Neurobiology of Disease

Altered Neuron Excitability and Synaptic Plasticity in the Cerebellar Granular Layer of Juvenile Prion Protein Knock-Out Mice with Impaired Motor Control

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Abstract

Although the role of abnormal prion protein (PrP) conformation in generating infectious brain diseases (transmissible spongiform encephalopathy) has been recognized, the function of PrP in the normal brain remains mostly unknown. In this investigation, we considered the effect of PrP gene knock-out (PrP0/0) on cerebellar neural circuits and in particular on granule cells, which show intense PrP expression during development and selective affinity for PrP. At the third postnatal week, when PrP expression would normally attain mature levels, PrP0/0 mice showed low performance in the accelerating rotarod and runway tests and the functioning of 40% of granule cell membrane currents, and also the EPSPs and EPSCs had slow kinetics. Overall, these alterations closely resembled an immature phenotype. Moreover, in slow-spiking PrP0/0 granule cells, theta-burst stimulation was unable to induce any long-term potentiation. This profound impairment in synaptic excitation and plasticity was associated with a protracted proliferation of granule cells and disappeared at P40–P50 along with the recovery of normal motor behavior (Büeler et al., 1992). These results suggest that PrP plays an important role in granule cell development eventually regulating cerebellar network formation and motor control.

Key words: cerebellum; granule cell; prion; LTP; action potential; motor control

Introduction

The prion protein (PrP) is a cell surface protein, whose abnormal conformation leads to transmissible spongiform encephalopathy (TSE), including “mad cow” disease in bovines, scrapie in sheep, and Creutzfeld–Jakob disease in humans (Prusiner, 1998; Harris and True, 2006). Brain alterations consist in vacuolization of neurons and synapses associated with PrP accumulation in polymeric aggregates. Prion infection causes a slowly progressive neurodegeneration impairing higher brain functions and motor control.

In face of the wealth of evidence on pathogenetic mechanisms, PrP physiological functions remain unclear. A first investigation on PrP knock-out mice (PrP0/0) (Büeler et al., 1992) did not reveal macroscopic developmental and behavioral alterations. However, subsequent analysis revealed several motor, cognitive, and emotional abnormalities (Roesler et al., 1999; Criado et al., 2005) and increased susceptibility to seizures (Walz et al., 2002). Recordings from brain slices of PrP0/0 mice showed reduced synaptic inhibition and long-term potentiation (LTP) (a neural correlate of learning) (Bliss and Collingridge, 1993) in the hippocampus (Collinge et al., 1994; Curtis et al., 2003; Maglio et al., 2004, 2006), but this observation was not confirmed (Lledo et al., 1996).

Several abnormalities have been observed in PrP0/0 neurons in culture, including reduced Ca2+-dependent K+ currents (Colling et al., 1996; Herms et al., 2001; Mallucci et al., 2002) and cytoplasmic Ca2+ levels (Herms et al., 2000). PrP is distributed on the neuron surface and in presynaptic terminals (Herms et al., 1999; Ferrer et al., 2000). Despite initial negative observations, there is now growing evidence that PrP regulates neuronal survival and differentiation (Steele et al., 2006) through its interaction with laminin and intracellular signaling cascades (Graner et al., 2000). In keeping with this, PrP expression is highly regulated during ontogenesis and by nerve growth factor, and susceptibility to prion diseases is maximal during development (Manson et al., 1992; Martins et al., 2002).

Although multiple structures of the CNS (including neocortex, thalamus, hippocampus, and cerebellum) are affected, PrP physiopathology is significantly related to the cerebellum and cerebellar ataxia appears to be one of the main clinical signs in a number of TSEs (Prusiner, 1998). Binding of prion protein to cerebellar granule cells may have a distinct role in the pathogenetic mechanism (Shmerling et al., 1998; Legname et al., 2002).
PrP is expressed in the cerebellum granular layer (Lainé et al., 2001) showing a characteristic developmental profile, with a steep increase during the second postnatal week (Sales et al., 2002). Interestingly, PrP expression parallels the major ontogenetic changes characterizing the mossy fiber–granule cell relay (Altman, 1972; Hámori and Somogyi, 1983; D’Angelo et al., 1993, 1994, 1997; Farrant et al., 1994; Rossi et al., 1994, 1998, 2002; Rumbaugh and Vicini, 1999; Cathala et al., 2000, 2003; Wall et al., 2002) and goes along with a marked improvement of supporting sensorimotor reactions (Altman and Sudarshan, 1975).

In this study, we investigated the effect of PrP knock-out at the cerebellar mossy fiber–granule cell relay (Eccles et al., 1967) in juvenile PrP<sup>0/0</sup> mice. PrP<sup>0/0</sup> mice showed low performance in sensorimotor tests. At the same time, a proportion of granule cells showed an immature electrophysiological phenotype and their synapses lacked long-term potentiation (D’Angelo et al., 1999; Armano et al., 2000; Hansel et al., 2001; Rossi et al., 2002). These alterations were associated with an abnormal duplication of granule cells in the germinal zone of the external granular layer (EGL) during the third postnatal week but fully recovered in the adulthood, suggesting an important role of PrP for cerebellar network formation and motor control during development.

Materials and Methods

Experimental animals

The prion protein knock-out (PrP<sup>0/0</sup>) mice were of the "Zurich-1" strain developed by Büeler et al. (1982). The wild-type mice (C57 × Sv129) are descendents of F<sub>2</sub> generation mouse produced by interbreeding the Goldfarb et al. (2007) to improve tissue viability. Kreb’s solution for cutting and recovery was modified as reported by Bu¨eler et al. (1992). The wild-type mice (C57BL/6J, which is the most common inbred strain used for experimen-
tal electrophysiology. Most experiments were performed on juvenile mice [postnatal day 17 (P17) to P22; day of birth, P0], when granule cells have just attained mature structural (Altman, 1972; Hámori and Somogyi, 1983), excitable, and synaptic properties (D’Angelo et al., 1993, 1994, 1997; Farrant et al., 1994; Rossi et al., 1994, 1998, 2002; Rumbaugh and Vicini, 1999; Cathala et al., 2000, 2003; Wall et al., 2002) and controls have also been performed at different ages.

