

# Chapter 2.1

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## **Motor Learning Requires Purkinje Cell Synaptic Potentiation through Activation of AMPA-Receptor subunit GluA3**

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### SUMMARY

Accumulating evidence indicates that cerebellar long-term potentiation (LTP) is necessary for procedural learning. However, little is known about its underlying molecular mechanisms. Whereas AMPA receptor (AMPA) subunit rules for synaptic plasticity have been extensively studied in relation to declarative learning, it is unclear whether these rules apply to cerebellum-dependent motor learning. Here we show that LTP at the parallel fiber to Purkinje cell synapse and adaptation of the vestibulo-ocular reflex do not depend on GluA1-, but on GluA3-containing AMPARs. In contrast to the classic form of LTP implicated in declarative memory formation, this form of LTP does not require GluA1-AMPARs trafficking, but rather changes in open-channel probability of GluA3-AMPARs mediated by an increase of cyclic AMP and activation of the protein directly activated by cAMP (Epac). We conclude that vestibulo-cerebellar motor learning is the first form of memory acquisition shown to depend on GluA3-dependent synaptic potentiation through an increase in channel conductance.

### INTRODUCTION

Plasticity mediated by synaptic trafficking of  $\alpha$ -amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid (AMPA) type glutamate receptors (AMPA) plays an important role in the acquisition of declarative memories (Kessels and Malinow, 2009). Ionotropic AMPARs drive fast excitatory neuronal activity and can consist of four different subunits named GluA1 to GluA4. In hippocampal pyramidal cells, most AMPARs are hetero-oligomers composed of either GluA1/GluA2 or GluA2/GluA3 subunits and the subunit composition dictates which role AMPARs play in synaptic plasticity (Shi et al., 2001). In the hippocampus, cortex and amygdala, both LTP and learning depend on the trafficking of GluA1-containing AMPARs to synapses (Makino and Malinow, 2011; Nedelescu et al., 2010; Rumpel et al., 2005; Mitsuhashi et al., 2011), whereas GluA3-containing AMPARs contribute relatively little to synaptic currents, synaptic plasticity or learning (Adamczyk et al., 2012; Meng et al., 2003; Humeau et al., 2007). To what extent GluA1-plasticity and GluA3-plasticity play a role in adaptive motor behavior, remains to be established.

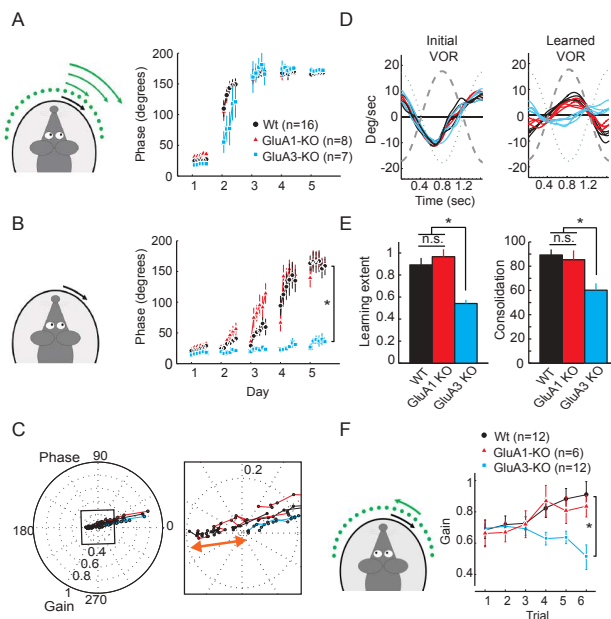
Here, we sought to unravel the potential role of GluA1- and/or GluA3-containing AMPARs in cerebellar motor learning. Unlike the rich insight in the role of AMPARs in declarative memory formation in the hippocampus, relatively little is known about their role in procedural memory formation in the cerebellum. AMPAR plasticity occurs at parallel fiber to Purkinje cell (PF-PC) synapses reflecting the expression of LTP or long-term depression (LTD) (Kakegawa and Yuzaki, 2005; Steinberg et al., 2006), but the full functional significance of and the precise molecular pathways underlying this plasticity remain to be further elucidated (Gao et al., 2012). In addition, the roles of GluA1- and/or GluA3-containing AMPARs in plasticity of PCs have hardly been studied (Bats et al., 2013; Douyard et al., 2007; Kakegawa and Yuzaki, 2005).

We found that adaptation of compensatory eye movements, which is one of the most widely studied forms of cerebellar motor learning serving to stabilize gaze (Anzai et al., 2010; Nguyen-Vu et al., 2013; Schonewille et al., 2011), depends on GluA3-containing AMPARs, but not GluA1-containing AMPARs. The GluA3-containing AMPARs in PCs are critical for the induction and expression of PF-PC LTP, not by trafficking of receptors, but by a change in conductance and open probability of the channel. This form of plasticity requires activation of Epac through an increase of cyclic AMP. Together, these findings do not only show that GluA3 is crucial for cerebellar potentiation and learning, but also that it evokes its actions of plasticity through a novel mechanism.

## RESULTS

## Cerebellar motor learning depends on GluA3, but not on GluA1

Unlike global knock-out mice of GluA2, which suffer from severe motor performance deficits including ataxia [Gerlai et al., 1998; Jia et al., 1996], mice that lack AMPAR subunit GluA1 or GluA3 (GluA1-KO and GluA3-KO) displayed intact basic motor behavior (Figure S1). Indeed, they were able to stabilize the images on their retina and/or gaze with respect to a moving visual field (i.e. optokinetic reflex or OKR; Figure S1A), with respect to their head movements (i.e. the vestibulo-ocular reflex in the dark or VOR: Figure S1B), or with respect to a combination of both as occurs in daily life (i.e. VOR in the light or VORL; Figure S1C). None of the comparisons between GluA1-KO, GluA3-KO and their control littermates showed a significant difference in any of these paradigms [for p-values, see Figure S1 and corresponding legends]. Far more challenging is the test for VOR phase-reversal adaptation, which involves cerebellum-dependent motor learning [Gutierrez-Castellanos et al., 2013]. During this paradigm mice learn to shift the phase of their VOR following sinusoidal visuovestibular mismatch stimulation, in which the visual stimulus moves in the same direction as the vestibular stimulus (i.e. in-phase), yet at a greater amplitude [Figure 1A]. After 5 days of visuovestibular training wild-type mice moved their eyes during table stimulation in the dark in the same direction as the body, rather



**Figure 1. GluA3 is required for oculomotor learning.**

(A) When adult (3-5 months of age) wild-type (black), GluA1-KO (red) and GluA3-KO mice (blue) are subjected to a visuo-vestibular mismatch training paradigm, in which the visual screen rotates sinusoidally in the same direction as the turntable, but at an increasingly greater amplitude (also referred to as phase-reversal task), they show a similar ability to follow the training signal over time as long as the light is on. Eye movement signals are expressed as phase values [in degrees] with respect to those of the turntable, which also rotates in a sinusoidal fashion [i.e., 360 degrees represents one sinusoidal cycle].

(B) However, when the light is turned off, but the turntable stimulus continues [i.e. the VOR adaptation catch trials of the phase-reversal task], the phase values of the GluA3-KO mice show significantly impaired motor learning compared to those of GluA1-KO and WT mice.

(C) Polar plot showing the trajectory of VOR gain and phase change during adaptation for WTs (black line), GluA3-KOs (blue), and GluA1-KOs (red). Gain [i.e. amplitude of the eye movement divided by that of the stimulus] is represented as distance from the center and phase as the angle relative to perfect compensation at 0 degrees. The data reveal a common learning trajectory and comparable initial gain, yet a difference in learning extent between the groups. Inset: the final VOR reached after 5 days of training is amplified to visualize the magnitude of the gain difference (red arrow) between the groups tested.

(D) GluA3-KOs (blue line) were unable to reverse their VOR phase compared with WT (black) and GluA1-KOs (red). Four representative eye velocity traces per group compare the initial VOR before (left) and after (right) the mismatch training (left).

(E) Both learning extent and consolidation during the phase-reversal task are significantly smaller in GluA3-KO mice compared to WT and GluA1-KO mice (T2-test  $p < 0.05$ ).

(F) Gain-increase learning also reveals deficits in GluA3-KO, but not in GluA1-KO, as compared to WTs.

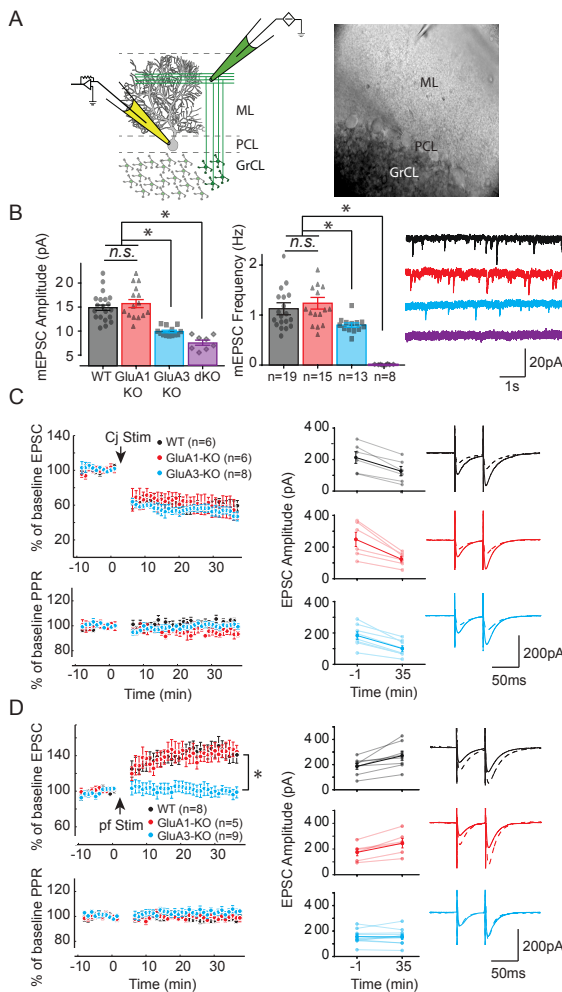
Error bars indicate SEM, \* indicates  $p < 0.05$ .

than the opposite direction as they used to do before the training (i.e. they normally show an innate contraversive compensation). More specifically, the mature wild-type mice learned to shift their VOR in the dark by  $159^\circ$  out of the perfect  $180^\circ$  after the training (Figure 1B). Likewise, when we subjected littermate GluA1-deficient mice to this phase-reversal adaptation paradigm, they reached final average phase shifts of  $162^\circ$  (GluA1-KO vs. WT:  $p=0.13$ ; Figure 1B), indicating that GluA1-containing AMPARs are dispensable for VOR adaptation. In contrast, GluA3-deficient mice showed striking deficits in shifting the phase of their VOR in the dark; they showed a final phase shift of only  $35^\circ$  after 5 training sessions (GluA3-KO vs. WT:  $p=0.001$ ; Figure 1B). When we looked not only at the oculomotor phase but also at the learning trajectory extent (as explained in Figure S2A), we observed that, although the initial performances of VOR catch learning trials were not significantly different for any of the 3 groups tested ( $p=0.3$  and  $p=0.11$  for GluA1-KO and GluA3-KO, respectively), the final performances of GluA3-KO after 5 days of training were significantly different from those of both wild-type littermates ( $p=0.01$ ) and GluA1-KO mice ( $p=0.01$ ) (Figure 1C). Accordingly, the vector of total learning extent per mouse, which equals the distance between the initial (1st recording, 1st day) and final VOR performance coordinate throughout the 5-day spanning phase-reversal paradigm (Figure S2), was significantly smaller for GluA3-KO mice compared to wild-type and GluA1-KO mice ( $p=0.001$  and  $p=0.0002$ , respectively) (Figures 1D and 1E). Moreover, the consolidation rate of learning, which equals the ratio between the total learning extent and the ideal learning extent with no overnight memory loss between training days (Figure S2A), was also significantly lower in GluA3-KO compared to both GluA1-KO and wild-type littermates ( $p=0.0008$  and  $p=0.001$ , respectively) (Figure 1E). Importantly, all groups of mice performed equally well during the visually-driven vestibular training trials over the 5 days of training (all  $p$  values  $>0.05$ ; Figure 1A), indicating that the learning deficits in GluA3-KO as measured in the dark during the catch trials did not directly result from a poor response to the visuo-vestibular training signal, but rather an impaired ability to maintain this learned vestibular response in the absence of the visual cue (Figure 1B).

In addition to phase modulation, we also investigated gain modulation of vestibulo-ocular movements after either in-phase [gain-down] or out-of-phase [gain-up] visuovestibular training paradigms that aim to reduce or increase the amplitude of the eye movement response to a constant vestibular input, respectively. GluA3-KO showed severe learning deficits in both the gain-down ( $p=0.001$  for final catch trials) and gain-up paradigms ( $p=0.009$  for final catch trials), whereas GluA1-KO and wild-types performed again equally well in both training paradigms ( $p=0.11$  and  $p=0.2$  for gain-down and gain-up final catch trials, respectively) (Figure 1F and S1D). These experiments indicate that GluA3-containing AMPARs contribute to cerebellum-dependent motor learning.

