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An a2-Na/K ATPase/a-adducin complex in astrocytes triggers non-cell autonomous neurodegeneration

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Abstract

Perturbations of astrocytes trigger neurodegeneration in several diseases, but the glial cellintrinsic mechanisms that induce neurodegeneration remain poorly understood. We found that a protein complex of α 2-Na/K ATPase and α -adducin was enriched in astrocytes expressing mutant superoxide dismutase 1 (SOD1), which causes familial amyotrophic lateral sclerosis (ALS). Knockdown of α 2-Na/K ATPase or α -adducin in mutant SOD1 astrocytes protected motor neurons from degeneration, including in mutant SOD1 mice *in vivo*. Heterozygous disruption of the α 2-Na/K ATPase gene suppressed degeneration *in vivo* and increased the lifespan of mutant SOD1 mice. The pharmacological agent digoxin, which inhibits Na/K ATPase activity, protected motor neurons from mutant SOD1 astrocyte–induced degeneration. Notably, α 2-Na/K ATPase

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AUTHOR CONTRIBUTIONS

A.B. directed and coordinated the project. G.G. designed and performed or participated in all experiments. J.B. performed mouse husbandry and survival studies. J.B.L. provided α2-Na/K ATPase knockout mice. H.S. performed mass spectrometry analysis. J. Ravits, T.S. and J. Robertson provided human tissue samples. The manuscript was written by G.G. and A.B. and commented on by all authors.

Astrocytes represent the most abundant cell type in the CNS and have diverse functions in the developing and mature CNS^{1,2}. Astrocytes and neurons share a common lineage during development, and both cell types often express disease genes that trigger neurodegeneration in the CNS. Notably, astrocytes are beginning to emerge as critical targets of CNS disorders that were once thought to selectively afflict neurons. In particular, mounting evidence suggests that astrocytes have a fundamental role in the progression of diverse neurodegenerative diseases^{3,4}. Expression of mutant proteins in astrocytes in ALS, Huntington's disease and spinocerebellar ataxias induce non–cell autonomous neurodegeneration ^{5–11}. However, with few exceptions¹², the cell-intrinsic mechanisms operating in mutant astrocytes that trigger non–cell autonomous neurodegeneration remain largely unknown.

The cellular basis of non–cell autonomous neurodegeneration has been best characterized in ALS^{9,13–15}. ALS is the most common motor neuron disease in adults and is characteristically fatal within 5 years of onset. Approximately 5–10% of patients with ALS are familial with an autosomal dominant pattern of inheritance¹⁶. Mutations in the gene encoding SOD1 account for 20% of familial ALS with over 140 distinct mutations identified to date^{16,17}.

Transgenic mice expressing the G93A mutation in SOD1 (SOD1^{G93A}) have been an invaluable model for studies of neurodegeneration, as these mice recapitulate the pathological features of ALS, including reactive gliosis, ubiquitin aggregates, loss of motor neurons and lethality^{13,18}. The degeneration of motor neurons in SOD1^{G93A} mice is thought to result in part from cell-autonomous mechanisms¹⁹. In addition, expression of mutant SOD1 in astrocytes induces the degeneration of motor neurons in a non–cell autonomous fashion^{9,14,15,20}. Notably, astrocytes from mutant SOD1 mice and astrocytes derived from post-mortem spinal cords of patients with either SOD1 mutations or sporadic ALS induce toxicity in primary motor neurons²⁰. Thus, mutant SOD1 mice provide an excellent model for elucidation of the glial cell–intrinsic mechanisms of non–cell autonomous neurodegeneration.

We found that a complex composed of the ion pump α 2-Na/K ATPase and the protein α adducin in SOD1^{G93A} astrocytes triggers the non–cell autonomous degeneration of motor neurons. Knockdown of α 2-Na/K ATPase or α -adducin in SOD1^{G93A} astrocytes markedly inhibited their ability to induce degeneration in co-cultured primary motor neurons. In addition, *in vivo* knockdown of the α 2-Na/K ATPase/ α -adducin complex by lentiviralmediated RNAi in the spinal cord of SOD1^{G93A} mice protected motor neurons from degeneration *in vivo*. Notably, inactivating one allele of *Atp1a2* (the gene encoding α 2-Na/K ATPase) in SOD1^{G93A} mice suppressed motor neuron degeneration and substantially increased mouse lifespan. In mechanistic studies, we found that mitochondrial respiration and inflammatory gene expression were induced in SOD1^{G93A} astrocytes, and removal of

one allele of *Atp1a2* reversed these effects, suggesting that the upregulation of α 2-Na/K ATPase stimulates mitochondrial respiration and expression of secreted inflammatory factors in SOD1^{G93A} astrocytes. The Na/K ATPase small molecule inhibitor digoxin, which has been widely used in the treatment of congestive heart failure²¹, blocked the degeneration of co-cultured primary motor neurons. Finally, α 2-Na/K ATPase and α -adducin were substantially upregulated in the spinal cord in individuals with familial ALS bearing distinct SOD1 mutations as well as in sporadic ALS. Together, our findings suggest that the α 2-Na/K ATPase/ α -adducin complex is critical role for the pathology of non–cell autonomous neurodegeneration and provides a potential drugable target in the treatment of neurodegenerative diseases.

RESULTS

a-adducin induces non-cell autonomous motor neuron degeneration

Using an antibody that recognizes phosphorylation events in cells following exposure to oxidative stress²², we unexpectedly identified a 105-kDa immunoreactive protein band enriched in lysates of spinal cord from symptomatic SOD1^{G93A} mice at 120 d of age as compared with age-matched wild-type littermate mice and non-symptomatic 60-d-old SOD1^{G93A} mice (Fig. 1a). Following incubation of symptomatic SOD1^{G93A} spinal cord lysates with λ -phosphatase, the immunoreactive 105-kDa protein band was eliminated (Fig. 1a). Mass spectrometry analysis following immunoprecipitation assays led to the identification of α -adducin as the putative phosphorylated protein in SOD1^{G93A} spinal cords (data not shown). We validated our mass spectrometry analysis by immunoprecipitating aadducin and immunoblotting with our phospo-antibody, confirming the identity of aadducin in symptomatic SOD1^{G93A} mice (Fig. 1b). Notably, immunoblotting for total aadducin protein levels in symptomatic SOD1G93A mice at 120 d and control age-matched wild-type littermate mice revealed that a-adducin protein was upregulated in spinal cord of symptomatic SOD1^{G93A} mice (Fig. 1c). The upregulation of α -adducin in the spinal cord of SOD1^{G93A} mice was evident as early as 90 d of age, which represents the time of disease onset (Supplementary Fig. 1a). In other experiments, we identified Ser436 as the site of α adducin phosphorylation in lysates of SOD1^{G93A} spinal cords (data not shown).

We next determined the cellular origin of α -adducin in SOD1^{G93A} mice. In immunoblotting analysis of primary SOD1^{G93A} glial cells and motor neurons, α -adducin and Ser436phosphorylated α -adducin were predominantly expressed in astrocytes rather than motor neurons (Fig. 1d). In complementary immunohistochemical analyses, Ser436phosphorylated α -adducin colocalized with the astrocyte marker glia fibrillary acidic protein (GFAP) in spinal cord of symptomatic SOD1^{G93A} mice (Fig. 1e). The Ser436phosphorylated α -adducin immunoreactivity was specifically blocked by a phosphorylated-Ser436 α -adducin peptide, validating the specificity of the phosphorylated-Ser436 α adducin immunoreactivity in astrocytes (data not shown). In control analyses, Ser436phosphorylated α -adducin did not colocalize with the motor neuron marker SMi32 in the spinal cord of symptomatic SOD1^{G93A}. In addition, Ser436-phosphorylated α -adducin colocalized with GFAP, but not SMi32, in the spinal cord of wild-type mice (Supplementary Fig. 1b,c). α -adducin forms heterodimers or heterotetramers with β -adducin or γ -adducin²³.

The adducins function as barbed-end actin-capping proteins that regulate actin filaments length²³. Notably, the absence of α -adducin leads to the downregulation of the other subunits, suggesting that α -adducin is the limiting subunit in the formation of adducin heteromers²⁴. The phosphorylation of α -adducin has been described in mutant SOD1 mice and sporadic ALS patients^{25,26}, but the role of α -adducin in neurodegeneration is not yet known. The localization and abundance of α -adducin in astrocytes in symptomatic SOD1^{G93A} mice raises the question of whether α -adducin might be involved in the toxic gain of function in SOD1^{G93A} astrocytes.

