GTP, 5 mM sucrose, and 0.4 mM EGTA (pH 7.25, 295 mOsm/kg) for long-term depression (LTD) recordings. The solution used for the slice storage and recording consisted of 125 mM NaCl, 2.5 mM KCl, 2 mM CaCl2, 1 mM MgCl2, 1.25 mM NaH2PO4, 26 mM NaHCO3, and 10 mM D-glucose, bubbled continuously with a mixture of 95% O2 and 5% CO2. Picrotoxin (100 μM, Sigma Aldrich) was always present in the saline to block the inhibitory inputs. To evoke the PF-EPSCs, square pulses were applied at various stimulus intensities (10 μs, 0 to 200 μA) through a stimulating electrode placed on the molecular layer (−50 μm)
away from the pial surface). Selective stimulation of the PFs was confirmed by paired-pulse facilitation (PPF) of EPSC amplitudes at a 50-ms inter-stimulus interval. Current responses were recorded with an Axopatch 200B amplifier (Molecular Devices), and pClamp software (v9.2, Molecular Devices) was used for data acquisition and analysis. Signals were filtered at 1 kHz and digitized at 4 kHz.

**CPTX injection**

### Acute intrahippocampal injection

We used a digitally controlled infusion system (UltraMicroPump III with Micro4 Controller, World Precision Instruments) fed with a 10 μl Hamilton syringe and a NanoFil (35 gauge) beveled needle. The mice were first anesthetized with 1 to 3% isoflurane and placed into the stereotaxic frame (Narishige). 1 ml CPTX (140 mg/ml) or vehicle (HBS buffer) was bilaterally injected at a rate of 3 nl/s. We used the following coordinates for bilateral injection: AP = ±2.0 mm from Bregma and L = ±1.5 mm; DV = 2.0 mm from the brain surface according to the mouse brain atlas ([85](#)).

### Acute intracerebellar injection

Mice were anesthetized with an intraperitoneal injection of ketamine/xylazine (80/20 mg/kg body weight, Daichi-Sankyo or Sigma Aldrich). A small hole in the occipital bone was made with a dental drill, and the dura matter was ablated as described previously ([36](#)). A glass needle was inserted into the vermis of lobule VI/VII at ± 0.7 mm from the midline at a depth of ~200 μm. Approximately 7 μl of vehicle, Cbln1-HIS or CPTX-HIS (7.7 μM) was bilaterally injected at a rate of 10 to 40 μl/hour.

### Acute intrahippocampal injection through guide cannulas

Intrahippocampal guide cannulas were implanted as described previously ([86-88](#)) with minor changes. In brief, mice were anesthetized with 1 to 3% isoflurane mixed with O2 through a vaporizer (Matrix VIP 3000, Midmark). Prior to any surgical manipulation, the mice were given the analgesic carprofen (5 mg/kg b.w., s.c., Rimadyl, Pfizer Pharma) as a postoperative analgesic. The intrahippocampal injections and behavioral tests were performed after the mice had fully recovered at least 5 to 7 days after surgery. Injections were done using a 10 μl Hamilton syringe as described in the **Acute intrahippocampal injection** section but being inserted in the guide cannula under brief sedation with 1% isoflurane.

### Acute intraspinal injection to SCI mice

A volume of 0.5 μl of CPTX (1 μg/μl), Cbln1 (1 μg/μl), Chondroitinase ABC (0.5 U/μl; Sigma Aldrich cat. no. C2905) or vehicle (HBS buffer) were injected in the proximity of the injury site by electrical micro-injector (BJ-110, BEX Co.) through glass capillaries (Drummond Scientific Company, cat. no. 3-000-203-G/X). Following injection, the muscle layers and skin were again closed by suturing.

### In utero electroporation

In utero electroporation (IUE) was performed at E14.0 in the cortex of ICR mice as described previously ([89](#)). Briefly, plasmid DNA was injected into the third ventricle by a glass pipette and a set of electrical pulses was applied 5 times. Positive electrodes were placed onto the cortical side. Plasmid DNA was dissolved in 21 mM HEPES, 137 mM NaCl, 5 mM KCl and 0.7 mM Na2HPO4 at a concentration of 1 mg/ml.

### Surgical procedure for spinal cord injury (SCI)

Mice (9 to 11 weeks old) were subjected to hemisection- or compression-induced SCI. After the anesthesia, the spinal cord was surgically exposed and the dorsal column at the 10th thoracic vertebra (T10) was dissected by micro-scissors and a scalpel. The muscle layers and skin were closed by sutures. Animals were recovered from the anesthesia by the administration of the antagonist. Mice were subjected to compression-induced damage using an SCI impactor device (Infinite Horizon Impactor; Precision Systems and Instrumentation) ([90](#)) and using a 70-kdyn impact force. We monitored the strength and the duration of the impact compression through the automated recording system in the SCI impactor device.

