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Sodium Valproate Exerts Neuroprotective Effects In Vivo through CREB-Binding Protein-Dependent Mechanisms But Does Not Improve Survival in an Amyotrophic Lateral Sclerosis Mouse Model

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Amyotrophic lateral sclerosis (ALS) is characterized by motoneuron (MN) degeneration, generalized weakness, and muscle atrophy. The premature death of MNs is thought to be a determinant in the onset of this disease. In a transgenic mouse model of ALS expressing the G86R mutant superoxide dismutase 1 (mSOD1), we demonstrated previously that CREB (cAMP response element-binding protein)-binding protein (CBP) and histone acetylation levels were specifically decreased in nuclei of degenerating MNs. We show here that oxidative stress and mSOD1 overexpression can both impinge on CBP levels by transcriptional repression, in an MN-derived cell line. Histone deacetylase inhibitor (HDACi) treatment was able to reset proper acetylation levels and displayed an efficient neuroprotective capacity against oxidative stress in vitro. Interestingly, HDACi also upregulated CBP transcriptional expression in MNs. Moreover, when injected to G86R mice in vivo, the HDACi sodium valproate (VPA) maintained normal acetylation levels in the spinal cord, efficiently restored CBP levels in MNs, and significantly prevented MN death in these animals. However, despite neuroprotection, mean survival of treated animals was not significantly improved (≤5%), and they died presenting the classical ALS symptoms. VPA was not able to prevent disruption of neuromuscular junctions, although it slightly delayed the onset of motor decline and retarded muscular atrophy to some extent. Together, these data show that neuroprotection can improve disease onset, but clearly provide evidence that one can uncouple MN survival from whole-animal survival and point to the neuromuscular junction perturbation as a primary event of ALS onset.

Key words: sodium valproate; HDAC inhibitor; CREB-binding protein; motor dysfunction; neuroprotection; acetylation

Introduction

Amyotrophic lateral sclerosis (ALS) is a fatal neurodegenerative disease that affects upper and lower motoneurons (MNs), leading to progressive muscle wasting, paralysis, and death within 2–5 years of diagnosis (Pasinelli and Brown, 2006). Although most of ALS cases occur sporadically, ~10% are inherited. Among these, 10–20% are attributable to mutations in the gene encoding the superoxide dismutase-1 (SOD1), one of the main free-radical scavenging enzymes that protect cells against oxidative stress (Rosen et al., 1993; Andersen et al., 2003). Different transgenic mice bearing mutations on the sod1 gene have been developed as ALS models, among which is the G86R mouse strain. These mice bear the equivalent G85R human mutation on the mouse sod1 gene. They die within ~120 d, presenting typical ALS symptoms: motoneuronal death, muscular atrophy, and paralysis (Ripps et al., 1995; Dupuis et al., 2000).

Premature death of MNs is believed to be determinant in the onset of ALS. However, the primary event leading to pathogenicity is still a matter of debate. All cell types implicated in the motor unit could be potential targets of toxicity (i.e., MNs, myocytes, astrocytes, microglia, and Schwann cells). Cell-specific knock-in or knock-down studies of the mutant sod1 gene have not so far brought a clear answer to that question (Gong et al., 2000; Boillée et al., 2006; Miller et al., 2006). Nevertheless, retraction of motor axon from synaptic connection to muscle and fragmentation of the neuromuscular junction are detected before any MN death, suggesting that ALS could be seen as a “dying-back” axonopathy (Fischer et al., 2004; Jokic et al., 2006). We have also shown that muscular metabolic defects and mitochondrial dysfunctions were early symptoms of the pathology (Dupuis et al., 2003, 2004). Furthermore, this idea that MN death could be dissociated from motor dysfunction has been recently strengthened by a study in...
which MN death was genetically prevented by Bax deletion (Gould et al., 2006).

Our recent work demonstrated that the cAMP response element-binding protein (CREB)-binding protein (CBP), a transcriptional coactivator displaying histone acetyltransferase (HAT) activity (Bannister and Kouzarides, 1996), is specifically lost in lumbar spinal cord MNs of G86R mice (Rouaux et al., 2003). Together with histone deacetylases (HDACs), HATs control gene transcription by finely tuning acetylation homeostasis (Verdene et al., 2005). Consistent with this, we also found decreased histone acetylation levels in the degenerating MNs (Rouaux et al., 2003). Based on these findings, we undertook to correct aberrant gene transcription by interfering with the HAT/HDAC balance with HDAC inhibitors (HDACis) to prevent neuronal death in vivo, hypothesizing that maintaining proper acetylation levels would preserve the transcription of neuroprotective genes. Indeed, several studies have reported the beneficial effects of HDACis on different aspects of neurodegeneration (Langley et al., 2005). Here, we tested the effect of a chronic treatment of G86R mice with sodium valproate (VPA). We provide an in vitro and in vivo proof of concept that VPA is a potent neuroprotective molecule but found that the distal pathology progresses despite neuroprotection.