To characterize α -adducin function in neurodegeneration, we first employed a cell culture model in which SOD1^{G93A} astrocytes are co-cultured with primary spinal cord motor neurons, which recapitulates the non-cell autonomous degeneration of motor neurons in vivo^{14,15,20}. Using a plasmid-based method of RNA interference (RNAi), we induced efficient knockdown of α -adducin in SOD1^{G93A} astrocytes (Supplementary Fig. 2a). We assessed the effect of a-adducin knockdown in SOD1G93A astrocytes on measures of degeneration in co-cultured motor neurons. SOD1^{G93A} astrocytes, but not wild-type astrocytes, transfected with the control U6 RNAi plasmid induced cell death and a substantial reduction in total dendrite length in motor neurons (Fig. 1f-h), confirming that mutant SOD1 astrocytes trigger non-cell autonomous degeneration of motor neurons. Notably, we found that knockdown of a-adducin in SOD1^{G93A} astrocytes protected motor neurons against the non-cell autonomous induction of motor neuron cell death (Fig. 1f-h). Although control SOD1^{G93A} astrocytes induced cell death in 50% of co-cultured motor neurons, α-adducin knockdown SOD1G93A astrocytes induced cell death in only 23% of cocultured motor neurons (Fig. 1g). Likewise, knockdown of a-adducin in SOD1^{G93A} astrocytes prevented the ability of SOD1^{G93A} astrocytes to induce abnormalities in motor neuron dendrite morphology (Fig. 1f,h). In control analyses, knockdown of a-adducin in non-transgenic astrocytes had little or no effect on the survival or morphology of co-cultured motor neurons (Fig. 1f-h).

To determine the specificity of the α -adducin RNAi–induced neuroprotective phenotype in SOD1^{G93A} astrocytes, we performed a rescue experiment. We expressed an RNAi-resistant form of α -adducin (Add-Res) in the background of α -adducin RNAi in SOD1^{G93A} astrocytes. Expression of Add-Res in SOD1^{G93A} astrocytes reversed the ability of α -adducin RNAi to protect co-cultured motor neurons from cell death and impairment of dendrite morphology (Supplementary Fig. 2). These data indicate that the α -adducin in SOD1^{G93A} astrocytes rather than off-target effects of RNAi. Together, our data suggest that α -adducin in SOD1^{G93A} astrocytes is critical for the non–cell autonomous degeneration of motor neurons.

We next assessed the effect of α -adducin knockdown on neurodegeneration in the spinal cord of SOD1^{G93A} mice *in vivo*. We injected lentivirus encoding α -adducin short hairpin RNAs and GFP (LV-Addi) or the corresponding control lentivirus (LV-U6) unilaterally in the lumbar spinal cord in SOD1^{G93A} mice (Fig. 2a). The *in vivo* RNAi approach allowed comparison of surviving motor neurons in the injected ventral horn with the non-injected contralateral ventral horn in the same spinal cord sections. We injected lentivirus in the

spinal cord of SOD1^{G93A} mice at 90 d, a time at which gliosis had set in, thereby maximizing the targeting of reactive astrocytes. Following injection of the lentivirus encoding GFP into the spinal cord of SOD1^{G93A} mice, 88% of the GFP-positive cells expressed the astrocyte marker GFAP, 8% expressed the motor neuron marker SMi32 and 3% expressed the microglia marker Iba1 (Supplementary Fig. 3a,b). Injection of lentivirus expressing GFP into wild-type mice at 90 d revealed 6% colocalization of GFP with SMi32, 3% with Iba1 and 91% with GFAP (Supplementary Fig. 3c,d). Thus, injected lentivirus into the spinal cord of SOD1^{G93A} and control mice largely targeted astrocytes *in vivo*.

Injection of control lentivirus in SOD1^{G93A} mice (LV-U6 SOD1^{G93A}) had little or no effect on the survival of motor neurons in the GFP-labeled injected ventral horn when compared with the contralateral non-injected ventral horn *in vivo* (Fig. 2b,c). By contrast, α -adducin kncockdown in SOD1^{G93A} mice (LV-Addi SOD1^{G93A}) strongly suppressed motor neuron degeneration *in vivo* (Fig. 2d,e). The α -adducin knockdown mice harbored 7.07 \pm 0.98 motor neurons in the GFP-labeled ventral horn injected with α -adducin RNAi virus, whereas the contralateral non-injected ventral horn contained only 3.26 \pm 0.56 motor neurons (Fig. 2d,e). In control analyses, we confirmed that α -adducin RNAi induced the knockdown of α adducin in the GFP-labeled injected ventral horn (Supplementary Fig. 4). Knockdown of α adducin had little or no effect on the presence of gliosis in the ventral horn (Supplementary Fig. 5). Likewise, α -adducin knockdown did not alter the migration or presence of microglia in the ventral horn *in vivo* (Supplementary Fig. 6). Collectively, our data suggest that α adducin mediates non–cell autonomous degeneration of motor neurons in the spinal cord of SOD1^{G93A} mice *in vivo*.

a2-Na/K ATPase triggers motor neuron degeneration

The identification of a critical role for α -adducin in non–cell autonomous motor neuron degeneration led us to determine the mechanism underlying the function of α -adducin in neurodegeneration. We performed immunoprecipitation of α -adducin followed by mass spectrometry in lysates of spinal cord from symptomatic SOD1^{G93A} mice. These analyses revealed that the ion pump α 2-Na/K ATPase interacted with α -adducin in symptomatic SOD1^{G93A} spinal cord lysates (data not shown). The Na/K ATPases are composed of two subunits, a catalytic α subunit and a non-catalytic β subunit²⁷. The four catalytic isoforms, α 1- α 4, display a unique tissue expression pattern^{27,28}. Notably, the α 2 catalytic subunit is selectively expressed in astrocytes in the CNS²⁸. These observations led us to characterize the role of α 2-Na/K ATPase in non–cell autonomous degeneration of motor neurons in SOD1^{G93A} mice.

We first validated the interaction of α -adducin with α 2-Na/K ATPase by immunoprecipitating α -adducin followed by immunoblotting with an α 2-Na/K ATPase antibody in spinal cord lysates of SOD1^{G93A} and littermate control mice (Supplementary Fig. 7a). In immunoblotting analyses of primary astrocytes and motor neurons, α 2-Na/K ATPase was predominantly expressed in astrocytes rather than motor neurons (Supplementary Fig. 7a). In immunohistochemical analyses, α 2-Na/K ATPase was predominately expressed in astrocytes in the spinal cord of both presymptomatic and symptomatic SOD1^{G93A} mice (Supplementary Fig. 7b,c). Notably, immunoblotting analyses

revealed that the levels of α 2-Na/K ATPase were upregulated in symptomatic SOD1^{G93A} mice at 120 d (Fig. 3a). Consistent with these results, the increase in α 2-Na/K ATPase protein levels was also evident in primary SOD1^{G93A} astrocytes (Fig. 3b). Knockdown of α -adducin in SOD1^{G93A} astrocytes reduced the levels of α 2-Na/K ATPase in these cells (Fig. 3b). These data suggest that upregulation of α 2-Na/K ATPase might act in concert with upregulated α -adducin to promote the toxic gain of function in SOD1^{G93A} astrocytes.

We determined the role of α 2-Na/K ATPase in the toxic gain of function of SOD1^{G93A} astrocytes. Knockdown of α 2-Na/K ATPase in non-transgenic control glia had little or no effect on motor neuron survival or dendrite morphology (Fig. 3c–e). Notably, knockdown of α 2-Na/K ATPase in SOD1^{G93A} astrocytes protected co-cultured primary motor neurons against non–cell autonomous cell death and impairment of dendrite morphology (Fig. 3c–e). These data suggest that knockdown of α 2-Na/K ATPase phenocopies the neuroprotective effects of α -adducin knockdown in SOD1^{G93A} astrocytes.