Electrophysiological recordings

All experiments were performed according to the guidelines laid down by the institution’s animal welfare committee. We performed whole-cell patch-clamp recordings from granule cells in acute cerebellar as reported previously (D’Angelo et al., 1999) from P17–P22 mice. Briefly, mice were killed by decapitation after anesthesia with halothane (1 ml in 2 L administered for 1–2 min). The cerebellum was gently removed, and the vermis was isolated and fixed on the stage of a vibroslicer (Dosaka) with cyano-acrylic glue. Acute 250-μm-thick slices were cut in the parasagittal plane in cold Kreb’s solution and maintained at room temperature before being transferred to a 1.5 ml recording chamber mounted on the stage of an upright microscope (Olympus). The slices were perfused with Kreb’s solution and maintained at 32°C with a Peltier feedback device (HCC-100A; Dagan Corporation). The Kreb’s solution contained the following (in mM): 126 K-glutonate, 4 NaCl, 5 HEPES, 15 glucose, 1 MgSO<sub>4</sub>·7H<sub>2</sub>O, 0.1 BAPTA-free and 0.05 BAPTA-Ca<sup>2+</sup>, 3 Mg-ATP, pH adjusted to 7.2 with KOH. Just after obtaining the cell-attached configuration, electrode capacitance was carefully cancelled to allow for electronic compensation of pipette charging during subsequent current-clamp recordings, and the amplifier was set to the “fast” operating mode (D’Angelo et al., 1995, 1998). At the beginning of each recording, passive transients were elicited to monitor passive cell properties (see below) and a series of depolarizing steps was applied in voltage clamp to measure the total voltage-dependent current of the granule cell (see Fig. 2). Leakage and capacitive currents were subtracted using hyperpolarizing pulses delivered before the test pulse (P/4 protocol). After switching to current clamps, intrinsic excitability was investigated (see Fig. 4) by setting resting membrane potential at −80 mV and injecting 2 or 800 ms steps of current from (from −8 to 48 pA in 2 or 4 pA increments).

Membrane potential during 2 s step current was estimated as the average value between 1.7 and 2 s, whereas membrane potential during 800 ms step current was estimated as the average value between 500 and 800 ms.

| Table 1. Properties of whole-cell recordings in mice granule cells |
|------------------|------------------|------------------|------------------|
|                  | C57BL/6J (n = 22) | C57 × Sv129 (n = 23) | PrP<sup>0/0</sup> fast (n = 12) | PrP<sup>0/0</sup> slow (n = 8) |
| R<sub>c</sub> (GΩ) | 1.9 ± 0.2*        | 1.6 ± 0.2*        | 1.9 ± 0.2*        | 3.0 ± 0.5        |
| C<sub>m</sub> (pf) | 3.8 ± 0.2         | 3.9 ± 0.2         | 4.0 ± 0.4         | 4.6 ± 0.4         |
| R<sub>m</sub> (MΩ) | 21.2 ± 1.6        | 20.5 ± 1.7        | 19.8 ± 3.3        | 17.7 ± 2.2        |
| f<sub>c</sub> (kHz) | 2.3 ± 0.2         | 2.2 ± 0.1         | 2.3 ± 0.2         | 2.2 ± 0.2         |
| γ<sub>c</sub> (µs)  | 81.6 ± 8.1        | 76.0 ± 5.0        | 75.1 ± 8.9        | 77.2 ± 7.3        |

The data were obtained using K<sup>+</sup>-gluconate intracellular solution and analyzing current transients elicited by −10 mV voltage-clamp steps delivered from the holding potential of −70 mV. The number of observations is indicated, and statistical significance is reported in comparisons with slow-spiking PrP<sup>0/0</sup> granule cells.

* p < 0.05, unpaired t test.

Whole-cell recording properties. The cerebellar granule cell is electrotonically compact and can be treated as a simple RC system, in which relevant parameters can be extracted by analyzing passive current relaxation induced by step voltage changes. In each recording, once in the whole-cell configuration, the current transients elicited by 10 mV hyperpolarizing pulses from the holding potential of −70 mV in voltage-clamp mode showed a biphasic relaxation, with a major component (−95%) related to somatodendritic charging (see supplemental material, available at www.jneurosci.org). According to previous reports (D’Angelo et al., 1995, 1999; Silver et al., 1996), the major component was analyzed to extract basic parameters useful to evaluate the recording conditions and to compare different cell groups. Membrane capacitance (C<sub>m</sub>) was measured from the capacitive charge (the area underlying current transients) and series resistance (R<sub>s</sub>) was calculated as R<sub>s</sub> = γ<sub>c</sub>/C<sub>m</sub>. The membrane resistance (R<sub>m</sub>) was computed from the steady-state current flowing after termination of the transient. The −3 dB cutoff frequency of the electrode-cell system, f<sub>c</sub>, was calculated as f<sub>c</sub> = (2πτ<sub>c</sub>)<sup>−1</sup>. The data are reported in Table 1. The electrotonic properties were further analyzed and simulated using a multicompartmental model, as explained in the supplemental material (available at www.jneurosci.org).

Granule cell excitability. Patch pipettes had of 8–12 MΩ resistance before seal formation with a filling solution containing the following (in mM): 126 K-gluconate, 4 NaCl, 5 HEPES, 15 glucose, 1 MgSO<sub>4</sub>·7H<sub>2</sub>O, 0.1 BAPTA-free and 0.05 BAPTA-Ca<sup>2+</sup>, 3 Mg-ATP, pH adjusted to 7.2 with KOH. Just after obtaining the cell-attached configuration, electrode capacitance was carefully cancelled to allow for electronic compensation of pipette charging during subsequent current-clamp recordings, and the amplifier was set to the “fast” operating mode (D’Angelo et al., 1995, 1998). At the beginning of each recording, passive transients were elicited to monitor passive cell properties (see below) and a series of depolarizing steps was applied in voltage clamp to measure the total voltage-dependent current of the granule cell (see Fig. 2). Leakage and capacitive currents were subtracted using hyperpolarizing pulses delivered before the test pulse (P/4 protocol). After switching to current clamp, intrinsic excitability was investigated (see Fig. 4) by setting resting membrane potential at −80 mV and injecting 2 or 800 ms steps of current from (from −8 to 48 pA in 2 or 4 pA increments).

Membrane potential during 2 s step current was estimated as the average value between 1.7 and 2 s, whereas membrane potential during 800 ms step current was estimated as the average value between 500 and 800 ms.
ms. Membrane potential during TBS was estimated as the mean of average values measured in the central 70 ms of each burst (tracings were filtered at 100 Hz). Action potentials in granule cells show a prepotential followed by the upstroke (D’Angelo et al., 1997; Armano et al., 2000). Action potential threshold was measured along the raising phase of membrane potential responses to step current injection. The prepotential threshold was identified at the flexus starting the regenerative process (a procedure that was improved by taking the first time derivative of membrane potential), whereas the upstroke threshold was identified by taking the second time derivative of the signal. Spike duration at half-amplitude was evaluated using the first spike generated with just-threshold current stimulation and measured at the midpoint between the threshold of spike upstroke and peak. The amplitude of the action potential overshoot was estimated as the difference between the threshold of spike upstroke and the maximum reached potential. The amplitude of the action potential afterhyperpolarization (AHP) was estimated as the difference between spike threshold and the lowest potential after the upstroke. Action potential frequency was measured by dividing the number of spikes by the current injected duration.

