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Sox2 conditional mutation in mouse causes ataxic symptoms, cerebellar vermis hypoplasia, and postnatal defects of Bergmann glia

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Supplementary Table 3

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Sox2 conditional mutation in mouse causes ataxic symptoms, cerebellar vermis

hypoplasia, and postnatal defects of Bergmann glia

Running title: Sox2 requirement in Bergmann glia

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Dedication. We dedicate this paper to the dear memory of our colleague and friend Ferdinando Rossi, whose insight and enthusiasm live on by our side.

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Abstract

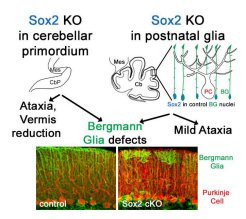
Sox2 is a transcription factor active in the nervous system, within different cell types, ranging from radial glia neural stem cells to a few specific types of differentiated glia and neurons. Mutations in the human SOX2 transcription factor gene cause various central nervous system (CNS) abnormalities, involving hippocampus and eye defects, as well as ataxia. Conditional Sox2 mutation in mouse, with different Cre transgenes, previously recapitulated different essential features of the disease, such as hippocampus and eye defects. In the cerebellum, Sox2 is active from early embryogenesis in the neural progenitors of the cerebellar primordium; Sox2 expression is maintained, postnatally, within Bergmann glia (BG), a differentiated cell type essential for Purkinje neurons functionality and correct motor control. By performing Sox2 Cre-mediated ablation in the developing and postnatal mouse cerebellum, we reproduced ataxia features. Embryonic Sox2 deletion (with Wnt1Cre) leads to reduction of the cerebellar vermis, known to be commonly related to ataxia, preceded by deregulation of Otx2 and Gbx2, critical regulators of vermis development. Postnatally, BG is progressively disorganized, mislocalized, and reduced in mutants. Sox2 postnatal deletion, specifically induced in glia (with GLAST-CreERT2), reproduces the BG defect, and causes (milder) ataxic features. Our results define a role for Sox2 in cerebellar function and development, and identify a functional requirement for Sox2 within postnatal BG, of potential relevance for ataxia in mouse mutants, and in human patients.

Key words:

Sox2, cerebellum, ataxia, Bergmann glia, mouse genetic models, transcription factors

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Main points

- -Deletion of Sox2 in the cerebellar primordium leads to ataxic symptoms, with vermis hypoplasia and Bergmann glia dysmorphology and displacement
- -Postnatal Sox2 deletion in glia causes mild ataxia with Bergmann glia defects

Introduction

The Sox2 transcription factor is widely expressed in the developing central nervous system (CNS), from early embryogenesis (Bertolini, 2016; Pevny & Nicolis, 2010). In humans, Sox2 heterozygous mutations lead to a spectrum of CNS defects, including eye defects, hippocampal reduction, seizures, as well as motor control defects and ataxia (Bertolini, 2016; Dennert et al., 2017; Ragge et al., 2005; Sisodiya et al., 2006). In mouse, complete Sox2 loss is early embryonic lethal (Avilion et al., 2003); by conditional knock-outs and hypomorphic mutants, we and others demonstrated that Sox2 loss leads to important CNS defects, affecting the eye, the developing ventral forebrain, and the hippocampus. Most of these defects arise as a consequence of the loss of neural stem cells (represented by radial glia), although in some cases specific glial or neuronal types have been involved (Cavallaro et al., 2008; Favaro et al., 2009; A. Ferri et al., 2013; A. L. Ferri et al., 2004; Pevny & Nicolis, 2010).

Ataxia is a motor control defect, depending on cerebellar abnormalities, and is often the consequence of inherited genetic defects observed in man (Basson & Wingate, 2013; Leto et al., 2016). While the genetic defects primarily operates in neurons in most types of ataxias, at least in two cases (SCA1, SCA7) Bergmann glia (BG), a specialized cerebellar glia essential for Purkinje neurons functionality, critically contributes to the defect; the BG dysfunction affects, in a non-cell-autonomous way, the functions of the adjacent Purkinje neuronal cells (PC), which are responsible for the functional defects (Buffo & Rossi, 2013; Custer et al., 2006; Shiwaku et al., 2010). While the primary genetic defect is well defined, the network of gene regulation and protein interactions downstream to it are not yet well understood.

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Previous studies of hypomorphic Sox2 mutants demonstrated motor abnormalities (Ferri 2004); in this work, we investigated the possibility that Sox2 conditional ablation in mouse reproduces some of the ataxic features demonstrated in human patients. To this end, we ablated Sox2 either prenatally, or postnatally, within the cerebellum, a CNS region essential in the control of movement, whose disfunction leads to ataxias (Basson & Wingate, 2013; Leto et al., 2016). We report that prenatal Sox2 loss in the developing midbrain-hindbrain region (via Wnt-1-Cre-mediated deletion) leads to a reduction of the cerebellar vermis at birth, and, postnatally, to progressive disorganization, mislocalization and reduction of BG, functionally accompanied by ataxia. BG retains Sox2 expression postnatally, and we find that postnatal deletion of Sox2 in glia (via GLAST-CreERT2) also leads to BG disorganization, mislocalization and reduction. Our findings point to a novel role for Sox2 in the postnatal establishment of the correct phenotype, layering and functions of this important cerebellar cell type.

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Materials and methods

Experimental animals

Experiments were performed on different mouse lines: Sox2^{flox} (Favaro et al., 2009), Wnt1Cre (Danielian, Muccino, Rowitch, Michael, & McMahon, 1998), GLASTCreErt2 (always used as heterozygous) (Buffo et al., 2008; Mori et al., 2006). Either sex was included. The day of vaginal plug detection was defined as embryonic day zero (E0) and the day of birth was considered as postnatal day zero (P0). Mice were maintained on a 12 hrs day / night cycle with adequate food and water. All procedures were in accordance with the European Communities Council Directive (2010/63/EU and 86/609/EEC), the National Institutes of Health guidelines, and the Italian Law for Care and Use of Experimental Animals (DL26/14).

They were approved by the Italian Ministry of Health and the Bioethical Committees of the Universities of Milan and Turin.

Histological, in situ hybridization and immunohistochemical procedures

In situ hybridization was performed as in (Favaro et al., 2009; A. Ferri et al., 2013). Briefly, embryos were fixed in 4% paraformaldehyde (PFA) in 0.12 M phosphate buffer (PB), pH7.2-7.4, cryoprotected, embedded in OCT and sectioned on a cryostat at 20 µM. Antisense probes for Sox2 (Avilion et al., 2003), Otx2 (Acampora, Avantaggiato, Tuorto, & Simeone, 1997) and Gbx2 (Wassarman et al., 1997), a kind gift (Otx2; Gbx2) from Drs. L. Muzio and V. Broccoli, were in vitro transcribed in the presence of DIG-UTP and hybridized to sections at 65°C. The signal was visualized by incubation with BM-purple (Sigma). For postnatal brain analysis, animals were transcardially perfused under anesthesia (hypothermia for pups up to postnatal day -P-7 and intraperitoneal injection of a mixture of ketamine -100 mg/kg; Ketavet, Bayern, Leverkusen, Germany- and xylazine -5 mg/kg; Rompun, Bayer, Milan, Italy- for older mice) with an appropriate volume of 4% PFA in 0.12 M phosphate buffer (PB), pH7.2-7.4. Brains were removed, stored overnight (o/n) in the same fixative at 4 °C, washed in Phosphate Buffered Saline (PBS) and finally cryoprotected in 30% sucrose in 0.12M PB. The cerebella were then embedded and frozen over dry ice in OCT (TissueTEK), sectioned in the parasagittal plane at 30 µm using a cryostat and collected in PBS (P30 juvenile and adult cerebella) or placed directly onto glass slides (E13-P7 cerebella). For immunolabelling, sections were incubated o/n at room temperature with the appropriate primary antibodies dissolved in PBS with 1.5% donkey or goat serum (Jackson ImmunoResearch) and 0.25% Triton X-100 (Sigma-Aldrich): anti-S100BETA (S100B, 1:1000, mouse, monoclonal; Sigma), anti-glial fibrillary acidic protein (GFAP, 1:1500, rabbit, polyclonal; Dako), anti-brain lipid Page 9 of 64 GLIA

binding protein (BLBP, 1:500; rabbit, polyclonal; Abcam); anti-SOX2 (1:500, rabbit, polyclonal; Merck Millipore; 1:50, mouse, monoclonal, R&D Systems); anti CALBINDIN(CALB, 1:1500, rabbit, monoclonal; Swant); anti-PARVALBUMIN (1:1500, mouse monoclonal; Swant); anti-myelin basic protein (MBP, 1:1000, mouse, monoclonal; BioLegend); anti-NG2 (1:250, rabbit, polyclonal; Merck Millipore); anti-Glutathione S-Transferase, Pi isoform (GSTPi, 1:500, rabbit, polyclonal; Eppendorf); anti-OLIG2 (1:500, rabbit, polyclonal; Merck Millipore). Sections were then exposed for 2h at room temperature to secondary species-specific antibodies conjugated with Alexa Fluor 488, Alexa Fluor 555 or Alexa Fluor 647 (1:500; Invitrogen), or Cy3 (1:500; Jackson ImmunoResearch). Cell nuclei were visualized using 4',6-diamidino-2-phenylindole (DAPI; Fluka). After processing, sections were mounted on microscope slides with Tris-glycerol supplemented with 10% Mowiol (Calbiochem). Statystical detail on Immunofluorescence cell counting is in Supplementary Table 2 and Figure legends.

Image processing and morphometric analyses

Histological specimens were examined using an E-800 Nikon microscope connected to a color CCD camera or a Leica TCS SP5 confocal microscope. Adobe Photoshop 6.0 (Adobe Systems) was used to adjust image contrast and assemble the final images. On the basis of confocal images, ectopia, densities, lengths and distances were mapped and quantified by Neurolucida 7.0 (MBF Bioscience) or by NIH ImageJ (http://rsb.info.nih.gov/ij/) softwares. Numbers of analysed animals for each time point or experimental condition are shown in the figure legends. Measurements were derived from at least three sections per animal. For evaluation of oligodendrocyte precursors and mature oligodendrocytes densities, cell counts

were performed in two areas of lobules IV-V and IX, on confocal stacks of about 25 μ m. Three sections per animal were examined.

The thickness (t) of the granular cell layer (GL) was determined on DAPI labelled slices according to the formula t = 2 A/b1 + b2, as in Carulli et al (2002). "A" was given by the area defined by the outer and inner profile of the granular layer, and "b1" and "b2" were the lengths of the two profiles. At least three slices/animal and three animals/genotype were analyzed. Values were pooled to yield the final average thickness.

For morphometric measurements in adult mice, DAPI-stained sections next to the midline of the entire cerebellar vermis were analysed. The whole PCL length, the extension along the Rostro-Caudal (R-C) and Ventro-Dorsal (V-D) axes, the PCL length along each lobule and the extension of the fissures between lobules VIa-VIb and IXb-IXc were evaluated using a fluorescent microscope and traced by means of the Neurolucida software (MicroBrightfield). For the of R-C extension measurement, the points of maximal convexity of the crowns of lobules V and IXb were chosen as points of reference, whereas the length of the V-D axis was evaluated by measuring the distance from the invagination between lobules I and X (maximal concavity) and the midpoint in the crown of lobule VIb. The length of the PCL along each lobule was measured as the line connecting the midpoint of the basis of the two fissures preceding and following the folium of interest. The extension of the fissures was evaluated by measuring the length of PCL between the related inward indentation points. For evaluation of BG ectopia, BG densities and BG to PC ratios, cell counts were performed in three cortical areas of lobules III, IV-V and IX, on confocal Z stacks of about 15 µm. Three sections per animal were evaluated.