We next assessed the role of α 2-Na/K ATPase in SOD1^{G93A}-dependent neurodegeneration *in vivo*. We used a lentiviral approach to induce knockdown of α 2-Na/K ATPase in the lumbar spinal cord in SOD1^{G93A} mice. Just as in the α -adducin experiments *in vivo*, injection of control lentivirus in SOD1^{G93A} mice (LV-U6 SOD1^{G93A}) had no effect on motor neuron survival (Supplementary Fig. 8a,b). By contrast, knockdown of α 2-Na/K ATPase in SOD1^{G93A} mice by lentivirus (LV-ATPi SOD1^{G93A}) suppressed the degeneration of spinal cord motor neurons *in vivo*. The GFP-labeled injected ventral horn in the α 2-Na/K ATPase knockdown mice harbored 6.7 ± 0.28 motor neurons, whereas only 4.33 ± 0.31 motor neurons were present in the non-injected contralateral ventral horn in these mice (Fig. 3f,g). Notably, α 2-Na/K ATPase knockdown had little or no effect on the presence or abundance of astrocytes or microglia in the ventral horns of SOD1^{G93A} mice (Supplementary Figs. 5 and 6). These data indicate that α 2-Na/K ATPase knockdown in SOD1^{G93A} mice suppresses motor neuron degeneration *in vivo*.

Heterozygous a2-Na/K ATPase suppresses motor neuron degeneration

We also used a genetic knockout approach to define the role of α 2-Na/K ATPase in neurodegeneration in SOD1^{G93A} mice. Although the complete absence of α 2-Na/K ATPase leads to embryonic lethality, heterozygous-null mice expressing approximately 50% of normal levels of α 2-Na/K ATPase protein display no gross abnormalities²⁹. We measured the ability of astrocytes from heterozygous-null *Atp1a*2^{+/-}; SOD1^{G93A} mice (*Atp1a*2^{+/-}; SOD1^{G93A}) to induce cell death of co-cultured motor neurons. Control SOD1^{G93A} astrocytes (*Atp1a*2^{+/+}; SOD1^{G93A}) induced non–cell autonomous cell death in 53% of cocultured motor neurons (Fig. 4a,b). In contrast, heterozygous-null *Atp1a*2^{+/-}; SOD1^{G93A} astrocytes (*Atp1a*2^{+/-}; SOD1^{G93A}) induced non–cell autonomous cell death in only 14% of motor neurons (Fig. 4b). Likewise, *Atp1a*2^{+/-}; SOD1^{G93A} astrocytes failed to induce dendrite abnormalities in motor neurons as compared to control SOD1^{G93A} astrocytes (Fig. 4a,c). These data corroborate the results of our knockdown analyses and buttress the conclusion that α 2-Na/K ATPase is critical for non–cell autonomous degeneration of motor neurons.

The genetic knockout approach facilitated analysis of the role of α 2-Na/K ATPase in the motor neuron disease phenotype of mutant SOD1 mice. We characterized disease onset, progression and lethality in heterozygous-null Atp1a2; SOD1^{G93A} mice and control SOD1^{G93A} littermates. Disease onset, defined as the first day of weight loss, was significantly delayed in $Atp1a2^{+/-}$; SOD1^{G93A} mice ($Atp1a2^{+/-}$) as compared with control $Atp1a2^{+/+}$; SOD1^{G93A} mice ($Atp1a2^{+/+}$) (P = 0.0009; Fig. 4d). Accordingly, disruption of Atp1a2 in SOD1^{G93A} mice delayed the age at which 10% weight loss was reached, which is a measurement of early disease (Fig. 4e). Early phase disease progression, as measured from the first day of weight loss to 10% weight loss, was not significantly altered between Atp1a2^{+/-}; SOD1^{G93A} and control SOD1^{G93A} littermates (P = 0.2023; Fig. 4f). In contrast, late-phase disease progression, as measured from 10% weight loss to end stage, was significantly delayed in Atp1a2^{+/-}; SOD1^{G93A} mice compared with control SOD1^{G93A} littermates (P = 0.0005; Fig. 4g). Because expression of SOD1^{G37R} in astrocytes drives latephase motor neuron degeneration progression in mice⁹, the suppression of late-phase disease progression in $Atp1a2^{+/-}$; SOD1^{G93A} mice supports the conclusion that α 2-Na/K ATPase mediates astrocyte-dependent mechanism of neurodegeneration. Notably, the overall survival of SOD1^{G93A} mice was increased after reducing the expression of a2-Na/ATPase to an average lifespan of 171.0 ± 2.5 d compared with 151.5 ± 2.7 d in control SOD1^{G93A} mice (Fig. 4h). In addition, the $Atp1a2^{+/-}$; SOD1^{G93A} mice were substantially more mobile at the time at which the control SOD1^{G93A} mice were at the endstage of the disease (Fig. 4i and Supplementary Movies 1 and 2). Thus, reducing the expression of a2-Na/K ATPase in SOD1^{G93A} mice delays the onset and slows the progressive process of neurodegeneration, thereby substantially increasing lifespan.

We next determined if the improvement in the lifespan of SOD1^{G93A} mice that are heterozygous for *Atp1a2* is associated with suppression of motor neuron degeneration *in vivo*. We quantified motor neurons at the endstage of disease in control SOD1^{G93A} mice and aged-matched SOD1^{G93A} littermates heterozygous-null for *Atp1a2*. Control SOD1^{G93A} mice harbored 3.39 ± 0.43 motor neurons per ventral horn (Supplementary Fig. 9a,b). By contrast, littermate *Atp1a2*^{+/-}; SOD1^{G93A} mice had more than twice the number of motor neurons at 7.27 ± 0.63 per ventral horn (Supplementary Fig. 9a,b). These results demonstrate that reducing the expression of α 2-Na/K ATPase in SOD1^{G93A} mice delays motor neuron degeneration.

We also determined the effect of disruption of Atp1a2 in SOD1^{G93A} mice on motor neuron innervation of muscles *in vivo*. We quantified neuromuscular junction (NMJ) innervation of the gastrocnemius muscle in 120-d SOD1^{G93A} mice heterozygous for the null Atp1a2 allele and age-matched control SOD1^{G93A} mice using published parameters³⁰. We categorized NMJs as completely innervated (as reflected by complete overlap of synapsin with α bungarotoxin staining), completely denervated (displaying no overlap staining) or partially denervated (displaying partial overlap). In control SOD1^{G93A} mice, 75% of NMJs were completely denervated, 23% were partially denervated and less than 1.5% were completely innervated (Fig. 4j,k). By contrast, in $Atp1a2^{+/-}$; SOD1^{G93A} mice, 47% of NMJs were completely denervated, 45% were partially denervated and 7.6% were completely innervated NMJs (Fig. 4j,k). These analyses demonstrate that knockout of one allele of

Atp1a2 substantially improves the integrity of motor neuron innervation of peripheral muscles.

Na/K ATPase stimulates mitochondria and inflammatory genes

The elevated levels of the a2-Na/K ATPase/a-adducin complex in SOD1G93A astrocytes raised the question of whether the activity of a2-Na/K ATPase per se has a pathogenic role in the toxic gain of function of SOD1G93A astrocytes. We used the Na/K ATPase small molecule inhibitors ouabain and digoxin, the latter of which is used widely as a therapeutic drug in treatment of heart failure²¹, to assess the role of Na/K ATPase activity in the toxic effects of SOD1^{G93A} astrocytes. Following the co-culturing of motor neurons with control and mutant SOD1^{G93A} astrocytes, we added ouabain, digoxin or control vehicle at a final concentration of 1 µm, which is sufficient to inhibit the a2-Na/K ATPase in primary glia cells³¹. We found that inhibition of Na/K ATPase with ouabain or digoxin substantially reduced motor neuron cell death induced by SOD1^{G93A} astrocytes to 22% and 19%, respectively, as compared with 56% motor neuron cell death in cultures treated with vehicle (Fig. 5a,b). Likewise, both ouabain and digoxin prevented impairment of dendrite morphology in motor neurons induced by SOD1^{G93A} astrocytes (Fig. 5a,c). In control analyses, co-cultures of motor neurons and control astrocytes exposed to ouabain and digoxin did not alter survival or morphology of motor neurons (Fig. 5a-c). These results support the conclusion that the catalytic activity of a2-Na/K ATPase/a-adducin complex in SOD1^{G93A} astrocytes triggers non-cell autonomous degeneration of motor neurons.

Because Na/K ATPase is the major consumer of ATP in the CNS, we would anticipate that an increase in the levels and activity of Na/K ATPase would alter demand for cellular metabolism and ATP. Mitochondria generate the majority of the cellular demand for ATP. We reasoned that elevated levels and activity of α 2-Na/K ATPase in SOD1^{G93A} astrocytes might reduce basal ATP levels and influence mitochondrial respiration. Measurement of oxygen consumption revealed that mutant SOD1^{G93A} astrocytes had significantly higher levels of basal oxygen consumption (P = 0.0001) and significantly increased maximum oxidative capacity (P = 0.0001), the latter of which was induced by the electron transport uncoupler p-triflouromethoxyphenylhydrazon (FCCP) (Fig. 5d,e). By contrast, $Atp1a2^{+/-}$; SOD1^{G93A} astrocytes had reduced basal and maximum oxidative capacity (Fig. 5d,e). These data suggest that α 2-Na/K ATPase in SOD1^{G93A} astrocytes enhance mitochondrial respiration, presumably secondary to increased ATP demand. Following the downregulation of α 2-Na/K ATPase in $Atp1a2^{+/-}$; SOD1^{G93A}, demand for ATP production is attenuated, thereby reducing basal and maximum oxidative capacity.