Materials and Methods

**HDAC inhibitors and antibodies.** VPA, sodium butyrate (NaBu), trichostatin A (TSA), and α-bungarotoxin (α-BGT) were purchased from Sigma (St. Louis, MO). Anti-CBP, anti-acetylated histone 3 (AcH3), and anti-p75 antibodies were obtained from Millipore (Billerica, MA). Anti-human synaptophysin antibody was obtained from Dako (Trappes, France). HRP-conjugated goat-anti-rabbit antibody was from Jackson Immunoresearch (West Grove, PA). Alexa Fluor 488 donkey-anti-rabbit antibody was from Invitrogen (Eugene, OR). Goat anti-choline acetyltransferase (ChAT) and biotinylated goat-anti-rabbit antibody were from Millipore.

**NSC34 cell culture.** NSC34 cell line was kindly provided by Dr. Cashman (Cashman et al., 1992). Cells were seeded at a density of 1 × 10⁶ cells/ml in 96-well plates for cell survival measurements, 24-well plates for transfection experiments, 12-well plates for Western blot assays, and six-well plates for quantitative PCR (Q-PCR) analysis. Cells were grown in DMEM (Invitrogen) supplemented with 10% fetal bovine serum (Invitrogen), 100 U/ml penicillin, and 100 µg/ml streptomycin (Eurobio, Les Ulis, France) at 37°C and 5% CO₂. After 24 h, cells were serum starved for 12 h before being exposed or not to a 15 min pulse of hydrogen peroxide (H₂O₂) at a final concentration in the millimolar range. The HDAC inhibitors tested were added to media during and after the H₂O₂ pulse. MTT [3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl-2-H-tetrazolium bromide] assays (cell survival) were performed on 12 h, and lysed in a Promega (Madison, WI) lysis buffer. Luciferase activity was read in a single tube luminometer (Berthold Detection Systems, Pforzheim, Germany). Expression vectors used for cotransfection experiments are pRC-CMV wild-type (WT)-SOD1 and pRC-CMV G86R-SOD1. One microgram of each expression vector and 0.5 µg of reporter vector were used. Luciferase activity was read 20 h after transfection.

**Western blot analysis.** Western blots were performed as described previously (Rouaux et al., 2003) with typically 50 µg of total cell extracts run on 7, 10, or 13% SDS-acrylamide gels for CBP, actin, or histone detection, respectively. Specific bands were detected by enhanced chemiluminescence (GE Healthcare, Buckinghamshire, UK). Size and intensity of specific bands were quantified using an NIH Image 1.62 software to allow the representation of the results as histograms.

**Quantitative and classical (reverse transcription)-PCR analysis.** Total RNA was extracted from animal tissues or NSC34 cells using Trizol (Invitrogen) according to the manufacturer’s instructions. cDNA syntheses were performed using 1 µg of total RNA (iScript CDNA Synthesis kit; Bio-Rad, Hercules, CA) with the following primers: cbp, forward, 5'-ggg gta acc agc cag ctt ta-3'; reverse, 5'-cag gga cat tgt gtt aat tga-3'; bcl2, forward, 5'-ctg cca atg ctc gac taa-3'; reverse, 5'-tct act tcc tcc gca atg ct-3'; smm, forward, 5'-aag gca cag cca gaa aaa-3'; reverse, 5'-tca cag gtc ggg gaa agt ag-3'; 18S, forward, 5'-ctg cct cac tat ccc ttc-3'; reverse, 5'-tct ctt gga tgt ggt agc cg-3'.

Q-PCR analysis was performed on iCycler System (Bio-Rad). The reactions were performed in 25 µl, in 96-well plates, using iQSYBR Green Supermix (Bio-Rad). A specific standard curve for each gene was performed in parallel to the analysis. Each sample was analyzed in duplicate. PCR conditions were 3 min at 94°C, followed by 40 cycles of 45 s at 94°C and 10 s at 60°C. Results were analyzed by using the iCycler software (Bio-Rad) and normalized for the reference gene encoding the 18S ribosomal subunit. Semiquantitative reverse transcription (RT)-PCR analysis was performed as described previously (Dupuis et al., 2004).

**Animals.** Transgenic male mice with the G86R murine SMN1 mutation (Ripps et al., 1995) were obtained from the animal facility at the University of Pirmasens, Pirmasens, Germany. Heterozygous mice were bred with female littermates, and transgenic animals were genotyped by PCR amplification of DNA extracted from tail tissue. Nontransgenic age-matched male littermates served as control. Mice received water and regular rodent chow ad libitum. HDAC inhibitors tested were diluted in 0.09% NaCl [vehicle (VEH)], and administration was performed intraperitoneally. The VPA dose to use was evaluated on 16 adult WT mice. Chronic injections were performed daily, from 60 to 105 d of age. Three doses of VPA were tested: 150, 250, and 400 mg·kg⁻¹·d⁻¹. Acute injections of TSA (2 mg/kg), VPA (250 mg/kg), NaBu (640 mg/kg), or VEH were performed on symptomatic G86R (n = 6 in each group). Six WT littermates were used as control. All animals were killed by decapitation. Animal manipulations followed current European Union regulations.

**Survival and clinical assessment.** Clinical assessment was estimated daily, according to a clinical rating scale going from score 4 to 0. Score 4 is attributed to asymptomatic G86R mice, relative to their WT littermates. Score 3 corresponds to an alteration in hindlimb extension when the animal is hung by the tail. Appearance of score 3 is considered the disease onset. Score 2 is attributed when any slight alteration in the locomotion is observed. Score 1 is assigned when a complete paralysis of the limbs. Score 0 corresponds to the stage at which animals are unable to roll over within 10 s after being pushed on their back. Mice were killed when they reached score 0.