Systemic administration of Tamoxifen

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To activate the Cre recombinase in GLASTCreERT2 Sox2 mice Tamoxifen (Tx) (20 mg/ml solution dissolved in corn oil) was administered via subcutaneous injection for pups (20 µg/g body weight) or via oral gavage for adults (5 mg/40 g body weight). Mice were given either a single Tx injection at P2 or administered a double treatment including a first single injection at P8 and a further treatment at P60. A distinct cohort of adult mice was treated at 4 months of age to assess the effects of Sox2 deletion after the completion of cerebellar maturation. However, at 2 months after treatment no changes in BG layering were observed, suggesting that Sox2 is dispensable to preserve BG morphology and layering at adult stages, when cerebellum maturation is fully accomplished. As for experiments performed in Wnt1-Cre Sox2 mice, animals with mixed genotypes were used as controls (Sox2F/+;CreERT2, Sox2FF or Sox2F/+) for some analyses. In other instances, Sox2F/F;CreERT2 mice with no Tx treatment were also employed as controls, as indicated in Supplementary Table 1.

Behavioural Assays

All behavioral experiments were conducted in a dedicated dimly illuminated (2 x 40 W indirect) testing room during the standard light phase, usually between 9:00 AM and 2:00 PM (rotarod from 10:00 AM to 12:00 AM, beam test from 10:00 AM to 1:00 PM, open field from 10:00 AM to 2:00 PM). In all tests, the cages containing mice were moved into the testing room 1 hour before the first trial began. For Wnt1-Cre Sox2 mutants of either sexes were examined and pooled together, as no difference was appreciated between genders. For GLASTCreERT2 Sox2 mutants, in cases of relevant differences data for males and females are presented separately.

Rotarod test. Motor coordination, balance and motor learning were assessed using an accelerating rotarod (Ugo Basile, Varese, Italy; (Fuca et al., 2017)). The rotarod apparatus had five flanges dividing the five 5.7 cm lanes, the cylinder diameter was 3.5 cm. Mice were placed on the rotating bar and given a habituation period of 2 min at constant velocity (5.5 rpm), after which the rotating bar accelerated from 5.5 to 60 rpm over 300 seconds. Mice were put on the rotarod until the latency to fall off reached the total time of 300 s. Three trials per day with 3-5 min intervals, on three consecutive days were performed; we considered the averages of the three sessions. During pauses between trials, mice rested in their home cages

Beam walking test. In order to assess balance capabilities, beam walking test has been used (adapted from (Hilber & Caston, 2001)). Testing was performed once for month for mice of each group. The apparatus consisted of a motionless wooden beam with a rectangular section (1 cm x 3 cm), 1 m in length, placed 80 cm above a foam carpet. At the onset of the single trial, each animal was placed at one extremity of the beam, its body axis being perpendicular to the beam long axis. We recorded the time spent to complete each cross and the number of paw faults or slips made during the passages to the wooden bar, for a final number of 5 crossings of the bar. Then, for each animal, the frequency of slip and the time spent to cross the beamwere calculated. After 5 consecutive, unsuccessful trials to stay on the bar, the test was considered failed.

Open field. Animals were tested in a forex arena ($50 \times 50 \text{ cm}$), which was divided into a central ($25 \times 25 \text{ cm}$) and a peripheral zone (Cupolillo et al., 2016). The animal was placed in the corner of the open field and was allowed to explore the arena for 10 min. Distance

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travelled and time spent in the central zone was recorded and analyzed by Ethovision XT video track system (Noldus Information Technology, Wageningen, Netherlands).

Electrophysiology

The animals were anesthetized with isoflurane (Isoflurane-Vet, Merial, Italy) and decapitated. The cerebellar vermis was removed and transferred to an ice-cold artificial cerebrospinal fluid (ACSF) containing (in mM); 125 NaCl, 2.5 KCl, 2 CaCl2, 1 MgCl2, 1.25 NaH2PO4, 26 NaHCO3, 20 glucose, which was bubbled with 95% O2/5% CO2 (pH 7.4). Parasagittal cerebellar slices (200 µm thickness) were obtained using a vibratome (Leica Microsystems GmbH, Wetzlar, Germany) and kept for 1 h at 35°C and then at 25°C. Single slices were placed in the recording chamber, which was perfused at a rate of 2–3 ml/min with ACSF bubbled with the 95% O2/5% CO2. All recordings were performed at room temperature (22–25°C). The data derive from 5 to 7 animals.

Whole-cell patch-clamp recordings were made from PCs of adult animals of both genotypes using an EPC-9 patch-clamp amplifier (HEKA Elektronik, Lambrecht/Pfalz, Germany). We performed recordings in the posterior lobules of cerebellum. Pipettes of borosilicate glass with resistances between 2.5 and 3.0 M Ω were used. Patch pipettes were filled with an internal solution containing (in mM): 130 CsCl, 4 MgCl₂, 10 HEPES, 4 Na₂ATP, 0.4 Na₃GTP, 10 EGTA, 5 N-(2,6-dimethylphenyl)acetamide-2-triethylammonium bromide (QX-314) and the pH was adjusted to 7.3 with CsOH and filtered at 0.2 μ m. Cs⁺ blocks most outward currents through K⁺ channels while QX-314 blocks voltage-gated Na⁺ channels. Recordings were accepted only if the series resistance was less than 9.0 M Ω (range: 5.0–9.0 M Ω), and if it did not vary by >20% during the experiment. Data were filtered at 3 kHz and sampled at 10 kHz.

For all the experiments, digitized data were stored on a personal computer using the Patch Master software (HEKA Elektronik, Lambrecht/Pfalz, Germany) and analized off-line. PF-EPSCs were evoked by stimulating the PFs in the molecular layer while the PC was kept at a holding potential of -90 mV. The stimulation was done with a glass pipette, pulled from sodalime glass to a tip diameter of 10–15 μ m, containing the saline solution. Negative current pulses ranging from 3 to 15 μ A with a duration of 100 Ξ μ s were delivered at 20s intervals. Paired pulse facilitation was elicited by applying double pulses of 9 μ A of intensity at different time intervals (50, 100, 150, 200 ms), and the ratio of the amplitude of the second PF-EPSC over the first one was calculated. All PF recordings were performed in the presence of gabazine (SR 95531, 20 μ M) in the saline solution.

Gliosome purification

Animals were euthanized and the whole brain rapidly removed. The cerebellar cortex was dissected on ice at 4°C; care was taken to dissect away the white matter, containing fibrous (parenchymal) astrocytes, as much as possible, to obtain an astrocyte preparation highly enriched in BG; glial perisynaptic processes (gliosomes) were then prepared from control and Sox2F/F;Wnt1Cre mutant mice essentially as previously described (Milanese et al., 2010; Raiteri et al., 2008), with minor changes. Briefly, the tissue was homogenized in 0.32 M sucrose, buffered at pH 7.4 with Tris-HCl, using a glass-teflon tissue grinder (clearance 0.25 mm - Potter-Elvehjem VWR International). The homogenate was centrifuged (5 min, 1,000 x g) to remove nuclei and debris, the supernatant centrifuged at 12,000 x g for 10 min and the pellet resuspended in Tris-buffered 0.32 M sucrose and gently layered on a discontinuous Percoll® (Sigma-Aldrich, St Louis, Missouri, USA) gradient (2, 6, 10 and 20% v/v in Tris-buffered 0.32 M sucrose). After centrifugation at 33,500 x g for 5 min, the layer between 2 and 6% Percoll® (gliosomal

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fraction) was collected and washed by centrifugation at 20,000xg for 15 min with physiological medium (PM) having the following composition (mM): NaCl, 140; KCl, 3; MgSO₄, 1.2; NaH₂PO₄, 1.2; NaHCO₃, 5; CaCl₂, 1.2; 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES), 10; glucose, 10; pH 7.4. Gliosomes were re-suspended in PM for uptake experiments or lysed in Ultrapure water (Milli-Q; Millipore, Billerica, MA, USA) for endogenous glutamate/glutamine determination. All the above procedures were conducted at 4°C. Protein content was measured according to (Bradford, 1976) using bovine serum albumin (Sigma-Aldrich, St Louis, Missouri, USA) as a standard.

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Uptake experiments

Uptake was measured in gliosomes purified from BG-enriched glia of control (Sox2FF/F+) and mutant (Sox2F/F;Cre) mice as previously described (Milanese et al., 2015). Briefly, gliosomes were re-suspended in PM and aliquots (500 μl, corresponding to 3-5μg protein) were incubated for 5 min at 37 °C and the 20μl of a solution of [³H]D-Aspartate ([³H]D-Asp)(specific activity: 16.5 Ci/mmol, Perkin Elmer Italia, Milan, Italy), a non-metabolizable analogue of glutamate (Glu), used to mimic the endogenous neurotransmitter as a substrate of Glu transporters (Raiteri et al., 2007) was used, to obtain the final concentrations of 0.3, 3, 5, 30, and 100μM. After 2 min incubation at 37°C, [³H]D-Asp uptake was blocked by rapid vacuum filtration on glass fiber filters (GF-B, Millipore, Billerica, MA, USA) and washed three times with 5 ml of standard medium to remove the excess of radioactivity. Aliquots of gliosomes were exposed to the same concentrations of [³H]D-Asp at 0–4 °C in the presence of 50μM of the broad spectrum Glu uptake inhibitor *DL*-threo-beta-benzyloxyaspartate (DL-TBOA; Tocris Bioscience, Bristol, UK; Shimamoto et al., 1998) to detect the non-specific binding to the gliosomal membranes. Radioactivity was determined by liquid scintillation counting. The

specific uptake was calculated by subtracting non-specific binding from the total uptake. To calculate the kinetic parameters Vmax and Km for the two experimental groups, data were fitted by the Michaelis-Menten equation V=Vmax*[S]/Km+[S], using the SigmaPlot 10.0 and SigmaStat fitting setup (Systat Software, Inc., San Jose, CA, USA).

Endogenous glutamate and glutamine determination

BG cells gliosomes from control (Sox2FF/F+) and mutant (Sox2F/F;Cre) mice were lysed in hypotonic ultrapure water, and the cytosolic extracts separated from membrane debris by centrifugation (10 min 20,000xg). The endogenous glutamate and glutamine content were quantified in supernatants by previously described high performance liquid chromatography analysis (Milanese et al., 2011). The methods consisted in a pre-column derivatization of the samples with o-phthalaldehyde, followed by discontinuous gradient separation using a C18 reverse-phase chromatographic column (Agilent MicroSpher C18, S100x4.6, 3 μ m; CPS Analitica, Milano, Italy) at 30 °C, and fluorometric detection (excitation wavelength 350 nm; emission wavelength 450 nm;). Homoserine (Sigma-Aldrich, St. Louis, MO, USA) was used as an internal standard.

Statistical analyses

Data elaboration and statistical analyses were conducted by means of GraphPad software Prism version 7.0 (GraphPad software, San Diego; SPSS,IBM) In all instances, *P<0.05 was considered as statistically significant, **P<0.01 very significant and ***P<0.001 extremely significant. Graphs and data are represented as means ± standard errors. If data sets passed normality test and equal variance test, we performed Student's t-test or One/Two Way Anova

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followed by Bonferroni post hoc tests. When data did not satisfy the normality or equal variance tests, corresponding non-parametric statistics was used. To compare different percentages, we transformed the values in radiants according to the arcsin transformation. Cumulative frequency plots were analyzed by Kolmogorov-Smirnov (KS) test. The graphs of electrophysiology results were designed using Igor Pro (WaveMetrics, Lake Oswego, Oregon, USA). Details about statistical assessment and numerical values obtained are provided in Supplementary Tables 1 and 2.