In a specific set of recordings, synaptic responses were elicited at 0.1–0.33 Hz to activate the mossy bundle while measuring EPSPs from a resting membrane potential of −70 mV. EPSPs were averaged and analyzed off-line to measure peak amplitude and decay time constant through a simple exponential fitting. LTP was induced by TBS delivered 10 min after establishing the whole-cell recording configuration (time = 0) from the membrane potential of −50 mV. In LTP recordings in current clamp, stability was monitored by measuring the resting membrane potential and the input resistance in a low membrane potential range (less than −70 mV).

Synaptic currents. Patch pipettes had 5–8 MΩ resistance before seal formation with a filling solution containing the following (in mM): 81 Cs2SO4, 4 NaCl, 2 MgSO4, 1 QX-314 (lidocaine N-ethyl bromide), 0.1 BAPTA-free and 0.05 BAPTA-Ca2+, 15 glucose, 3 Mg-ATP, 0.1 GTP, and 15 HEPES, pH adjusted to 7.2 with CsOH. The EPSCs were elicited at 0.33–0.1 Hz by interleaving periods at +60 mV and at −70 mV to modify the NMDA/AMPA current ratio. EPSCs were averaged and digitally filtered at 1.5 kHz off-line. EPSC peak amplitude and EPSC amplitude 25 ms after stimulation were taken at −70 and +60 mV to measure the non-NMDA and NMDA currents, respectively. The decay phases of EPSCs at either potentials were fitted with double exponentials of the following form by a nonlinear least-squares method using pClamp software (Molecular Devices): \[ A(t) = A_1 \exp(-t/\tau_1) + A_2 \exp(-t/\tau_2), \] where \( A(t) \) is the amplitude of the decay phase of the EPSC or EPSP as a function of time \( t \), \( A_1 \) and \( A_2 \) are the amplitudes of the fast and slow decay components, and \( \tau_1 \) and \( \tau_2 \) are their respective decay time constants. For a single exponential decay, \( A_2 \) was set to zero. The weighted decay time constant (Rumbaugh and Vicini, 1999) was estimated as follows: \( \tau = \tau_1 \times [A_1/(A_1 + A_2)] + \tau_2 \times [A_2/(A_1 + A_2)] \). Fittings were performed from peak at −70 mV and from 25 ms after the stimulus at +60 mV.

In some experiments, after a 10 min control period, LTP was induced by TBS while stepping membrane potential from −70 to −30 mV. In LTP recordings in voltage clamp, \( R \), stability was monitored throughout the experiment.

Behavioral tests
The behavioral analysis was performed on naive, juvenile PrP<sup>−/−</sup> and wild-type (C57 × Sv129) mice. At the first day of the test the animals were P19. The animals were housed in groups of three to four in clear plastic cages maintained in a temperature- and humidity-controlled room on a 12 h light/dark schedule with food and water provided ad libitum. All experiments were conducted in the light phase of circadian cycle between 9:00 A.M. and 4:00 P.M. Every day, animals were sequentially subjected to the runway and accelerating rotarod test. The study was approved by the Institutional Ethical Committee of the School of Medicine, Université Libre de Bruxelles, Belgium.

In the runway test, mice had to run along an elevated runway with low obstacles intended to impede their progress. The runway was 100 cm long and 0.7 cm wide, and obstacles of 1 cm diameter were placed every 10 cm along the runway. Mice were placed on one brightly illuminated
were deeply anesthetized (pentobarbital, 60 mg/kg) and transcardially perfused with 4% paraformaldehyde plus 0.05% glutaraldehyde in cold 0.1 M phosphate buffer. The dissected cerebellum was further postfixed in 4% paraformaldehyde plus 15% sucrose for 2 hours at 4°C. Its vermal part was cut in 80-μm-thick parasagittal sections with a vibrotome. Sections were then cryoprotected and permeabilized by freeze-thawing. Preembedding was performed with a standard free-floating immunocytochemical procedure using 0.1 M saline phosphate buffer as detergent. Briefly, aldehyde quenching in 0.1 M glycine was followed by preincubation at 4°C in SAF 32, a monoclonal antibody directed against the N-terminal region of PrP (Demart et al., 1999). After incubation in 1/200 biotinylated horse anti-mouse IgG (Vector Laboratories), a standard avidin–biotin complex revelation (Vectorstain Elite; Vector Laboratories) was applied, and characterized as recognizing the human oc-tarepeat sequence 52–92 (Demart et al., 1999). During this training session, every mouse was placed on the rod rotating at a constant speed (4 rpm). As soon as the animals were placed on the rod, the rod started to accelerate continuously from 4 to 40 rpm over 300 s. The latency to fall off the rotarod was recorded. Animals staying during 300 s were taken from the rotarod and recorded as 300 s.

Strains were compared using a two-way ANOVA and Bonferroni’s post hoc test. Results are expressed as mean ± SEM and were considered significant if p < 0.05.

**Morphological techniques Immunoelectron microscopy.** P20 C57 × Sv129 mice were deeply anesthetized (pentobarbital, 60 mg/kg) and transcardially perfused with 4% paraformaldehyde plus 0.05% glutaraldehyde in cold 0.1 M phosphate buffer. The dissected cerebellum was further postfixed in 4% paraformaldehyde plus 15% sucrose for 2 hours at 4°C. Its vermal part was cut in 80-μm-thick parasagittal sections with a vibrotome. Sections were then cryoprotected and permeabilized by freeze-thawing. Preembedding was performed with a standard free-floating immunocytochemical procedure using 0.1 M saline phosphate buffer as detergent. Briefly, aldehyde quenching in 0.1 M glycine was followed by preincubation at 4°C in SAF 32, a monoclonal antibody directed against the N-terminal region of PrP (Demart et al., 1999). After incubation in 1/200 biotinylated horse anti-mouse IgG (Vector Laboratories), a standard avidin–biotin complex revelation (Vectorstain Elite; Vector Laboratories) was applied, using 0.05% diaminobenzidine as the chromagen. For electron microscopy (EM), after 2% OsO4 postfixation and 2% uranyl acetate en bloc staining, selected sections were dehydrated in graded acetone and finally embedded in Durcupan (Fluka) resin. Un-counterstained ultrathin sections were examined with a Philips CM100 electron microscope, operated at 60 kV.