### **GluA3 is required to induce LTP, but not LTD, at PF-PC synapses**

Purkinje cells (PCs) form the sole output of the cerebellar cortex. It has previously been shown that synaptic plasticity at their parallel fiber afferents crucially contributes to motor learning (Schonewille et al., 2010). To investigate the contribution of GluA1- and GluA3-containing AMPARs to basal synaptic transmission, we recorded spontaneous miniature excitatory synaptic currents (mEPSCs) of PCs in cerebellar slices from 4-6 week old mice (Figure 2A), an age at which GluA3-KO mice showed similar motor learning deficits as during adulthood (Figure S2D). The average amplitude and frequency of mEPSCs in GluA1-deficient PCs were not significantly different ( $p=0.4$  and  $p=0.2$ , respectively) from those in wild-type PCs (Figure 2B). In PCs of GluA3-KO mice the average amplitude ( $p=0.0003$ ) and frequency ( $p=0.02$ ) of mEPSCs were significantly lower than those in wild-type PCs. The low basal transmission in the GluA3-KO mice was neither reflected in structural changes at the level of spine densities of proximal or distal PC dendrites ( $p=0.7$  and  $p=0.2$  for proximal and distal, respectively)



**Figure 2. GluA3 is required for PF to PC LTP, but not LTD.** (A) Scheme of cerebellar cortical circuitry (Left) and representative picture of the in-vitro preparation (right) showing positions of recording electrode (yellow) at PC soma and stimulus electrode (green) at parallel fiber beam. ML, PCL and GrCL indicate molecular layer, Purkinje cell layer, and granule cell layer, respectively.

(B) mEPSC amplitude (left panel) and frequency (middle panel) of both single GluA3-KO PCs (blue bar) and double GluA1/3-KO PCs (purple bar) were significantly reduced compared to those in WT PCs (black bar) [for amplitude and frequency WT vs. GluA3-KO  $p=0.0003$  and  $p=0.023$ , respectively; for WT vs. GluA1&3-dKO  $p<0.0001$  and  $p<0.0001$ , respectively] and single GluA1-KO PCs (red bar) [for amplitude and frequency GluA1-KO vs. GluA3-KO,  $p<0.0001$  and  $p=0.0032$ , respectively]. In contrast, GluA1-KO and WT PCs presented comparable basal transmission [for amplitude and frequency WT vs. GluA1-KO  $p=0.37$  and  $p=0.16$ , respectively]. Right panel shows corresponding raw traces of mEPSCs.

(C) Both GluA1-KOs (red) and GluA3-KOs (blue) show similar cerebellar synaptic weakening after LTD induction compared to WT littermates (black) (top left panel) with unchanged PPR over time (bottom left panel). EPSC magnitude was held in a comparable range for all cases to prevent potential bias due to differential basal synaptic strength (middle panels). Representative traces of paired EPSCs before (solid lines) and after LTD induction (dashed lines) (right panels, matched genotype color code). Cj Stim indicates conjunctive stimulation (so as to induce LTD).

(D) GluA3-KO PCs show severe deficits in PF to PC LTP compared with WT and GluA1-KOs with no changes in PPR or baseline EPSC magnitude. Representative traces of paired EPSCs before (solid lines) and after LTP induction (dashed lines) (same configuration as in B). pf Stim indicates parallel fiber-only stimulation (so as to induce LTP).

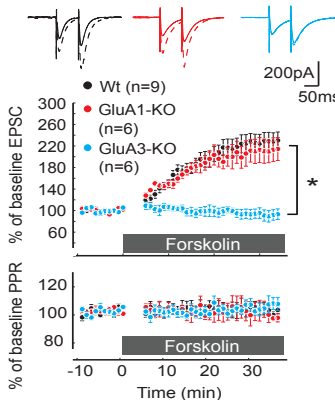
Error bars indicate SEM, \* indicates  $p<0.05$ .

(Figure S3) nor was it compensated for by increased synaptic currents from kainate receptors (for details see Figure S4 and corresponding legends). In PCs of mice that lack both GluA1 and GluA3, mEPSC events were virtually absent (Figure 2B), suggesting that the far majority of synaptic currents in PCs are derived from either GluA1- or GluA3-containing AMPARs.

A reduced basal transmission in GluA3-deficient PC synapses can either be a cause or a consequence of impaired synaptic plasticity. We therefore investigated both LTD and LTP at the PF to PC synapse, using whole-cell recordings. LTD was induced either by pairing PF stimulation with a depolarizing voltage-clamp step, mimicking climbing fiber (CF) input (Linden, 2001; Figure 2C), or by pairing PF stimulation with CF stimulation (Schonewille et al., 2011; Figure S3C-E). The magnitudes of LTD in PCs of GluA1-KO and GluA3-KO were indistinguishable from that in the PCs of wild-type littermates with either induction protocol [with somatic depolarization, for GluA1-KO vs. wild-type  $p=0.4$  and for GluA3-KO vs. wild-type  $p=0.2$ ; with direct CF stimulation, for GluA1-KO vs. wild-type  $p=0.9$  and for GluA3-KO

vs. wild-type  $p=0.8$ ). These data are in line with other studies showing that GluA2 is the key subunit for AMPAR internalization and therefore for LTD induction (Steinberg et al., 2006; Schonewille et al., 2011). Next, we induced LTP in PCs by 1 Hz tetanic stimulation of PF input alone (Lev-Ram et al., 2002) (Figure 2D). This stimulus protocol reliably produced significant LTP in both wild-type PCs ( $p=0.0005$ ) and GluA1-KO PCs ( $p=0.0003$ ) with a similar magnitude ( $p=0.5$ ), and without significant changes in paired-pulse facilitation (PPF) of the evoked EPSCs after LTP induction ( $p=0.11$  and  $p=0.6$  in GluA1-KO and wild-type PCs, respectively). In contrast, with the same stimulation protocol LTP could not be induced in GluA3-KO PCs ( $p=0.7$ ) (Figure 2D). These experiments demonstrate that PF to PC LTP requires GluA3-containing AMPARs.

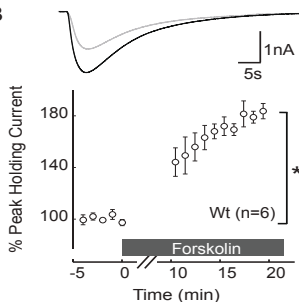
A



### GluA3-dependent synaptic potentiation involves a cAMP-driven change in channel conductance

What is the molecular mechanism underlying GluA3-dependent LTP of PF-PC synapses? To test whether GluA3-dependent synaptic plasticity in PCs depends on cAMP-signaling we administrated the adenylyl cyclase activator forskolin (FSK) to PCs of GluA3-KO brain slices and compared the effects to those in wild-type slices and GluA1-KO slices (Figure 3A). Whereas FSK produced on average a two-fold potentiation in PF-evoked EPSCs in both wild-type and GluA1-KO PCs ( $230 \pm 25\%$  and  $215 \pm 35\%$ , respectively), it failed to induce synaptic potentiation in PCs that lack GluA3 ( $95 \pm 10\%$ ;  $p=0.005$  GluA3-KO vs. wild-type). Importantly, AMPAR potentiation also occurred in wild-type PCs when local stimulation with  $1 \mu\text{M}$  AMPA was used, while blocking PFs with TTX ( $189 \pm 17\%$ ,  $p=0.001$ ; Figure 3B), highlighting its postsynaptic nature (Chen and Regehr, 1997). These data indicate that a GluA3-dependent synaptic potentiation at PF-PC synapses can occur upon a rise in the cellular level of cAMP.

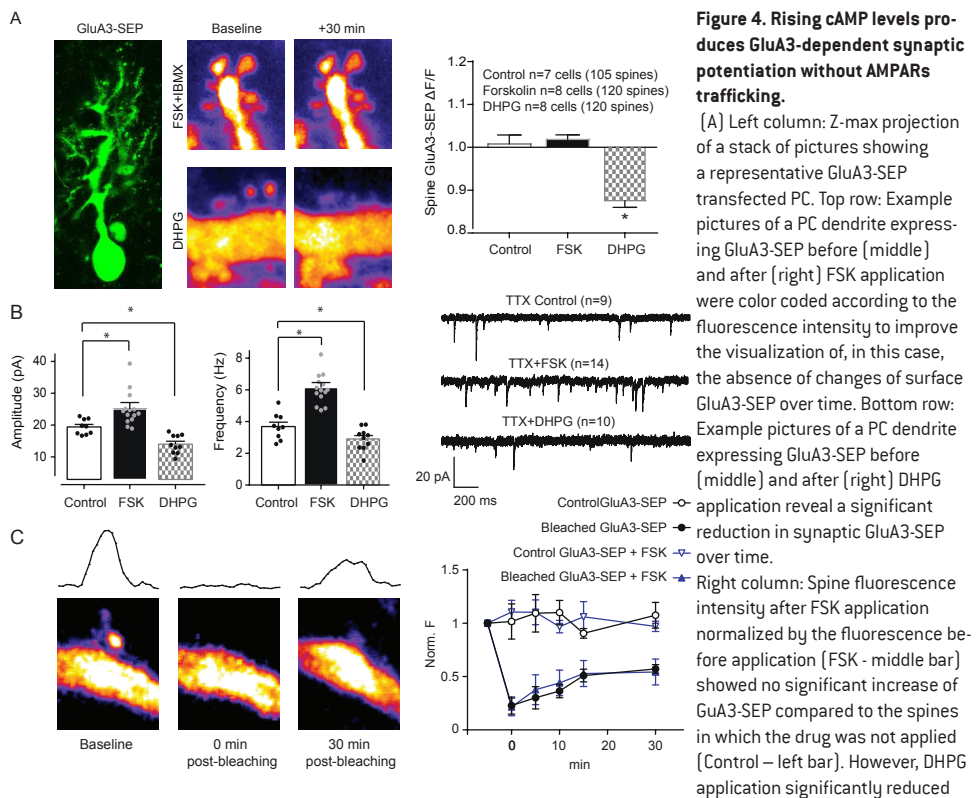
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**Figure 3. Rising cAMP levels produces GluA3-dependent post-synaptic potentiation.** [A] Wash-in of  $50 \mu\text{M}$  FSK causes synaptic potentiation at wild-type PCs (black) and GluA1-KO PCs (red), but not at GluA3-KO PCs (blue). Top, middle and bottom panel show example traces, normalized EPSC amplitude and paired pulse ratio (PPR), respectively. [B] Enhancement of currents evoked by local puffs of  $1 \mu\text{M}$  AMPA at the molecular layer following FSK application can also occur in the presence of TTX blocking PF input. Error bars indicate SEM, \* indicates  $p < 0.05$ .

We next examined whether cAMP-driven synaptic potentiation is a result of synaptic trafficking of GluA3-containing AMPARs. To assess whether FSK increases GluA3 levels on the cell surface of spines, we performed time-lapse 2-photon imaging of PCs in cultured organotypic cerebellar slices infected with Sindbis virus to acutely express GluA3 subunits fused to Super Ecliptic pHluorin (SEP). SEP is a pH-sensitive variant of GFP that shows a reduction in fluorescence upon rapid application of acidic (pH 5) ACSF (Figure S4F) (Makino and Malinow, 2009). To test whether GluA3 trafficking can be detected with this method, we first triggered LTD chemically by adding the metabotropic mGluR1 receptor agonist DHPG to induce internalization of AMPARs (Linden, 2001). Indeed, application of DHPG to wild-type PCs expressing SEP-GluA3 produced a significant decrease in SEP fluorescence at spines ( $p < 0.0001$ ; Figures 4A) and in synaptic strength ( $p=0.004$  for amplitude and  $p=0.04$  for frequency; Figure 4B), which is in line with the endocytosis of AMPARs that occurs during the expression of LTD at the PF-PC synapse

(Wang and Linden, 2000). In contrast, washing in FSK failed to induce any change in SEP-GluA3 fluorescence at PC spines ( $0.03 \pm 0.015\%$  change,  $p=0.4$ ) (Figures 4A), even though FSK significantly increased synaptic currents in the SEP-GluA3 expressing PCs ( $p=0.04$  for amplitude and  $p<0.0001$  for frequency) (Figure 4B). These data suggest that the cAMP-driven synaptic potentiation does not require an insertion of GluA3-containing AMPARs at the surface of spines. To assess whether FSK promotes lateral mobility of GluA3 receptors instead of an increase in receptor externalization, we performed fluorescence recovery after photobleaching (FRAP) experiments of single spines of PCs expressing SEP-GluA3 (Figure 4C). After 80% photobleaching, the SEP signals recovered to 50%, suggesting that a proportion of SEP-GluA3 is immobilized at synapses (Makino and Malinow 2009). The SEP fluorescence intensity recovered at a similar pace in the presence or absence of FSK ( $p=0.9$ ), indicating that the lateral mobility of GluA3-containing AMPARs is not influenced by a rise in cAMP.



GluA3-SEP in PC spines in accordance to the observed synaptic depression. (E) Lowering the extracellular pH produced a drastic reduction of GluA3-SEP fluorescence intensity corroborating its pH sensitivity.