RNAseq from wild type and mutant neural stem cell-derived in vitro differentiated cells Neurosphere cultures were derived from wild type and Sox2-mutant mouse forebrain cells. Sox2 ablation was obtained at E11.5 by Nestin-Cre deletion; at P0, neural stem cells populations from three wild type and three Sox2 mutant forebrains, were individually placed in culture (in the presence of EGF and FGF) to generate neurospheres (as in Favaro et al. 2009). For differentiation, neurospheres from the individual cultures were dissociated to single cells, and plated onto MATRIGEL (Becton-Dickinson)-coated tissue culture dishes at 1,5 x 10⁶ cells/10cm dish (Gritti et al., 1996; Zappone et al., 2000), with bFGF only as mitogen. After 4 days, the medium was changed to neural stem cell medium without bFGF, and supplemented with 1% fetal calf serum (FCS). At day eleven, cells were collected for RNA analysis (at this stage, over 90% of the cells are glial cells in wild type cells, and about 95% in mutant cells). RNA extraction was performed on the three independent NSC populations for both wild type and mutant cells, using Trizol and RNeasy Kit (Qiagen). 1/5 volume of chloroform was added to one volume of Trizol. The aqueous phase was transferred into a new tube. 1.5x volume of ethanol was added and thoroughly mixed. The mixture was filtered through Rneasy (Qiagen) columns; the columns were washed with Buffer RW1. On-column DNase treatment was performed as described by the RNase-Free DNase Kit (Qiagen). Post treatment columns were cleaned up with Buffer RW1 and two washes of RPE buffer. Columns were then dried and total RNA was eluted with RNase-free water. PolyA Stranded Truseq Libraries were generated using the Truseq Stranded mRNA Sample Preparation Kit (Illumina). First, mRNA was purified from 1µg of total RNA using magnetic beads containing poly-T oligos. mRNA was then fragmented and reverse transcribed using Superscript II (Invitrogen), followed by second strand synthesis. Double stranded cDNA was treated with end-pair, A-tailing, adapter ligation and 8 cycles of PCR amplification.

RNA-Seq was performed on all individual samples (three wild type and three mutant) studied, to yield single-end reads. The number of sequences obtained in each sample ranged from 7.5 to 12.5 millions.

Read counts and transcript levels for each sample were computed with the RSEM software package version 1.17 (B. Li & Dewey, 2011), on the RefSeq gene annotation available at the UCSC Genome Browser for mouse genome assembly mm9 (24,148 genes). For downstream analyses, expression levels measured as Transcripts per Million (TPM) were employed. Differential expression analysis was performed with the NOISeqR package (Tarazona et al., 2015). Differentially expressed genes were selected by employing a q-value threshold of 0.99, corresponding to an empirical false discovery rate (FDR) of 0.01.

Results

Conditional Sox2 mutants develop ataxia

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To address the consequences of Sox2 loss in the developing cerebellum, we used mice carrying a Sox2flox conditional knockout (cKO) allele, previously generated in our laboratory (Favaro et al., 2009); to delete Sox2, we used Wnt1-Cre, active from embryonic day 9.5 (E 9.5) in the developing midbrain-hindbrain region, where the cerebellum will arise (Danielian et al., 1998). In situ hybridization at E9.5 in wild type embryos shows Sox2 mRNA expression in the neuroepithelium (consisting, at this stage, of undifferentiated progenitors of neurons and glia), throughout the developing neural tube, including the midbrain-hindbrain region where the cerebellum will arise; in the mutant, Sox2 mRNA was lost in the developing mesencephalon and anterior rhomboencephalon by E9.5 (Fig. 1a).

Wnt1-Cre Sox2 mutant mice (Sox2F/F;Cre hereafter) are born in normal mendelian ratios; most survive to adulthood, though a few are lost before weaning (when mutants represent 20% of the offspring, instead of the expected 25%).

We initially observed clear motor abnormalities in a number of adult mutant mice, suggesting possible cerebellar defects. To precisely address functional cerebellar abnormalities in Sox2 mutants, we subjected them to behavioural tests of motor coordination and equilibrium, used to detect alterations that arise following cerebellar damage: the rotarod and balance beam tests (Fuca et al., 2017; Hilber & Caston, 2001). We compared motor performances of mutants and control littermates between 1 and 8 months of life (Fig. 1b-d). By the rotarod test, latency to fall (whose shortening is indicative of motor control damage) was significantly shorter from 2 months of age (Fig. 1b). Heterozygous (Sox2F/+;Cre) mice also showed a moderately reduced latency, though the defect was less pronounced than that of homozygous mutants (Fig. 1b). The beam-walking test revealed a significant impairment of the Sox2F/F;Cre mutants as compared to controls, not only in the execution of the test, with a significantly higher number of slips (Fig. 1c), but also in the time needed to complete the crossings on the

beam bar, especially at the earliest time points (Fig. 1d). Thus, Sox2 deletion during cerebellar development leads to defects in fine motor coordination and balance.

Wnt1-Cre-mediated Sox2 deletion leads to morphological cerebellar abnormalities

To understand the causes of the observed ataxic defects, we studied the mutant cerebellum during development and postnatally. Sox2 deletion at E9.5 via Wnt1-Cre results in early posterior shifts of the expression boundaries of two genes, Otx2 and Gbx2, and an early thinning of the neuroepithelium in the medial, prospective vermis region (Supporting Information Fig. S1). Otx2 and Gbx2 encode transcription factors required for the precise spatial definition of the isthmic organizer, a signalling center marking the boundary between prospective mesencephalon and cerebellum, and for subsequent vermis development (Joyner, Liu, & Millet, 2000; J. Y. Li, Lao, & Joyner, 2002; Martinez, Andreu, Mecklenburg, & Echevarria, 2013; V. Y. Wang & Zoghbi, 2001); the correct relative positioning of Otx2 and Gbx2 expression territories is critical for the appropriate development of the isthmic organizer (Broccoli, Boncinelli, & Wurst, 1999).

At birth, abnormalities of cerebellar morphology were observed (Fig. 1e, showing postnatal day 5, P5). The most prominent defects were seen in the medial region, the vermis, which was underdeveloped, in particular along its antero-posterior axis (Fig. 1e,f). Different degrees of severity were observed: from severe reduction (Fig. 1e, far right; "severe" mutants, 11% of total) to milder (though visible) effects, in all other mutants (Fig. 1e left; "moderate" phenotype).

We addressed cerebellar vermis morphology in more detail in three adult mutants (8 months, moderate phenotype), by morphometric analyses (Fig. 2). Vermis alterations were observed in all three mutants: measurements of the total length of the Purkinje cell layer (PCL), and of

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the granule layer (GL) (Fig. 2e-h). Consistent with this result, also PCL lengths in various specific lobules were shorter compared to controls; this reduction was most prominent in lobules IV/V, VIb, and IX and X (20-60% reduction; these lobules are particularly important for equilibrium and posture) (Fig 2i-k). More subtle alterations in foliation were also detected; indeed, the late-born secondary fissure separating lobule IXb/IXc was not observed in mutants, and the VIa/VIb fissure showed a trend to decrease in length (Fig. 2a,b,c,l,m).

Sox2 mutants show alterations of Bergmann glia monolayer formation arising postnatally

To further investigate possible mechanisms for the mutant phenotype, we looked for abnormalities at the cellular level within the postnatal cerebellum, where neurons and glia complete their terminal differentiation and acquire their function.

We first looked at Sox2 expression in the wild type postnatal cerebellum, by double immunofluorescence for SOX2 and cell-type-specific antibodies. At early postnatal ages, SOX2 was clearly expressed in S100B-positive BG (Fig. 3a-b"), though apparently not in adjacent, CALBINDIN-positive PCs (Fig. 3c-c"), in agreement with previous reports (Shiwaku et al., 2010; Sottile, Li, & Scotting, 2006). SOX2 was also expressed by parenchymal astrocytes of the GL and prospective white matter (PWM; Fig. 3a-b), as expected (Sutter et al., 2010). The same pattern of expression was maintained in adult cerebella at 8 months of age (Fig. 3d-d"). In Sox2F/F; Cre mutant mice, Sox2 was not detected, as expected (Fig. 3e-f').

We then looked at possible alterations, in mutants, within specific postnatal cell types, by immunofluorescence with cell type-specific markers, at P7, P19 and 8 months. At 8 months, we did not detect major changes in the density, nor morphology, of PCs, nor of interneurons

(by CALBINDIN and PARVALBUMIN immunofluorescence, respectively) in mutants compared to controls (Supporting Information Fig. S2). As Sox2 is expressed in oligodendrocyte progenitors during development (Zhao et al., 2015), we also examined oligodendrocytes by immunofluorescence using anti-NG2 (recognizing immature oligodendrocyte progenitors), GSTPi (post-mitotic differentiating oligodendrocytes) and Olig2 (the entire oligodendrocyte lineage): no significant differences in the density or distribution of labelled cells was observed at 10 weeks of age (Supporting Information Fig. S3a-c"'). In addition, overall myelination, as detected by MBP immunofluorescence, was not altered in mutant as compared to control mice (Supporting Information Fig. S3d-e"). We then investigated glial cells, which have essential roles in the maintenance of cerebellar neuronal function. In particular, BG cells, which express Sox2 (Fig. 3), surround with their cell bodies the PC, enwrap their cell dendrites, and are critical for their function (Buffo & Rossi, 2013; Custer et al., 2006; Miyazaki et al., 2017). We performed immunofluorescence with antibodies against BLBP (recognizing BG progenitors at P7), and S100B and GFAP (recognizing fully differentiated glia cells, in particular BG, at later stages) (Fig. 4a-r). At P7, mutant cerebella did not reveal overt differences in BLBP- or S100Bpositive glial cell number, morphology, or localization, when compared to the two control genotypes (Sox2F/F, Sox2F/+, or Sox2F/+; Cre; 'controls')(Fig. 4, compare m-o, mutant, with a-c and g-I, controls). At P19, however, abundant S100B/GFAP-positive cell bodies suggestive of mislocalized BG cells were observed in the molecular layer (ML) of the mutant, (Fig. 4p-p"), in contrast with the orderly alignment of \$100BETA/GFAP-positive cell bodies along the PCL, with parallel radial projections extending across the ML, observed in controls (Fig. 4d-d", j-j"). GFAP immunofluorescence (staining the cytoplasm more intensely than S100B) further showed that ectopic cells had a non-radial, abnormal morphology (Fig. 4p", arrowheads). These ectopic cells were still abundant at 8 months (Fig. 4, compare q-r, mutant, to e-f and k-l,

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controls; quantitation in s), and GFAP immunofluorescence highlighted a characteristic aberrant, multipolar morphology, with dense arborizations (Fig. 4r, arrowheads). The abnormal cell morphology was particularly evident in the ectopic cells (Fig. 4r', inset), but also occasionally observed on the cells aligned along the PCL (Fig. 4r, arrows). Overall, the number of non-ectopic BG in the mutant PCL was significantly decreased compared to control animals (Fig. 4t,u). Importantly, by contrast with the abnormalities in mutant BG, parenchymal astrocytes of the GL and WM appeared unchanged among genotypes (Supporting Information Fig. S2). We conclude that the localization and morphology of glial cells (BG, based on the location) are progressively altered during postnatal life in Wnt1Cre-Sox2 mutant mice.