**5-Bromodeoxyuridine immunostaining.** At different postnatal days (8, 12, 15, 20, 25, 40), PrP0/0 and control C57BL/6J mice were injected intraperitoneally with 5-bromodeoxyuridine (BrDU) (Sigma-Aldrich) (50 μg/g body weight) dissolved in ethanol 25% in PBS. Three hours after treatment, animals were killed by decapitation, and brains were excised and placed into Carnoy’s fixative (ethanol absolute/chloroform/acetic acid, 6:3:1) overnight. Subsequently, brains were dehydrated through a series of graded ethanols and embedded in Paraplast X-tra. Sagittal 10 μm sections were cut and mounted onto gelatin-coated slides. The slides were deparaffinized in xylene, rehydrated, and processed for immunohistochemistry for BrDU. Sections were treated with proteinase K (20 μg/ml; Sigma-Aldrich) for 7 min at room temperature and then with 2N HCl in 0.1 M PBS for 45 min to enhance binding of the primary antibody. After that, sections were incubated in 3% H2O2, in 10% methanol in 0.1 M PBS for 7 min, to block endogenous peroxidase, and then sections were exposed to 5% nonfat milk in 0.1 M PBS containing 0.5% Triton X-100 and 0.1% NaN3. Thereafter, the sections were incubated overnight with an anti-BrDU antibody (clone B44; BD Biosciences) diluted 1:50 in 0.1 M PBS containing 0.5% Triton X-100 and 0.1% NaN3. The primary antibody was detected by the biotin–streptavidin technique according to the procedure suggested by the supplier (Vectastain ABC Elite kit; Vector

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**Figure 2.** Granule cell electrosensitivity. Voltage responses were elicited from −80 mV using step current injection (intensities are indicated below tracings) in wild-type and PrP0/0 mice. A, Examples of C57 × Sv129, C57BL/6J, and PrP0/0 granule cells generating fast repetitive spike discharge. Note also the fast inward rectification in the subthreshold voltage response. B, A slow-spiking PrP0/0 granule cell generates a broad nonrepetitive action potential and shows no inward rectification. C, Threshold-aligned wild-type and PrP0/0 action potentials. Note smaller spike overshoot, smaller AHP, and greater half-width in the slow-spiking PrP0/0 granule cell. D, Subthreshold steady-state depolarization (left) and discharge frequency (right) in the different groups of granule cells. In the plot at the left, relative membrane potentials were calculated by dividing the actual values by −70 mV. The straight dashed line interpolates the points more negative than −70 mV in slow-spiking PrP0/0 granule cells, evidencing the absence of inward rectification, which conversely characterizes the other three cell groups. In the plot at the right, note that slow-spiking PrP0/0 granule cells have very low frequency of discharge on current injection (in the case that only one spike was observed, the frequency was set to 0). Data are mean ± SEM (C57BL/6J, n = 22; C57 × Sv129, n = 23; PrP0/0 fast, n = 12; PrP0/0 slow, n = 8). Note in the inset that slow-spiking PrP0/0 granule cells were evenly distributed in the P17–P22 age window used for the present recordings (gray, slow-spiking PrP0/0; hatched, fast-spiking PrP0/0).
granule cells, a voltage command of Figure 3. Voltage-activated inward and outward currents in granule cells. In slow-spiking PrP0/0 granule cells, the outward steps from the holding potential of and inward currents are smaller than in wild-type granule cells. slow-spiking granule cell) and n

<table>
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<tr>
<th>V&lt;sub&gt;rest&lt;/sub&gt; (mV)</th>
<th>CS7BL/6J (n = 22)</th>
<th>CS7 × Sv129 (n = 23)</th>
<th>PrP&lt;sup&gt;0/0&lt;/sup&gt; fast (n = 12)</th>
<th>PrP&lt;sup&gt;0/0&lt;/sup&gt; slow (n = 8)</th>
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<tr>
<td>K&lt;sub&gt;out&lt;/sub&gt; (mV)</td>
<td>−60.7 ± 2.8*</td>
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<td>Aν0 (mV)</td>
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<td>49.1 ± 4.0***</td>
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<td>Aν0 (mV)</td>
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<td>Aν0 (mV)</td>
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<td>(−36.0 ± 2.2*)</td>
<td>(−36.6 ± 0.7***)</td>
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<td>(−29.1 ± 1.8)</td>
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<td>Aν1 (ms)</td>
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<td>1.4 ± 0.2***</td>
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<td>EPSP&lt;sub&gt;0&lt;/sub&gt; (mV)</td>
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<td>EPS&lt;sub&gt;0&lt;/sub&gt; (ms)</td>
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<td>183.4 ± 22.7***</td>
<td>170.8 ± 13.9***</td>
<td>198.2 ± 36.9***</td>
<td>107.5 ± 10.8</td>
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The table also reports values for the maximum transient inward current (I<sub>fast,peak</sub>) and the persistent outward current at +20 mV (I<sub>out,hold</sub> + 20). The number of observations is indicated and statistical significance is reported in comparisons with slow-spiking PrP<sup>0/0</sup> granule cells.

*p < 0.05, **p < 0.001, ***p < 0.001, unpaired test.

Figure 3. Voltage-activated inward and outward currents in granule cells. In slow-spiking PrP<sup>0/0</sup> granule cells, the outward and inward currents are smaller than in wild-type granule cells. A. Voltage-dependent currents evoked by depolarizing voltage steps from the holding potential of −70 mV. The inset shows the VC protocol (note that, to elicit inward currents in the PrP<sup>0/0</sup> granule cells, a voltage command of −10 mV rather than −30 mV was needed). Traces were leak-subtracted (see Materials and Methods). B. The scatter plot shows inward and outward current amplitudes measured at −30 mV (except for −10 mV in the slow-spiking granule cell) and +20 mV in experiments like those shown in A, C. I-V relationships for the peak inward current and the steady-state outward current in the different groups of granule cells. Note smaller currents in slow-spiking PrP<sup>0/0</sup> granule cells, in which the inward current attains its negative peak at higher potential. Data are mean ± SEM (CS7BL/6J, n = 22; CS7 × Sv129, n = 23; PrP<sup>0/0</sup> fast, n = 12; PrP<sup>0/0</sup> slow, n = 8).

Laboratories), using 3,3'-diaminobenzidine tetrahydrochloride as chromogen. BrdU-positive cells were counted over 100 strips of 150 μm length of external granule cell layer in each of three wild-type and PrP<sup>0/0</sup> mice.