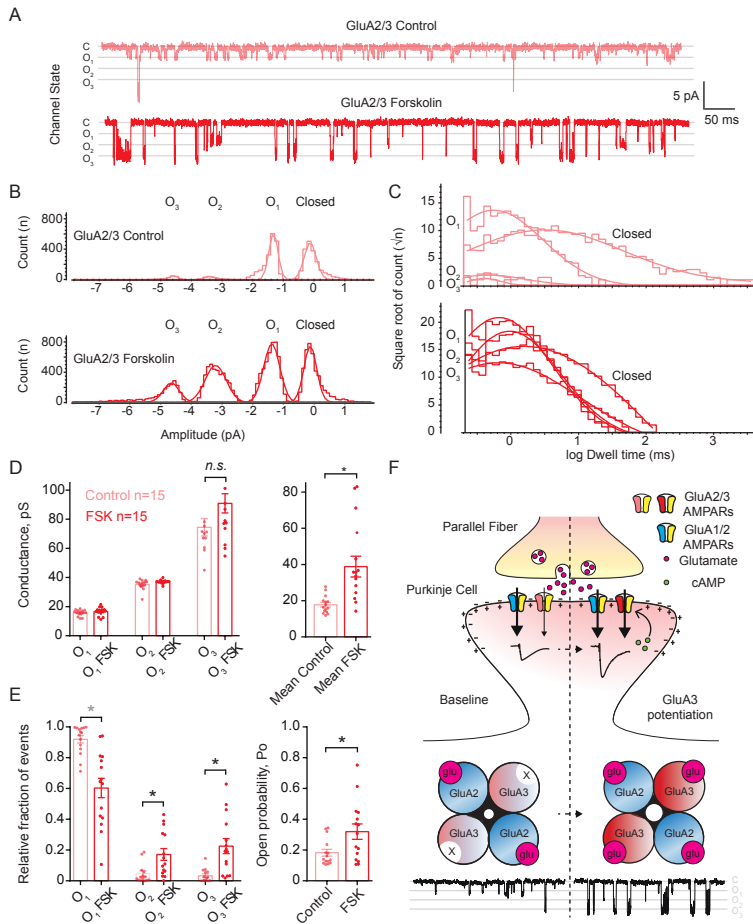
(B) Despite the lack of a detectable increase in surface GluA3-SEP, FSK produced a significant increase in mEPSCs amplitude and frequency in GluA3-SEP transfected PCs in organotypic slices. DHPG induced a significant decrease in mEPSC amplitude and frequency in these neurons.

(C) Left: Example baseline max projection Z-stack ( $3 \mu\text{m}$ , 6 optical planes) of a dendrite transfected with GluA3-SEP obtained with two-photon microscopy before, immediately after and 30 minutes after photobleaching of the spine. The black traces above the pictures represent quantifications of SEP fluorescence across the spine and parallel to the dendrite.

Right: Overall quantification of spine FRAP dynamics over time for PCs transfected with GluA3-SEP, either with (n=5) or without (n=4)  $50 \mu\text{M}$  forskolin added after the moment of bleaching (0 min). SEP fluorescence intensity is normalized to baseline intensity (-5 min). No changes in SEP intensity were observed over time in spines neighboring the bleached spines.



To assess whether GluA3-plasticity involves a change in channel properties, we resolved single-channel AMPA mediated currents by clamping a single AMPAR in cell-attached mode at the cell body of either GluA1-KO or GluA3-KO PCs with the recording pipette containing near-saturating concentrations of AMPA [Armstrong et al., 2002]. GluA1-containing AMPARs at the surface of GluA3-KO PC cell bodies stochastically reached open states 1, 2 and 3 (indicating binding of 2, 3 and 4 glutamates per receptor complex, respectively) and displayed similar conductance levels and open-channel probability in the presence or absence of FSK application (Figure S5). In contrast, GluA3-containing AMPARs on cell



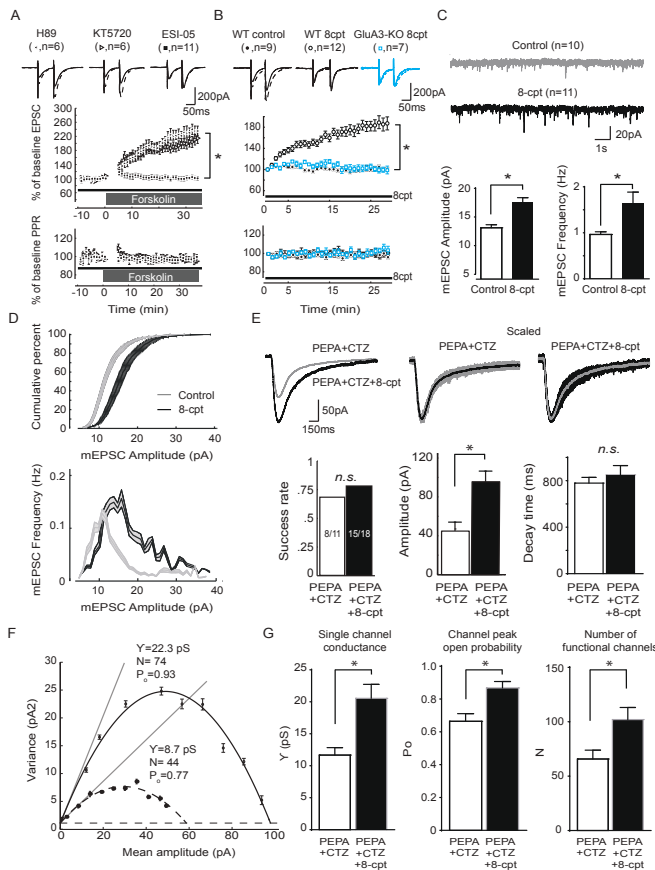
**Figure 5. GluA3-plasticity occurs through cAMP-dependent changes of single channel conductance and open probability.**

(A) Example traces of cell attached single channel recordings of GluA2/3 AMPARs in PCs of GluA1-KO mice. Under basal conditions (light red) the vast majority of the openings of GluA2/3 AMPARs occur at the low conductance level (O<sub>1</sub>), but in the presence of FSK the amount of openings in the higher conductance levels (O<sub>2-3</sub>) increase (red). (B) Count versus amplitude histograms of the events detected in the representative recordings shown in A illustrate the uneven distribution of events across the different conductance levels in the absence (light red) or presence (red) of FSK. (C) The opening durations [dwell time] of the same events shown in A and B were unchanged after FSK application. However, the duration of the closed state times were reduced suggesting a net increase in the total number of openings

produced by GluA2/3 channels in the presence of FSK. (D) Overall quantification of GluA2/3 single channel recordings show that the conductance significantly increased in the presence of FSK, yet the conductance per open state remained unchanged. (E) FSK significantly changed the distribution of GluA2/3 AMPARs events as revealed by a significant decrease of events in O<sub>1</sub> state and a significant increase in events at O<sub>2</sub> and O<sub>3</sub> states. The reduction of the closed state time shown in C was translated into a significant increase of the open channel probability. (F) The classical model of GluA1-subunit dependent LTP in pyramidal cells [see introduction] does not hold valid at PF to PC synapses. Note that the absolute numbers of subsets of both GluA1/2 and GluA2/3 AMPARs are unchanged upon LTP induction, whereas the GluA2/3 AMPARs are activated by cAMP signaling, enhancing their channel conductance and thereby increasing the current generated in potentiated synapses. This model describes for the first time a form of GluA3-dependent LTP.



bodies of GluA1-KO PCs produced the vast majority of their openings in the first and lower conductance state [O1] under basal conditions (Figure 5A,B), indicating that only two out of the four ligand binding domains (LBDs) present in the AMPAR tetramer are activated by AMPA. After application of FSK the behavior of GluA3-containing AMPARs changed strikingly and produced a significantly higher amount of openings in state O2 and O3 similar to GluA1-containing AMPARs (compare Figures 5A, B and E with Figure S5B). The average duration of the openings was unchanged ( $p=0.4$ ,  $p=0.13$  and  $p=0.09$  for O1, O2 and O3, respectively) (Figure 5C), but an increase of the absolute frequency of the openings caused shortening of the closed state dwell-time and thus a significant net increase in open probability ( $p<0.0001$ ; Figure 5E). Although FSK did not significantly change the conductance level of any of the open states ( $p=0.7$ ,  $p=0.14$  and  $p=0.15$  for O1, O2 and O3, respectively; Figure 5D), the higher relative fraction of events in the highly conductive open states O2 and O3 caused a significant increase ( $p<0.0001$ ) in the overall conductance of cAMP-stimulated GluA3 channels (Figures 5B and 5D). These experiments suggest that a rise in intracellular cAMP produces synaptic potentiation by increasing the open-channel probability of the GluA3 subunit, indicating a novel mechanistic model for GluA3-dependent synaptic plasticity.



noise analysis (NSNA) was done by fitting a parabolic equation to this distribution in order to estimate conductance, open probability and number of active receptors. (G) NSNA performed on the PC recordings in E revealed a significantly increased single channel conductance (left panel) and peak open channel probability upon 8CPT application (middle panel), which in turn led to an increased number of functional channels responding to the local AMPA application (right panel).

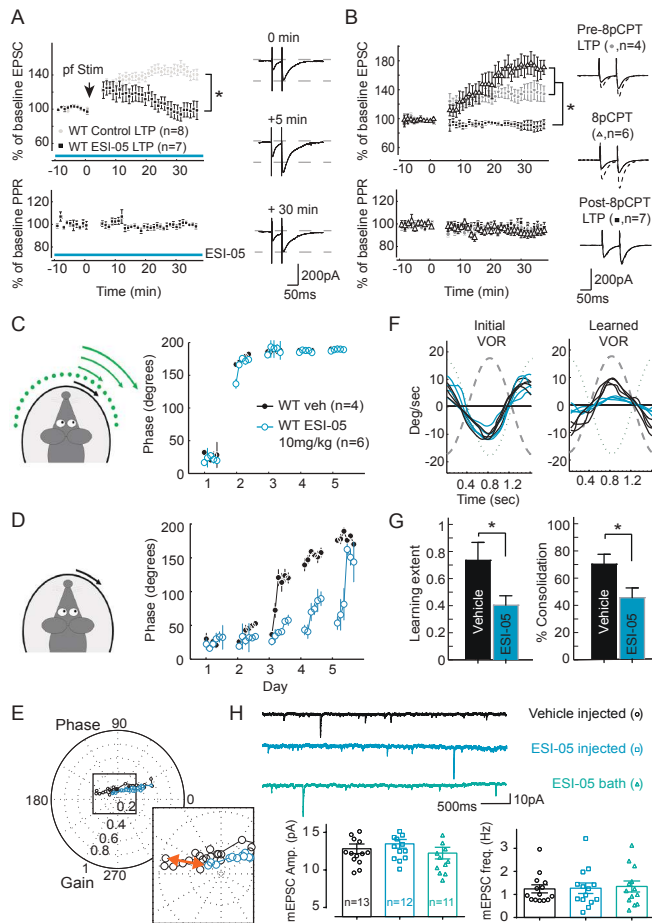
### GluA3-mediated plasticity is induced via cAMP-mediated Epac activation

To further elucidate the molecular mechanism underlying GluA3-dependent plasticity, we aimed to identify the intermediary factor that translates a rise in cAMP into synaptic potentiation of GluA3-containing AMPARs. Protein kinase A (PKA) is activated by a rise in cAMP and exerts cAMP-dependent synaptic effects [Lev-Ram et al., 2002; Sokolova et al., 2006]. However, incubating wild-type PCs with PKA antagonists KT5720 or H89 did not have any significant effect on synaptic potentiation induced by FSK ( $215 \pm 20\%$  with KT5720 and  $235 \pm 19\%$  with H89;  $p=0.7$  and  $p=0.9$ , respectively) (Figure 6A) indicating that PKA is not involved in mediating GluA3-plasticity. We next assessed the involvement of Epac (exchange proteins directly activated by cAMP, a.k.a. Rap guanine-nucleotide-exchange factor) as an alternative cAMP-dependent pathway that can trigger synaptic changes [Gekel and Neher, 2008; Woolfrey et al., 2009]. The blockade of Epac with its selective antagonist ESI-05 [Tsalkova et al., 2012] did not reduce basal transmission at PF-PC synapses (Figure 7H), but it effectively prevented the FSK-induced synaptic potentiation in wild-type PCs ( $p=0.9$  vs. baseline and  $p<0.0001$  vs. without ESI-05) (Figure 6A). To assess whether Epac activation is not only necessary, but also sufficient to cause GluA3-dependent synaptic potentiation, we investigated the impact of the selective Epac activator 8-CPT-2Me-cAMP (8CPT). Adding  $20 \mu\text{M}$  8CPT to the intracellular recording solution produced synaptic potentiation by  $185 \pm 17\%$  compared to baseline in wild-type PCs ( $p=0.0004$ ; Figure 6B), but not in GluA3-deficient PCs ( $100 \pm 5\%$ ;  $p=0.8$ ; Figure 6B). In addition, the postsynaptic application of 8CPT increased the amplitude and frequency of PC mEPSCs ( $p=0.0005$  and  $p=0.001$ , respectively; Figures 6C and 6D) and did not change the PPF ratio ( $104 \pm 5\%$ , Figure 5B). These experiments indicate that a rise in cAMP triggers synaptic potentiation through Epac-mediated activation of postsynaptic GluA3-containing AMPARs. This Epac-driven activation of GluA3-containing AMPARs was not limited to those located at synapses. Outside-out patches excised from wild-type PC somata produced a peak current of approximately 10 pA in response to puffs of  $100 \mu\text{M}$  AMPA (Figure S4E). When the Epac activator 8CPT was added to the internal solution of the patch pipette, the peak current obtained under the same conditions was increased 2.5-fold in the absence of a presynaptic component ( $25 \pm 3$  pA,  $p<0.0001$  vs. control). This difference in peak current was largely maintained in the presence of AMPAR desensitization blockers PEPA and cyclothiazide ( $45 \pm 8$  pA without vs.  $97 \pm 10$  pA with 8CPT,  $p<0.0001$ ; Figure 6E), indicating that cAMP-driven GluA3-plasticity does not depend on a change in desensitization properties of AMPAR channels. As expected from our single channel results, non-stationary noise analysis of these non-desensitizing AMPAR responses showed a significant increase in conductance and open probability (Figures 6F-G). This analysis revealed how in a mixed pool of GluA1- and GluA3-containing AMPARs Epac-dependent GluA3 potentiation was translated into an increase in current amplitude without altering the dynamics of the response (Figure 6E), highlighting the consistent results with miniature and evoked EPSC recordings.