**Weight assessment and motor function testing.** Mice were weighed twice a week. Grip strength was assessed on forelimbs, twice a week, using a grip strength meter (Bioseb, Chaville, France). Measurements were performed from 60 d of age to the end of mouse life. Animals were placed over a metallic grid that they instinctively grabbed to try to stop the involuntary backward movement performed by the manipulator until the pulling force overcame their grip strength. The strength meter scored the peak pull force. Three measurements were collected at each time, and the best one was retained.

**Tissue preparation.** Lumbar spinal cords, cerebellum, and soleus and gastrocnemius muscles were carefully dissected from G86R or WT mice and immediately frozen in liquid nitrogen for additional biochemical analysis or postfixed for 24 h in 4% paraformaldehyde (PFA) and cryoprotected in 20% sucrose during 24 h at 4°C for immunohistochemistry.

**Histone acetylation measurement.** Tissue lysates from spinal cord or cerebellum were obtained by homogenizing each sample in 300 µl of ice-cold lysis buffer for 30 s at 30 Hz in a Qiagen (Haan, Germany) Tissuelyser. and histones were acid extracted as described by Rouaux et al. (2003). Five micrograms of purified histones were separated on 13% polyacrylamide gels and further processed for Western blot analysis.

Rouaux et al. • Uncoupling of Neuroprotection and Motor Dysfunction in ALS
**Immunohistochemistry.** Serial lumbar spinal cord tissue sections from L3–L5 spinal cord segments were used for motoneuronal analysis. The sections were mounted onto gelatin-coated slides and kept at −80°C. Briefly, slides were dried for 45 min on a hot plate (70°C), rehydrated for 10 min in PBS, and fixed for 10 min with 4% PFA. For CBP and AcH3 immunostaining, endogenous peroxidases were inactivated by 10 min of incubation in 1% H2O2/PBS. Antigenic sites were then uncovered in a citrate buffer (1.8 mm citric acid and 8 mm sodium citrate), warmed for 10 min in a microwave oven, and cooled down on ice. Sections were permeabilized for 10 min in 1% Triton X-100/PBS before being blocked for 30 min in 5% goat serum/0.1% Triton X-100/PBS. Primary antibodies were applied overnight, and secondary antibodies were applied for 1 h. Revelation was performed using the VectaStain ABC kit (Vector Laboratories, Burlingame, CA). For ChAT immunostaining, the method was adapted to a FITC-conjugated secondary antibody. Pictures were taken with a Nikon (Tokyo, Japan) digital camera DXM 1200. ChAT-positive cells were counted in the ventral horn from six spinal cord sections per animal as described previously (Dupuis et al., 2004). For counting innervated synapses, 40-μm-thick longitudinal sections of PFA-fixed, sucrose-embedded, and frozen soleus muscles were cut and stained with antibodies against synaptophysin and α-BGT. The same protocol was used for p75 immunostaining. Usually 5–6 sections were analyzed per animal, which represents 100–150 nicotinic acetylcholine receptor (nAChR) clusters. Immunostaining was analyzed confocally (Laser Scanning System 510; Zeiss, Oberkochen, Germany).

**Myelinated axon area measurements.** Lumbar spinal cords were dissected from animals and postfixed in 4% PFA for 48 h. The fourth lumbar ventral roots (L4 VRs) were dissected and stained by 1% osmium tetroxide (H9262) and 0.5 mg/kg xylazine. A monopolar needle electrode (diameter, 0.3 mm; 9013S0011; Medtronic). Gastrocnemius muscles were monitored on both sides with a concentric needle electrode (diameter, 0.3 mm; 9013S0011; Medtronic). Gastronemius muscles were transversally cryosectioned (10 μm) and stained with 0.1% toluidine blue, and fiber areas were measured with the LuciaG software in two square fields. At least 40 fibers were analyzed in each field.

**Carnitine assay.** Total plasma t-carnitine concentrations were determined by a spectrophotometric enzymatic assay using carnitine acetyltransferase (CAT; EC 2.3.1.7) and 5,5-dithiobis (2-nitrobenzoic acid) (DTNB) as thiol group color reagent. Total carnitine was quantified after deesterification by alkaline hydrolysis as described previously (Wan and Hubbard, 1998). Diluted sample (50 μl) was mixed with 200 μl of the primary reagent (200 μM DTNB and 600 μM acetyl-CoA in 200 μM phosphate buffer, pH 7.5), and the reaction was started with 25 μl of diluted start reagent (CAT from pigeon muscle, 123 U/mg protein) to give a final activity of 1.53 U/sample. All specific reagents used were from Sigma. The reaction mixture was incubated for 15 min at 37°C, and absorbance was read at 415 nm. Calibrators containing 10–100 μM t-carnitine were prepared by dilution of the t-carnitine stock solution. The assay was linear for carnitine concentrations up to 200 μM. All specimens were assayed in replicate in the same analytical run.