Results

In the cerebellum granular layer, PrP attains high expression levels by the third postnatal week (Sales et al., 2002). By using EM immunostaining, at P17–P22 we observed PrP expression on granule cell somata and dendrites and on mossy fibers of C57 × Sv129 mice (Fig. 1). At high magnification, PrP demonstrates a punctate immunoprecipitation on the plasma membrane. The extensive expression of PrP on the plasma membrane of glomerular structures suggests a role for the appropriate functioning of the cerebellar circuit. Conversely, in PrP knock-out mice (PrP<sup>0/0</sup>) (Bueller et al., 1992), PrP is absent from all structures forming the mossy fiber–granule cell relay (Lainé et al., 2001). We therefore investigated whether the electrophysiological properties of the mossy fiber–granule cell relay were altered by PrP knock-out at the end of the third postnatal week.

Altered electroresponsiveness in PrP<sup>0/0</sup> granule cells

As reported in previous investigations on rats and mice cerebella (D’Angelo et al., 1995, 1998; Brickley et al., 1996; Rossi et al., 1998; Chadderton et al., 2004), wild-type granule cells in the present recordings were characterized by fast repetitive spike discharge (Fig. 2 A,C, Table 2). Similar properties were also observed in 60% (n = 12 of 20) of PrP<sup>0/0</sup> granule cells. However, the remaining 40% (n = 8 of 20) of PrP<sup>0/0</sup> granule cells showed a single or a few slow spikes with marked adaptation, high threshold, small overshoot, and small AHP (Fig. 2 B,D, Table 2), and were therefore called “slow-spiking.” Slow-spiking PrP<sup>0/0</sup> granule cells showed very low discharge frequency (often just one spike was generated) compared with fast-spiking (either wild-type or PrP<sup>0/0</sup>) granule cells (Fig. 2 D, right). Slow-spiking PrP<sup>0/0</sup> granule cells did not show inward rectification, which was evident in fast-spiking (either wild-type or PrP<sup>0/0</sup>) granule cells (Fig. 2 B,D, left), and showed a slight reduction in resting membrane potential (Table 2).
The slow-spiking PrP<sup>0/0</sup> granule cells were distributed over the 6 d on which recordings were performed (Fig. 2D, inset), and their properties did not depend on series resistance, which was similar regardless of the recorded cell group (compare Table 1). Both in their fast-spiking and slow-spiking pattern, the granule cells had typical membrane capacitance (Table 1) confirming the normal size and homogeneous shape observed in EM recordings (Fig. 1) (Laine et al., 2001). Moreover, regardless of firing pattern, current relaxations elicited by voltage-clamp steps were biphasic with almost identical decay components (the major fast component is related to somatodendritic charging and a smaller slow component is related to axon). The current relaxations were reproduced with a mathematical model supporting axon integrity in all recorded granule cells (see supplemental material, available at www.jneurosci.org).

In the same experiments, whole-cell currents differed when recorded from slow-spiking compared with fast-spiking granule cells (Fig. 3, Table 2). The “transient inward current” (corresponding to a fast Na<sup>+</sup> current) (Magistretti et al., 2006) was significantly smaller in slow-spiking PrP<sup>0/0</sup> compared with fast-spiking (either wild-type or PrP<sup>0/0</sup>) granule cells and reached its inward peak at higher potentials (Fig. 3C). The “transient and persistent outward currents” (corresponding to A-type, delayed rectifier, and calcium-dependent K<sup>+</sup> currents) (Bardoni and Belluzzi, 1993) were also significantly smaller in slow-spiking PrP<sup>0/0</sup> compared with fast-spiking (either wild-type or PrP<sup>0/0</sup>) granule cells. Thus, the impairment of intrinsic electroresponsiveness in slow-spiking PrP<sup>0/0</sup> granule cells was correlated with the abnormal expression of voltage-dependent membrane currents.

In the same recordings, the EPSPs elicited by mossy fiber stimulation showed comparable size but slower kinetics in slow-spiking PrP<sup>0/0</sup> granule cells compared with fast-spiking (either wild-type or PrP<sup>0/0</sup>) granule cells (Fig. 4, Table 2). Therefore, not just intrinsic electroresponsiveness but also synaptic transmission was altered in slow-spiking PrP<sup>0/0</sup> granule cells.

**Altered synaptic transmission at the PrP<sup>0/0</sup> mossy fiber–granule cell relay**

In specific experiments, glutamate receptor-mediated synaptic currents were measured both at −70 and +60 mV to identify the differential contribution of the AMPA and NMDA components (Fig. 4C). Actually, whereas the NMDA current is relatively slower and shows a marked increase with depolarization, the AMPA current is much faster and generates large transient responses at negative membrane potentials (D’Angelo et al., 1995, 1999; Cathala et al., 2003). These recordings were performed using Cs<sup>+</sup>-based intracellular solutions to improve voltage clamp at depolarized potentials, so that PrP<sup>0/0</sup> cells could not be sorted based on their spike shape.

The AMPA current [estimated at the EPSC peak at −70 mV: C57BL/6J, −62.2 ± 11.3 pA (n = 13); C57 × Sv129, −54.2 ± 14.1 pA (n = 20); PrP<sup>0/0</sup>, −46.1 ± 5.7 (n = 26)] and the NMDA current [estimated 25 ms after the stimulus: C57BL/6J, 59.5 ± 11.5 pA (n = 13); C57 × Sv129, 60.7 ± 13.9 pA (n = 20); PrP<sup>0/0</sup>, 43.6 ± 8.5 (n = 26)] showed similar amplitude in the different strains (Fig. 4C). However, EPSCs were on average slower in PrP<sup>0/0</sup> than in wild-type granule cells (Fig. 4D). The rate of decay was estimated by performing double-exponential fittings and calculating the weighted time constant (τ<sub>w</sub>) (see Materials and Methods). At −70 mV, PrP<sup>0/0</sup> EPSCs often showed a large late component significantly increasing τ<sub>w</sub> values compared with wild-type mice [C57BL/6J, 5.8 ± 0.5 ms (n = 13), p < 0.01; C57 × Sv129, 5.0 ± 0.5 ms (n = 20), p < 0.001; PrP<sup>0/0</sup>, 8.5 ± 0.8 ms (n = 26)]. At +60 mV, EPSC decay was also slower in PrP<sup>0/0</sup> compared with wild-type mice, causing a significant increase in τ<sub>w</sub> [C57BL/6J, 79.0 ± 6.3 ms (n = 13), p < 0.001; C57 × Sv129, 70.9 ± 6.3 ms (n = 20), p < 0.001; PrP<sup>0/0</sup>, 133.1 ± 12.8 ms (n = 26)].