### Epac activation is required for LTP and motor learning

We next tested whether PF-PC LTP depends on Epac activation. Incubation of slices with Epac-inhibitor ESI-05 significantly inhibited synaptic potentiation induced by tetanic PF stimulation ( $102 \pm 13\%$  versus  $140 \pm 8\%$  in control conditions,  $p<0.0001$  after 15 min; Figure 7A). In addition, LTP was fully occluded when brain slices were pre-incubated with the membrane-permeable analog of the Epac activator [8pCPT] (GluA3-KO vs. WT,  $p=0.008$ ; Figure 7B). Together, these data indicate that Epac2 activation is responsible for postsynaptic LTP at the PF to PC synapse through activation of GluA3-containing AMPARs.

To investigate the involvement of Epac activation in cerebellar synaptic plasticity in-vivo, we



**Figure 7. Pharmacological manipulation of Epac activity impairs LTP in-vitro and motor learning in-vivo without affecting synaptic transmission.**

(A) Epac antagonist ESI-05 prevents PF to PC LTP induced by tetanic PF stimuli. (B) Epac activation through incubation with 8pCPT potentiates AMPAR currents (open triangles). As a consequence, a minimal 30 minute incubation with 8pCPT fully occludes LTP induction (black squares) compared with LTP induction in the absence of 8pCPT (gray circles). (C) Eye movement phase values in wild-type mice that are injected with ESI-05 10 mg/kg (open blue circles) or injected with vehicle only (black circles) are virtually identical during the visuo-vestibular mismatch training when the light is on.

(D) During the catch trials in the dark, the phase shift of VOR adaptation in the mice injected with ESI-05 10 mg/kg is significantly delayed compared with those in their littermates injected with vehicle only.

(E) Polar plot of the combined gain and phase data shows a common learning trajectory and comparable initial gain, yet different final outcome, for both groups. Inset: the final VOR reached after 5 days of training is amplified to visualize the magnitude of the gain difference (red arrow) between ESI-05 and vehicle-injected mice.

(F) Four representative eye velocity traces of the VOR before (left) and after (right) phase-reversal training show that, whereas both ESI-05 and vehicle-injected mice show equal baseline performance and that both are able to flip the phase of the VOR, the magnitude of the VOR reached after the training is substantially different.

(G) Both learning extent and consolidation during the phase-reversal task are significantly smaller in the mice injected with ESI-05 compared to those injected with vehicle only (T2-test  $p < 0.05$ ).

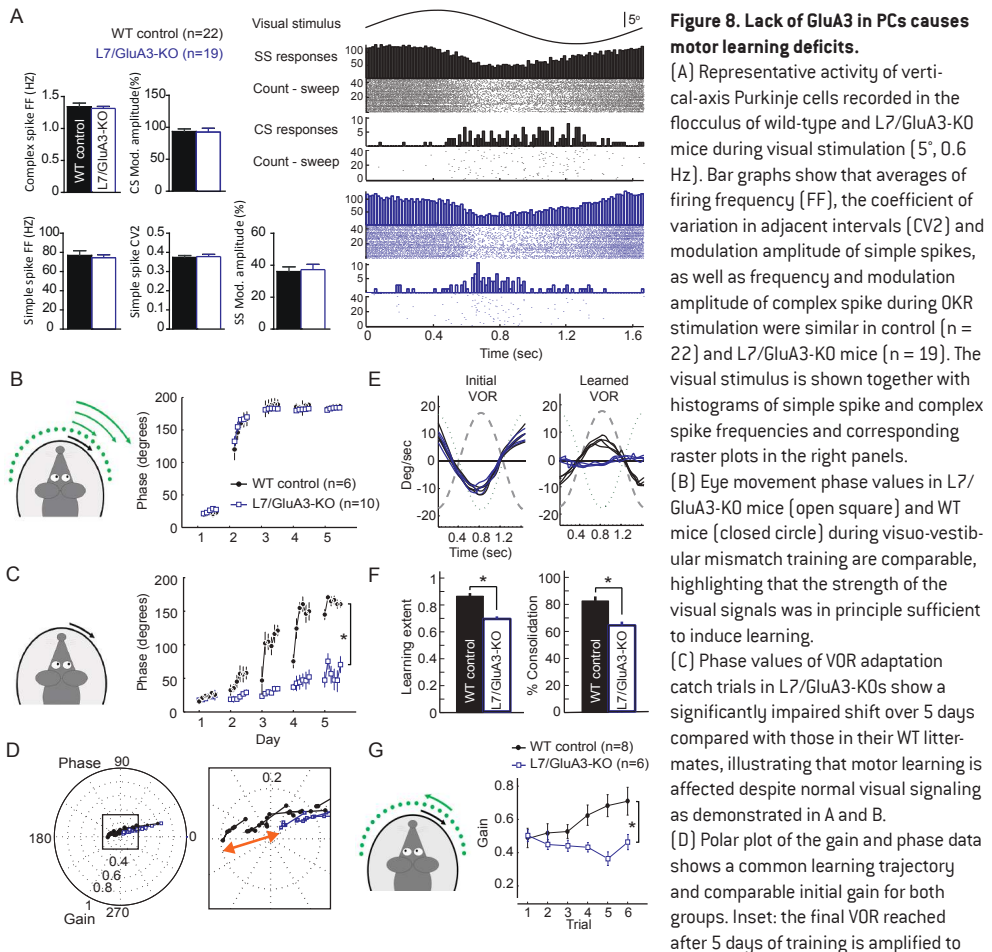
(H) Impaired motor learning after ESI-05 injections does not correlate with a decreased transmission at PF to PC synapse. The PC mEPSC amplitude and frequency of WT mice injected with ESI-05 are similar compared with those in vehicle-injected mice. Error bars indicate SEM, \* indicates  $p < 0.05$ .

performed phase-reversal adaptation in wild-type mice that received daily IP injections with either Epac antagonist ESI-05 (0.2-0.3 ml at 10 mg/kg) or with vehicle alone 30 minutes prior to the training protocol. Mice administrated with ESI-05 were not affected in basal eye reflex behavior, but performed significantly worse in the phase-reversal task compared to vehicle injected animals (Figure 7D). Although both groups eventually reached a reversal of the VOR phase (Veh:  $157 \pm 2\%$ , ESI-05:  $148 \pm 15\%$ ; Figure 7D), its magnitude was significantly lower in the ESI-05 injected mice compared to vehicle controls ( $p = 0.01$ ; Figure 7E-F). This difference reached after training could not be explained by a poor basic response to the training stimuli (Figure 7C), but only by a significantly reduced learning extent ( $p = 0.01$ ) and consolidation rate ( $p = 0.03$ ). Importantly, systemic ESI-05 injections produced learning

deficits without a change in basal synaptic transmission compared to vehicle-injected mice ( $p=0.5$  and  $p=0.9$  for mEPSCs amplitude and frequency, respectively) (Figure 7H), suggesting that absence of Epac-dependent synaptic potentiation without any change in basal transmission is sufficient to impair learning capabilities.

### GluA3 expression in Purkinje cells is required for VOR learning

We showed that VOR learning depends on global expression of GluA3 and that LTP at PF-PC synapses requires GluA3-plasticity, but does VOR learning depend on GluA3 specifically in PCs? To address this question, we generated and tested a PC-specific GluA3-KO mouse (referred to as L7/GluA3-KO; Figure



S6) by crossbreeding mice expressing Cre-recombinase under the PC-specific promotor L7-pcp2 with mice in which the GluA3 gene is flanked by loxP sites [Barski et al., 2000; Sanchis-Segura et al., 2006]. After establishing the single-unit identity of floccular vertical-axis Purkinje cells in adult L7/GluA3-KO mice by demonstrating a climbing fiber pause in their simple spike activity as well as a preferred modulation tuning-curve during extracellular recordings in-vivo (Figure 8A) [Hoebeek et al., 2005], we investigated the action potential generation of their simple spike activity in the awake state. In the absence of visual stimulation, both the firing frequency and regularity (i.e. coefficient of variation of adjacent interspike intervals or CV2) of the simple spike activity of the L7/GluA3-KO mice did not differ significantly from those of wild-types (Figure S7A), which is consistent with the similar I-V relationships recorded in-vitro for wild-type and GluA3-lacking PCs (Figure S7B). Next, we provided visual stimulation at the frequency that was used for the training paradigm (0.6 Hz). Again the firing frequency and regularity (i.e. CV2) of the simple spike activity of the L7/GluA3-KO mice did not differ from those of wild-types ( $p=0.7$  and  $p=0.8$  respectively) (Figure 8A). Moreover and most importantly, the amplitude of the simple spike modulation during visual stimulation did not differ ( $p=0.8$ ; Figure 8A), suggesting that the PF output is in effect sufficient to mediate the visual training signals in the L7/GluA3-KO mice despite the reduced PF to PC synaptic transmission (Figure S7C). Finally, the firing frequency and modulation amplitude of the complex spikes did not show any significant difference either ( $p=0.7$  and  $p=0.9$ , respectively; Figure 8A). Together, these data indicate that the in-vivo excitability and spike generation of PCs are intact in L7/GluA3-KO mice.

We then tested 3 to 5 month-old L7/GluA3-KO and control littermates for their ability to adapt eye reflexes. The baseline OKR and VOR performances of these L7/GluA3-KO mice were indistinguishable from those in controls (Figure S1). Similarly to global GluA3-KOs, VOR motor learning was prominently affected in L7-GluA3-KO mice (Figure 8C-G). The absence of GluA3 in PCs showed significant deficits throughout the phase-reversal paradigm (all  $p$  values  $< 0.01$  after day 2), including a significantly different learning extent at the end of it ( $p=0.0005$ ; Figure 8B-F). Moreover, consolidation during the phase-reversal paradigm was significantly lower ( $p=0.0006$ , Figure 7F). Gain modulation was also impaired as shown by a significant difference between the final eye movement gain of L7/GluA3-KO mice compared to control littermates after either gain-down or gain-up training sessions ( $p=0.04$  and  $p=0.006$  for gain-down and gain-up, respectively) (Figure 8G).

In contrast to the single L7/GluA3-KO as well as single global GluA1-KO and GluA3-KO mice, mice that lack both GluA1 and GluA3 receptor subunits specifically in Purkinje cells (L7-GluA1/3-KO) showed significant aberrations in baseline eye movement performance ( $p=0.0001$  for OKR and  $p=0.03$  for VORD) (Figures S1A and S1B). Together with the findings presented above (see also Figure 1), these data suggest that the presence of GluA3 in the GluA1-KO can compensate for their lack of GluA1 during both baseline and learning behavior, but that the presence of GluA1 in the global GluA3-KO and single L7-GluA3-KO is not sufficient to fully compensate for the lack of GluA3 during adaptation of compensatory eye movements, highlighting the putative impact of GluA3-dependent synaptic plasticity in PCs.

## DISCUSSION

It is widely believed that LTP- and LTD-type synaptic plasticity mechanisms act in concert to mediate several types of learning in brain regions such as the hippocampus, amygdala and cerebral cortex [Malinow and Malenka, 2002; Makino and Malinow, 2011; Nabavi et al., 2014; Nedelescu et al., 2010; Rumpel et al., 2005; Takahashi et al., 2003]. For cerebellar learning, LTD at the PF to PC synapse has historically been considered the dominant plasticity mechanism [Ito, 2002; Linden and Connor, 1995].

Although the simple spike suppression observed at early stages of some forms of motor learning *in-vivo* may suggest LTD occurrence (ten Brinke et al., 2015; Yang and Lisberger, 2014), an increasing amount of studies suggest that LTD is not a strict requisite for motor learning (Hesslow et al., 2013; Schonewille et al., 2011). In the present study we show that LTP at PF to PC synapses is a required mechanism for cerebellar motor learning. We show that LTP, but not LTD, at the PF to PC synapse requires plasticity of GluA3-containing AMPARs and that both the selective removal of GluA3 in PCs and the pharmacological blockade of the pathway leading to GluA3 plasticity *in-vivo* severely impair the ability to adapt the vestibulo-ocular reflex. Combined, these findings provide the first correlative link between GluA3-dependent LTP and behavioral learning in general.