Cellular stresses, including increased metabolic rates, induce the production of mitochondrial reactive oxygen species (ROS) that in turn activate an inflammatory response associated with the secretion of cytokines and other inflammatory factors that promote chronic inflammatory diseases^{32–34}. Notably, both ROS and inflammation have been linked to the pathogenesis of ALS^{20,35}. Consistent with this possibility, whereas conditioned medium from SOD1^{G93A} astrocytes induces degeneration of motor neurons, conditioned medium from $Atp1a2^{+/-}$; SOD1^{G93A} astrocytes failed to induce motor neuron degeneration (Supplementary Fig. 10a,b). These results suggest that the upregulation of α 2-Na/K ATPase

in SOD1^{G93A} astrocytes mediates the non-cell autonomous degeneration of motor neurons via secreted factors.

We next assessed whether the upregluation of α 2-Na/K ATPase in SOD1^{G93A} astrocytes might stimulate the expression of specific inflammatory genes^{20,36}. In quantitative RT-PCR (qRT-PCR) analyses, we characterized a panel of 20 inflammatory genes reported to be increased in SOD1^{G93A} astrocytes^{20,36}. We found that expression of a set of cytokines, chemokines and related inflammatory astrocytic genes was increased in SOD1^{G93A} astrocytes compared with control wild-type astrocytes (Fig. 5f). Among this panel, more than half were downregulated in *Atp1a2*^{+/-}; SOD1^{G93A} astrocytes (Fig. 5f). These data support the conclusion that the upregulation of α 2-Na/K ATPase in SOD1^{G93A} astrocytes promotes an inflammatory response, which may in turn induce motor neuron degeneration.

a2-Na/K ATPase and a-adducin are upregulated in ALS spinal cord

To determine the clinical relevance of the a2-Na/K ATPase/a-adducin mechanism in noncell autonomous neurodegeneration, we characterized the expression of the a2-Na/K ATPase/ α -adducin complex in spinal cord lysates from both familial ALS expressing distinct mutations in SOD1 and sporadic ALS patients and controls. Familial ALS cases demonstrated autosomal dominant inheritance. All of these patients had definite ALS by the El Escorial criteria³⁷, with a mean age of 42 years (range of 21-65). The mean age of sporadic ALS patients was 60 years (range 46-69). All patients with sporadic ALS met modified El Escorial criteria for probable or definite ALS, with their spinal cords displaying gliosis, demyelination and long-term motor neuron degeneration³⁷. The mean age of control patients was 64 years (range 54-70). The CNS had normal appearance in samples of control patients. Immunoblotting of lysates of familial ALS, sporadic ALS and control patients revealed that the levels of α 2-Na/K ATPase were significantly increased in lysates of spinal cord in both familial (P = 0.0302) and sporadic (P = 0.0467) ALS patients as compared with controls (Fig. 6). Likewise, the protein levels of α -adducin were also significantly increased in the spinal cord of ALS patients (familial, P = 0.0336; sporadic, P = 0.0181; Fig. 6a,d). Quantification of α 2-Na/K ATPase and α -adducin immunoreactivity as a continuous variable revealed that the levels of these two proteins doubled in both familial and sporadic ALS (Fig. 6b,c,e,f). Together, these data suggest that the abundance of α 2-Na/K ATPase and α -adducin in familial and sporadic ALS mimics the elevated levels of the α 2-Na/K ATPase/a-adducin complex in SOD1^{G93A} mice and may therefore contribute to neurodegeneration.

DISCUSSION

Our results reveal a glial cell-intrinsic mechanism of non-cell autonomous neurodegeneration. We found that α 2-Na/K ATPase and α -adducin form a complex that is highly abundant in mutant SOD1 astrocytes. Knockdown of α 2-Na/K ATPase or α -adducin by RNAi or by introducing a null allele of *Atp1a2* in mutant SOD1 astrocytes protects motor neurons from non-cell autonomous degeneration. Notably, introducing a null allele of *Atp1a2* in mutant SOD1 mice suppressed motor neuron degeneration and muscle denervation *in vivo* and thereby substantially increased mouse lifespan. Mechanistic studies

revealed that mitochondrial respiration and inflammatory genes were induced in mutant SOD1 astrocytes and that these alterations were suppressed in mutant SOD1 *Atp1a2*^{+/-} astrocytes. We have also found that the therapeutic drug digoxin, which inhibits Na/K ATPases and is commonly used to treat heart failure²¹, protected motor neurons from non– cell autonomous degeneration. Finally, we found that α 2-Na/K ATPase and α -adducin were upregulated in both familial ALS patients expressing distinct SOD1 mutations and sporadic ALS patients, suggesting a pathogenic role for this protein complex in human disease. Taken together, our results define the α 2-Na/K ATPase/ α -adducin complex as a key glial cell–intrinsic regulator of non–cell autonomous neurodegeneration.

Although neurodegenerative diseases were once thought to be selectively neuronal, growing evidence suggests that expression of neurodegenerative disease genes in astrocytes triggers neuronal degeneration non–cell autonomously^{3,4,7,10}. However, the glial cell-intrinsic mechanism that mediate neurodegeneration have remained poorly understood. Our study reveals that upregulation of α 2-Na/K ATPase/ α -adducin complex stimulates mitochondrial respiration and an inflammatory response in mutant SOD1 astrocytes, leading to the non–cell autonomous degeneration of motor neurons. The finding that the α 2-Na/K ATPase/ α -adducin complex is upregulated in familial ALS patients expressing distinct mutations of SOD1 and in sporadic ALS supports the conclusion that the α 2-Na/K ATPase/ α -adducin complex may represent a common mechanism of non–cell autonomous degeneration in motor neuron disease. Identification of a critical pathogenic role for the glial α 2-Na/K ATPase/ α -adducin complex in non–cell autonomous neurodegeneration in ALS suggests that it will be important to determine whether chronic activation of this complex in astrocytes might contribute to the pathogenesis of other neurodegenerative diseases.

Our findings shed light on the role of Na/K ATPase activity in motor neuron disease. The levels of Na/K ATPase isoforms have been reported to be downregulated in mutant SOD1 spinal cord³⁸. However, consistent with our findings, the activity of Na/K ATPase has been shown to be elevated in spinal cord lysates from symptomatic mutant SOD1 mice³⁹. We found that α 2-Na/K ATPase was upregulated in the spinal cord of symptomatic mutant SOD1 mice³⁹. We found that the upregulation of α 2-Na/K ATPase occurred specifically in glia in these mice. Notably, consistent with our findings in mutant SOD1 mice, we also found that the levels of α 2-Na/K ATPase were elevated in spinal cord of familial ALS patients harboring distinct SOD1 mutations as well as sporadic ALS. Our distinct genetic approaches of knockdown and knockout analyses both in primary cultures and in mice *in vivo* suggest a requirement for α 2-Na/K ATPase in motor neuron degeneration.

Recent studies have suggested that elevated levels of the $\alpha 1$ isoform of Na/K ATPase may induce hippocampal morphological abnormalities and memory deficits in Angelman syndrome, a neurodevelopmental disorder that features developmental delay, speech impairment and seizures^{40,41}. Attenuation of the $\alpha 1$ Na/K ATPase levels suppresses the hippocampal morphological abnormalities and memory deficits in a mouse model of Angelman syndrome. Whether the $\alpha 1$ isoform of Na/K ATPase acts non–cell autonomously in glial cells or cell autonomously in neurons and whether $\alpha 1$ -Na/K ATPase operates in a complex with α -adducin in Angelman syndrome remains unknown. In view of our findings, it will be interesting to determine whether the glial $\alpha 2$ -Na/K ATPase/ α -adducin complex

contributes to the pathogenesis of other neurological diseases beyond neurodegeneration, including developmental disorders of cognition.