Alteration of synaptic transmission at parallel fiber-Purkinje cell synapse Given the important role of BG in modulating the excitatory synapse transmission in PCs, we examined synaptic transmission between parallel fibers (PFs) and PCs by means of whole-cell voltage clamp. We performed our analysis in adult mice, in the posterior lobules of the cerebellum, which had revealed the most prominent reduction of the PCL length (Fig. 2). We electrically stimulated PFs using a stimulation electrode placed in the molecular layer and recorded excitatory glutamatergic synaptic currents (PF-EPSCs) in the PC. PF-EPSCs had comparable amplitudes at every intensity of stimulation in both mutant (n = 16 cells) and control mice (n = 17 cells) (P> 0.05, Two- way ANOVA repeated measures –RM-) (Fig 5a-b). However, while most EPSCs of control mice had amplitudes smaller than 250 pA with stimulus strength of 12 μ A (Fig 5c), a large fraction of mutant mice EPSCs exceeded this value (Fig 5d). Cumulative plots revealed significant differences between genotypes in the distribution of EPSC amplitudes, with mutant values shifted to the right towards larger values (KS, P<0.0001) (Fig 5e). To evaluate possible alterations in glutamate release from

presynaptic terminals we stimulated PFs with paired stimuli at different intervals. The analysis of paired pulse facilitation of the PF-PC synapse did not show differences between genotypes, suggesting a preserved presynaptic release of glutamate by PFs (P>0.05, Two way Anova RM) (Fig 5f-g). Alterations of BG are associated with changes in the kinetics of PF-EPSCs (Takayasu et al., 2005). To address this issue we measured the decay time constant of PF-EPCSs. We did not find changes between genotypes in the decay time constant (for controls: 19.26±1.45ms; for mutant mice: 18.36±2.25ms P>0.05, Student's t-test). However, the analysis of the distribution of decay time constants of mutant mice EPSCs revealed the presence of two populations with faster and slower decays (data not shown), while the data of control mice were better described as a single population. The heterogeneity of the synaptic properties of the EPSCs is in line with the presence of ectopic BG, which might alter synaptic transmission at nearby synaptic contacts.

Sox2 postnatal deletion in radial glia leads to ectopic, abnormal and disorganized Bergmann glia and to mild motor control defects

The expression of SOX2 in postnatal BG, together with the defects in BG observed in mutants, raised the possibility that SOX2 could be required postnatally, within the BG itself, to maintain these cells and their characteristics. To test this hypothesis, we deleted Sox2 in BG postnatally, by means of a tamoxifen (Tx)-inducible Cre encoded by a GLAST^{CreERT2} transgene (Buffo et al., 2008; Mori et al., 2006). Tx was administered at P2 or P8, and cerebella were analysed at 1 or 8 months of age (Fig. 6). Cre induction leads to efficient ablation of SOX2 from S100B-positive BG, reaching 60-80% (Fig. 6c-d, mutant, compare to a-b, controls; quantitation in e). In controls, at P30, the S100B-positive BG (red), largely Sox2-positive (green), form an ordered array of cells with their cell bodies aligned in the PCL (at the bottom

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of the molecular layer)(Fig. 6a-a",b-b"); by contrast, in the mutant, S100B-positive cells show wide disorganization, with their cell bodies localized throughout the molecular layer (Fig. 6c-c",d-d", quantitation in f). Sox2 immunofluorescence confirms absence of SOX2 from most S100B-positive cells in the mutant (Fig. 6c-c",d-d"), to be compared to the wide Sox2-positivity of S100B-positive cells in the control (Fig. 6a-a",b-b"). S100B/GFAP immunofluorescence at 8 months confirmed that S100B-positive cells in mutant were widely disorganized and mislocalized, and showed highly abnormal morphology by GFAP immunostaining, together with a disruption of the normal, radial organization of the BG, and abundant ectopic BG (Fig. 6g-i, compare h, mutant, to g, control). Similar to what found in Wnt1-Cre Sox2 mutants, deletion of Sox2 also leads to a decreased amount of BG in the PCL, as indicated by a similar reduced BG/PC ratio in both mutant lines (Sox2F/F;CreERT2, 2.939±0.098 vs Sox2F/F;Cre 2.606±0.1487) compared to controls (not shown).

These findings indicate that Sox2 is required during postnatal life to maintain BG alignment along PC cell bodies, and their characteristic radial morphology and organization.

We further asked if postnatal Sox2 deletion would lead to motor control defects, together with BG abnormalities. Since Tx injection in P2 animal significantly affected newborn survival and for behavioural studies numerous litters are needed, we postponed the treatment to P8 and combined it with a second administration at P60 (see Materials & Method section). This protocol determined BG ectopia to an extent similar to Tx treatment at P2 (Fig. 6i). By the beam-walking test, a significant difference was found between mutants and controls, with no differences between sexes (not shown), similarly to what observed with Wnt1-Cre mutants, although the extent of the motor alterations was much smaller (Fig. 7a,b). By the rotarod test, mutants did not perform significantly different from controls (Fig. 7c). However, when male

and female mutant and control mice were separately compared, mutant male mice, that initially performed comparably to controls (month 1), later (month 2) showed a significantly worse performance than controls, though the difference was clearly smaller than that found for Wnt1-Cre mutants (see Fig. 1b-d)(Fig. 7d). We wished to check if the abnormalities detected in the motor control tests (small, though statistically significant) could be affected by an increased anxiety in the mutants influencing motor performances. We thus performed, on the same mice previously evaluated in the motor control tests, a test evaluating both anxiety and general locomotor activity, the Open field test. No significant difference was observed between mutants and controls. (Fig. 7d). We conclude that postnatal Sox2 deletion in glia leads to mild motor control defects, based on both the beam-walking and rotarod tests.

Abnormalities of glutamate transport in Sox2 mutant BG

BG expresses the excitatory amino acid - glutamate transporter GLAST (encoded by the Slc1a3 gene), important for the removal of glutamate from the synaptic cleft, and mutations in human SLC1A3 were found in patients affected by episodic ataxia (Jayadev & Bird, 2013). Further, BG was shown to be essential for synaptic organization and electrophysiological function of PCs through the action of GLAST, in studies of Slc1a3 knockout mice (Miyazaki et al., 2017). Glutamatergic transmission requires the involvement of glia cells to efficiently clear the synaptic cleft and to recycle the transmitter through a Glutamate to Glutamine (Glu/Gln) metabolic shuttle, in a multicomponent regulated process. We thus performed Glu uptake experiments in gliosomes purified from BG-enriched cell preparations of 6 wild type and 5 Sox2-Wnt1Cre mutant mice. Gliosomes represent a well-recognised subcellular preparation that recapitulates molecular and functional features of the astroglial perisynaptic processes (Carney et al., 2014; Stigliani et al., 2006). As shown in Fig. 8, Sox2-Wnt1Cre gliosomes

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showed a significant increase of uptake Vmax, compared to age matched control mice, with no significant changes in the Km values (Supporting Information Figure S4). In parallel we measured the gliosomal content of endogenous Glu and Gln, and calculated the Glu/Gln ratio: these ratios were statistically comparable in gliosomes from Sox2-Wnt1Cre and WT BG mice (Supporting Information Figure S4). These data demonstrate a clear biochemical alteration of the Sox2-mutant BG cells, namely, Glu uptake abnormalities, without modification of Glu/glutamine conversion.

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Sox2 deletion in neural stem cells causes widespread deregulation of gene expression in in-vitroderived glial cells

Previous work (Koirala & Corfas, 2010) identified genes expressed in BG at P6, or P30, or at both stages. To gain insight into the effect of Sox2 deletion on gene expression in glia, we differentiated in vitro neural stem cells obtained, at P0, both from wild type newborn mice and from littermates embryonically deleted for Sox2 (at E11.5, see (Favaro et al., 2009)). We then determined genes deregulated at day 11 of in vitro differentiation; at this stage, cells are postmitotic, and 90-95% of them are glial cells, with a small proportion of oligodendroglia (1-2%) and neurons (about 5%). Finally, we compared the list of deregulated genes in differentiated mutant glia with those of genes expressed in BG at P6, P30, or both. Over 450 genes, out of 1183 genes expressed at both P6 and P30 in BG, were found to be deregulated in Sox2-mutant glia (Supplementary Table 3); furthermore, a highly significant fraction (266 genes) is likely to be direct SOX2 targets, based on published SOX2 ChIP-seq data (Lodato et al., 2013) in which SOX2 target genes were defined as being SOX2-bound in regulatory regions (Supplementary Table 3). These results point to an important role of Sox2 in the control of gene expression in differentiated glia, and possibly BG. Interestingly, among SOX2-bound, BG-

expressed genes downregulated in Sox2-mutant cells (SOX2_UP WT_UP BOTH in Supplementary Table 3), several (highlighted) are potentially related to ataxia, based on previous publications. Among these genes, Slc1a3 (GLAST) was Sox2-bound and moderately reduced, at statistically significant values, in *in vitro* differentiated glia, implying a potential effect of Sox2 deficiency on glutamate uptake; however, Slc1a3 expression, as detected by immunofluorescence in BG in tissue sections, was somewhat variable between different mutant mice, and overall not significantly different from controls (not shown). On the other hand, glutamate uptake by BG is a complex phenomenon, affected by many different components (i.e. gene products) in addition to the Slc1a3 protein amount. Future work will better define the relations between the altered gene expression pattern caused by Sox2 deficiency and the increased glutamate uptake that we detected in mutant BG, as well as their relation with PC dysfunction and ataxia.

Discussion

Sox2 mutations in man cause genetically dominant defects in the central nervous system, including hippocampus hypoplasia, cognitive defects, motor control abnormalities, seizures, retina and eye defects; some of these abnormalities have been reproduced in mouse models of conditional Sox2 ablation (Dennert et al., 2017; Favaro et al., 2009; A. Ferri et al., 2013; Pevny & Nicolis, 2010; Ragge et al., 2005; Sisodiya et al., 2006). In previously studied Sox2 mutant mouse models, the observed defects develop prenatally, or immediately after birth, and are related mainly to neural stem/progenitor cells defects. An exception is represented by abnormalities of GABAergic interneurons morphology and postmitotic migration (Cavallaro et al., 2008) in the forebrain, as well as defects of differentiated glia types: Mueller glia, a

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differentiated glial type of the eye, and oligodendroglia (Bachleda, Pevny, & Weiss, 2016; Hoffmann et al., 2014; Pevny & Nicolis, 2010).

In our Wnt1Cre-Sox2flox mouse model, we report that Sox2 is important for the correct prenatal morphogenesis of the cerebellum, and in particular of the vermis. In addition, Sox2 controls the correct morphology, organization/localization and development of the BG in the first three weeks after birth. As the BG population is mainly postmitotic after the first postnatal week, when BG defects arise, this finding points to important roles of Sox2 within a specifically differentiating, non-dividing, glia cell type.

The cerebellar defects we observed lead to ataxic features, which are more pronounced when Sox2 deletion is performed prenatally, but are still observed with (partial) postnatal deletion within BG.

Genetic mechanisms contributing to the development of ataxic features

The significant reduction of the vermis observed in newborn Wnt1Cre Sox2 mutants is one plausible contributor to the ataxic features appearing shortly after birth. In fact, vermis reduction, together with ataxia, was previously observed, in several mutant mouse models and human inherited diseases. In mouse, an experimental posterior shift (by Otx2 knock-in into the En1 locus) of the caudal expression boundary of the Otx2 transcription factor, a critical regulator of the cerebellum/mesencephalon boundary, leads to impaired vermis development and to ataxia (Broccoli et al., 1999). Our Sox2 mutant mimics, to a certain extent, the expression changes of Otx2 (and of Gbx2) observed in this mutant mouse model (see Supporting Information Fig. 1), suggesting that vermis reduction in the Sox2 mutant is due, at least in part, to Otx2/Gbx2 dysregulation. In humans, vermis reduction is observed, together with ataxia, in CHARGE syndrome, an autosomal dominant disorder caused by mutation of

CHD7 (Yu et al., 2013), a chromatin regulatory protein interacting with SOX2 (Engelen et al., 2011). In mouse models carrying Chd7 mutations, Otx2 and Gbx2 expression boundaries are shifted posteriorly, similarly to what we find in Sox2 mutants (Supporting Information Fig.S1), and CHD7 directly binds to the Otx2 and Gbx2 regulatory regions (Yu et al., 2013), on sites at least partially overlapping those bound by SOX2 (Engelen et al., 2011; Lodato et al., 2013; Yu et al., 2013). This suggests that a shared SOX2 and CHD7-controlled gene regulatory network, involving early Otx2 and Gbx2 regulation, may contribute to pathological features shared between the Sox2-deficiency and the CHARGE human syndromes. Vermis hypoplasia is also characteristic of Joubert and Meckel syndromes (JMS), caused by mutation of several different genes (Aguilar et al., 2012); interestingly, SOX2 ChIPseq experiments on cultured neural stem cells (NSC) (Engelen et al., 2011; Lodato et al., 2013) detect SOX2 binding to the promoter or to other gene sequences of various JMS genes (8 out of 17 JMS genes, mentioned in Aguilar et al., 2012, carry SOX2-bound promoters), a finding that was confirmed in studies of mouse brain-derived NSC (our unpublished observations), suggesting that Sox2 may be involved in a wider gene regulatory network regulating vermis development. Future experiments will address whether deregulation of these genes could also contribute to the cerebellar abnormalities in our mutants.