Previous studies proposed a role for cerebellar LTP in the context of bidirectional gain modulation (Boyden et al., 2006). This work suggested that gain-down modulation of the eye movements might require PF to PC LTP, and conversely, gain-up modulation would require LTD. According to this hypothesis one would expect that GluA3-KO mice show impaired gain-down modulation with intact gain-up adaptation. However, our data show that the specific absence of GluA3 in Purkinje cells most prominently impairs gain-up and phase modulation, supporting an opposite kind of role for GluA3-dependent LTP in oculomotor learning. Whereas the role of GluA3 in PC plasticity and cerebellar motor learning is becoming more clear now, that of GluA1 is still largely obscure. The presence of GluA1 in PCs was neither essential for the induction of LTD or LTP nor were there overt signs of deficits in motor performance or motor learning in the GluA1-KO. Its possible role became only indirectly apparent, when we observed that, in contrast to the single GluA1-KO, the double GluA1/GluA3-KO virtually completely lacked glutamatergic currents in PCs and that the double L7-GluA1/A3-KO showed significant signs of ataxia and deficits in motor learning. Given that the single GluA3-KO did not show any sign of motor performance deficit, these findings indicate that GluA1-containing AMPARs in PCs do contribute to cerebellar motor performance, but that their absence can be compensated for by GluA3-containing AMPARs.

The possible role of LTP at the PF to PC synapse in cerebellar motor learning has been suggested before by various other cell-specific mouse mutant studies (Andreescu et al., 2007; Schonewille et al., 2010; Peter et al., 2016). However, these studies tackled more upstream PC processes, which involved the nuclear estrogen receptor, cytosolic protein phosphatase calcineurin and subsynaptic protein shank2, and as a consequence they suffered from various side-effects that prevented definitive conclusions (Gao et al., 2012). In the current study, in which we tackled PF to PC LTP more downstream by targeting GluA3-containing AMPARs at the level of the synapse itself, we did not find any evidence for structural changes or firing differences in PCs of awake behaving mice. We did find that the basal transmission was reduced in PCs lacking GluA3 (both in the global and the cell-specific KO mice), but this deficit was probably not the reason for the impairment of LTP or of motor learning, because acutely inhibiting GluA3-plasticity through blockade of Epac prevented both LTP and motor learning without affecting basal transmission. We therefore propose that the reduced basal transmission in GluA3-KO mice is the consequence of a prolonged deficit in LTP.

GluA1-dependent synaptic plasticity is mediated by active trafficking (Makino and Malinow, 2011; Shi et al., 2001) as well as by changes in conductance and open probability at the single receptor level (Benke et al., 1998; Derkach et al., 1999). Here, we present evidence that, at least at the short-term scale of tens of minutes, synaptic potentiation through activation of GluA3-plasticity does not involve trafficking but mainly a prominent increase in open-channel probability of GluA3-containing receptors, suggesting that in the case of GluA3, changes in receptor properties is the predominant mechanism to produce synaptic potentiation. These findings imply that PF to PC LTP is mechanistically not just



the reverse of LTD at this synapse [Jorntell and Hansel, 2006]. Linden and colleagues have shown that PF to PC LTD is largely expressed by endocytosis of GluA2-containing AMPARs [Linden, 2001; Schonewille et al., 2011], thus mainly dependent on AMPAR trafficking. Given the current findings on GluA3-mediated LTP, it may be worthwhile to find out whether changes in AMPAR unitary conductance or glutamate affinity also play a minor role in early LTD expression at the PF to PC synapse, as interference with clathrin-mediated endocytosis did not produce a total attenuation of LTD expression [Wang and Linden, 2000].

Full genetic ablation of GluA2 subunits produces, in contrast to that of GluA3, an ataxic, hardly viable phenotype [Gerlai et al., 1998]. Interestingly, the remaining, mainly GluA1- and GluA3-containing, AMPARs in these knock-out mice have an unusual subunit composition and are abnormally distributed at the synapse [Sans et al., 2003]. In this respect, it should be noted that GluA3 is an obligatory heteromeric subunit: GluA3 homomers are energetically unfavorable [Rossman et al., 2011] and form intracellular aggregates that don't reach the cellular surface efficiently [Coleman et al., 2016]. Genetic mouse models in which GluA2 trafficking is blocked reveal an impairment in LTD induction at their PF to PC synapses, whereas LTP is normal [Schonewille et al., 2011; Steinberg et al., 2006], which is in line with our finding that LTP can be induced without trafficking of GluA2/A3-containing AMPARs. Differently from these GluA2 mutants, GluA3-KO mice prominently express surface GluA2-containing AMPARs (heteromerized with GluA1), but lack a cAMP-dependent synaptic LTP. These data highlight the differential roles of GluA2 and GluA3 in the structural dynamics and localization of AMPARs and the related forms of synaptic plasticity. In contrast to GluA3, GluA2 is unlikely to be directly involved in cAMP-dependent plasticity, since its expression coupled to GluA1 does not compensate for the absence of GluA3 subunits. We propose that GluA2 expression is a structural requisite for GluA3 plasticity, as it appears necessary for proper expression and location of GluA3-containing AMPARs. GluA1/2 heteromers in PCs may then serve to maintain basal synaptic currents when cAMP levels are low.

The finding that an Epac-mediated change in single channel conductance and open probability of GluA2/3-containing AMPARs may underlie LTP at the PF to PC synapse raises the question how this change in configuration comes about. Interestingly, the distribution of GluA3-containing AMPARs openings does not seem to respond to a stochastic probability distribution of four ligand binding domains [LBDs] "catching" glutamate with equal probability. Instead it is biased towards the lowest conductance state opening, in which only two out of four LBDs bind glutamate. Since GluA3-containing AMPARs predominantly consist of two GluA3 and two GluA2 subunits, only the GluA2 LBDs effectively may bind glutamate under basal conditions. Our observation that enhancing cAMP levels exerts GluA3-containing receptors to produce higher conductance openings [resembling the behavior of GluA1-containing receptors] may suggest that Epac activation triggers a conformational change in the two GluA3 subunits present in each tetramer, such that they become responsive to glutamate binding at the LBD [Figure 5F].

It is widely accepted that intracellular calcium signaling is a key mechanism for LTP induction in PCs [Coesmans et al., 2004; van Woerden et al., 2009]. In the present study we show that postsynaptic LTP depends on cAMP-dependent activation of GluA3-containing receptors. How low calcium signals in PCs are transduced into activation of adenylyl cyclase to raise cAMP levels remains to be elucidated. Interestingly, it has been shown that the tetanic activity of PFs required for LTP induction produces local calcium increases dependent on low-threshold CaV3.1 T-type calcium channels [Hildebrand et al., 2009] and that global deletion or blockage of these channels prevents LTP induction and motor learning [Ly et al., 2013]. In this respect, the calcium/calmodulin-dependent adenylyl cyclase *Adcy1*

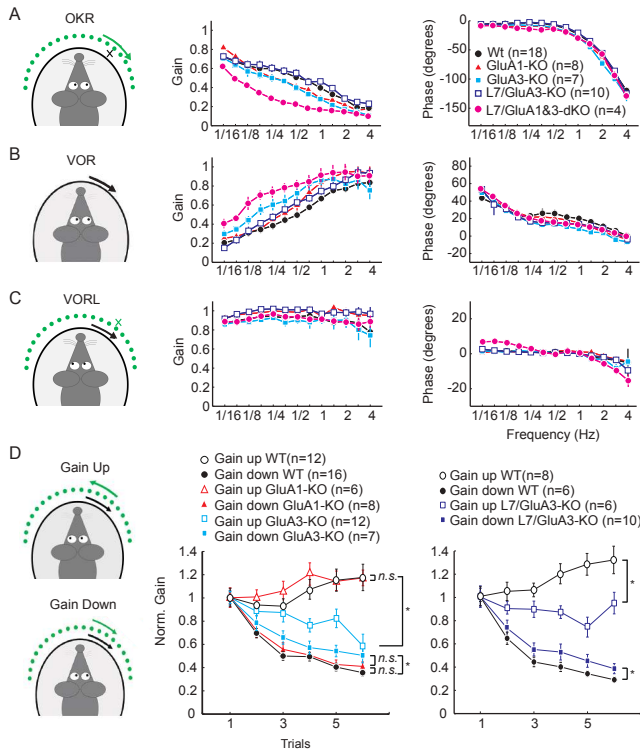


[Masada et al., 2012] could be an interesting candidate to convert local calcium activity in a rise in cAMP.

We have shown here that postsynaptic GluA3-dependent synaptic potentiation depends on a rise in cAMP. Therefore, this study expands the repertoire of forms of PC plasticity already known to depend on cAMP, such as presynaptic plasticity [Chen and Regehr, 1997; Kaneko and Takahashi, 2004; Lev-Ram et al., 2002], intrinsic plasticity [Belmeguenai et al., 2010] or plasticity at inhibitory synapses [Mitoma and Konishi, 1996]. Epac2 has recently been reported to also have a role in presynaptic plasticity in that it may modify glutamate release probability [Gekel and Neher, 2008]. This raises the interesting possibility that Epac2 and/or cAMP in their pre- and postsynaptic domains operate in a synergistic fashion to control synaptic plasticity [Le Guen and De Zeeuw, 2010; Wang et al., 2014]. Likewise, the induction protocol of LTP produces an increase in intrinsic excitability in PCs, via cAMP-mediated PKA modulation of SK potassium channels [Belmeguenai et al., 2010]. Thus, since this change in intrinsic excitability occurs at least partly as a secondary process following tetanic PF stimulation, LTP at the PF to PC synapse may act as a feed-forward amplifier of synaptic inputs to modulate firing rate in PCs via cAMP production. Finally, it should be noted that rebound potentiation at the molecular layer interneuron to PC synapse, which occurs following PC depolarization, is also mediated by cAMP-mediated PKA modulation [Hirano and Kawaguchi, 2012]. Together, these findings point towards a central role of cAMP following induction of PF to PC LTP, regulating multiple forms of plasticity with different identities and natures in a synergistic fashion [Gao et al., 2012].

Synapses are highly dynamic structures and early removal of synaptic proteins can lead to compensatory mechanisms to occur in order to overcome unbalanced synaptic function. However, no compensatory mechanism is able to overcome the declarative memory deficits observed in GluA1 KO mice [Feyder et al., 2007; Humeau et al., 2007]. In contrast, GluA1-KO learned to adapt their vestibulo-ocular reflexes virtually identically to wild-type littermates. This finding suggests two possible scenarios: either PCs synapse are capable to compensate for the absence of GluA1 through a mechanism that is not present in hippocampal pyramidal or amygdalar cells, or GluA1 is not involved in this form of learning at all. With the evidence presented here, none of these possibilities can be unequivocally discarded. Yet, these findings in the GluA1-KO emphasize the insufficiency of compensation in the GluA3-KO; the fact that their PCs could not compensate for the absence of GluA3 to overcome the lack of LTP and the learning deficits, highlights the importance of GluA3 for PC synaptic plasticity and motor learning. Taken together, the picture emerges that the learning rules for AMPAR-mediated plasticity in PCs are inverted compared with those in the hippocampus: cerebellar LTP and learning do not require GluA1, but depend on plasticity of GluA3-containing AMPARs.

## Supplementary information



**Figure S1. Basic compensatory eye movements in the single global KO mice for GluA1 or GluA3 as well as in PC-specific KO for GluA3 are relatively normal, whereas those in the double PC-specific KO for GluA1 and GluA3 show pronounced gain deficits.**

(A) The optokinetic reflex (OKR), which stabilizes gaze with respect to a moving visual field [Stahl et al., 2000], showed a normal baseline gain in GluA1-KO and GluA3-KO mice [ $F(2,29)=2.361$ ,  $p = 0.11$ ; Repeated measures ANOVA and Tukey post hoc analysis], whereas phase values in both mutants presented a small, but consistent, delay across the entire frequency range tested [ $F(2,29)=14.86$ ,  $p < 0.01$ , Tukey's multiple comparisons test revealed differences for the 95% confidence intervals of both mutants with respect to wild-type controls, but not between them]. L7/GluA3-KO mice presented intact gain and phase values compared to controls [ $F(1,26) = 0.21$ ,  $p = 0.64$  and  $F(1,26) = 1.24$ ,  $p = 0.27$ , respectively]. In contrast, PC-specific double GluA1 and GluA3 KO mice [L7/GluA1&3-dKO] showed a highly significantly impaired OKR response [ $F(1,20) = 21.30$ ,  $p = 0.0001$ ]

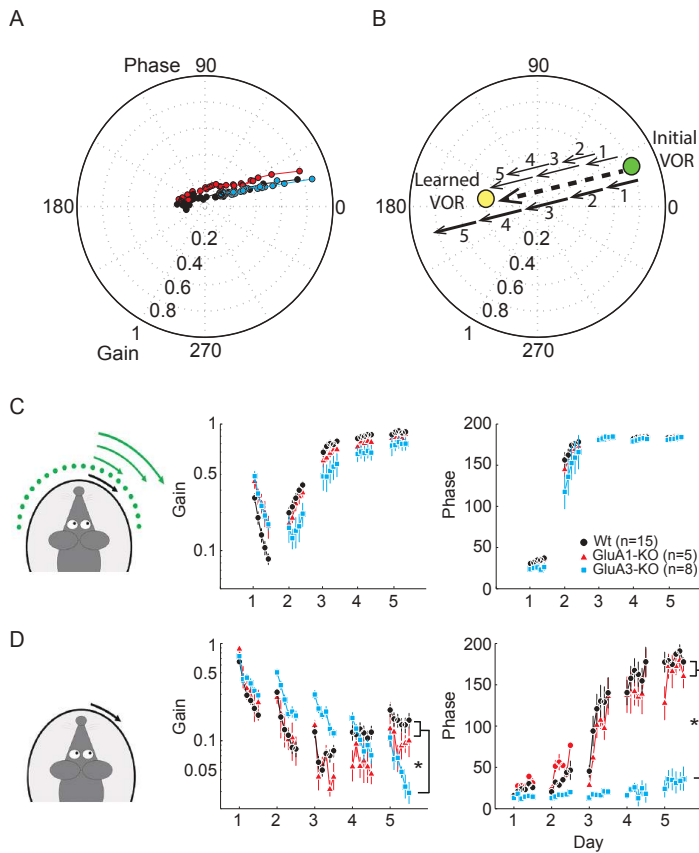
(B) During VOR compensation GluA1-KO

and GluA3-KO showed both a normal gain [ $F(2,29)=1.745$ ,  $p = 0.17$ ] and normal phase [ $F(2,29)=1.382$ ,  $p = 0.26$ ]. In addition, L7/GluA3-KO mice also showed a normal basic eye movement performance [ $F(1,26) = 1.65$ ,  $p = 0.21$  and  $F(1,26) = 1.53$ ,  $p = 0.22$ , for gain and phase, respectively]. Interestingly, L7/GluA1&3-dKO showed significantly improved VOR performance when compared to control mice [ $F(1,20) = 5.245$ ,  $p = 0.033$ ], most likely as a compensation for their impaired OKR.