Our study also illuminates a mechanism by which the α 2-Na/K ATPase/ α -adducin complex elicits neurodegeneration non–cell autonomously. The upregulation of α 2-Na/K ATPase is anticipated to reduce ATP levels and increase demand for mitochondria function. Accordingly, we found that the upregulation of α 2-Na/K ATPase stimulates mitochondrial respiration in mutant SOD1 astrocytes. Mitochondrial respiration may increase ROS in mutant SOD1 astrocytes, which in turn may activate the observed induction of inflammatory factors leading to non–cell autonomous degeneration of motor neurons.

The role of α -adducin and Na/K ATPase activity in the renal epithelium may be instructive for the study of neurodegeneration. Consistent with our findings in mutant SOD1 astrocytes, α -adducin promotes the stability and activity of Na/K ATPase in renal epithelial cells^{42,43}. Notably, polymorphisms in α -adducin may enhance Na/K-ATPase activity in the renal epithelium, leading to abnormal renal ionic handling and hypertension in humans^{42–44}. Furthermore, in models of renal interstitial fibrosis, chronic activation of Na/K ATPase in proximal tubule cells and macrophages leads to the release of pro-inflammatory cytokines that facilitate the development of chronic inflammation, oxidative stress and fibrosis^{45,46}. Thus, our findings that α 2-Na/K ATPase/ α -adducin complex in astrocytes triggered non– cell autonomous degeneration suggest the exciting idea of parallels in the pathogenesis of hypertension and neurodegenerative diseases.

The unexpected finding that chronic activation of α 2-Na/K ATPase in astrocytes is critical for neurodegeneration suggests that α 2-Na/K ATPase might represent an attractive target for the identification of new therapies for neurodegenerative diseases. We found that the therapeutic drug digoxin, which has been widely used to treat heart failure²¹, protects motor neurons against degeneration. Consistent with our findings, cardiac glycosides appear to be neuroprotective in models of ischemic stroke, prevents polyglutamine-induced cell death, and inhibits SOD1 and TDP-43 aggregation in cells^{47–50}. Whether chronic activation of the α 2-Na/K ATPase/adducin complex in astrocytes has an active role in other neurodegenerative diseases and in brain injury remains to be determined. The enormous experience with Na/K ATPase inhibitors in the treatment of heart disease should prove useful in the development of inhibitors selective for the α 2 isoform of Na/K ATPase in glial-dependent neurodegeneration.

ONLINE METHODS

Human tissue

All human spinal cord tissue samples were acquired by way of an Investigational Review Board and Health Insurance Portability and Accountability Act compliant process. The SOD1 A4V, V148G and E100G nervous system were confirmed by DNA testing. All patients who had been followed during the clinical course of their illness had met El Escorial criteria for definite ALS. After death, autopsies were performed immediately by an on-call tissue acquisition team. Control spinal cord tissue samples were from patients from the hospital's critical care unit when life support was withdrawn or patients on hospice. Tissue

samples were completed within 4–6 h of death and the entire motor system was dissected and elaborately archived for downstream applications by creating two parallel tissue sets from alternating adjacent regions. For biochemical studies, segments were embedded in cutting media, frozen on blocks of dry ice and stored at -80 °C.

Animals

We used transgenic mice overexpressing mutant G93A human superoxide dismutase 1 (Jackson Laboratory stock# 002726), α 2-Na/K ATPase null heterozygous (provided by J.B.L.) and C57BL6 (Charles River). Mouse husbandry with timed mating and analysis of age-dependent phenotypes was carried out as described⁵¹. Two or three males or two to four females per cage were housed under a 12-h/12-h light-dark cycle. Age-matched littermates of both males and females were used in the experiments. No animals were excluded from analysis. All analysis was done with littermate mice derived from SOD1G93A heterozygous mating with non-transgenic wild-type and heterozygous α 2-Na/K ATPase mice. All animal experiments were conducted under the institutional guidelines and were approved by the Institutional Animal Care and Use Committees of Harvard Medical School and Washington University School of Medicine.

Plasmids

RNAi plasmids were designed as described previously⁵². The U6/ α -adducin plasmids were cloned using the following primers: 5'-agt cga aga cta agt gga ctt-3' and 5'-agt cga aga cta agt gga cta-3'. The U6/ α 2-Na/K ATPase plasmids were cloned using the following primers: 5'-gca tca tat cag agg gta acc-3' and 5'-gtg gca aga aga aac aga aac-3'. α -adducin and α 2-Na/K ATPase were cloned from C57BL/6J mice cDNA and inserted into the pcDNA3 vector (Invitrogen) at the ECOR1 and Xho1 sites. The primer for site direct mutagenesis for RNAi-resistant form of α -adducin was 5'-aac gga agc agt ccc aaA tcA aaA acA aaA tgg acA aaa gag gat gga cat ag-3.

Antibodies

Antibodies to α -adducin (1:1,000, H-100, Santa Cruz), phospho-Ser436-adducin (1:1,000, Ser436, Santa Cruz), α 2-Na/K ATPase (1:1,000, A-16, Santa Cruz), α 2-Na/K ATPase (1:5,000, a5, Developmental Studies Hybridoma Bank), α 2-Na/K ATPase (1:1,000, ATP1A2, Thermo Fisher Scientific), Islet-1 (1:1,000, 39.4D5, Developmental Studies Hybridoma Bank), MAP2 (1:1,000, ab5622, Fisher), GFAP (1:1,000, AB5804, Millipore), Iba-1 (019-19741, WACO), 14-3-3 β (1:2,000, H-8, Santa Cruz), ERK (1:2,000, Molecular Probes), superoxide dismutase (1:1,000, SD-G6, SIGMA), ERK (4685, Cell Signaling), GFP (1:1,000, Molecular Probes) and synapsin (1:500, AB1543, Millipore) were purchased.

Protein biochemistry

To determine the relative amounts of total proteins, whole spinal cords from aged match littermate mice were homogenized in 50 mM Tris-buffer (pH 7.2), 100 mM NaCl, 1 mM EDTA, 1 mM PMSF and a cocktail of protease and phosphatase inhibitors. Equal concentrations from each sample were solubilized in SDS-loading buffer and boiled for 10 min. For analyses of α 2-Na/K ATPase protein levels, equal concentrations (1 mg ml⁻¹) from

each sample were solubililzed in SDS-loading buffer. Using an ultrasonic water sonicator protein samples in SDS-loading buffer were sonicated in an ice bath to prevent oligomerization of the α 2-Na/K ATPase (6 pulses for 30 s with 30-s intervals). Protein quantification was carried out by immunoblotting using Amersham ECL plus and analyzed with Bio-Rad ChemiDoc gel imaging system. Signals were normalized for internal controls run on the same blots. For immunoprecipitation of α -adducin, proteins were extracted under non-denaturing (1% Triton X-100 for 1 h at 4 °C, vol/vol) and denaturing conditions (1% SDS and 0.5% DOC for 1 h at 4 °C, vol/vol) and the insoluble material was removed by centrifugation (18,000 g for 10 min). Following the denaturing conditions the supernatant was diluted to a final concentration of 0.33% SDS and 0.16% Triton. The supernatants were precleared with protein A for 30 min. at 4 °C before the addition of α -adducin antibody for 1 h at 4 °C. For co-immunoprecipitation of α -adducin and α 2-Na/K ATPase, immunoprecipitated α -adducin antibody was eluted from protein A with 0.2 M glycine pH 2.5 on ice for 15 min followed by quenching with 1 mM Tris pH 8.8.

Primary astrocyte cultures

We prepared monolayer astrocytes culture from P2 SOD1^{G93A} and non-transgenic littermate mice that were previously described^{14,15}. Astrocytes were plated in DMEM (GIBCO) supplemented with 10% FBS (vol/vol) containing 100 U ml⁻¹ penicillin and 100 mg ml⁻¹ streptomycin for 2 weeks or until confluent. Once confluent, cells were replated onto glass cover slips pre-coated with poly-d-lysine at a density of 40,000 cells per well in a 24-well dish. For RNAi experiment, astrocyte cultures were transfected with RNAi or control U6 plasmid using Lipofectamine 2000 (Invitrogen) according to the guidelines of the manufacturer 4 d before co-culturing with motor neurons.