**Electrophysiological recording of spontaneous activity.** All recordings were made with a standard EMG apparatus (Dantec, Les Ulis, France) in accordance with the guidelines of the American Association of Electrodiagnostic Medicine. Mice were anesthetized with 1 mg/kg ketamine hydrochloride and 0.5 mg/kg xylazine. A monopolar needle electrode (diameter, 0.3 mm; 9013R0312; Medtronic, Minneapolis, MN) was inserted into the tail of the mouse to ground the system. Recordings were made with a concentric needle electrode (diameter, 0.3 mm; 9013S0011; Medtronic). Gastronemius muscles were monitored on both sides (right and left) for at least 2 min. Only spontaneous activity with a peak-to-peak amplitude of at least 50 μV was considered to be significant. Fibrillations were graded from 0 to 4: no fibrillations (0), persistent single small strains of potentials (longer than 2–3 s) in at least two areas of the muscle (1), moderate number of potentials in three or more areas (2), many potentials in all areas (3), or full interference pattern of potentials (4).

**Statistical analysis.** Data are expressed as the mean ± SEM. All statistical analysis were performed on PRISM version 4.0b software (GraphPad, San Diego), using an ANOVA followed by the post hoc Newman–Keuls multiple-comparisons test for classical analysis, a two-way ANOVA followed by a Bonferroni post-test for the grip strength analysis or by means of Kaplan–Meier curves for survival, score onsets, and weight loss onset.

**Results**

**HDACi protects from oxidative stress-induced MN death**

Oxidative stress is described as one of the possible major causes of motoneuronal death in ALS (Liu et al., 1998). To test whether HDACi treatment could be used as a therapeutic approach to prevent MN death in ALS mice, we first used a simplified cellular model, the NSC34 cell line, a hybridoma of spinal cord motoneurons and neuroblastoma (Cashman et al., 1992), in which cell death can be induced by oxidative stress (Cookson et al., 1998). Treated with a pulse of H2O2, NSC34 cells died in a dose-dependent manner (Fig. 1A). Interestingly, death was accompanied by a progressive decrease of both CBP protein and histone H3 acetylation levels (Fig. 1B), two features that we already observed in MNS of symptomatic SOD1(G86R) mice (Rouaux et al., 2003). Several HDACis have been reported to promote cell survival both in vitro and in vivo, and we tested three HDACis (VPA, TSA, and NaBu) for their ability to counteract H2O2-induced NSC34 cell death. All of the three HDACis were able to reverse oxidative stress-induced histone deacetylation in a dose-dependent manner (Fig. 1C). In addition, HDACi treatment could also rescue NSC34 cells from death, with different efficiencies (Fig. 1D). Altogether, these results show that HDACis efficiently restored histone acetylation and counteracted cell death in a cellular model that mimics oxidative stress-induced motoneuronal death.

**Oxidative stress and mutant SOD1 induce transcriptional repression of the cbp gene, which can be upregulated by HDACi treatment**

Taking advantage of this homogenous cellular model of MNs (in contrast to the whole spinal cord), we further investigated which mechanism could underlie CBP downregulation under oxidative stress conditions. Figure 2A shows that a 15 min pulse of H2O2 induced a transcriptional repression of the cbp gene within 4 h. The same result was observed with a reporter luciferase gene driven by the cbp promoter (Fig. 2B). Interestingly, overexpression of SOD1(G86R) strongly repressed cbp promoter activity, whereas the WT SOD1 had no effect (Fig. 2C). This result suggests that expression of SOD1(G86R) in ALS transgenic animals could intrinsically exert a transcriptional repression of cbp gene in the lumbar spinal cord, thus triggering MN death. By blocking histone deacetylation, HDACis act as transcriptional activators; therefore, we next tested whether they could directly induce cbp gene expression. Figure 2D shows that HDACis were able to increase cbp promoter-driven luciferase activity in a dose-dependent manner, both under control and oxidative stress conditions. Except for NaBu, the efficient doses in reversing oxidative stress-induced cbp transcriptional repression (1 mM VPA and 10 mM TSA) corresponded to those exhibiting neuroprotective effects (Fig. 1D).

**Chronic VPA treatment in vivo prevents histone deacetylation in the spinal cord of symptomatic ALS mice**

We then undertook to treat ALS mice with an HDACi to see whether protecting MNs could provide beneficial effects in the pathology. We chose to treat mice with VPA, a Food and Drug
Chronic VPA treatment ensures neuroprotection and restores normal CBP levels in vivo

We then assessed the effects of VPA on survival of lower MNs in these mice. MNs were counted in the ventral horns of lumbar spinal cord sections from 105-d-old (score 2) (Fig. 4A, left) and end-stage (score 0) mice (right) (for clinical score assessment, see Material and Methods and Fig. 5A). We found that at 105 d of age, when MN loss was ~30% in G86R mice compared with age-matched WT mice, chronic VPA treatment completely prevented neuronal death. In the end-stage mice, MN loss was much more important (77.7%), but VPA effect was still significant, and the number of MNs was twofold higher in treated than in untreated transgenic mice (Fig. 4A). These results clearly indicate that VPA treatment efficiently delayed motoneuronal death in vivo. To get more convincing information on the extent of neuroprotection by VPA, we verified the occupancy of myelinated axons in L4 VRs from the spinal cords. Typical sections of WT and G86R mice at the onset of locomotor symptoms (score 2) are shown in Figure 4B, and focus images made on the different groups of mice (treated or not) are shown in Figure 4C. We found a reduction of ~20% of occupancy of myelinated L4 VR axons in score 2 G86R mice [day 105 (D105)] compared with WT (49.9% vs 71.5%) (Fig. 4D). Here again, the chronic VPA treatment helped in protecting from the loss of myelinated fibers at that age (15% improvement). At the end stage of the disease, the myelinated axons occupancy in VPA-treated animals was still larger than in untreated animals (~10%) (Fig. 4D), a stage at which we previously counted twice as many large lumbar MN cellular bodies (Fig. 4A). These data further support the neuroprotective role of VPA.