On closer inspection, PrP<sup>0/0</sup> τ<sub>w</sub> exceeded the value of wild-type EPSCs in 42% of cases at −70 mV and in 38.5% of cases at +60 mV (Fig. 4D). Thus, the proportion of granule cells with slow τ<sub>w</sub> was similar to that of slow-spiking granule cells identified in current-clamp recordings, suggesting that slow EPSCs explained slow EPSP decay kinetics in PrP<sup>0/0</sup> granule cells.

**Impaired long-term potentiation at the PrP<sup>0/0</sup> mossy fiber–granule cell relay**

At the mossy fiber–granule cell relay of rat cerebellum, high-frequency stimulation is known to induce a composite form of LTP, in which both synaptic transmission and intrinsic excitability are enhanced (D’Angelo et al., 1999; Armano et al., 2000; Sola...
et al., 2004). In certain conditions in which granule cell activation is weakened, long-term depression (LTD) has been reported (Gall et al., 2005; Mapelli and D’Angelo, 2007).

In current-clamp recordings, LTP was induced by delivering TBS from −50 mV to improve the depolarizing action of EPSPs (Fig. 5, Table 3). In fast-spiking granule cells (all wild-type and one PrP0/0), robust action potential discharge was generated during low-frequency test stimulation increased, precluding EPSP amplitude from being measured (Fig. 5C). In all of the four slow-spiking PrP0/0 granule cells, although TBS had a strong depolarizing action (even stronger than in controls) (Table 3), just a few spikes could be elicited in response to TBS (Fig. 5A), in line with the alterations in intrinsic firing reported above (compare Fig. 3). After TBS, the probability of action potential generation did not increase and EPSPs remained measurable in most responses (Fig. 5A, right) without showing any net increase (−4.7 ± 7.6%; n = 4).

Intrinsic excitability was investigated in the same recordings reported in Figure 6. In all fast-spiking granule cells (including the PrP0/0 one), TBS enhanced action potential generation and decreased the current needed to reach threshold (Ith) (Fig. 6A, Table 3). Conversely, in the four slow-spiking PrP0/0 cells, no changes were observed. The apparent granule cell input resistance (Fig. 6B, C, Table 3) was measured from membrane potential changes caused by small current steps in the 10 mV potential range either below −70 mV (Rm-low) or above −70 mV (Rm-high). After TBS, Rm-high rapidly increased in all fast-spiking (including the PrP0/0 one) but not in PrP0/0 slow-spiking granule cells. It should be noted that Rm-low remained unchanged in all mice strains, providing an internal control for recording stability (Fig. 6B, C, Table 3).

In a different group of recordings (Fig. 7), granule cells were voltage clamped with a Cs+-based solution, and LTP was induced by TBS while depolarizing the granule cell at −30 mV. At this potential, Mg2+ block is mostly removed from NMDA channels (D’Angelo et al., 1993). As for experiments shown in Figure 4, spikes cannot be monitored and PrP0/0 data were pooled. After TBS, the AMPA EPSC in wild-type mice increased, following a similar time course in C57 × Sv129 and C57BL/6J strains, and remained potentiated throughout the recordings (at least 20 min after TBS; average time courses are shown in Fig. 7A, B). At 20 min after TBS, the EPSC increase was 27.5 ± 5.9% (n = 7) in C57 × Sv129 and 32.8 ± 10.4% (n = 7) in C57BL/6J mice. However, by using the same induction protocol, no LTP was observed in the PrP0/0 mice group (1.9 ± 5.6; n = 9; p < 0.01 vs either wild-type strains, unpaired t test). At a closer inspection, a single PrP0/0 recording showed clear LTP and had EPSCs as fast as in wild type. All the other PrP0/0 recordings showed either no LTP or even LTD and had an EPSC decay time constant, τm, remarkably larger than normal (compare Fig. 4D). As a whole, LTP was present in 85.7% (six of seven) granule cells in wild-type mice but just in 22% (two of nine) granule cells in PrP0/0 mice (Fig. 7C). The low probability of having LTP in PrP0/0 granule cells is consistent with the absence of LTP in PrP0/0 slow-spiking granule cells.

Juvenile PrP0/0 mice show impaired motor control

Given the marked abnormalities in mossy fiber–granule cell transmission and plasticity and considering the essential role played by the granular layer for cerebellar functioning, we asked whether the PrP0/0 mice had impaired sensorimotor control. Motor learning and performance of PrP0/0 mice were assessed at P19–P25 and compared with age-matched C57 × Sv129 control mice (n = 11 in both cases) (Duham and Miya, 1957; Goldowitz et al., 1992; Kashiwabuchi et al., 1995). In the accelerating rotarod test (Fig. 8A), PrP0/0 were already worse than C57 × Sv129 on the first day of training, and then both PrP0/0 and C57 × Sv129 mice improved performance with training. However, learning was incomplete in PrP0/0 compared with C57 × Sv129 mice. The greatest difference was observed during the first 4 d of training (p < 0.001 to p < 0.01). At days 5–7, when learning tended to plateau, PrP0/0 remained stably below the performance of C57 × Sv129 (p < 0.05). In the runway test (Fig. 8B), PrP0/0 were worse than C57 × Sv129 mice on days 1 and 2 (p < 0.05), but there was no statistical difference on days 3–5 (saturation probably occurred...
because the difficulty of test was low). The worse performances of PrP^0/0 mice was not attributable to a different body development, because PrP^0/0 and C57 × Sv129 mice had similar body weight (P19: 8.1 ± 1.2 g for PrP^0/0 mice; 7.6 ± 1.2 g for C57 × Sv129 mice) (P23: 8.4 ± 1.2 g for PrP^0/0 mice; 8.05 ± 0.8 g for C57 × Sv129 mice) (P25: 9.4 ± 1.5 g for PrP^0/0 mice; 9.4 ± 1.1 g for C57 × Sv129 mice). Preliminary experiments with C57BL/6J mice revealed aspecific differences, which could be attributed to the attitude to attentive, emotional, and fear responses in this strain (Bearzatto et al., 2005) and to developmental differences, because C57BL/6J are still relatively smaller than C57 × Sv129 and PrP^0/0 mice at P19.