Real-time PCR reactions were performed using Light cycler 480 SYBR Green 1 Master (Roche). The following primers were used: ATPase forward, tcttagcctatggtatcc; ATPase reverse, gcatcttttctccctctc; GFAP forward, ggaccagcttacggccaacagtgc; GFAP reverse, ccagcgattcaacctttctctcc; SPP1 forward, ccctcccggtgaaagtgactgattc; SPP1 reverse, ggtctccatcgtcatcatcgtc; LCN2 forward, ggcccaggactcaactcagaacttg; LCN2 reverse, cctgaccaggatggaggtgacattg; CLM1 forward, attgctggtcccctttctct; CLM1 reverse, ggaactccttggcaccagta; WNT11 forward, caggatcccaagccaataaa; WNT 11 reverse, gacaggtagcgggtcttgag; CCL8 forward, gcttctttgcctgctgctcatag; CCL8 reverse, ccatgtactcactgacccacttc; CCL11 forward, gagctccacagcgcttctattcc; CCL11 reverse, ccacttcttcttggggtcagcac; CXCL1 forward, gcttctttgcctgctcatag; CXCL1 reverse, gatetccatgtactcactgaccc; CXCL12 forward, getetgcatcagtgacggtaaacc; CXCL12 reverse, cgggtcaatgcacacttgtctgttg; AIMP1 forward, gctgttctgaagaggctggagca; AIMP1 reverse, agttcgtctgcagtggagtactc; CCR4 forward, tgaccctcggaggatgtacacaa; CCR4 reverse, accatgtttccgagttctgccgg; CCR10 forward, atctggcagccagacgaactacc; CCR10 reverse, agagetegtgcgatggccacata; CEBP forward, aaggccaaggacgagacggt; CEBP reverse, tcgggcagctgcttgaacaagtt; II1b2 forward, ggctgcttccaaacctttgacct; II1b2 reverse, catcatcccatgagtcacagagg; Itgb2 forward, attcaatgtgactttccggcggg; Itgb2 reverse, caggcetteteettgttgggaca; Il10ra forward, gaagtggeeetcaaacagtaegg; Il10ra, gatgccgtccattgctttcagag; II1R1 forward, ggggctcatttgtctcatggtgcc; II1R1 reverse, gctgatgaatcctggagtcccggtc; II1R2 forward, cgcatcccactgtgagcaaatgt; II1R2 reverse,

tgtggggtctcctgcactcagaa; TNFr1 forward, acttggtgagtgactgtccgagc; TNFr1 reverse, ggaagtgtgtctcactcaggta; IFN γ forward, ggctttgcagctcttcctcatgg; IFN γ reverse, cgcttatgttgttgctgatggcc.

Primary motor neuron cultures and morphological analyses

We performed spinal motor neuronal cultures from E12.5 wild-type rodents that were described previously⁵³. We plated at 12,000 cells per well onto astrocyte monolayer in Neurobasal-A medium (Gibco) supplemented with 2% B-27 serum free supplement (vol/ vol) and 0.5 mM glutamine; DMEM/F12 (3:1, vol/vol ratio) with a cocktail of trophic factors composed of 0.5 ng ml⁻¹ glia-derived neurotrophic factor, 1 ng ml⁻¹ brain-derived neurotrophic factor and 10 ng ml⁻¹ ciliary neurotrophic factor (trophic factor cocktail, R&D Systems).

Motor neurons were fixed after 7 d of co-cultures with astrocytes and immunostained with motor neurons marker Islet1 and dendrite marker MAP2. Cell death of motor neurons was assessed on the basis of the integrity of neuritis and the morphology of the nucleus (condensation or fragmentation) as determined using DNA dye Hoechst 33258 (Sigma) previously described²². Approximately 150 cells were counted per cover slip per condition. For motor neuron morphology, images were taken using a NIKON eclipse TE2000 epifluorescence microscope using a digital CCD camera (DIAGNOSTIC instruments) and imported into the SPOT imaging software. Dendrites length was measured by tracing the dendrites emanating from the soma in a manner that produced the longest continuous path. Total dendrite length was quantified as a sum of all dendrites emanating from the soma. Approximately 40 cells were analyzed per cover slip.

Condition medium studies were performed as previously described¹⁴. For pharmacological inhibition of Na/K ATPase in co-cultures, 1.0 μ M of ouabian or digoxin was added 4 h following the co-culturing of motor neurons with control or G93A astrocytes (d *in vitro* (DIV) 0). This was followed by administration of 0.5 μ M of ouabain or digoxin every 48 h (DIV2, DIV4 and DIV6).

Seahorse XF-24 metabolic flux analysis

Oxygen consumption rates and extracellular acidification rate were measured at 37 °C using an XF24 extracellular analyzer (Seahorse Bioscience). Astrocytes were plated in 24-well plates for 24 h (50,000 cells per well) in DMEM 10% FBS containing 100 U ml⁻¹ penicillin and 100 mg ml⁻¹ streptomycin. Cells were changed to unbuffered DMEM (DMEM supplemented either 25 mM glucose, 1 mM sodium pyruvate, 31 mM NaCl, 2 mM GlutaMax, pH 7.4) and incubated at 37 °C in a non-CO₂ incubator for 60 min. Four baseline measurements were taken before sequential injection of mitochondrial inhibitors. Four measurements were taken following the addition of 1.0 μ M oligomycin, 1.0 μ M FCCP, 1.0 μ M rotenone/antimycin A. Oxygen consumption rate and extracellular acidification rate were automatically calculated by Seahorse XF-24 software. Every point represents an average of *n* = 3.

Morphological studies

All morphological studies were performed as previously described⁵¹. Briefly, mice were deeply anesthetized and perfused with 4% paraformaldehyde (vol/vol) in phosphate buffer. Spinal cords were post-fixed in the same fixative for 4 h and processed for cryoprotective embedding. All immunohistochemistry analyses were carried out on 30-µm cryosections from spinal cords with the indicated antibodies. For Na/K ATPase, immunohistochemical analyses antigen retrieval was performed using 1 mg ml⁻¹ pepsin in 0.2 M HCl for 10 min followed by blocking in 3% BSA (vol/vol) for 30 min. For measurement of motor neurons survival, alternating GFP-positive sections were either immunostained or Nissl stained and images were taken using a NIKON eclipse TE2000 epifluorescence microscope or brightfield using a digital CCD camera (DIAGNOSTIC Instruments) and imported into the SPOT imaging software. For lentiviral-mediated RNAi, a minimum of 20 sections were quantified for surviving motor neurons in the injected and non-injected ventral horns from Nissl-stained and immunofluorescence-labeled sections.

Lentiviral injections

Lentivirus was produced in 293T cells and concentrated by ultracentrifuge. All surgical procedures were performed as described⁵⁴. Briefly, 90-d-old mice were anesthetized with ketamine/xylazine intraperitoneally (90–200 mg per kg of body weight ketamine/10 mg per kg xylazine). A 2-cm longitudinal skin incision was made above the lumbar region under a dissecting microscope. Using a dental drill, a small (1 mm) hole was made into the spinal cord. The lentiviral solution was injected into the L3–L4 region using a stereotaxic frame (Stoelting) at 2 mm unilaterally. The viral solution (50 nl) was injected 25 times per animal with 45-s intervals using a fine micropipette (Drummond 30- μ l microcapillaries pulled with P-9 capillary puller, Sutter Instruments) and a Nanoject II (Drummond). The micropipette was then left for an additional 5 min and gently withdrawn.

Statistics

Statistical analyses were done using GraphPad Prism 4 software. Bar graphs are presented as the mean \pm s.e.m. For experiments in which only two groups were analyzed, the two-tailed unpaired *t* test was used. No statistical methods were used to pre-determine sample sizes, but our sample sizes are similar to those generally employed in the field. Data distribution was assumed to be normal, but this was not formally tested. Blinding and randomization was performed on weight and survival data analyses.

A Supplementary Methods Checklist is available.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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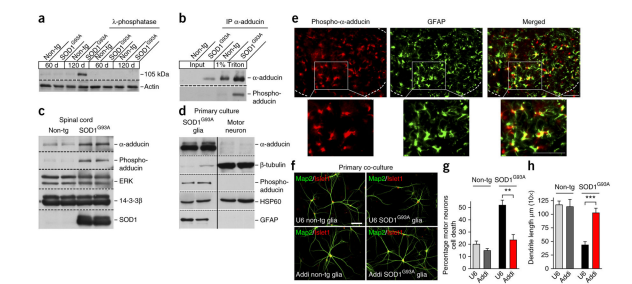


Figure 1.