We also verified that chronic VPA administration had no effect on the expression of the sod1(G86R) gene by Q-PCR experiments performed on lumbar spinal cord extracts (data not shown), suggesting that the neuroprotective effect of VPA does not result from a repression of the mutated sod1 transgene.

Most importantly, we found that CBP levels were maintained in MN nuclei from VPA-treated mice (Fig. 4E), suggesting that, as seen in NSC34 cells, VPA could directly upregulate CBP at the transcriptional level. Together, these results suggest that chronic VPA administration could reverse not only histone deacetylation in MNs but also CBP protein loss, two events that could account for the neuroprotective effect mediated by VPA in vivo. Additionally, we checked by Q-PCR performed on whole lumbar spinal cord extracts whether CBP loss in G86R mice, whereas they were maintained in other cell types (Fig. 3C). In contrast, histone acetylation levels in VPA-treated transgenic mice were significantly restored to levels comparable with those of WT littermates (Fig. 3C).
We then tested the effects of chronic VPA treatment (250 mg·kg⁻¹·d⁻¹) on disease onset and survival. To this aim, we first established a rating scale representing disease progression and ranging from score 4 (no visible symptoms) to score 0 (dying mouse) (see Material and Methods). Figure 5A shows the mean ages of appearance and duration of each score, as assessed on a cohort of untreated G86R mice. Treated and untreated mice were then evaluated for their onset in each of these scores (Fig. 5B–E). Statistical analysis of Kaplan–Meier curves showed that onset of score 3 (i.e., first ALS symptoms) was significantly delayed in VPA-treated G86R mice by 10% (Fig. 5B) (p = 0.027). Onsets of scores 2 and 1 were also significantly delayed but to a lesser extent (Fig. 5C,D). Overall and despite the neuroprotective effect observed on lumbar MNs, we found no significant effect of chronic VPA treatment on the lifespan of G86R mice (110.5 ± 1.46 d for untreated animals (n = 21) vs 114.3 ± 2.34 d for VPA-treated animals (n = 15) (Fig. 5E). A lower dose of VPA (150 mg·kg⁻¹·d⁻¹) did not modify the mean lifespan of animals (108.9 ± 1.96 d for untreated animals (n = 12) vs 108.6 ± 4.48 d for VPA-treated animals (n = 9)), whereas increasing VPA concentrations to 400 mg·kg⁻¹·d⁻¹ was toxic to G86R mice (data not shown). We also conducted a survival study with another HDACi, sodium butyrate, at a dose that has previously been shown to display in vivo neuroprotective effect on a Huntington’s disease mouse model (Ferrante et al., 2003), but we found no beneficial effect on G86R mice (supplemental material 1, available at www.jneurosci.org). Together, these data show that protecting against MN death could only provide mild beneficial effects at the onset of the disease and did not prevent mice from dying with the same clinical symptoms typical of ALS. We thus evaluated more carefully different criteria of motor dysfunction in treated and untreated mice, to further analyze the lack of effect of VPA on mouse survival (more precisely, the muscular and the neuromuscular junction aspects).

**Chronic VPA administration prevents muscular alteration in mouse only at the first stages of ALS**

The muscle functions were then further evaluated, first by measuring muscle strength with the grip test. Two-way ANOVA analysis of the results showed that muscular strength of G86R mice was gradually reduced as the disease progressed (p < 0.0001) and that VPA-treated transgenic mice presented a significant increase in motor performances (p < 0.0001) compared with untreated mice (Fig. 6A). This difference was observed very early, after 2 weeks of treatment, at ~11 weeks of age, when mice did not present any motor symptoms (score 4). Altogether, these results show that VPA treatment induced a rapid and sustained increase in skeletal muscle strength. Second, we measured muscle fiber area in our group of experimental mice and found that, if the size of fibers was diminished as expected because of muscular atrophy during the course of the disease, VPA did prevent muscular atrophy at the early stage of the disease (score 2) (Fig. 6B). Nevertheless, muscle fiber area was not significantly different at the end

**Motoneuronal protection by chronic VPA treatment delays disease onset but does not increase the lifespan of transgenic ALS mice**

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stage. These results are in line with the previous observation, in that mice with muscle fibers that are preserved from atrophy by VPA also present a better score at the grip test, and underline that muscle function is protected only at the early stage of the disease.