(C) When we combined optokinetic stimulation with vestibular stimulation [i.e. VOR in the light or VORL] as occurs in daily life, all mutants also showed normal performances for both gain and phase compared to those in wild-type littermates [ $F(2,29) = 1.33$ ,  $p = 0.29$  for GluA1-KO, GluA3-KO and their WT littermates and  $F(1,26) = 1.51$ ,  $p = 0.23$  for L7/GluA3-KO vs. control littermates gain values].

(D) Oculomotor adaptation was assessed through paradigms aiming to either increase [i.e. gain-up paradigm, in which the visual stimulus moves with the same amplitude as the vestibular stimulus, but with opposite direction, resulting in improved VOR compensation] or decrease [i.e. gain-down paradigm, in which both visual and vestibular stimuli move with the same amplitude in the same direction, resulting in cancelation of compensatory eye movements] the amplitude of the VOR. Schematic drawings of the training stimuli are shown in the left column. Our results show that whereas GluA1-KO mice show a comparable increase [ $p=0.23$  for final catch trials] and decrease [ $p=0.11$ ] of gains compared to those in WT mice, GluA3-KO mice show significantly impaired gain-up ( $p=0.009$ ) as well as gain-down ( $p=0.001$ ) paradigms. The deficits observed in the global GluA3-KO mice were also present in the PC-cell specific KO [L7/GluA3-KO compared to WT littermates,  $p = 0.006$  and  $p = 0.04$  for gain-up and gain-down, respectively].

Error bars indicate SEM, \* indicates  $p < 0.05$ .

**Figure S2.**

(A) The full range of the 2 variables that explain ocular movements (0 to 1 for the gain and 0 to 360 for the phase) determine a circular bi-dimensional Cartesian plane (shown as a polar plot), in which every eye movement can be defined. Given that phase-reversal learning takes place through a defined common learning trajectory over several days during which phase covariates with gain (left polar plot), we performed statistics on the Cartesian coordinates defining gain and phase using the paired Hotelling's T2-Test.

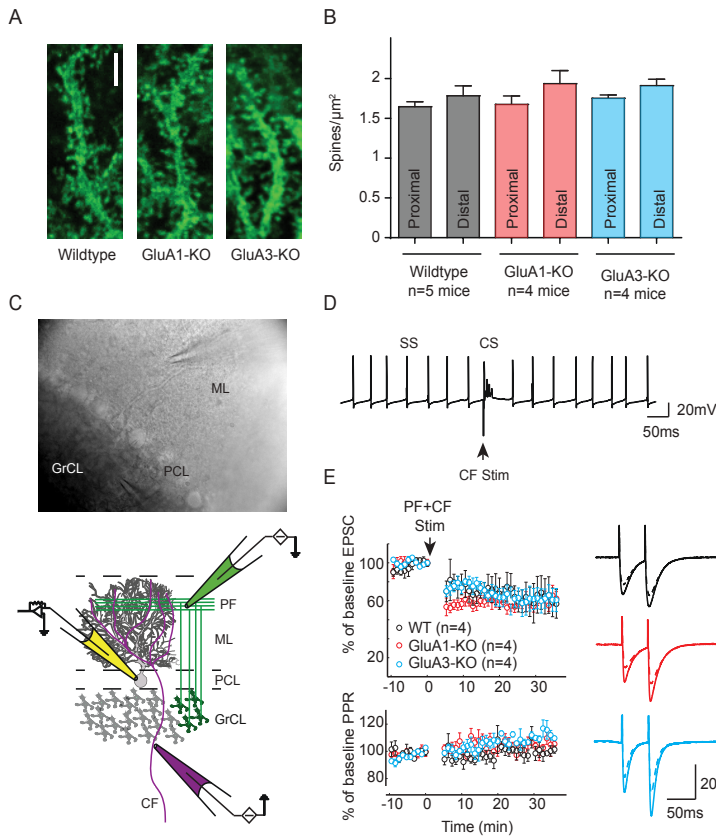
(B) Polar plots of gain and phase vectorial representation during phase-reversal VOR learning data to illustrate the data analysis procedure. The data are composed of 5 individual learning vectors (one per day) moving across a constant learning trajectory towards the target set by the training paradigm (Phase of 180 degrees; Gain of 1). Based on the raw gain and phase data (A), we first calculate the learning extent for each mouse as the

vectorial difference between the final performance and the initial performance (recording 6 of day 5 – recording 1 of day 1) and subsequently average these values per group. Between days of training there is partial retention of motor memories; to calculate the overall consolidation we calculate the ratio between the learning extent and the absolute summed extent of the learning vectors as if there was no memory loss overnight (100% consolidation). This ratio calculated per mouse is then also averaged across the mice, generating consolidation values for each group.

(C) Eye movement behavior of 4-6 week old GluA3-KO mice is virtually identical to that of 3-5 month old mice. Scatter plots of gain and phase values of 4-6 week old mice during the visuo-vestibular training for VOR phase-reversal shows no significant differences in the ability to follow the training signal ( $p > 0.05$  for last training recording on day 5 for comparison of GluA1-KO vs GluA3-KO and of WT vs GluA3-KO).

(D) Scatter plots of gain and phase values of the VOR catch trials show that WT and GluA1-KO mice, but not GluA3-KO mice, are able to reverse the phase of the VOR after training ( $p < 0.01$  for last catch recording on day 5 for comparison of GluA1-KO vs GluA3-KO and of WT vs GluA3-KO). For comparison with data in 3-5 month old animals see also Figure 1.

Error bars indicate SEM, \* indicates  $p < 0.05$ .



**Figure S3. PCs lacking GluA1 or GluA3 have comparable spine density and show comparable levels of LTD induction compared to those in wild type (WT) PCs.**

(A) Representative confocal optical planes ( $0.5\ \mu\text{m}$  thick) of PC distal dendrites used to quantify spine densities of WT, GluA1-KO or GluA3-KO PCs. The results show that neither the lack of GluA1 nor that of GluA3 yielded differences in spine densities of proximal or distal dendrites of PCs (bottom panel).

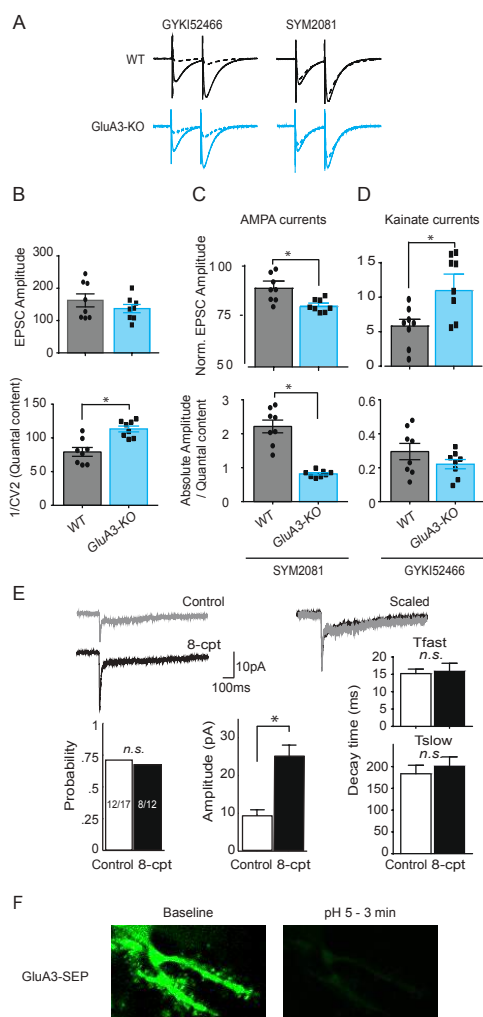
(B) Densities plotted for each genotype correspond to the average spine density of at least 20 dendritic branches of PCs in lobules V to X per animal. Scale bar =  $5\ \mu\text{m}$ .

(C) Scheme of cerebellar cortical circuitry (bottom panel) and representative picture of the in-vitro preparation (top panel) showing positions of recording electrode (yellow) at PC soma and stimulus electrodes (green and purple) at parallel fiber (PF) beam and climbing fiber (CF), respectively. ML, PCL and GrCL indicate molecular layer, Purkinje cell

layer, and granule cell layer, respectively.

(D) PCs were recorded in current clamp mode and the location of the stimulus pipettes were determined functionally by evoking responses to electrical stimulation at resting potential. Once the proper locations were identified, cells were kept in hyperpolarized state ( $-80\ \text{mV}$  approx.) and a conjunctive CF and PF stimulation protocol was applied to the cell for 5 minutes [see Suppl. Methods for details].

(E) Both GluA1-KOs (red) and GluA3-KOs (blue) show similar cerebellar synaptic weakening after LTD induction (top panel) compared to WT littermates (black) with unchanged PPR over time (bottom panel). Representative traces of paired EPSCs before (solid lines) and after LTD induction (dashed lines) (right panels; genotypes match the color codes in B).



**Figure S4. Kainate receptors do not compensate for weakening of glutamatergic transmission at PF to PC synapses in the absence of GluA3.**

(A) To assess possible compensatory components in the glutamatergic transmission of PCs in GluA3-KO mice we investigated the impact of blocking either AMPA-receptors with 30  $\mu$ M of GYKI-52466 [Cossart et al., 2002] or kainate receptors with 5  $\mu$ M of SYM2081 [Yan et al., 2013] after establishing a stable baseline of eEPSCs in WT and GluA3-KO PCs.

(B) PF stimulation intensity was manually adjusted to obtain comparable EPSC amplitudes between 100-200 pA in WT and GluA3-KO PCs ( $p = 0.3$ , GluA3-KO vs. WT). The average quantal content released to produce events of comparable amplitude (estimated as the inverse of the square coefficient of variation; Kerchner and Nicoll, 2008), was significantly higher in the GluA3-KO ( $p < 0.001$ ), indicating post-synaptic weakening.

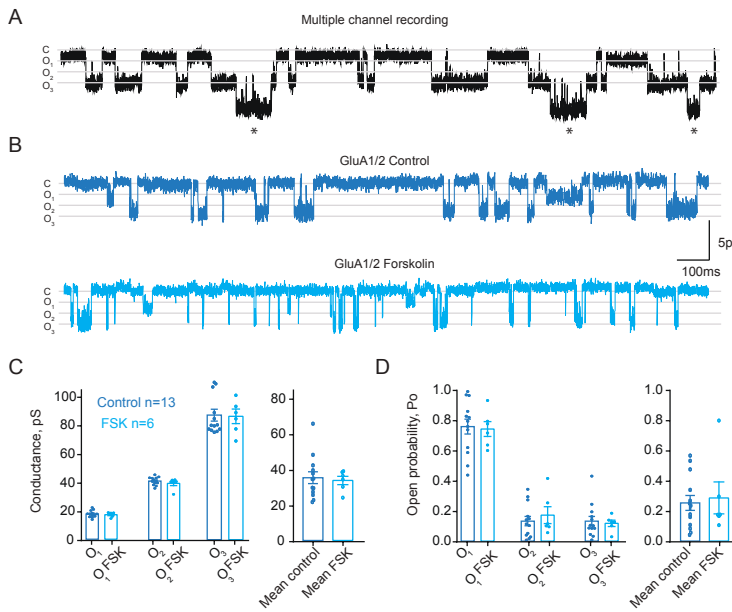
(C) Blocking AMPARs reduced the total glutamatergic transmission in GluA3-KO PCs by  $89 \pm 2\%$ , which was significantly less than that in wild-type PCs [ $94 \pm 2\%$ ;  $p = 0.01$  for GluA3-KO vs. WT, top panel]. However, this difference was exacerbated after normalizing the amplitude to the quantal content, revealing that in the absence of GluA3, PCs have about half the normal magnitude of AMPA-mediated current ( $p < 0.001$ , bottom panel).