Upregulation of α-adducin in SOD1^{G93A} astrocytes mediates non-cell autonomous degeneration of motor neurons. (a) Lysates of spinal cord from symptomatic SOD1^{G93A} transgenic mice and control non-transgenic mice were subjected to immunoblotting using an antibody that recognizes phosphorylation events in cells following exposure to oxidative stress. Lysates in lanes 5–8 were incubated with λ -phosphatase, which largely eliminated the 105-kDa immunoreactive band. (b) Lysates of spinal cords from symptomatic SOD1^{G93A} and control mice were subjected to immunoprecipitation using the α -adducin antibody followed by immunoblotting with our phospho-antibody. (c) Immunoblots from spinal cord lysates showed an increase in a-adducin and phosphorylated a-adducin relative to the internal control proteins ERK and 14-3-3ß in symptomatic SOD1G93A mice as compared with control wild-type littermates (120 d). (d) Immunoblots revealed that α -adducin and phosphorylated a-adducin were predominately expressed in primary glial cultures enriched with the astrocyte marker GFAP relative to primary motor neuron cultures enriched with the neuron marker β -tubulin. HSP60 was used as an internal control (lower panel). Blots shown in a-d are cropped. Full-length blots are presented in Supplementary Figure 11. (e) Immunohistochemistry of symptomatic SOD1^{G93A} lumbar spinal cord sections revealed that phosphorylated-Ser436 a-adducin (phospho-a-adducin) colocalized with the astrocyte protein GFAP. Dashed line indicates ventral horn. The boxed areas are shown at high magnification; scale bars represent 50 um. (f) Co-cultured astrocytes and motor neurons were subjected to immunocytochemistry with antibodies recognizing the motor neuron nuclear protein Islet1 (red) and the dendrite protein Map2 (green); scale bar represents 50 μ m. Wild-type astrocytes transfected with the control U6 or α -adducin RNAi plasmid had little or no effect on motor neuron morphology or survival (upper and lower left panels). Control U6 SOD1^{G93A} astrocytes induced non-cell autonomous motor neuron cell death and dendrite abnormalities (upper right panel). α-adducin knockdown in SOD1^{G93A} astrocytes protected motor neurons against the non-cell autonomous cell death and dendrite abnormality (lower right panel). (g,h) Quantification of motor neuron survival was derived from *n* 900 cells per condition and values presented are the average of three independent

experiments; two-tailed unpaired *t* test, P = 0.0095. Quantification of dendrite length was derived from *n* 240 images per condition and values presented are the average of three independent experiments; two-tailed unpaired *t* test, P = 0.0001. Data are presented as mean \pm s.e.m. **P < 0.01, ***P < 0.001.

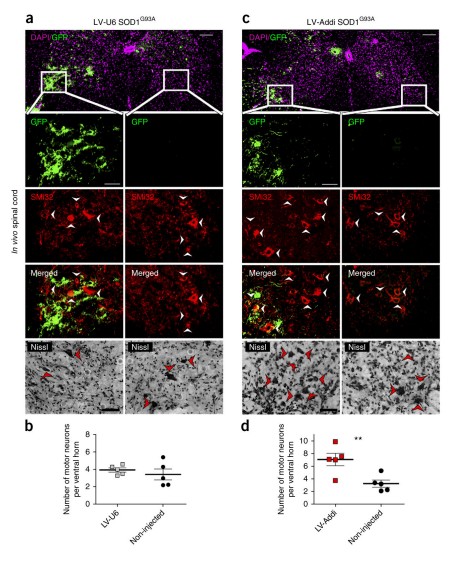
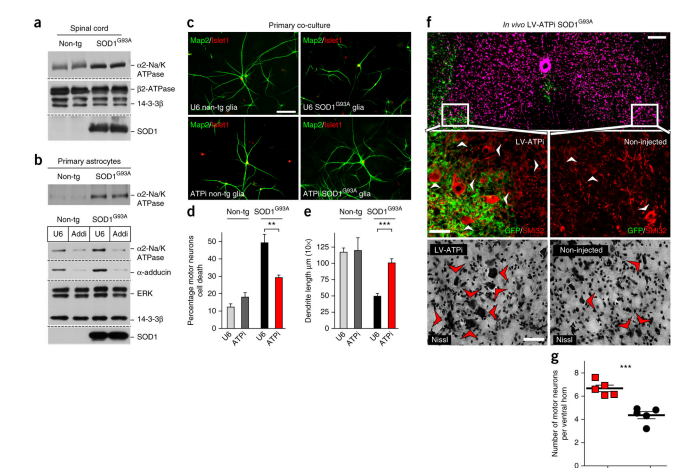


Figure 2.

Knockdown of α -adducin in SOD1^{G93A} mice suppresses motor neuron degeneration *in vivo*. (**a,c**) Spinal cords from SOD1^{G93A} mice injected intraspinally were subjected to immunohistochemistry at end stage. End stage was defined as a time point at which the animal was unable to upright itself within 30 s of placement on its side. Immunohistochemistry with GFP in sections of the spinal cord in SOD1^{G93A} mice lumbar revealed delivery of control virus (LV-U6; top, **a**) or α -adducin RNAi virus (LV-Addi; top, **c**) into the ventral horn; scale bars represent 100 µm. Alternating GFP-positive sections were subjected to immunohistochemistry using the GFP antibody and the neurofilament-SMi32 antibody (red), a motor neuron marker, or Nissl stained (bottom) for quantification of surviving motor neurons in GFP-labeled injected ventral horn and contralateral non-injected Ventral horn (n = 20 sections per animal); scale bars represent 50 µm. (**b**) Control LV-U6 SOD1^{G93A} mice (n = 5) displayed equivalent degeneration of motor neurons in injected GFP-labeled ventral horn and non-injected contralateral ventral horn. (**d**) α -adducin knockdown in SOD1^{G93A} mice (LV-Addi SOD1^{G93A}; n = 5) increased motor neuron

survival in GFP-labeled injected ventral horn as compared with non-injected contralateral ventral horn. Two-tailed unpaired *t* test: LV-U6, P = 0.340; LV-Addi, P = 0.0100. Data are presented as mean \pm s.e.m. **P < 0.01.

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LV-ATPi Non-injected

Figure 3.

Enrichment of the a2-Na/K ATPase/a-adducin complex in SOD1^{G93A} astrocytes triggers motor neuron degeneration. (a) Immunoblots from spinal cord lysates showed an increase in the protein levels of α 2-Na/K ATPase relative to the internal control proteins β 2-ATPase and 14-3-3β in symptomatic SOD1^{G93A} mice as compared with control wild-type littermates (120 d). (b) Immunoblots revealed that α 2-Na/K ATPase was upregulated in SOD1^{G93A} astrocytes as compared with non-transgenic controls. Knockdown of a-adducin in SOD1^{G93A} astrocytes attenuated a2-Na/K ATPase protein levels. Protein levels are relative to ERK and 14-3-3 β . Blots shown in **a** and **b** are cropped. Full-length blots are presented in Supplementary Figure 11. (c) Co-cultured astrocytes and motor neurons were subjected to immunocytochemistry with the motor neuron nuclear protein Islet1 (red) and the dendrite protein Map2 (green); scale bar represents 50 µm. Wild-type astrocytes transfected with the control U6 or a2-Na/K ATPase RNAi plasmid had little or no effect on motor neuron morphology or survival (upper and lower left panels). Control U6 SOD1^{G93A} astrocytes induced non-cell autonomous motor neuron cell death and dendrite abnormalities (upper right panel). a2-Na/K ATPase knockdown in SOD1G93A astrocytes protected motor neurons against the non-cell autonomous cell death and dendrite abnormalities (lower right panel). (\mathbf{d}, \mathbf{e}) Quantification of motor neuron survival was derived from n = 900 cells per condition and values presented are the average of three independent experiments; two-tailed unpaired t

test, P = 0.0080. Quantification of dendrite length was derived from n = 240 images per condition and values presented are the average of three independent experiments; two-tailed unpaired t test, P = 0.0001. Data are presented as mean \pm s.e.m. **P < 0.01, ***P < 0.001. (f,g) Alternating GFP-positive sections from SOD1^{G93A} mice injected intraspinally with lentivirus expressing α 2-Na/K ATPase RNAi were subjected to immunohistochemistry using the GFP and neurofilament-SMi32 (red) antibodies or Nissl stained (bottom panels) for quantification of surviving motor neurons in GFP-labeled injected ventral horn and contralateral non-injected ventral horn at end stage (n = 20 sections per animal); scale bars represent 100 (top) and 50 (middle, bottom) µm. End stage was defined as a time point at which the animal was unable to upright itself within 30 s of placement on its side. α 2-Na/K ATPase knockdown in SOD1^{G93A} mice (LV-ATPi SOD1^{G93A}, n = 5) increased motor neuron survival in GFP-labeled injected ventral horn as compared with non-injected contralateral ventral horn. Arrowheads indicate surviving motor neurons (quantified in g). Two-tailed unpaired t test, P = 0.0005. Data are presented as mean \pm s.e.m. ***P < 0.001.