Taken together, these previous results suggest that the ataxic features observed in our Wnt1-Cre Sox2-mutant mice are due, at least in part, to the abnormalities that we observe in the vermis at birth. However, it is important to note that functional motor abnormalities are observed, although in a milder form, also in mice in which Sox2 ablation was specifically obtained after birth, and were not therefore related to any vermis morphological abnormality. This point is further discussed below.

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The relation between ataxia and Bergmann glia abnormalities

The observed BG abnormalities raise the issue of their possible direct involvement in the ataxia found in Sox2 mutants. Sox2 is expressed in BG postnatally (Fig. 3). BG ectopic location and cellular dysmorphology is observed in both embryonic (Wnt1-Cre) and postnatal (GLASTCreERT2) Sox2 mutants, (Figs. 4,6), beginning after the first week of postnatal life; it is first detected at P19, and persists in adults (8 months and later). Similarly to embryonic Sox2 deletion, postnatal Sox2 deletion specifically in BG leads to ataxic features (Fig. 7), although the severity of the motor control defect is reduced compared to that observed in embryonic Sox2 mutants (Fig. 1e,f); note, however, that in postnatal mutants, the deletion of Sox2 in BG is incomplete (60-80%, Fig. 6), which may explain, in part, the milder phenotype. It is important to note that, in postnatal cerebellum, only BG and other glial cells, but neither PC nor other neurons, express Sox2; in addition, only BG, but not other types of glial cells, show obvious abnormalities. This points to BG as a likely contributor to the functional abnormalities observed in Sox2 mutants, both prenatal (Wnt1-Cre) and, in particular, postnatal (GLAST-CreERT2).

Mutation of other mouse genes leads to BG cytoarchitectural defects comparable to those in our mutants, accompanied by ataxia. Inactivation in BG (via a GFAP-Cre transgene) of the tumor suppressor adenomatous polyposis coli (APC) gene, encoding an inhibitor of the Wnt/beta-catenin pathway, disrupts BG architecture after P10, with translocation of their cell bodies to the molecular layer, loss of their pial contact and transformation into stellate-shaped cells, leading to motor coordination deficits (X. Wang, Imura, Sofroniew, & Fushiki, 2011). The Sox4 transcription factor gene is expressed in normal BG; intriguingly, its overexpression, in BG, via a hGFAP promoter leads to BG cytoarchitectural defects, resulting in ataxia (Hoser et al., 2007). Interestingly, ChIP studies (Lodato et al., 2013, and our

unpublished observations) show binding of Sox2 to the promoter regions of Sox4 and APC, suggesting a possible direct regulation. It is interesting to note that, in our GLAST^{CreERT2} Sox2 mutant, the Sox2 deletion occurred in a subpopulation of the glial cells, BG and astroglia; the displaced BG cells included both Sox2-null and Sox2-positive cells. This result implies that non-cell autonomous mechanisms might be involved in causing cells which retain Sox2 expression to migrate away from their normal location along the PCL. As shown in Supplementary Table 3, glial cells express many Sox2-bound genes encoding diffusible factors, whose deregulated expression, following Sox2 ablation in glia, might thus affect the localization of both Sox2-deleted and non-deleted BG cells.

BG is known to play critical roles for the functionality of PCs in the mature cerebellum, and its defects have been shown to lead to ataxia by non-cell-autonomous mechanisms, whereby BG defects caused PC dysfunction. Indeed, mutant ataxin 7 (causing the autosomal dominant disease spinocerebellar ataxia 7, SCA7) caused neuronal degeneration when expressed in mouse BG via a cell-type-specific transgene (Custer et al., 2006), and mutant ataxin-1 (causing SCA1), when expressed in BG, contributed to non-cell-autonomous toxicity affecting PCs (Shiwaku et al., 2010). In our experiments, a proportion of the PC show aberrant postsynaptic currents of the PF-PC synapse, as expected given the connection between regulatory circuits operating in BG, and PC physiology (Fig. 5); on the other hand, the majority of responses were normal, perhaps reflecting the patchy distribution of ectopic BG.

How BG alterations relate to the PC disturbed function in Sox2-mutant cerebellum, is difficult to speculate at the present time. One important fact might be the loss of the normally close apposition of many BG cells relative to PC in the mutant cerebellum. Additionally, the presence of aberrant postsynaptic responses at the PF-PC synapse, suggests that the dysregulation of BG-PC signalling might have impaired the normal homeostasis based on a

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balance between synaptic potentiation and depression, leading to the generation of improper cortical output signals by the affected PCs.

Collectively, our results demonstrate a novel role of Sox2 in the maintenance of morphology and localization of BG, as well as on at least one major biochemical activity (gliosome glutamate uptake) of these cells. Further, gene expression analysis of Sox2 mutant glia identifies a large number of potential direct Sox2 target genes (Supplementary Table 3), and point to a gene regulatory network that connects Sox2 to the regulation of other genes important in BG maintenance and in ataxia. Identifying the downstream targets of Sox2 which are responsible, in Sox2 mutants, for the observed BG abnormalities, will be important for a better understanding of mechanisms operating in BG cells to affect PC function.

Future studies will clarify whether Sox2 expression is somehow altered in BG in ataxias occurring in man; additionally, it is possible that some of the transcriptional targets of Sox2 are among the genes whose activity is altered in ataxias. The future characterization and understanding of the Sox2-dependent regulatory network in BG might hopefully provide novel targets and new perspectives for therapeutic approaches.

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Figure legends

Figure 1. Wnt1-Cre Sox2-mutant mice show defects in motor control and cerebellar morphology

- (a) Sox2 early conditional deletion in the midbrain-hindbrain. Sox2 in situ hybridization on E9.5 sagittal sections of control (Sox2F/F, left) and Sox2-deleted embryos (Sox2F/F;Cre, right) showing loss of Sox2 expression in the mid-hindbrain by E9.5. Fb: forebrain; Mes: mesencephalon; CbP: Cerebellar primordium.
- (b-d) Motor performances of Sox2F/F;Cre mutants were evaluated at rotarod test (b) and beam test (c,d) and compared to control littermates (Sox2FF/F+, Sox2F/+;Cre) from 1 to 8 months of age. At the rotarod test, both Sox2F/F;Cre and Sox2F/+;Cre animals showed a significantly lower latency to fall compared to controls (b; Two-way Anova, main effect of genotype, P=0.0001); at the beam test, Sox2F/F;Cre mutants overall made more slip errors (b;Two-way Anova, main effect of genotype, P=0.0001) and took more time to cross the bar compared to controls (d, Two-way Anova, main effect of genotype, P=0.0001; Time, Two-way Anova, main effect of genotype, P=0.0001, main effect of time, P=0.002). Details on statistical analysis are reported in Table 1.
- (e,f) Sox2 early conditional deletion leads to cerebellar morphological defects. (e) Cerebellar morphology at P5. Two Sox2F/F;Cre mutants are shown, with "moderate" (left) and "severe" (right) phenotype. Vertical arrows point to the antero-posterior extension of the vermis, reduced in the mutants. Some enlargement of the adjacent mesencephalic lobes is also seen in mutants, and in controls carrying the Cre transgene as previously described (Lewis,

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Vasudevan, O'Neill, Soriano, & Bush, 2013). The lateral arrow in the second mutant ("severe") points to a central "hole" at the mid/hindbrain boundary. The severe phenotype was observed in 4/44 mutant mice examined (11%). (f) Cerebellar morphology in adults, dorsal view; vermis lobules are numbered. In mutants, the vermis is reduced along the antero-posterior axis. Scale bars: A, 200μ M; B,C: 2mm.

Figure 2. Adult Wnt1Cre mutant mice show morphological alterations in the cerebellar vermis

(a-c) Sagittal sections of the cerebellar vermis of Sox2FF/F+ (a), Sox2F/+;Cre (b), and Sox2F/F;Cre (c) adult mice (8 months) display a reduction in size in the mutant accompanied by subtle alterations in defined fissures (asterisks in a',a",b',b",c',c"). (d) Morphometric measurements for each examined section comprised the evaluation of i) the PCL length along the entire section (green line); ii) the maximal extension along the Rostro-Caudal (R-C) axis axis, defined as the distance connecting the maximal convexity of the crowns of lobules V and IXb, and the Ventro-Dorsal (V-D) axis, corresponding to the distance connecting the invagination between lobules I and X and the midpoint in the crown of lobule VIb (dotted lines, see Methods section); iii) the thickness of the GL in each lobule (red dotted line); iv) the PCL length along each lobule (green line between the two green crosses) and the extension of the fissures separating lobules IVa/VIb (yellow line) and lobules IXb/IXc (purple line). (e) The extension of the entire PCL is significantly diminished in Sox2F/F;Cre compared to both Sox2FF/F+ and Sox2F/+; Cre mice (One-way Anova, P=0.0026). A reduction in size is also observed in the mutant R-C length (f; One-way Anova, P=0.0014), whereas no differences are detected in the V-D extension (g), as well as in GL thickness (h). (i-k) PCL length declines in defined lobules of the mutant compared to controls (k, lobule IV-V, One-way Anova, P=0.0195; i, lobule VIb, One-way Anova, P=0.0084; j, lobules IX and X, One-way Anova, P=0.0055 and P=0.0124, respectively). (k-m) Mutant mice show fine alterations in foliation, as revealed by the absence of the fissure between lobules IXb/IXc (a", b", c" - violet asterisk -, l) and by the poor development of the fissure separating lobules VIa/VIb (a', b', c' - yellow asterisk -, M; One-way Anova, P=0.0411). Dapi staining (blue in a-c") labels all cell nuclei and allows visualization of layering. Scale bars: a, b, c, 300 μ m; a', a", b', b", c', c", 100 μ m. Details on statistical analysis are reported in Table 1. * P< 0.05; ** P<0.01; *** P<0.001.

Figure 3. SOX2 expression in postnatal and adult cerebellum

(a-f') Immunofluorescence for SOX2 (green) and glial (S100B, red), or neuronal (Calbindin, Calb, red) markers in the control (a-d''') and mutant (e-f') postnatal cerebellum. (a-b'') SOX2 immunofluorescence (green) in the postnatal (P7) cerebellum shows SOX2 expression in S100B+ astrocytes(red in a-b'), i.e. BG cells in the PCL (arrowheads in b-b'') and parenchymal astrocytes of the GL and PWM (arrows in b-b''). (c-c''') At P7, SOX2labelling is not detected in Calbindin+ PCs (red). (e-f') Absence of Sox2 expression in mutant Sox2F/F;Cre cerebella at both P7 (e,e') and 8 months (f,f'). ML, molecular layer; PCL, Purkinje cell layer, GL, granular layer, PWM, prospective white matter. Scale bars: A, D, E-F', 50 μm; B-C, D'-D''', 20 μm; C'-C''', 10 μm.