**Behavioral Analysis**

### Spatial navigation in a labyrinth

The 3D-printed dry maze was developed at the DZNE. In short, before implantation of cannulas, mice were put on the light food restriction diet (LFRD) scheme where ad libitum food was removed from mice and was given daily manually at the same time in amounts comparable with normal consumption (4 to 5 g/day/mouse). The weight of all mice was monitored daily and kept within a 10 to 15% reduction. Before the cannulation of mice, labyrinth habituation and training was performed for 2-3 weeks. Once all mice were familiar with the labyrinth, the room was changed to introduce new landmarks and cues, and the configuration of the labyrinth was changed to be more complex ([Fig. 5A](#)). Testing in the labyrinth began on day 3 after CPTX or Mock injection. Mice were placed into a starting zone in the labyrinth and given 20 min to find a hidden colorless and odorless round pellet reward (AIN-76A 10 mg tablet, TestDiet, cat. no. 1811213). Two hours later the mice were placed again in the same labyrinth with the same room landmarks and cues for 10 min to find the reward. On day 4, the mice were placed again into the same labyrinth in which the reward zone was changed, and the mice had to relearn the new position of the reward within 10 min. Two hours later, during the retrieval session (5 min), the mouse had to find the new location of the reward. The labyrinth was cleaned with 60% ethanol between trials.

The position of mice in the labyrinth was recorded using a USB video camera (Microsoft) and special behavioral video acquisition and analysis software (ANY-maze). All recorded movies were analyzed using ANY-maze by a trained observer blind to the mouse genotype and treatments. ANY-maze was run in tracking mode to trace coordinates of mice (10 frames/s) and compute distance traveled from the starting zone to the hidden reward zone. The criteria for successful performance were that during encoding and retrieval sessions, the hidden rewards (7 to 8 pellets) were found and eaten. Travel distances in meters were computed as follows: mice had to find the reward and take at least 1 pellet. This time point, as a beginning of the association of the reward location in the labyrinth and all
To validate the action of CPTX in another hippocampus-dependent cognitive task, we chose contextual fear conditioning (CFC). Shortly after spatial navigation in the labyrinth, at day 5 all mice were first recorded in a neutral context (context B, 5 min) and then 1 hour later they were put into a conditioned context (context A, 5 min) and given three foot shocks, each with 0.5 mA and 1 s in duration and separated by 1 min. Contexts A and B were different in terms of walls (chess-like black-and-white pattern versus grey color), smell (different cleaning solutions) and floor (different cleaning solutions) and floor like black-and-white pattern versus grey color), and B were different in terms of walls (chess-like black-and-white pattern versus grey color), smell (different cleaning solutions) and floor like black-and-white pattern versus grey color). Then, mice were kept for 1 hour in context B, and then at least 1 hour later in the context B for 5 min for retrieval sessions. Context A served as the fraction of the time a mouse was freezing in contexts A and B. Freezing of mice was defined as total immobility of animals (except for breathing) with a characteristic tense fear posture. The analysis of data was done by using ANY-maze software. The fear conditioning system consisted of a touch-pad controller and a conditional sound-attenuated cabinet and chambers (Ugo Basile).

Rotor-rod test and gait analyses

To examine the skilled motor coordination in the mice, the rotor-rod test was performed. As this phenotype is clearly observed on the ICR genetic background (13, 16, 84), we used in 4- to 6-week-old Cbln1-null mice with this background. The rotor-rod test was performed 1 day before and 3 days after the injection. Six trials were performed at 20 rpm with a 30-s interval between trials, and the latency to fall from the rotor-rod was measured (maximum score; 120 s). For gait analyses, we used 4- to 6-month-old Cbln1-null and Ghδ2-null (C7BL/6J) mice. In addition, 4- to 6-week-old Cbln1-null mice (ICR) were used. Mice were habituated to the experimenter and the behavior room from 2 to 3 days before the test. Then, the mice were habituated to the gait analysis apparatus made of transparent plexiglas (6 cm × 85 cm) and trained to walk straight to their home cage. Each step during the walking was recorded by a video camera across the transparent floor at 30 frames per second. The time points and locations of the hind limb to attach and detach on the floor were manually determined by an experimenter blinded to experimental conditions, using custom software written in HSV (Hot Soup Processor, ONION software). These analyses were performed > 4 times for each mouse 1 day before and 3 days after Mock, Cbln1 or CPTX injection. Gait parameters were calculated as follows: (i) the coordinated step ratio was calculated as the number of alternate left/right steps divided by the number of total steps; (ii) the stride length of each step along the x-axis (plus direction to home cage); (iii) the stride speed and (iv) stride irregularity (the standard deviation of stride length during the single journey divided by the average stride length).