Altersations at the mossy fiber–granule cell relay disappear in adulthood

Because it was reported that motor performance in adult PrP^0/0 mice was normal (Büeler et al., 1992; Herms et al., 1995; Lipp et al., 1998), an obvious question was whether the alterations at the mossy fiber–granule cell relay of juvenile PrP^0/0 mice disappeared with adulthood. We therefore repeated the critical measurements on PrP^0/0 mice at P40–P50 (Fig. 9), when cerebellar morphological development is concluded (Altman, 1972; Hámori and Somogyi, 1983) [the intrinsic excitatory properties of adult granule cells were recently reported by Goldfarb et al. (2007)]. Interestingly, all adult PrP^0/0 granule cells (seven of seven) showed fast repetitive firing (Fig. 9A) and inward rectification (Fig. 9B). Moreover, all of the four cases in which TBS was applied demonstrated LTP, which was manifest both as an increase in the probability of activating EPSP–spike complexes (Fig. 9C,D) and as an increase in intrinsic excitability and cell input resistance (Fig. 9A,B). Thus, adult PrP^0/0 granule cells did not significantly differ in their major functional properties from fast-spiking neurons in the juvenile stage. It should also be noted that no significant difference was observed between adult PrP^0/0 mice and age-matched C57BL/6J mice used for control. Minor differences between adult and juvenile granule cells (both in PrP^0/0 and C57BL/6J mice) are summarized in the supplemental material (available at www.jneurosci.org).

Protracted mitosis in the germinal zone of EGL during the third postnatal week

The fact that granule cells recovered a normal phenotype at P40–P50 suggested that the deficit in PrP^0/0 mice was transient and related to the developmental period. Because PrP was reported to affect the early stages of neuronal development (Manson et al., 1992; Graner et al., 2000; Martins et al., 2002; Steele et al., 2006), we investigated granule cell proliferation in the germinal zone of the EGL by using BrdU immunostaining in sagittal sections of the cerebellar vermis taken at different ages (Fig. 10). As expected, the

| Table 3. Effect of TBS on granule cell response properties |

<table>
<thead>
<tr>
<th>PrP^0/0 slow (n = 4)</th>
<th>C57 × Sv129 (n = 8)</th>
<th>C57BL/6J (n = 5)</th>
</tr>
</thead>
<tbody>
<tr>
<td>ΔAPprob (%)</td>
<td>51.9 ± 2.3*</td>
<td>55.7 ± 1.7**</td>
</tr>
<tr>
<td>ΔAPlow (%)</td>
<td>-33.1 ± 21.2*</td>
<td>-24.1 ± 21.0*</td>
</tr>
<tr>
<td>ΔRth (%)</td>
<td>42.2 ± 15.8**</td>
<td>55.5 ± 11.8**</td>
</tr>
<tr>
<td>ΔRlow (%)</td>
<td>6.2 ± 7.9</td>
<td>0.39 ± 5.2</td>
</tr>
<tr>
<td>TBSdep (mV)</td>
<td>-44.3 ± 1.9*</td>
<td>-44.6 ± 1.8*</td>
</tr>
</tbody>
</table>

The table compares percentage changes in wild-type and PrP^0/0 mice caused by TBS in current-clamp recordings: variation in probability of generating action potentials during synaptic stimulation (ΔAPprob), variation in current needed to reach spike threshold from the holding potential (ΔAPlow), variation in apparent input resistance in the region above or below −70 mV (ΔRth), and average depolarization during TBS (TBSdep). The number of observations is reported in parentheses, and statistical significance is reported in comparisons with slow-spiking PrP^0/0 granule cells.

*p < 0.05, **p < 0.01, unpaired t test.
but not in pooled PrP0/0 granule cells. Data are mean PrP 0/0 granule cell. After TBS, EPSCs increased in the wild-type but not in the PrP0/0 granule cell. Control and 20 min after TBS application are shown superimposed in C57 of average EPSC amplitude changes in C57
The top plot correlates EPSC amplitude changes after TBS for the different mice strains with the EPSC decay time constant, 0.05, two-way ANOVA).

In PrP0/0 mice, a large proportion of granule cells (Altman, 1972). Conversely, duplication of calcium-dependent K-channels (Colling et al., 1996; Fuhrmann et al., 2006) (see also Korte et al., 2003), could reduce spike afterhyperpolarization by weakening activation of NMDA currents (D’Angelo et al., 1993; Farrant et al., 1994; Rumbaugh and Vaccini, 1999; Cathala et al., 2000; Rossi et al., 2002; Wall et al., 2002). In normal mice, the transition from immature to mature excitable patterns occurs soon after granule cell migrate from the external to internal granular layer and proceeds throughout postnatal weeks 1 and 2: in week 3, electrophysiological parameters are stabilized and immature slow-spiking granule cells disappear [for mice, see Rossi et al. (1998); for the rat, see D’Angelo et al. (1993, 1994, 1997)]. The observation of protracted mitosis in the germinal zone of the EGL suggests that, in PrP0/0 mice, immature granule cells are abnormally supplied to the granular layer all along the third postnatal week. These neurons will eventually mature explaining why only a part of granule cells at P20 are slow-spiking and why at P40–P50 slow-spiking PrP0/0 granule cells disappear (cf. Goldfarb et al., 2007). In addition to the arrival of new immature granule cells, other PrP-dependent mechanisms may be altered in granule cells once migration is concluded. A reduced L-type Ca2+-channel expression, which was observed in PrP0/0 granule cells in culture (Hermes et al., 2000; Fuhrmann et al., 2006) (see also Korte et al., 2003), could reduce spike afterhyperpolarization by weakening activation of calcium-dependent K-channels (Colling et al., 1996; Hermes et al., 2001) and contribute to alter repetitive spike discharge. LTP was altered only in granule cells with immature phenotype and could therefore also be dependent on the altered developmental program of the granule cells. However, the alteration of LTP could be more complex and involve additional mechanisms. In fact, membrane depolarization during TBS

wild-type granule cells actively duplicated until the end of the second postnatal week (Altman, 1972). Conversely, duplication in the PrP0/0 mice continued also during the third postnatal week, suggesting that PrP was involved in controlling granule cell proliferation.

Discussion
The central observation in this study is that PrP knock-out causes abnormal functioning of the cerebellar mossy fiber–granule cell relay and impairs motor control in juvenile mice. In PrP0/0 mice, a large proportion of granule cells (~40%) showed slow nonovershooting nonrepetitive action poten-

tials, slow EPSPs, and no inward rectification. Moreover, the slow-spiking PrP0/0 granule cells were unable to generate LTP with the TBS protocol. The disturbances were evident during the third postnatal week but disappeared thereafter, supporting a major role of PrP in granular layer formation and motor control (Salès et al., 2002; Nicolas et al., 2007).