Figure 4. Wnt1-Cre Sox2 mutants show postnatal BG alterations

(a-r) Glial cells were labelled with specific markers as BLBP, S100B and GFAP in control (a-f, Sox2FF/F+; g-l, Sox2F/+; CRE) and in mutant (m-r, Sox2F/F; Cre) cerebella at different ages. At P7 (a-c; g-i; m-o) no significant differences were detected in BG progenitors distribution, as BG cells were similarly labelled with BLBP and S100B in the arising PCL of the three

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genotypes. At P19 (d-d"; j-j"; p-p") ectopic S100B+ cells appeared in the mutant ML (p-p"; arrows in p') and these cells ectopic remained in the adult cerebella at 8 months (arrows in q). GFAP staining confirmed the presence of mislocalized BG in the mutant ML, at both P19 and 8 months, and highlighted theaberrant multipolar morphology of these cells (arrowheads in p" and r; r'). (s) Quantification of S100B+ cells in 8 months old mutant or control cerebella confirmed the significant presence of ectopic BG cells in the mutant ML (Student T test, P < 0.0001; Sox2F/F;Cre mice vs. Sox2FF/F+ and Sox2F/+;Cre mice). (t,u) Analysis of cell density (n. of cells/mm²) of S100B+ cells in the PCL and related ratio of S100B+ cells to PCs (u) showed a significant decrease of BG somata correctly aligned in the PCL (t) of 8 months old mutant mice (Student T tests, t, P=0.0179; u, P=0,0014; Sox2F/F;Cre vs. Sox2FF/F+ and Sox2F/+;Cre mice). (s-u) Being fully comparable, data from the two control genotypes were pooled together (see Table 2). Dapi staining (blue in a,e,g,k,m,q) labels all cell nuclei. ML, molecular layer; PCL, Purkinje cell layer, GL, granular layer, WM, white matter. Scale bars: a, g, m, 100 μm; c, d-d", f, hj", l, o-p", r, 50 μm; b, 20 μm; e,k,q, 30 μm; n, 10μm. Details on statistical analysis are reported in Table 1. * P< 0.05; ** P<0.01; *** P<0.001.

Figure 5. Alteration of the PF-EPSC in adult mutant mice

(a) Five superimposed representative traces for control (blue; Sox2FF/F+) and mutant mice EPSCs (red; Sox2F/F;Cre) obtained with stimulus strengths from 3 to 15 μ A. (b) Amplitudes of Parallel Fibre (PF)- Excitatory postsynaptic potentials (EPSCs) plotted as a function of stimulus intensity for control (Sox2FF/F+, blue circles, n = 17) and mutant (Sox2F/F;Cre; red triangles, n = 16) PCs. (c) Histogram distribution of EPSC amplitudes (obtained with stimulus strength of 12 μ A) for control (c; Sox2FF/F+) and for mutant (d) mice. (e) Cumulative distribution of EPSC amplitudes for control (Sox2FF/F+, blue trace) and mutant mice

(Sox2F/F;Cre, red trace) (P<0.0001, Kolmogorov-Smirnov test). (f) Representative traces of PF-EPSCs evoked by paired-pulse stimulation with an interpulse interval of 100 ms in control (blue; Sox2F/F;+) and mutant mice (red; Sox2F/F;Cre). (g) Time course of paired-pulse facilitation of PF-EPSCs in control (Sox2FF/F+, blue circles, n = 17) and mutant mice (Sox2F/F;Cre, red triangles, n = 16). The facilitation is expressed as the percentage of the second EPSC relative to the first one (mean \pm SEM) and is plotted as a function of interstimulus interval.

Figure 6. Postnatal Sox2 deletion in GLAST^{CreERT2}Sox2 mutants causes BG alteration

(a-d") Immunofluorescence analysis of Sox2F/F;CreERT2mutants after Tamoxifen (Tx) administration. P30 Sox2F/F;CreERT2 cerebella induced with Tx at P2 showed high numbers of mislocalized S100B+ BG in the ML (c-d"), most of which have lost Sox2 expression (arrows; arrowheads point to cells that maintained Sox2), compared to controls (a-b"). (d) A higher fraction of ectopic cells could be occasionally observed, within regions where Sox2 ablation had been particularly efficient (d-d"). (e) Quantification of Sox2-ablated cells. At P30, after tamoxifen administration at P2, a high percentage of S100B+ cells in Sox2F/F;CreERT2mutant cerebella does not express SOX2, in agreement with an efficient ablation of the gene. (f) Fraction of ectopic S100B+BGon total S100B+BGwas significantly higher in mutant Sox2F/F;CreERT2 cerebella induced at P2 with respect to control littermates at P30 (Student T test, P = 0,0011). (g-i) Ectopic S100B+ BG in the ML of mutant mice (arrows in h) was maintained at 8 months after Tx treatment at P2 orat P8 plus P60 (P8-60). Controls include genotypes as for Table 2.Tx, tamoxifen; ML, molecular layer; PCL, Purkinje cell layer. Scale bars: a-d", 50 μm; g,h, 20 μm. Details on statistical analysis are reported in Table 1.* P< 0.05; ** P<0.01; *** P<0.001.

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Figure 7. Postnatal Sox2 deletion in GLAST^{CreERT2} Sox2mutants causes motor control abnormalities

(a-d) Motor performances of Tx-treated Sox2F/F;CreERT2 mutants were evaluated with the beam (a,b) and rotarod tests (c,d) and compared to those of control littermates (Controls, untreated) from 1 to 4 months of age. In the beam test, mutants of either sex exhibited slight alterations of motor behaviour compared to controls, as revealed by an increase in the number of slip errors (a, Two-way AnovaRM, Main effect of genotype, P<0.0454) and in the time needed to cross the bar (b, Two-way AnovaRM, Main effect of time, P<0.0001). When all animals were pooled together, at the rotarod test no significant defects were detected in mutants, as showed by a similar latency to fall compared to controls (c). However, comparisons of mutant and control males highlighted that mutant males have a trend to perform in a worse way compared to controls (d, Two-way AnovaRM, Main effect of interaction of genotype per time, P<0.0001). (e,f) In the Open field test, both the distance travelled and the time spent in the center of the field were comparable in mutants and controls (Student T test, distance, P = 0.802; time, P = 0.7957). Further details on statistical analysis are reported in Supplementary Table 1.

Figure 8. Wnt1-Cre Sox2 mutants show gliosomal glutamate uptake alterations Gliosomes from control (Sox2FF/F+, blue circles) and mutant (Sox2F/F;Cre; orange circles) mice were exposed for 2 min at 37 °C to increasing concentration of [3 H]D-Asp (0.3, 3, 5, 30 and 100 μ M). The specific glutamate uptake for each concentration tested is expressed as nmol/2min. Values are expressed as nmol Glu/mg protein content / 2 min and are the mean \pm SEM of 5 independent experiments for each group. The kinetic parameters Vmax and Km

values were obtained by fitting the points with the Michaelis-Menten equation $(Vmax_{[Sox2FF/F^+]}=7.68\pm0.992 \quad nmol/mg/2min, \quad n=5; \quad Vmax_{[Sox2F/F;Cre]} = 12.60\pm1.16 \quad nmol/mg/2min, \quad n=5). \quad *P<0.05 \quad (P_{t=-3.195}=0.019) \quad for \quad Vmax_{[Sox2FF/F^+]} \quad vs. \quad Vmax_{[Sox2F/F;Cre]} \quad (two tailed Student's$ *t* $-test). No significant differences were registered for Km values <math>(Km_{[Sox2FF/F^+]}=25.70\pm9.57 \, \mu M \, n=5; \, Km_{[Sox2F/F;Cre]}=24.30\pm6.60 \, \mu M, \, n=5).$

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Supporting Information Figure and Table legends

Supporting Information Figure S1. In Sox2 mutants, Otx2 and Gbx2 show early alterations of expression boundaries

(a,b) Otx2 and Gbx2 in situ hybridization on adjacent sagittal (a) or coronal (b) sections at E10.5 of mutants and controls (images of the mid-hindbrain boundary region are shown). (a) sagittal sections are shown at a lateral or medial level. Arrowheads point to the expression boundary of Otx2 or Gbx2, shifted posteriorly in mutants, evident in the medial region. A thinner neuroepithelium in mutants is visible in medial sections, corresponding to the prospective vermis region; this is reminiscent of a similar observation in the Gbx2 knock-out (J. Y. Li et al., 2002), reported to cause vermis hypoplasia. (b) Coronal sections. Arrowheads point to the posteriorly shifted boundary of Otx2 expression. Embryos shown in A and B are representative of: for coronal and sagittal sections analysis, respectively, n=5 and n=4 Sox2F/F;Cre mutants, n=7 and n=3 Sox2F/+;Cre heterozygotes, and n=4 and n=4 Sox2F/F or F/+ homozygous wild type controls. Ant: anterior; Post: posterior; Mes: mesencephalon; CbP: cerebellar primordium; Rho: rhomboencephalon (in green in b), which includes the cerebellar primordium; IVv: fourth ventricle. Scale bars: 200μM.

Supporting Information Figure S2. Wnt1-Cre Sox2 mutants do not show alterations in the density of Purkinje cells, interneurons or astrocytes in deep cerebellar layers

(a,b) At 8 months of age, no significant differences were detected in the density of either CALBINDIN-positive Purkinje cells (a) or PARVALBUMIN-expressing molecular layer interneurons (b) between control (Sox2FF/F+ and Sox2F/+;Cre) and mutant animals

(Sox2F/F;Cre). (c,d) Similar results were obtained for the density of S100B-positive astrocytes in the GL (c) and WM (d) at P19. DAPI staining (blue) labels all cell nuclei. GL, granular layer, WM, white matter, PCL, Purkinje cell layer, Calb, CALBINDIN, PV, PARVALBUMIN. Scale bars: 30μm.

Supporting Information Figure S3. Wnt1-Cre Sox2 mutants do not show alterations in the density of cells in the oligodendroglial lineage or myelination defects

(a) At 10 weeks of age, staining for NG2 (a, red) did not reveal differences in the density of oligodendrocyte progenitors between mutant (Sox2F/F;Cre) and control (Sox2FF/F+ and Sox2F/+;Cre) mice, either in the molecular layer (ML, a'), or in the granular layer (GL, a") and white matter (WM, a""). (b) Similarly, as revealed by GSTPi immunostaining (b, red), postmitotic differentiating oligodendrocytes showed comparable densities in all layers in both mutant and control mice (b',b"). These results were further confirmed by Olig2 immunostaining (c-c"", red), which labels the entire oligodendrocyte lineage. (d-e) Anti-MBP immunofluorescence (red) was overall comparable in mutant (e) and control (d) mice, in both lobular (d',e') and deep (d", e") white matter. Dapi staining labels cell nuclei and allows layers visualization. ML, molecular layer, GL, granular layer, WM, white matter. Scale bars: a,b,c,d',d",e',e", 50µm; d,e, 200µm.

Supporting Information Figure S4. Endogenous glutamate and glutamine quantification. High performance liquid chromatography analysis of the cytosolic endogenous glutamate (a) and glutamine (b) content measured in gliosomes of BG cells purified from control (Sox2FF/F+, blue bars) and mutant (Sox2F/F;Cre; orange bars) mice. Endogenous glutamate (Glu) and glutamine (Gln) content are expressed as nmol/mg of

gliosomal protein content and represent the mean \pm SEM of 5 independent experiments for each experimental group ($Glu_{[Sox2FF/F+]} = 38.26\pm4.88$ nmol/mg, n=5; $Glu_{[Sox2F/F;Cre]} = 34.49\pm5.01$ nmol/mg, n=5; $Gln_{[Sox2FF/F+]} = 20.12\pm3.98$ nmol/mg, n=5; $Gln_{[Sox2F/F;Cre]} = 24.94\pm5.66$ nmol/mg, n=5). (c) Endogenous cytosolic glutamate/glutamine content ratio in gliosomes of BG cells of control and mutant mice ($Glu/Gln_{[Sox2FF/F+]} = 2.07\pm0.23$ nmol/mg, n=5; $Glu/Gln_{[Sox2F/F;Cre]} = 1.52\pm0.17$ nmol/mg, n=5). No statistically significant differences have been detected.