We wanted to further check whether VPA could have some toxic side effects on mice at the dose tested in vivo. It is known that VPA decreases the body mass in rodents (Wolden-Hanson et al., 1998), and we then wondered whether the lack of a significant effect on mice survival could result from a deleterious effect of the molecule on body mass. This question is of particular importance because it is now well established that a dramatic loss of body mass is one of the symptoms of the disease in G86R mice and is related to major metabolic dysfunction (Dupuis et al., 2004). Chronic VPA treatment indeed reduced the gain of body mass of WT animals (Fig. 6C). This effect was observed as soon as 1 week after the beginning of injections and was maintained during all of the time of treatment (p < 0.0001). Surprisingly, VPA had no significant effect on the body mass of transgenic animals. Indeed, although G86R mice presented a characteristic lighter body mass than their WT littermates (p < 0.0001), no difference was found between untreated and VPA-treated G86R mice (Fig. 6C). VPA treatment had no effect on onset of body weight loss either (data not shown), suggesting that variation of mouse body mass cannot account for the lack of beneficial VPA effect on G86R mouse survival. It might be noteworthy that prevention of muscular atrophy (Fig. 6B) may have accounted for the retardation of the appearance of symptoms, at least at the early stage of the disease.

The most frequent alterations reported for VPA, in cases of acute intoxication, are hepatotoxicity and alteration of carnitine metabolism (for review, see Lheureux et al., 2005). Moreover, a recent study showed that L-carnitine treatment of SOD1(G93A) mice significantly delayed the onset of signs of disease (including deterioration of motor activity) and extended the lifespan of the mice (Kira et al., 2006). We further checked whether VPA could have some toxic side effects on animals through modulation of plasmatic carnitine levels.

**Chronic VPA treatment delays but does not prevent neuromuscular denervation**

Progressive paralysis observed in ALS results from progressive denervation of skeletal muscles (Bjornskov et al., 1984). The delay of symptom onset and the increase in muscle strength seen under VPA treatment suggest that this molecule could retard skeletal muscle denervation. To test this hypothesis, we performed electromyographic analysis on score 3 transgenic mice and their WT age-matched littermates (100 d of age). Recording of the resting activity in the gastrocnemius muscle of untreated animals was done in vivo. A, Six 90-d-old WT mice received a single injection of VEH (0.09% NaCl) or 150 or 300 mg/kg VPA. Twenty-four hours later, cerebella and spinal cords (SC) were carefully dissected and processed for acid extraction of histones. Top, Five micrograms of purified histones were then analyzed by Western blot to detect specific AcH3. Bottom, Bands were quantified. #p < 0.05 and ##p < 0.01 compared with VEH condition; **p < 0.01. B, WT mice were injected once with a dose of 250 mg/kg VPA and killed 0, 2, 6, 12, or 24 h later. Left, Spinal cords were dissected and processed for Western blot analysis. Right, Bands were quantified. #p < 0.05 compared with time 0. C, Spinal cord sections from WT (n = 5) or end-stage G86R mice daily injected with VEH (n = 5) or 250 mg/kg VPA (n = 5) were taken, and sections were immunostained using an anti-AcH3 antibody followed by a peroxidase revelation. Representative pictures are shown. Arrowheads depict lumbar motor neurons. Scale bars: top, 200 μm; bottom, 40 μm.
G86R mice (Fig. 7A, middle) consistently revealed abnormal spontaneous activity with fibre potentials graded from 3 to 4 (see Material and Methods). Numerous fascinations were also observed. In contrast, recording the resting activity in the gastrocnemius muscle of chronically VPA-treated mice showed strongly reduced abnormal spontaneous activity with fibre potentials graded from 1 to 2 and without any fascication (right). As expected, neither fibrelation nor fascication were observed in WT animals (left).

To check whether VPA could have had a direct effect on membrane potential of myocytes (independently of motor innervation), we tested the electromyographic response to treatment of mice with two doses of VPA (250 and 400 mg/kg) after a right sciatic nerve amputation. All groups of mice, whether treated or not, presented with a full interference pattern of spontaneous potentials in the right gastrocnemius muscle 3 and 8 d after amputation of the right sciatic nerve (supplemental material 2, available at www.jneurosci.org). These results suggest that treatment with VPA does not interfere with the electrical activity of normal and denervated skeletal muscle.

Thus, it seems that VPA treatment may delay muscle denervation in ALS mice, and we further counted the number of innervated synapses in muscles. To this aim, the presynaptic terminals were identified on longitudinal soleus sections using an anti-synaptophysin antibody, and nACHRs were labeled by fluorescent α-BGT binding. Figure 7B represents typical photographs obtained after confocal analysis of a WT and a score 1 G86R mouse. We found that the number of innervated neuromuscular junctions (NMJs) decreased in nontreated animals and that VPA treatment slightly helped in preventing denervation but only at the early stage of the disease (Fig. 7C).

Finally, we performed an immunohistochemical labeling against p75, which is highly expressed in both MN presynaptic terminals and terminal Schwann cells after denervation (You et al., 1997). Figure 7D shows representative sections of each type of animal (WT and G86R, treated or not, D105 and end-stage mice) photographed using confocal microscopy. Quantification is presented on Figure 7E. We observed that most of the NMJs were p75 positive in end-stage G86R animals, either treated or not. VPA significantly delayed p75 expression at D105, although p75-positive NMJs tend to be more present in