The slow-spiking PrP0/0 granule cells bear a striking resemblance to an immature phenotype by showing nonovershooting nonrepetitive spikes, no inward rectification, and relatively high resting potential (D’Angelo et al., 1997). Like wild-type immature granule cells, slow-spiking PrP0/0 granule cells have small Na+, Ca2+, and K+ currents (D’Angelo et al., 1994; Rossi et al., 1998) and lack inward rectification (Rossi et al., 1998). Moreover, immature granule cells show slow EPSPs in association with slow NMDA currents (D’Angelo et al., 1993; Farrant et al., 1994; Rumbaugh and Vaccini, 1999; Cathala et al., 2000; Rossi et al., 2002; Wall et al., 2002). In normal mice, the transition from immature to mature excitatory patterns occurs soon after granule cell migrate from the external to internal granular layer and proceeds throughout postnatal weeks 1 and 2: in week 3, electrophysiological parameters are stabilized and immature slow-spiking granule cells disappear [for mice, see Rossi et al. (1998); for the rat, see D’Angelo et al. (1993, 1994, 1997)]. The observation of protracted mitosis in the germinal zone of the EGL suggests that, in PrP0/0 mice, immature granule cells are abnormally supplied to the granular layer all along the third postnatal week. These neurons will eventually mature explaining why only a part of granule cells at P20 are slow-spiking and why at P40–P50 slow-spiking PrP0/0 granule cells disappear (cf. Goldfarb et al., 2007). In addition to the arrival of new immature granule cells, other PrP-dependent mechanisms may be altered in granule cells once migration is concluded. A reduced L-type Ca2+-channel expression, which was observed in PrP0/0 granule cells in culture (Hermes et al., 2000; Fuhrmann et al., 2006) (see also Korte et al., 2003), could reduce spike afterhyperpolarization by weakening activation of calcium-dependent K-channels (Colling et al., 1996; Hermes et al., 2001) and contribute to alter repetitive spike discharge. LTP was altered only in granule cells with immature phenotype and could therefore also be dependent on the altered developmental program of the granule cells. However, the alteration of LTP could be more complex and involve additional mechanisms. In fact, membrane depolarization during TBS
was larger in slow-spiking PrP<sup>0/0</sup> than in fast-spiking granule cells, and the slow time course of NMDA EPSCs suggests that Ca<sup>2+</sup> influx, which is needed for LTP induction (D’Angelo et al., 1999; Armano et al., 2000; Rossi et al., 2002; Gall et al., 2005), might be even larger in PrP<sup>0/0</sup> than in wild type. Therefore, the mechanisms of dysfunction may involve alterations in intracellular Ca<sup>2+</sup> dynamics, which have been reported in PrP<sup>0/0</sup> cultured cerebellar granule cells (Hermes et al., 2000). A defect in nitric oxide signaling (Keshet et al., 1999), which is required for mossy fiber–granule cell LTP (Maffei et al., 2003), could explain the absence of presynaptic changes needed to generate EPSC potentiation (Sola et al., 2004). Therefore, PrP knock-out may directly affect the LTP mechanism, as originally proposed for hippocampal synapses (Collinge et al., 1994; Curtis et al., 2003; Maglio et al., 2004, 2006) (but see Lledo et al., 1996).

These results suggest a hypothesis on how PrP could interact with granular layer functions. The absence of a timely arrest of proliferation in the EGL germinal zone may reflect the absence of signals normally provided by PrP to the biochemical cascades regulating granule cell apoptosis (Nicolas et al., 2007). Thus, the mechanism of PrP action, at least in part, could reside in a protracted proliferation in the germinal zone of the EGL supplying the granular layer with immature granule cells. Moreover, because PrP expression proceeds in parallel with the development of granule cell excitatory functions (cf. D’Angelo et al., 1994, 1997) and Rossi et al. (1998) to Salès et al. (2002) and is abundantly expressed in mature granule cells and presynaptic mossy fiber terminals, the effects of PrP may continue in the granular layer. The coexistence of mature with immature granule cells at early stages and the full recovery of mature excitability and LTP at later stages indicate that, despite the altered developmental program in PrP knock-out mice, mature functions are finally recovered possibly through the engagement of compensatory mechanisms. It seems less likely that slow-spiking granule cells reflect ongoing degeneration, because no gross anatomical abnormalities are apparent in the adulthood (Lainé et al., 1998), as it would be expected with an extended granule cells loss [e.g., in the weaver mutation (Rossi et al., 1998)].

The multiple granular layer alterations in the PrP<sup>0/0</sup> mice could each contribute to impair cerebellar functioning in their own way. Slow EPSPs and action potentials are expected to reduce the velocity and temporal precision of mossy fiber–granule cell synaptic transmission (D’Angelo et al., 1995, 1997; Cathala et al., 2003), the absence of repetitive firing to prevent coding of afferent information into spike bursts (Chadderton et al., 2004; Jörntell and Ekerot, 2006; Nieus et al., 2006; Rancz et al., 2007), and the absence of LTP to limit granular layer changes by specific input patterns (Mapelli and D’Angelo, 2007). This could bring about a diffused impairment of synaptic transmission and plasticity throughout the cerebellar network (Casado et al., 2002; Brunel et al., 2004). Although PrP<sup>0/0</sup> functional alterations occur also in other parts of the brain [e.g., the hippocampus (Colling et al., 1996) (but see Lledo et al., 1996; Johnston et al., 1998)], the marked granular layer abnormalities are likely to play a relevant role in altering sensorimotor control in the runway and rotarod tests.

In conclusion, these results suggest an active role of PrP in the functional maturation of the mossy fiber–granule cell relay of cerebellum implying consequences for motor control and reinforcing the evidence for an active role of PrP in brain development (Manson et al., 1992; Graner et al., 2000; Martins et al., 2002; Steele et al., 2006). The reason why, in adult PrP<sup>0/0</sup> mice, behavioral tests appeared normal (Büeler et al., 1992; Herms et al., 1995; Lipp et al., 1998), is probably because slow-spiking granule cells eventually recover a normal fast-spiking phenotype and synaptic plasticity. The potential impact of the temporary disruption of the cerebellar circuit during a time window critical for sensorimotor development (Altman and Sudarshan, 1975) remains to be determined, but it may have a reflection in some of the late motor, cognitive, and emotional abnormalities of PrP<sup>0/0</sup> mice (Roesler et al., 1999; Walz et al., 2002; Criado et al., 2005). In this context, it should be noted that the late-onset cerebellar ataxia observed in different PrP<sup>0/0</sup> strains is a distinct phenomenon, which
arises much later (after 60 postnatal weeks) and depends on mutation in the homologous Dpl gene (Sakaguchi et al., 1996; Katamine et al., 1998; Moore et al., 2001; Behrens and Aguzzi, 2002).

References