(D) To investigate to what extent kainate receptors can compensate for an impairment in GluA3-dependent transmission in PCs [Yan et al., 2013], we investigated the impact of a blockage of kainate-receptors in both WT and GluA3-KO PCs. The contribution of kainate-receptor mediated events to EPSC amplitude normalized to baseline magnitude was significantly higher in PCs of GluA3-KO [ $21 \pm 1.5\%$ ] than that in WT PCs [ $16 \pm 3\%$ ;  $p = 0.024$  for GluA3-KO vs. WT, top panel]. However, when normalized to the quantal content, the absolute contribution of kainate receptors was comparable among genotypes ( $p = 0.19$ , bottom panel). Together, these data indicate that glutamatergic transmission in GluA3-KO mice can be largely explained by GluA1/GluA2-mediated AMPA-currents and to a lesser extent by kainate-currents, none of which is

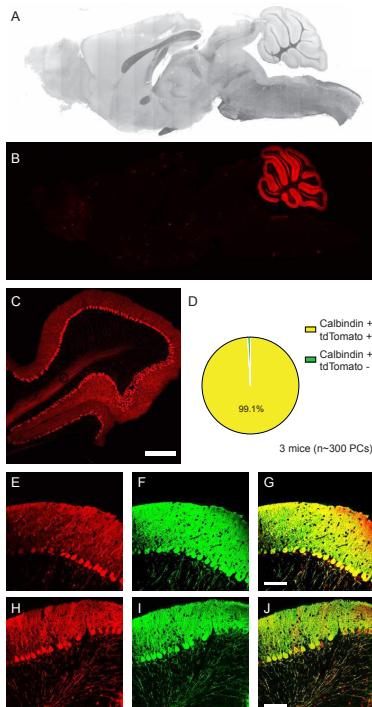
able to compensate for the synaptic weakening caused by the absence of GluA3.

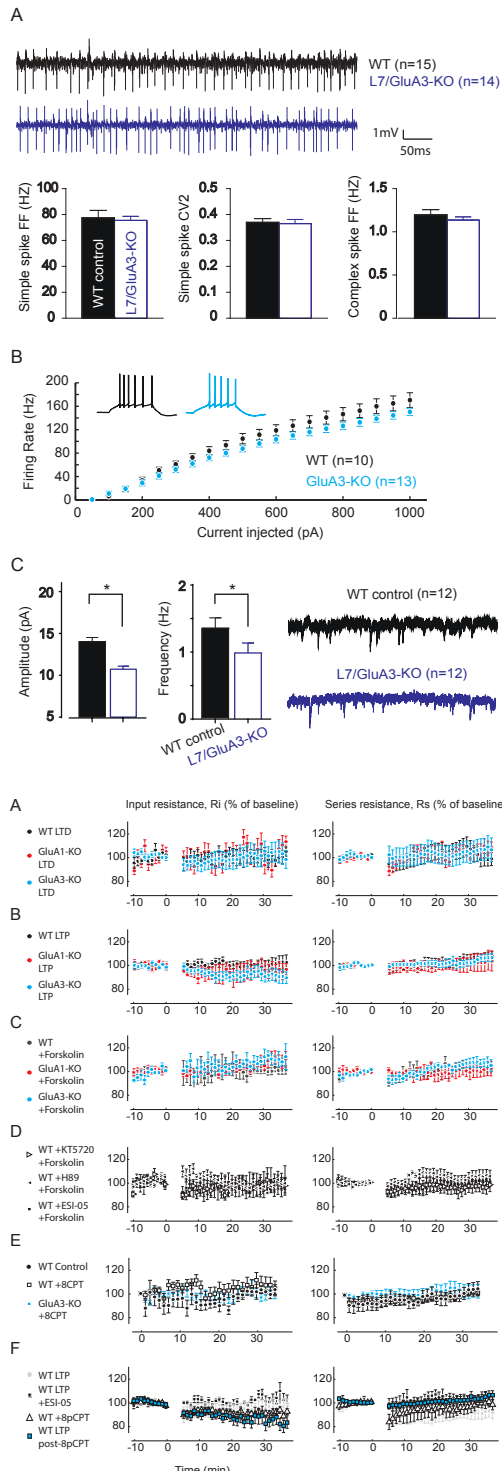
(E) Excised patches of PC somata that received puffs of 100  $\mu$ M AMPA generated significantly larger currents when 8-CPT-2Me-cAMP [8-CPT] was present in the internal solution. Note that the control patches showed the same probability of presenting AMPA events [left]. Fast desensitizing and slow decay time kinetics were also unchanged [right panels].

(F) Super-ecliptic pHluorin [SEP] fused to GluA3 AMPARs showed the expected pH sensitivity. Washing in of acidic ACSF (pH 5) produced a dramatic reduction in the fluorescence intensity of externalized GluA3-SEP receptors. This is in line with the fact that GluA3-SEP AMPARs internalized in acidic vesicles contribute marginally to the fluorescent signal imaged.



channels was unchanged in the presence of forskolin and also comparable to that of GluA3 channels. (D) The relative fraction of openings and overall open probability of GluA1 channels was also unchanged after forskolin application and it resembled that of cAMP-activated GluA3 channels.





**Figure S7. GluA3 lacking PCs show intact excitability in-vitro and in-vivo despite their reduced synaptic transmission.**

[A] In-vivo spontaneous firing of L7/GluA3-KO PCs show comparable firing frequency and regularity of simple spikes as well as comparable amount of complex spikes, suggesting once more that, despite weaker PF to PC synapses, PC excitability is unaffected. [B] Short square steps of increasing current injected into PCs of both wild-types and GluA3-KOs showed no differences in the I/V relationships between genotypes ( $F(1,21)=2.3$ ,  $p = 0.14$ , Repeated Measures ANOVA), showing that despite the weaker synaptic transmission in the absence of GluA3, PCs have unchanged excitability in vitro. [C] Synaptic transmission is also reduced in the PC specific KO for GluA3 (L7/GluA3-KO) tested in-vitro. Error bars indicate SEM, \* indicates  $p<0.05$ .

**Figure S8. Overview of membrane resistance ( $R_m$ ) and series resistance ( $R_s$ ) of every group of PCs used to generate the experimental figures of the current study.**

Data are plotted with the same color code as in main figures. All PCs that had a change in resistance bigger than 20% over a period longer than 2 minutes were discarded for further analysis.  $R_s$  and  $R_m$  for data shown in figure 2C (A), figure 2D (B), figure 3A (C), figure 6A (D), figure 6B (E) and figure 7A-B (F). Error bars indicate SEM.



## EXTENDED EXPERIMENTAL PROCEDURES

### Mice

Inbred breeding colonies were used to obtain the experimental knockout mice. GluA1-KO mice, kindly provided by Dr. R. Huganir [Kim et al., 2005], were generated by mating heterozygous c57bl6/129 mice; GluA3-KO and wild-type littermates were bred from c57bl6x129P2-Gria3tm1Dgen/Mmnc mutant ancestors (MMRRC, Davis, CA) at least 6 times backcrossed to c57bl6 mice; and Purkinje cell specific GluA3 knockout mice were generated by crossing floxed GluA3 mice [Sanchis-Segura et al., 2006] with L7-Cre mice [Barski et al., 2000]. All experiments were conducted in line with the European guidelines for care and use of laboratory animals [Council Directive 86/609/EEC]. The experimental protocol was approved by the Animal Experiment Committee (DEC) of the Royal Netherlands Academy of Arts and Sciences (KNAW). All data of the experiments described below were acquired and analyzed in a blinded fashion with respect to the genotype.

### Eye movement recordings and oculomotor learning tasks

Baseline performance of compensatory eye movements and VOR adaptation were first tested in three groups of male mice at the age of 4-6 weeks and 3-5 months. These included wild-type littermate mice (WT,  $n = 15 + 14$ , for both age categories, respectively), GluA1 knockout mice ( $n = 5 + 6$ ) and GluA3 knockout mice ( $n = 8 + 6$ ). Mice were surgically prepared for chronic head restrained experiments [de Jeu and De Zeeuw, 2012]. During the experiment the mouse was placed head-fixed in a holder tube on a vestibular motion platform (R2000 'Rotopod', Parallel Robotic Systems Corporation, Hampton, USA). Left eye orientation was measured using video pupil tracking with a table-fixed CCD camera (Pulnix TM-6710CL, 120 frames/s) and IR illumination (850 nm LED, 6.5 cm distance from the eye). Pilocarpine (2%) eye drops were applied before the experiment to limit pupil dilatation in darkness. Online image analysis was performed to extract the location of pupil edges and corneal light reflections using custom built software for Labview (National Instruments, Austin, TX, USA). Angular eye velocity was computed offline using custom software written for Matlab (The Mathworks Inc., Natick, MA) using the algorithm outlined elsewhere [Stahl et al., 2000]. Saccadic eye movements and quick-phases of the vestibular nystagmus were removed using a 50°/s velocity threshold and 200 ms margins at each threshold crossing. Each mouse was accustomed to the setup for a period of three training days before the experimental data were collected. The horizontal VOR was characterized in both darkness and light using sinusoidal rotation about the vertical axis, using frequencies ranging between 1/16th to 4 Hz, presented in a sequence of increasing order, holding constant peak velocity of 18.8°/s. The number of cycles ranged between 5 at 1/16Hz to 60 at 4 Hz. Mice were subjected to a VOR cancellation stimulus on the first day (in-phase sinusoidal movement at 0.6 Hz, 5° amplitude of both the table and the visual surround) and a VOR reversal stimulus on subsequent days (2-5), where the amplitude of the visual surround was increased to 7.5° (day 2) and 10° (days 3, 4, and 5). The amplitude of the turntable remained constant at 5° amplitude (18.8°/s peak velocity). Training sessions consisted of 6 VOR measurements (30 cycles, 50 seconds, in darkness) that were alternated with 5 periods of visuo-vestibular mismatch training (300 cycles, 500 seconds). Apart from the training sessions, animals were kept in total darkness during the consecutive training days. The eye movement response was expressed as gain and phase relative to head movement, which was calculated using multiple linear regression of eye velocity to in-phase and quadrature components of the turntable velocity trace. Gain of the eye movement response was defined as the ratio between the eye velocity and the table velocity magnitudes. Phase was expressed in degrees and offset by 180°, so that a phase of 0° indicates an eye movement that is in-phase with contraversive head movement; positive phase values indicate phase leads. Consolidation of the adapted VOR was assessed by computing the ratio between the long-term change in VOR and the cumulative sum of short-term changes in VOR of preceding training sessions. The long-term change was defined as the absolute difference between the ending VOR on day 5 and the naive VOR on day 1. The short-term change was defined as the absolute difference between the VOR at the beginning and end of a training session.

For a period of at least 10 days animals were allowed to rest between different VOR adaptation protocols. Bivariate 2-sample Hotelling's T2-test was used to compare gain and phase values between groups, and One Way ANOVA/ Tukey post-hoc test was used for cumulative consolidation values.

### Spine density quantification in Purkinje neurons

In order to calculate the spine density in PCs, 5 WT, 4 GluA1-KO and 4 GluA3-ko mice received an overdose of sodium pentobarbital via IP injection and were perfused intracardially with 10 ml of PB 0.1M (pH 7.6) followed by 60 ml of fixative [4% paraformaldehyde in 0.1 M PB, pH7.6] at a rate of 5.5 ml/min. Brains were carefully removed from the skull, post-fixed for a maximum of 2 hours in the same fixative solution at 4°C, immersed in 30% sucrose in PB at 4°C until they sank, and subsequently cut into 40µm thick frontal sections, which were collected as four matching series. For calbindin detection, the slices were incubated in blocking solution containing 10% horse serum in 0.1 M PB to minimize non-specific binding of the antibodies. After 1 hour, blocking solution was replaced by the primary antibody solution containing 5% horse serum in 0.1M PB, rabbit anti-Calbindin antibody (Chemicon, Millipore) at a concentration of 1:1000 for 12 hours at 4°C. After several rinses with 0.1 M PB slices were incubated for 4 hours in a solution containing 5% horse serum in 0.1 M PB and horse anti-rabbit combined with Alexa 488 secondary antibody at a concentration of 1:200. After several rinses, slices were mounted and covered with Dako mounting medium (Dako), and imaged under a confocal microscope (Leica SP5). All images were acquired with the same settings and the analysis was performed with ImageJ. Stacks of pictures across the Z-axis (10-30µm) were made to count total spine number in proximal (max 30µm away from the PC soma) and distal dendrites of PCs. The spine density was calculated for each dendrite dividing the dendrite's spine count by its length; all images were processed using ImageJ. All proximal and distal dendrites counted were averaged for each mouse and mice of the same genotype were averaged to obtain the final spine densities (Figure S3A-B).