Figure 4. Chronic VPA administration delays motoneuronal death, maintains high levels of CBP protein, and upregulates CREB/CBP-dependent transcription in vivo. WT and G86R mice were chronically injected with VEH (0.09% NaCl) or VPA (250 mg · kg⁻¹ · d⁻¹) from 60 d of age until score 2 (105 d; G86R-D105) or score 0 (end stage; G86R-ES). A, Nonadjacent spinal cord sections from WT (n = 5) or transgenic mice were immunostained using an anti-ChAT antibody followed by a FITC-conjugated antibody. ChAT-positive MNs with an area >600 μm² are represented as means ± SEM per section. The counting was performed on G86R-D105 and G86R-ES, white and gray histograms correspond to untreated and VPA-treated animals, respectively. *p < 0.05 and **p < 0.0001 compared with WT conditions; #p < 0.05 compared with G86R plus VEH condition. B, Representative semithin section of an L4 ventral root from a WT mouse or a G86R-D105. Scale bars, 50 μm. C, D, Representative pictures (C) and quantification (D) of myelinated fibers occupancy in the different groups of mice (a, WT; b, G86R-D105 plus VEH; c, G86R-D105 plus VPA; d, G86R-ES plus VEH; e, G86R-ES plus VPA). Scale bar, 20 μm. D, The percentage of occupancy of myelinated axons within the L4 VR is represented for each group. The data represent means ± SEM. The number of each animal tested is noted within the histogram. Statistical analysis were performed with ANOVA followed by the Newman–Keuls multiple-comparisons test (Pgrim). All situations are compared with WT, and in each group, VPA treatment is compared with VEH. *p < 0.05; **p < 0.001. E, Representative pictures of CBP protein immunostaining in lumbar spinal cord from WT and G86R mice (score 0) chronically injected with VEH or VPA (250 mg · kg⁻¹ · d⁻¹). Arrowheads depict lumbar motor neurons. Scale bars, 40 μm. F, Expression of bcl-2 and smn were analyzed by semiquantitative RT-PCR 24 h after injection in spinal cord extracts from 105-d-old WT or G86R (score 2) that received or not an acute injection of VEH, VPA (250 mg/kg), TSA (2 mg/kg), or NaBu (640 mg/kg). Agarose gels were scanned, and specific bands were quantified. The results are represented relative to the expression levels of the house-keeping 18S gene mRNA levels. *p < 0.05 and #p < 0.001 compared with the untreated G86R condition.
D105 VPA-treated animals than in WT or WT VPA-treated mice (27 vs 8–10%).

Thus, we can conclude that chronic VPA administration modestly delays life extent and motor performance in vivo.

**Discussion**

**Modulation of cbp gene transcription by oxidative stress and HDACis**

In a previous report, we evidenced a loss of CBP in MN nuclei from G86R mice spinal cord (Rouaux et al., 2003). Herein, we show that an acute oxidative stress is able to downregulate cbp gene transcription. Loss of CBP is a recurrent phenomenon observed in different neurological disorders. Interestingly, it has been shown to occur through a diversity of mechanisms depending on the pathological context [for review, see Rouaux et al. (2004) and Saha and Pahan (2006)], but so far, cbp transcriptional repression has only been reported in a presenilin knock-out mouse model presenting increased neurodegeneration and memory alterations (Saura et al., 2004). It is thought that MN death in ALS involves oxidative stress (for review, see Simpson et al., 2003). Consistent with this, we found that overexpression of the mutant sod1(G86R) strongly repressed cbp promoter-driven luciferase activity, suggesting that chronic SOD1 dysfunction in transgenic animals and subsequent oxidative stress could be responsible for the drop of CBP levels. Remarkably, cbp was found among the downregulated genes in a gene expression profiling performed on MNs microdissected from spinal cords of ALS patients (Jiang et al., 2005).

In a therapeutic view, if cbp gene transcription were repressed in MNs, then a drug that proves efficient at reactivating its transcription (and subsequent CBP-dependent transcription) might favor neuroprotection. For a few years, because of their capacity to activate transcription by blocking deacetylation, HDACis have been tested as potential neuroprotective agents in several cellular and animal models of neurodegenerative diseases (Rouaux et al., 2004; Langley et al., 2005; Saha and Pahan, 2006). It is believed that global deacetylation will induce transcriptional repressions associated with pathologies. Such genetic alterations are observed in the spinal cord of transgenic mice and ALS patients (Dupuis et al., 2000; Ishigaki et al., 2002; Malaspina and de Belleroche, 2004; Jiang et al., 2005). Thus, inhibiting HDAC activity could remove repressive blocks from promoters, thereby providing neuroprotection. Herein, we present for the first time evidence that HDACis can combat neurodegeneration through modulation of the cbp transcriptional pathway (cbp and its targeted genes), not only in simplified cellular models in response to oxidative stress, but also in vivo in the pathological context of ALS. Interestingly, HDACi treatment did not revert neuronal apoptosis induced by electrical activity depletion (Rouaux et al., 2004), a model in which the CBP protein is degraded by posttranslational caspase-6-dependent mechanisms (Rouaux et al., 2003). Together, these results suggest that the presence of CBP is required, or at least more effective than random reactivation with HDACi treatment, in counteracting neuronal death, either because its other functions (molecular scaffolding and transcriptional coactivator) are needed or because HDAC inhibition is not specific to CBP-dependent genes.
VPA delays MN death but does not significantly increase the mean survival of ALS mice