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Supplementary Table 1. Details of statistical analyses.

In most cases only comparisons leading to P<0.05 are reported; n = number of animals (unless differently stated). RM, repeated measures.

Supplementary Table 2. Average numerical values obtained in immunofluorescence experiments, and statistical analysis.

Supplementary Table 3. Genome-wide analysis of genes deregulated in glia following Sox2 loss, and their overlap with BG-expressed genes, and direct SOX2 binding

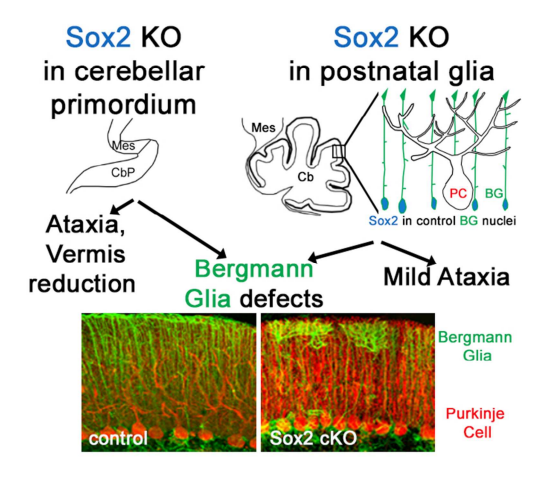
(a) "Expression" sheet: Output of the analysis performed by NOISeq (see Methods) on RNA-Seq data. Expression values, defined as Transcripts per Million (TPM) in three replicates of wild-type (WT 1-3) and Sox2-deleted cells (MUT 1-3). Genes are ordered according to significance of expression variation, from high to low. For each gene, the table reports: q-value (1-FDR) output by NOISeq; the MUT/WT fold ratio; the significance of expression

variation at a false discovery rate (FDR) threshold of 0.01 (SIGP, +1 over-expressed in MUT, -1 over-expressed in WT); whether or not the gene is included in the genes expressed in BG at postnatal day 6 (P6) (UP P6 column), at P30 (UP P30), or both (UP BOTH) (datasets from Koirala and Corfas 2010); whether or not the gene is SOX2-bound on the promoter or proximal enhancer (SOX2 in PROMOTER or SOX2 in ENHANCER) (dataset in Lodato et al., 2013). The significance of each overlap and corresponding p-values (columns Y to AE in "Expression" sheet) were computed with a hypergeometric test.

Note that, in Koirala and Korfas (2011), UP P6 are genes expressed, in BG at P6, at levels at least 3 times higher that at P30; UP P30 are genes expressed at P30 at levels at least 3 times higher that at P6; UP BOTH are genes expressed at both stages.

(b) "Gene lists" sheet: Lists of genes deregulated (UP MUT or UP WT) in Sox2 deleted cells (columns A,B), and lists of genes belonging to the different classes defined by overlap of UP WT or UP MUT genes and genes expressed in BG (UP P6, UP P30, or UP BOTH), and/or SOX2-bound genes (SOX2). Genes of known potential relevance for ataxia are in bold.

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TOCI 56x50mm (300 x 300 DPI)

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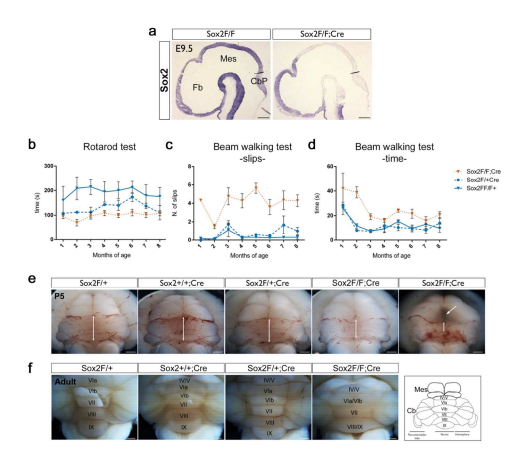


Figure 1 160x136mm (300 x 300 DPI)

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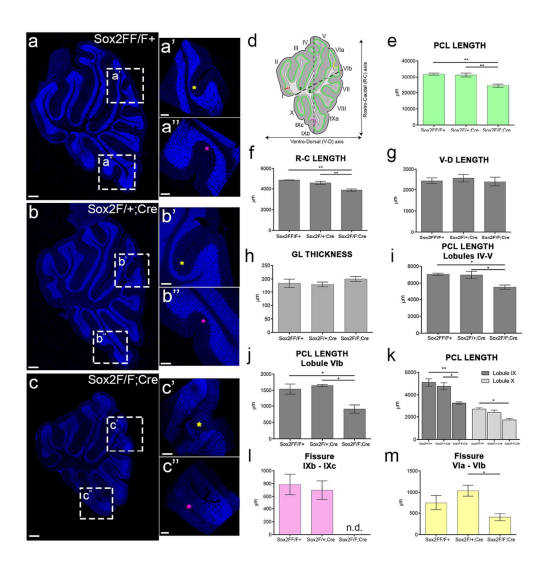


Figure 2 109x117mm (300 x 300 DPI)

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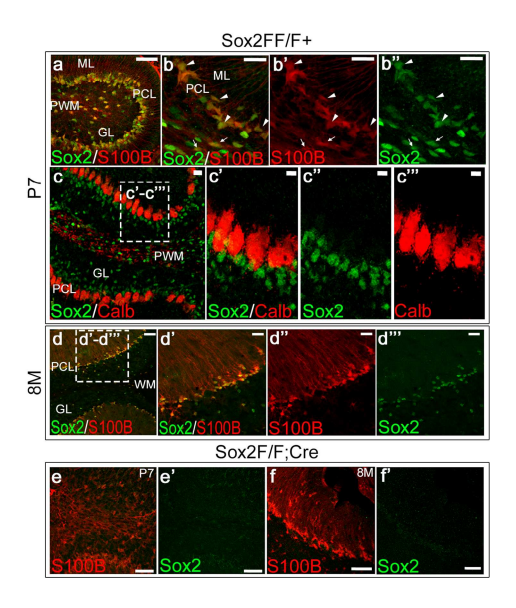


Figure 3 115x129mm (300 x 300 DPI)

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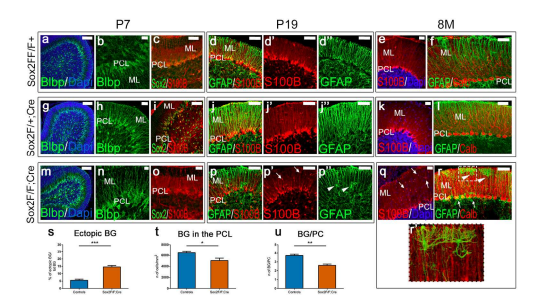


Figure 4 172x98mm (300 x 300 DPI)

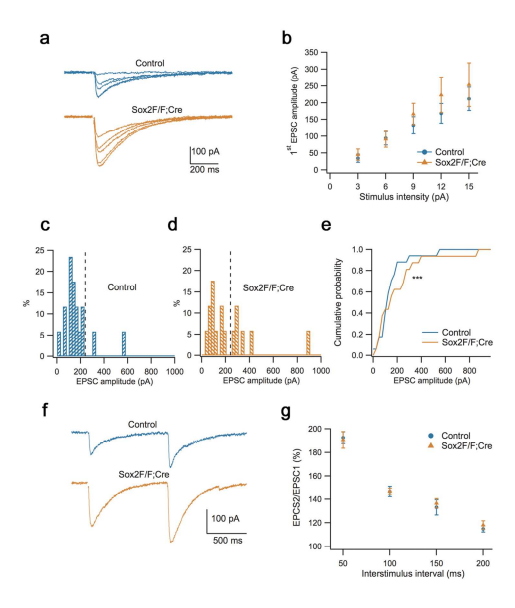


Figure 5 115x132mm (300 x 300 DPI)

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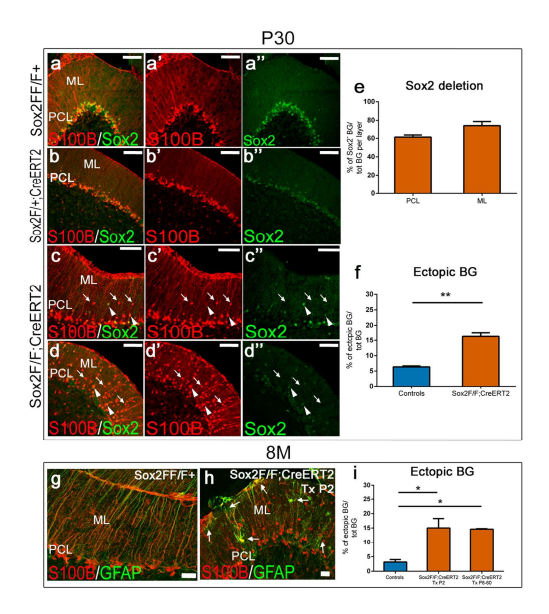


Figure 6 115x131mm (300 x 300 DPI)

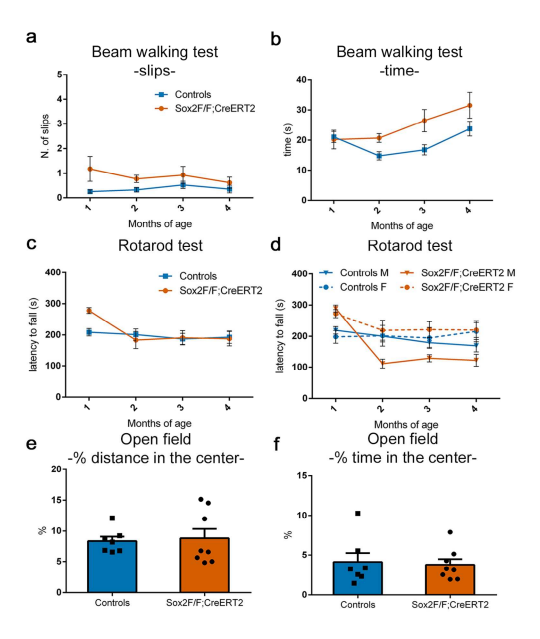


Figure 7 113x134mm (300 x 300 DPI)

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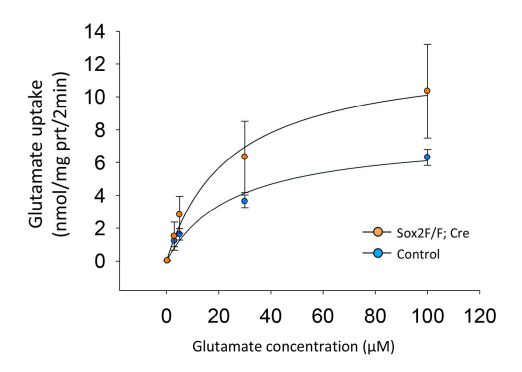
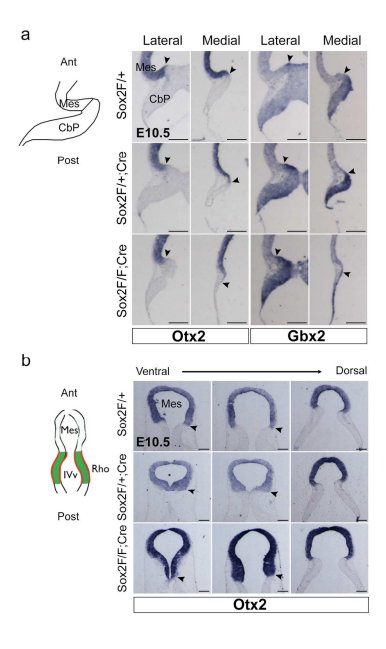
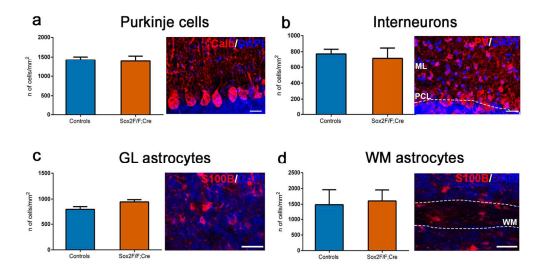