Locomotor recovery after SCI

The locomotor recovery assessment was performed using video recording as previously reported (90). BMS (Basso Mouse Scale) open-field scoring and footfall tests were performed weekly during the 6 to 8 weeks following SCI. The BMS score was evaluated by at least two independent investigators who were blinded to the experimental groups. Mice were excluded if they had an incomplete injury (BMS score > 0) on day 2 after SCI. To quantify the recovery effect, the increase in BMS score between the 1st and 2nd week after SCI was calculated as ABMS (week 1 − BMS score (2nd week)) = BMS score (2nd week) − BMS score (1st week).

For footfall tests, mice were placed on a wiremesh grid and videotaped for 5 min while on the grid. Mice that walked longer than 3 min with more than 70 steps were subjected to scoring by three independent examiners. The total number of footfalls from the bars during the total walking time was counted.
W.K.), the Takeda Science Foundation (to M.Y. and W.K.), the Marie-Curie Actions postdoctoral fellowship (grant 328531 to J.E.), the University of Bordeaux Initiative of Excellence (IdEx) fellowship (to J.E.), the European Research Council (ERC Starting Grant 850820 SynLink to J.E.), the European Union 7th Framework Programme Initial Training Network (EU FP7 ITN: grant 606950 EXTRABRAIN to A.D.), and the Bundesministerium für Bildung und Forschung (BMBF: EnergI project, TP5, 01GQ1421A to A.D.).

**Author contributions:** K.S. performed cell biological experiments, immunohistochemical experiments of hippocampus, cerebellum, and spinal cord tissues, and behavioral studies of cerebellar function; A.J.C., J.E. and A.R.A. collected x-ray data and solved the NP1PTX crystal structure; A.R.A. conceived the CPTX organizer; J.E. performed SPR and MALS experiments; W.K. performed electrophysiological experiments in the cerebellum; I.S. performed electrophysiological experiments in the hippocampus; J.E., V.T.C., and A.J.C. prepared recombinant proteins; O.S. developed a 3D-printed labyrinth and performed behavioral studies of hippocampal function; M.F.-F. and S.S. performed spine analysis; E.M. performed electron microscopic analyses and immunohistochemical experiments of the spinal cord; R.K. performed immunocytochemical and immunohistochemical analyses of hippocampal neurons; K.M. performed immunocytochemical, immunohistochemical and biochemical analyses; K.S. and T.S. performed cell-based binding assays; S.O. developed analytical tools for gait performance; H.S., M. I., Y.M., Y.T. and K.T. generated the experimental models of spinal cord injury and performed behavioral analyses. M.W. raised new antibodies against neuronal pentraxins; A.D., A.R.A., and M.Y. designed and supervised the project; K.S., J.E., A.D., A.R.A., and M.Y. wrote the paper.

**Competing interests:** K.S., K.T., and M.Y. are inventors on Japan patent application 2020-2879 (10 January 2020) describing the use of synthetic synapse connector for spinal cord injury. **Data and materials availability:** All data are available in the main text or the supplementary materials. All the plasmids and antibodies generated in this study are available from the authors upon request. Structure factors and coordinates of NP1PTX are deposited in the Protein Data Bank (PDB code 6YPE).

**SUPPLEMENTARY MATERIALS**

science.sciencemag.org/content/369/6507/eabb4853/suppl/DC1

Figs. S1 to S22

Table S1

Movies S1 to S8

View/request a protocol for this paper from Bio-protocol.

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A synthetic synaptic organizer protein restores glutamatergic neuronal circuits


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Synthetic excitatory synaptic organizer

The human brain contains trillions of synapses within a vast network of neurons. Synapse remodeling is essential to ensure the efficient reception and integration of external stimuli and to store and retrieve information. Building and remodeling of synapses occurs throughout life under the control of synaptic organizer proteins. Errors in this process can lead to neuropsychiatric or neurological disorders. Suzuki et al. combined structural elements of natural synaptic organizers to develop an artificial version called CPTX, which has different binding properties (see the Perspective by Salinas). CPTX could act as a molecular bridge to reconnect neurons and restore excitatory synaptic function in animal models of cerebellar ataxia, familial Alzheimer's disease, and spinal cord injury. The findings illustrate how structure-guided approaches can help to repair neuronal circuits.

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