The capacity of HDACis to modulate SOD1(G86R) mice lifespan has recently been investigated in several other studies. Sugai et al. (2004) showed that a chronic preonset VPA treatment significantly increased mice survival of SOD1(G93A) mice. Our results thus contrast with this study, although both reports showed a neuroprotective effect of VPA in vivo. However, some differences are found in the experimental design between studies. Whereas we treated G86R mice at 250 mg·kg⁻¹·d⁻¹ intraperitoneally, Sugai and coworkers administrated VPA in drinking water at 0.26% w/v, which corresponds to approximately twice our dose (530 mg·kg⁻¹·d⁻¹). It is thus possible that at a higher dose, VPA may have affected other genes and cellular pathways. Indeed, gene-profiling experiments showed that different doses of VPA induced different panels of genes (Massa et al., 2005), so they might also have different therapeutic effects. However, in our hands, a higher dose of VPA (400 mg·kg⁻¹·d⁻¹) injected intraperitoneally was lethal to the G86R mice population. It is noteworthy that not only the mode of injection is different in the two studies, but also the mouse strains used (i.e., SOD1(G93A) vs SOD1(G86R)), which do not display the same pathological time course and may not have the same sensitivity/metabolism for VPA. Nevertheless, VPA therapy was also established as ineffective in a retest performed on the G93A mice strain (n = 18 nontreated and n = 19 treated animals), given at preonset of disease (60 d of age) by subcutaneous pump [131.6 d control group vs 125.9 d VPA-treated group (Scott et al., 2007) (see also details on http://www.als.net/)]. Overall, the only currently available FDA-validated drug to treat ALS is riluzole (Bensimon et al., 1994). In the ALS mouse model, this molecule has been found to increase survival by 10% (Gurney et al., 1996, 1998), a threshold that can be considered clinically significant. In this respect, our data are obtained with a sufficient number of treated mice (n = 15) to assess with a power of >80% that we can reach 10% survival improvement. As we only observed 5% (not statistically significant) increase in lifespan, we concluded that VPA, although protecting MNs efficiently, does not display clinically significant improvement in survival.

Nevertheless, it has been shown recently that treatment with another HDACi, phenyl butyrate (PBA), resulted in significant enhancement of survival and improvement in clinical and neuropathological phenotypes of the SOD1(G93A) mice (Ryu et al., 2005; Petri et al., 2006). PBA was shown to increase histone acetylation (Petri et al., 2006), block apoptotic signaling, upregulate bcl-2 gene (Ryu et al., 2005), and increase MN survival, life extent, and motor performances in vivo (Ryu et al., 2005; Petri et al., 2006). Although here again, the strategy was aimed at resetting global histone acetylation levels, the 22% survival improvement under PBA treatment observed by Ryu et al. (2005) was associated with a significant increase in body weight (~20%), which is a major difference with our study, in which we observed that VPA had no effect on diseased animal body weight. This discrepancy seems of prime interest, because we previously showed that increasing adiposity and body weight of G86R mice by feeding them with a high-energetic diet significantly increased
their survival (Dupuis et al., 2004). Thus, the lack of beneficial effect of VPA on energy deficit of G86R mice may be an explanation why it failed to improve mice survival although an efficient neuroprotection was observed.

Overall, these data indicate the poor efficiency of HDACi treatment for ALS therapy despite the neuroprotective efficiency of HDACiS (see below). The recent results obtained in other laboratories with HDACiS (i.e., PBA) may be reconsidered in light of these studies, as they may have revealed beneficial effects on ALS mice by other means than only a strict neuroprotection.

Motor dysfunctions are not counteracted by VPA treatment in ALS mice

It is noteworthy that, despite neuroprotection, VPA-treated animals died with the neuropathological criteria typical of ALS. Interestingly, several recent studies question the origin of ALS, suggesting that biochemical and molecular events occurring in the muscle fibers (such as hypermetabolism, oxidative stress, and mitochondrial dysfunction) could increase NMJ integrity and spinal MN vulnerability (for review, see Gonzalez de Aguilar et al., 2007). It is thus conceivable that protecting against MN death might not be sufficient to cure ALS. We found that preonset treatment with VPA delayed disease onset in G86R mice and induced a significant amelioration of muscle strength, as well as decreased muscle denervation (EMG studies). This is reminiscent of the fact that the MNs as well as their proximal axons were spared in the treated animals: protecting MN from death may have retarded the disease onset. Recently, in a series of elegant experiments using genetic elimination of programmed cell death in SOD1 mice by crossings with Bax knock-out mutants, Gould et al. (2006) provided evidence that damage to the distal motor axon was an event dissociated from MN death. More recently, Dewil et al. (2007) found that the antiapoptotic effect of inhibition of p38 MAPK (mitogen-activated protein kinase) activation rescued MNs and to a lesser degree the proximal axon but that the distal denervation still progressed. In fact, our experiments aimed at counting the number of innervated motor synapses reached the same conclusions: NMJs were already damaged at D105 in VPA-treated animals, whereas the number of MNs was similar to that of WT animals. Moreover, we found that the low-affinity neurotrophin receptor p75 was highly expressed in NMJs of control and VPA-treated animals at the end stage of the disease. Both of these results suggest that the distal pathology (i.e., denervation) was not prevented by neuroprotection.

Altogether, these data show that at the clinical level, neuroprotection of lower MNs is not sufficient for ALS therapy, because it does not prevent MN denervation during the pathology. These data strongly suggest that MN death activation is not the required pathogenic event of ALS and point instead to disruption of MN presynaptic terminals.

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Rouaux et al. • Uncoupling of Neuroprotection and Motor Dysfunction in ALS

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