### In-vitro electrophysiology

Sagittal slices of the cerebellar vermis (250 to 400 µm thick) from 4 to 6 weeks old mice were obtained in ice-cold oxygenated "slicing" solution containing (in mM) 2.5 KCl, 1 CaCl<sub>2</sub>, 3 MgCl<sub>2</sub>, 25 NaHCO<sub>3</sub>, 1.25 NaH<sub>2</sub>PO<sub>4</sub>, 240 sucrose, 25 D-glucose and 0.01 kynurenic acid. Slices were transferred to the same slicing solution at 34°C for 5-10 minutes and then transferred to oxygenated ACSF at 34°C containing (in mM) 124 NaCl, 5 KCl, 1.25 NaH<sub>2</sub>PO<sub>4</sub>, 1 MgSO<sub>4</sub>, 2 CaCl<sub>2</sub>, 26 NaHCO<sub>3</sub>, 20 D-glucose. Subsequently, the slices were allowed to recover for at least 30 minutes until they were moved to the recording chamber containing the same oxygenated ACSF with 100 µM picrotoxin to prevent GABAergic transmission at near physiological temperature of 30±2°C. Whole-cell patch-clamp recordings of Purkinje cells located in lobules VI to X were performed using an EPC-10 amplifier (HEKA, Lambrecht). 3-5 MΩ resistance patch pipettes were filled with (in mM) K-Gluconate 122.5 mM, NaATP 4, NaGTP 0.4, HEPES 10, NaCl, KCl 9 and 0.6 mM EGTA (Sigma) at pH 7.25 for all the recordings that required current clamp mode (including LTP) or with (in mM) 115 mM cesium methanesulfonate, 20 mM CsCl, 10 mM Hepes, 2.5 mM MgCl<sub>2</sub>, 4 mM Na<sub>2</sub>ATP, 0.4 mM Na<sub>3</sub>GTP, 10 mM sodium phosphocreatine (Sigma), and 0.6 mM EGTA (Sigma), at pH 7.25 for the experiments that were exclusively done in voltage clamp mode. For both voltage-clamp and current-clamp recordings, PC membrane potential was held at -70mV to prevent spontaneous firing. Series resistance (5-10 MΩ) was measured before the experiment and compensated with standard procedures. During the experiment series and membrane resistances were monitored by applying a 100 ms hyperpolarizing pulse (-10 mV). Only cells with stable membrane and series resistance (change < 20% of the last 5 minutes of recordings compared to the last 5 minutes of baseline) were included in the analysis (Figure S8). Whole-cell recordings were digitized at 40 kHz and filtered with a Bessel filter at 4 kHz for voltage clamp recordings (8 kHz for current clamp mode). PF to PC LTD was induced by pairing PF stimulation at 1Hz for 1 minute with a 100 ms somatic depolarization from -70 mV to 0 mV, mimicking climbing fiber input (Linden, 2001; Saab et al., 2012), or by pairing PF stimulation at 1Hz for 5 minutes with real climbing fiber stimulation at 1Hz (Schonewille et al., 2011). Instead, PF to PC LTP was induced

by PF stimulation alone at 1 Hz for 5 min. To monitor EPSC amplitude over time, two test responses to a PF pulse (with 50 ms interval) were evoked every 20s in voltage-clamp mode. In LTP experiments, cells were switched to current-clamp mode for tetanization. Paired-pulse ratio (PPR) was calculated as the ratio of the amplitude of the second evoked excitatory postsynaptic current (eEPSC) to that of the first. eEPSC amplitudes and PPR were averaged per minute and normalized for final representation. For the experiments on intrinsic excitability recordings were performed in current-clamp mode, again using an EPC-10 amplifier (HEKA Electronics). Intrinsic excitability was monitored through injection of brief steps (550 ms) of increasing depolarizing current (20 steps from 50 to 1000pA). The spike count was taken as a measure of excitability. Input resistance ( $R_i$ ) was measured by injection of hyperpolarizing test currents (200 pA; 100 ms) and was calculated from the voltage transient toward the end of current injection. Recordings were excluded if the input resistance varied by  $> 20\%$ .

Single channel activity was measured in cell attached configuration with pipettes between 6-8 M $\Omega$  of resistance, containing the same intracellular solution used for whole cell recordings but containing 100  $\mu$ M of S-AMPA (Tocris). After reaching a patch resistance above 2 G $\Omega$ , the patch voltage clamp was decreased from close to resting potential (-60 mV approx.) to twice as negative (-120 mV approx.). In this configuration the ionic driving force across the channel was reversed and therefore the openings produced depolarizing events in the patch pipette. To determine the actual driving force across the AMPAR we broke into whole cell mode after the single channel recording was acquired and measured the cell resting potential. The driving potential, resulting from subtraction of the resting potential and clamped voltage, was used to calculate the receptor conductance. To further corroborate that the openings observed were caused by AMPARs, a subset of channels was also recorded close to resting potential voltages (-60 mV) and at 0 mV. When clamped close to cell resting potential, the driving force across the channel was minimal and the openings were no longer visible. When clamped at 0 mV the events detected by the pipette were of similar size, but the driving force was reversed, consistent with AMPARs behavior. For the outside-out patches of AMPA responses, pipettes with 4-6 M $\Omega$  resistance were used to establish a Giga-seal (1 G $\Omega$  resistance) with PC somata. After compensating the capacitance artifact, we let the seal rest until it reached a resistance above 2 G $\Omega$ . After breaking into whole cell mode, the pipette was slowly retracted until both the cell and the outside-out patch were re-sealed again. Every 20 seconds a 100 ms puff of 100  $\mu$ M AMPA was delivered with a Picospritzer III (Parker, Hollis, USA) to generate an AMPA-dependent response. In each sweep, a 100 ms depolarizing test pulse (-10 mV) was applied in order to test series resistance and membrane capacitance. Only patches with a constant resistance over 1 G $\Omega$  were considered for analysis. Membrane capacitance was used to control for outside-out patch size, assuming a specific membrane capacitance of 0.01 pF per 1  $\mu$ m<sup>2</sup> (Schmidt-Hieber and Bischofberger, 2010). Our patches presented comparable estimated areas of  $12.1 \pm 0.9$  and  $11.8 \pm 0.8$   $\mu$ m<sup>2</sup> in control and 8-CPT containing patches, respectively ( $p=0.42$ ).

### Drugs and pharmacology

For mEPSC recordings, tetrodotoxin (TTX, 1  $\mu$ M, Sigma) was added to the bath solution to block network activity in order to only measure excitatory spontaneous release. In order to isolate the specific contribution of AMPA and kainate receptors to glutamatergic transmission in WT and KO mice, the AMPA specific blocker GYKI52466 (30  $\mu$ M, Sigma) or the kainate specific blocker SYM2081 (5  $\mu$ M, Sigma) were added to the extracellular bath solution. For pharmacological investigation of the cAMP-GluA3 dependent pathway the following membrane permeable drugs were added to the bath of ACSF: 50  $\mu$ M Forskolin (adenylyl cyclase activator, Sigma), 20  $\mu$ M H89 (PKA antagonist, Tocris), 5  $\mu$ M KT5720 (PKA antagonist, Sigma), and 10  $\mu$ M ESI-05 (EPAC antagonist, BioLog). In addition, we applied the membrane non-permeable agonist for EPAC, 8-CPT-2Me-cAMP (20  $\mu$ M, Tocris Bioscience) to the intracellular whole cell recording solution to investigate the postsynaptic impact of EPAC. In order to obtain a monophasic time decay of the AMPA-evoked responses in outside-out patches we added a final mixture of 80  $\mu$ M PEPA (AMPA flop splice variant desensitization blocker, Tocris bioscience) and 100  $\mu$ M cyclothiazide (CTZ, AMPAR flop splice variant desensitization blocker, Tocris bioscience) to the bath solution.

### Analysis of cell physiological data

Spontaneous mEPSC and evoked EPSC recordings were analyzed with MiniAnalysis software (Synaptosoft) and ClampFit (Molecular Devices), respectively. To calculate  $T_{fast}$  (fast desensitizing component) and  $T_{slow}$  (slow non-desensitizing component) of AMPA evoked currents in outside-out patches a double exponential function was fitted using ClampFit with DC offset set at 0. The decay of the averaged current was fitted to the following equation:

$$I = A_1 e^{-t/\tau_1} + A_2 e^{-t/\tau_2}$$

In this equation  $T_1$  represents  $T_{fast}$ . The percentage of the decay represented by the slow component ( $\%_{slow}$ ) was calculated by the function  $A_1/(A_1+A_2)$ , as described elsewhere [Christian et al., 2013]. The weighted decay time constant for AMPA evoked currents in outside-out patches in the presence of desensitization blockers was calculated by dividing the total charge transfer (in fC) by the peak amplitude (in pA). Non-stationary fluctuation analysis of outside-out patches traces was carried out following previously described methods [Alvarez et al., 2002; Benke et al., 2001; Hartveit and Veruki, 2007]. In short, peak aligned AMPA evoked currents recorded over 10-15 sweeps per outside-out patch were binned in 10 equally sized bins of 150 ms each and for each bin the mean amplitude and variance were calculated. The data distribution resulting after plotting amplitude versus variance was fitted with the following equation:

$$\sigma^2 = iI - \frac{I^2}{N} + \sigma_b^2$$

Where the variance ( $\sigma^2$ ) of the amplitude of the current ( $I$ ) obtained at each time point is explained as a function of the single unitary current ( $i$ ) and the number of functional conducting channels ( $N$ ) with an offset given by the variance of the baseline noise ( $\sigma_b^2$ ). The number of functional channels was extracted from the derivative at  $I = 0$ , and the single channel conductance was calculated by dividing the unitary current by the applied voltage with respect to the reversal potential ( $V_{holding} - E_{reversal}$ , -70 mV and 0 mV, respectively). The peak open probability ( $P_o$ ), which corresponds to the fraction of available functional channels open at the time of the peak current ( $I_{peak}$ ), was calculated from the following equation:

In this equation  $N_{max}$  represents the theoretical maximum of available channels opened at the point where the theoretical maximum amplitude reaches the minimum variability ( $\sigma_b^2$ ) in the given parabola fit.

Single channel activity was analyzed using ClampFit (Molecular Devices). Three detection thresholds were used to detect O1 (1.5 pA), O2 (3 pA) and O3 (4.5 pA) openings in single channel AMPA receptors in steady baseline recordings (no holding current fluctuations). Events with a latency shorter than 0.3 ms were ignored to prevent noise to be recognized as openings.

### Statistics

For statistical analysis of behavioral and in-vitro electrophysiological data we used either Matlab statistical toolbox (The MathWorks Inc., Natick, MA, 2000) or GraphPad Prism 6 (La Jolla, California, USA). Although Matlab always reports exact p-values, GraphPad Prism 6 does not report exact values when  $p < 0.0001$ . Thus we have reported exact p-values when possible, taken into consideration the limitation explained above.

### In-vitro two-photon imaging

Organotypic cerebellar slices were made from P7-9 mice using a protocol adapted from previous studies [Hurtado de et al., 2011; Stoppini et al., 1991] and kept in culture 4-7 days prior to the experiments. Slices were then transfected with sindbis virus expressing rat flip GluA3 AMPAR fused to the pH sensitive version of GFP Super Eccliptic pHluorophor (GluA3-SEP) for a period of 24-48 hours prior to the imaging session. Electrophysiological

recordings of PC mEPSCs were performed in this preparation. In our hands, mEPSC amplitudes and frequencies were consistently higher in organotypic cultured PCs than in acute (e.g. Fig 2B and 4B, WT-Acute vs. WT-Organotypic  $p = 0.0002$  and  $p < 0.0001$  for amplitude and frequency, respectively), using the same concentration of TTX and PTX. For imaging, slices were transferred from the incubation solution to the recording chamber containing ACSF [same composition as mentioned before but with  $4\ \mu\text{M}$  calcium and  $4\ \mu\text{M}$  magnesium]. Three-dimensional images were collected on a custom-built two-photon microscope based on a Fluoview laser-scanning microscope (Olympus). The light source was a mode-locked Ti:sapphire laser (Chameleon, Coherent) tuned at 850 nm using a 60x objective. Optical sections were captured every  $0.5\ \mu\text{m}$  from transfected PC dendrites. Fluorescence intensity was quantified from projections of stacked sections using ImageJ software (NIH). For single spine bleaching in the FRAP experiments, a ROI was selected covering the surface of a single spine, which was used to target the laser for 20-30 seconds [with the same intensity as for regular imaging].

### **In-vivo electrophysiology**

Mice (males, 4-6 month old) were prepared for chronic experiments as described previously (Wulff et al., 2009). In short, under general anesthesia a pedestal with a magnet was placed on the frontal and parietal bones of the animal, and a recording chamber was constructed around a small craniotomy in the left occipital bone. After 2 days of recovery, animals were habituated in the setup for 20 min for two days. During the experiments, the animals were alert and immobilized in a custom restrainer. Extracellular activities were recorded with glass micropipettes filled with 2M NaCl solution and advanced into the cerebellar cortex from the surface of Crus I and II. Electrode signals were filtered, amplified and stored for off-line analyses (Spike2, CED, and Cambridge, UK). PCs were identified by the occurrence of both simple spikes and complex spikes, and single-unit activity was confirmed by a brief pause in simple-spike firing following each complex spike (i.e. climbing fiber pause; see De Zeeuw et al., 2011). The whole field visual stimulation was presented by rotating a cylindrical screen (diameter 63 cm) with a random-dotted pattern (each element  $2^\circ$ ) at 0.6 Hz with an amplitude of  $5^\circ$ . Offline analysis was conducted in Matlab (Mathworks, Natick, MA, USA). CV2 of simple spikes was calculated as the mean value of  $[2 \times (IS_{In+1} - IS_{In})] / (IS_{In+1} + IS_{In})$  (Wulff et al., 2009). Modulation of simple spikes and complex spikes was calculated as the amplitude of the sine wave fitted to the histogram of spike rate. Statistical analysis was done using Student's t-test with SPSS (IBM Corporation, Armonk, NY, USA).

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