Figure 8 149x109mm (300 x 300 DPI)

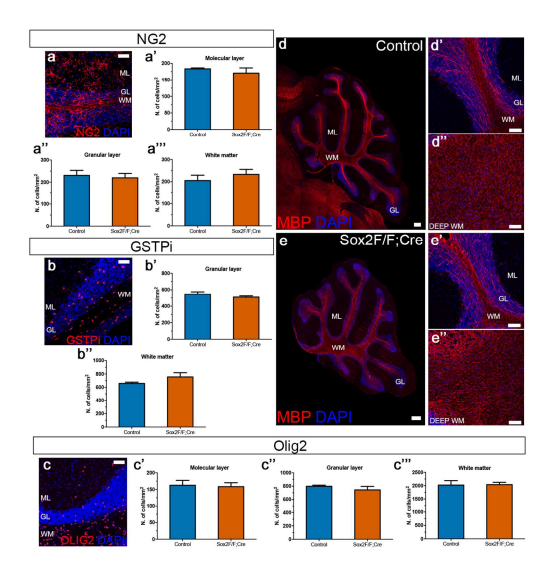


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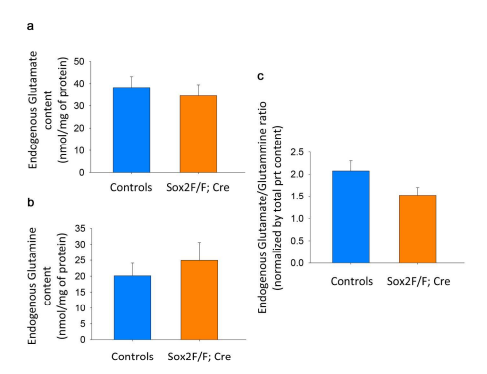
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128x66mm (300 x 300 DPI)



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199x169mm (300 x 300 DPI)

Figure	Applied Test	P value	F/t value	Post hoc analyses	Post hoc results
Fig. 1, rotarod Sox2FF/F+, n=4 Sox2F/+;Cre, n=3 Sox2F/F;Cre, n=10	Two-way Anova	Genotype: P<0.0001	Genotype (2, 14)= 70.54	Bonferroni's Multiple Comparison Test	Sox2F/F;Cre vs Sox2FF/F+: P<0.001 2,3 months P<0.05 4,7 months P<0.001 5,6 months Sox2F/+;Cre vs Sox2F/F;Cre P<0.001 2,3 months
Fig. 1, beam (slips) Sox2FF/F+, n=4 Sox2F/+;Cre, n=3 Sox2F/F;Cre, n=10	Two-way Anova	Genotype: P<0.0001	Genotype F(2,14)= 73.49	Bonferroni's Multiple Comparison Test	Sox2F/F;Cre vs Sox2FF/F+: P<0.01 1,4,7,8 months P<0.05 3,6 months P<0.001 5 months Sox2F/F;Cre vs Sox2F/+;Cre: P<0.01 1,4 months P<0.05 6,8 months P<0.001 5 months
Fig. 1, beam (time) Sox2FF/F+, n=4 Sox2F/+;Cre, n=3 Sox2F/F;Cre, n=10	Two-way Anova	Genotype: P<0.0001 Time: P=0.002	Genotype F(2,14)= 18.40 Time F(7,14)= 6.16	Bonferroni's Multiple Comparison Test	Sox2F/F;Cre vs Sox2FF/F+: and Sox2F/F;Cre vs Sox2F/+;Cre: P<0.01 2 months
Fig. 2e Sox2FF/F+, n=3 Sox2F/+;Cre, n=3 Sox2F/F;Cre, n=3	One-way Anova	Main effect of genotype, P=0.0026	(2, 6) = 18.93	Bonferroni's Multiple Comparison Test	Sox2FF/F+ vs. Sox2F/F;Cre, P=0.0045 Sox2F/+;Cre vs. Sox2F/F;Cre, P=0.0065
Fig. 2f Sox2FF/F+, n=3 Sox2F/+;Cre, n=3 Sox2F/F;Cre, n=3	One-way Anova	Main effect of genotype, P=0.014	(2, 6) = 24.14	Bonferroni's Multiple Comparison Test	Sox2FF/F+ vs. Sox2F/F;Cre, P=0.016 Sox2F/+;Cre vs. Sox2F/F;Cre, P=0.0078
Fig. 2i Lobule IV-V Sox2FF/F+, n=3 Sox2F/+;Cre, n=3 Sox2F/F;Cre, n=3	One-way Anova	Main effect of genotype, P=0.0195	(2, 6) = 8.137	Bonferroni's Multiple Comparison Test	Sox2FF/F+ vs. Sox2F/F;Cre, P=0.0344 Sox2F/+;Cre vs. Sox2F/F;Cre, P=0.0442
Fig. 2j Lobule VIb Sox2FF/F+, n=3 Sox2F/+;Cre, n=3 Sox2F/F;Cre, n=3	One-way Anova	Main effect of genotype, P=0.0084	(2, 6) = 11.76	Bonferroni's Multiple Comparison Test	Sox2FF/F+ vs. Sox2F/F;Cre, P=0.0259 Sox2F/+;Cre vs. Sox2F/F;Cre, P=0.0125
Fig. 2k Sox2FF/F+, n=3 Sox2F/+;Cre, n=3 Sox2F/F;Cre, n=3	One-way Anova	IX, Main effect of genotype, P=0.0055 X, Main effect of genotype, P=0.0124	IX, (2, 6) = 14.02 X, (2,6)=9.965	Bonferroni's Multiple Comprison Test	IX, Sox2FF/F+ vs. Sox2F/F;Cre, P=0.0076 Sox2F/+;Cre vs. Sox2F/F;Cre, P=0.0196 X, Sox2FF/F+ vs. Sox2F/F;Cre, P=0.00144
Fig. 2n VIa-VIb Sox2FF/F+, n=3 Sox2F/+;Cre, n=3 Sox2F/F;Cre, n=3	One-way Anova	Main effect of genotype, P=0.0411	F(2,6)=5.693	Bonferroni's Multiple Comprison Test	Sox2F/+;Cre vs. Sox2F/F;Cre, P=0.0452
Fig. 4s BG ectopia 8 months Sox2F/F;Cre, n= 5 Sox2F/+;Cre, n= 2 Sox2FF/F+, n=3	Student T test two tailed	P<0.0001	t(8)=7.338		
Fig. 4t BG density in PCL mutants: Sox2F/F;Cre, n=3 controls: Sox2F/+;Cre, n=3	Student T test two tailed	P = 0.0179	t(7)=3.075		

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Sox2FF/F+, n=3					
Fig. 4u BG/PCL mutants: n= 3 Sox2F/F;Cre controls: n= 2 Sox2F/+;Cre n=3, Sox2FF/F+	Student T test two tailed	P = 0.0014	t(7)=5.080		
Fig. 5e Electrophysiology Sox2F/F;Cre, cells, n =16 Sox2FF/F+, cells, n = 17	Kolmogorov- Smirnov test	P < 0.0001	D = 0.68		
Fig. 6k BG ectopia Tx P2, 30 days survival mutants: Sox2F/F;CreERT2, n=3 controls: Sox2F/+;CreERT2, n=1, Sox2FF or Sox2F/+, n=2	Student T test two tailed	P = 0,0011	t(4)=8.478		
Fig. 6n ectopia Tx P2 vs Tx P8-60 vs controls, 8 months survival mutants: Sox2F/F;CreERT2-P2, n=4 Sox2F/F;CreERT2- P8/60, n=3 controls: Sox2F/+;CreERT2-P2, n=2, Sox2FF or Sox2F/+P2, n=2	One way Anova	P = 0,0066	Genotype (2, 8)= 10.03	Bonferroni's Multiple Comparison Test	Sox2F/F;CreERT2-P2 vs controls, P=0.0114 Sox2F/F;CreERT2-P8/60 vs controls, P=0.0209
Fig. 7a beam (slips), both sexes Sox2F/F;CreERT2, TxP8-60, n=9 Sox2F/F;CreERT2, no Tx, n=14	Two-way RM Anova	Main effect of genotype, P<0.0454	F(1,21)=4.526	Bonferroni's Multiple Comparison Test	month 1, P=0.0128
Fig. 7b beam (time) Sox2F/F;CreERT2, TxP8-60, n=9 Sox2F/F;CreERT2, no Tx, n=14	Two-way RM Anova	Main effect of genotype, P<0.0333 Main effect of time, P<0.0001	F _{senotype} (1,21)=5.192 F _{time} (3,63)=8.330	Bonferroni's Multiple Comparison Test	Genotype, month 3, P=0.0288 Time, controls: month 2 vs 4, P=0.0050; month3 vs 4, P=0.049 mutants: month 1 vs 4, P=0.0042; month 2 vs 4, P=0.0069
Fig. 7d Rotarod, males Sox2F/F;CreERT2, TxP8-60, n=3 Sox2F/F;CreERT2, no Tx, n=7	Two-way RM Anova Student T test	Main effect of interaction genotype x time, P<0.0001	F(3,24)=17.05 t(8)=-3.195	Bonferroni's Multiple Comparison Test	month 2, P=0.0131
Fig. 8 Vmax	two tailed	P=0.019	ι(ο)3.193		

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Supplementary Table 1. Details of statistical analyses. In most cases only comparisons leading to P<0.05 are reported; n = number of animals (unless differently stated). RM, repeated measures.

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Dataset	Time	Genotype	Mean ± SEM	n, animals	Student t test
0/ of optomic DC	8 months	Sox2F/+;Cre	5.179 ± 1.298	2	P=0.6445
% of ectopic BG	8 months				t(3)=0.5111
		Sox2FF/F+	6.081 ± 0.7485	3	
BG density in PCL	8 months	Sox2F/+;Cre	6141 ± 396.4	3	P=0.2233 t(4)=1440
		Sox2FF/F+	6781 ± 201.1	3	
BG/PC	8 months	Sox2F/+;Cre	3.518 ± 0.1780	3	P=0.1252 t(4)=1.935
		Sox2FF/F+	3.939 ± 0.1256	3	
% of ectopic BG Tx P2	P30	Sox2FF/F+	6.215 ± 0.4674	2	n/a
		Sox2F/+;CreERT2	6.63	1	
% of ectopic BG Tx P2	8 months	Sox2FF/F+	2.118 ± 0. 6345	2	P=0.2992 t(2)=1.389
		Sox2F/+;CreERT2	4.207 ± 1.363	2	
PCs density	8 months	Sox2FF/F+	1423 ± 56,26	3	P=0.9798 t(3)=0.02749
		Sox2F/+;Cre	$1428 \pm 222,2$	2	
ML interneurons density	8 months	Sox2FF/F+	859,7 ± 71,81	2	P=0.1412 t(2)=2.370
		Sox2F/+;Cre	$685,7 \pm 15,35$	2	
GL astrocytes density	P19	Sox2FF/F+	821,4 ± 84,98	2	n/a
		Sox2F/+;Cre	761,1	1	
WM astrocytes density	P19	Sox2FF/F+	$1743 \pm 692,5$	2	n/a
		Sox2F/+;Cre	974,8	1	

Supplementary Table 2. Average numerical values obtained in immunofluorescence experiments, and statistical analysis.