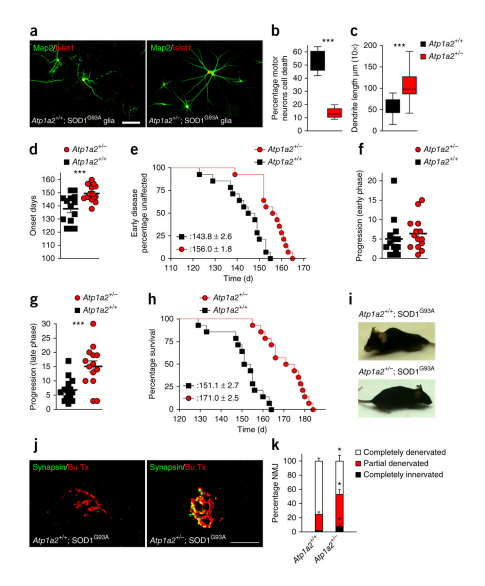


Figure 4.

Heterozygous disruption of Atp1a2 in SOD1^{G93A} mice suppresses motor neuron degeneration and enhances mouse lifespan. (**a**) Downregulation of α 2-Na/K ATPase in SOD1^{G93A} astrocytes by crossing SOD1^{G93A} mice with $Atp1a2^{+/-}$ (right) protected motor neurons from non–cell autonomous cell death and dendrite abnormalities induced by control SOD1^{G93A} astrocytes (left). Scale bar represents 50 µm. (**b**,**c**) Quantification of motor neuron survival is derived from n = 900 cells per condition and values presented are the average of three independent experiments; unpaired t test, P = 0.0003. Quantification of dendrite length is derived from n = 240 images per condition and values presented are the average of three independent experiments performed in duplicates; two-tailed unpaired ttest, P = 0.0001. Data are presented as mean \pm s.e.m. ***P < 0.001. (**d**) Disease onset, that is, initial day of weight loss, was significantly delayed in $\alpha Atp1a2^{+/-}$; SOD1^{G93A} mice ($Atp1a2^{+/-}$, n = 14) as compared with control SOD1^{G93A} littermates ($Atp1a2^{+/+}$, n = 14); two-tailed unpaired t test, P = 0.0009. (**e**) Early disease process, that is, age at which 10% of weight loss is reached, was significantly delayed in $Atp1a2^{+/-}$; SOD1^{G93A} mice ($Atp1a2^{+/-}$;

SOD1^{G93A}, red circles, n = 14) as compared with control SOD1^{G93A} littermates (Atp1a2^{+/+}; SOD1^{G93A}, black squares, n = 14) (P = 0.0001). (f) Early phase disease progression, that is, days from onset to 10% weight loss, displayed no change between $\alpha Atp1a2^{+/-}$; SOD1^{G93A} mice and control SOD1^{G93A} littermates; two-tailed unpaired t test, P = 0.2023. (g) Latephase disease progression, that is, from 10% weight loss to end stage, displayed significant delay in $Atp1a2^{+/-}$; SOD1^{G93A} mice ($Atp1a2^{+/-}$, n = 14) as compared with control SOD1^{G93A} littermates ($Atp1a2^{+/+}$, n = 14); two-tailed unpaired t test, P = 0.0005. (h) Kaplan-Meier survival plots showed substantial and significant increase in lifespan for $Atp1a2^{+/-}$; SOD1^{G93A} mice ($Atp1a2^{+/-}$; SOD1^{G93A}, red circles, n = 14) as compared with control SOD1^{G93A} littermates ($Atp1a2^{+/+}$; SOD1^{G93A}, black squares, n = 14) (P = 0.0001). (i) Control SOD1^{G93A} mice displayed morbidity of reduced mobility and the inability to upright at endstage ($Atp1a2^{+/+}$; SOD1^{G93A}), whereas age-matched SOD1^{G93A} littermates that harbor a heterozygous null Atp1a2 allele displayed increased mobility and health (Supplementary Movies 1 and 2). (j) Immunohistochemistry of NMJs from gastrocnemius muscle using the presynaptic marker synapsin (green) and postsynaptic marker abungarotoxin (red, Bu Tx) at 120 d showed that control SOD1^{G93A} mice had denervated NMJs (left), whereas $Atp1a2^{+/-}$ displayed an increased NMJ integrity (right); scale bar represents 20 µm. (k) Quantification of NMJs from n = -300 NMJs from three animals per group (two-tailed unpaired t test: complete denervated, P = 0.0196; partial denervated, P =0.0203; completely innervated, P = 0.0370). Data are presented as mean \pm s.e.m. *P < 0.05.

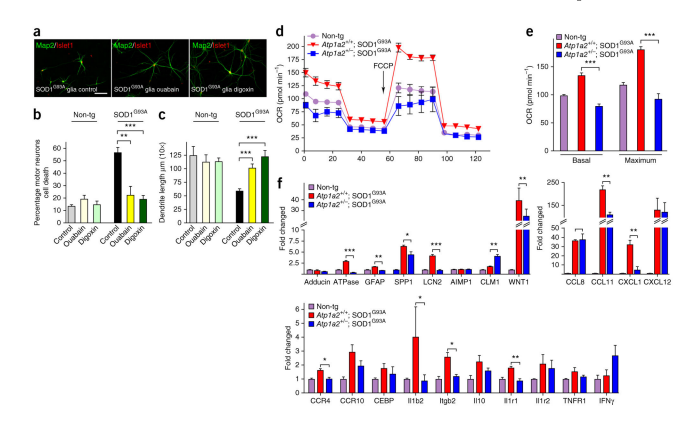


Figure 5.

Na/K ATPase stimulates mitochondrial respiration and expression of inflammatory genes in SOD1^{G93A} astrocytes. (a) Control SOD1^{G93A} astrocytes (left) induced non-cell autonomous motor neuron cell death and dendrite abnormalities. Pharmacological inhibition of Na/K ATPase with ouabain (middle) and digoxin (right) was neuroprotective from the non-cell autonomous cell death and dendrite abnormalities induced by SOD1^{G93A} astrocytes (left); scale bar represents 50 µm. Pharmacological inhibition of Na/K ATPase in control cocultures non-transgenic astrocytes (Non-tg) and motor neurons with ouabain and digoxin did not alter motor neuron survival or morphology. (b,c) Quantification of motor neuron survival was derived from n = 900 cells per condition and values presented are the average of three independent experiments (two-tailed unpaired t test: ouabain, P = 0.0024; digoxin, P = 0.0001). Quantification of dendrite lengths was derived from n = 240 images per condition and values presented are the average of three independent experiments (two-tailed unpaired t test: ouabain, P = 0.0002; digoxin, P = 0.0001). Data are presented as mean \pm s.e.m. **P < 0.01, ***P < 0.001. (d) A representative plot of oxygen consumption measured in astrocytes from control non-transgenic, SOD1G93A and heterozygous-null SOD1G93A mice using Seahorse Bioscience XF Analyzer. Arrow indicates time when the mitochondrial uncoupler FCCP was added. (e) Basal and maximum oxygen consumption were quantified in control non-transgenic (n = 3), SOD1^{G93A} (n = 3) and heterozygous-null SOD1^{G93A} mice (n = 3) astrocytes using Seahorse Bioscience XF Analyzer (two-tailed unpaired t test: basal consumption, P = 0.0001; maximum consumption, P = 0.0001). Data are presented as mean \pm s.e.m. ****P* < 0.001. (f) Total RNA of astrocytes from control non-transgenic (*n* = 3), SOD1^{G93A} (n = 3) and heterozygous-null SOD1^{G93A} mice (n = 3) were subjected to qRT-

PCR analyses using primers to a panel of pro-inflammatory genes. Gene expression was normalized to GAPDH expression. The expression of 18 inflammatory genes was upregulated in SOD1^{G93A} astrocytes. Downregulation of α 2-Na/K ATPase in SOD1^{G93A} astrocytes significantly decreased expression of half of the upregulated inflammatory genes (two-tailed unpaired *t* test: SPP1, *P* = 0.0253; LCN2, *P* = 0.0013; CLM1, *P* = 0.0052; WNT1, *P* = 0.0198; CCL11, *P* = 0.0387; CXCL1, *P* = 0.0246; CCR4, *P* = 0.0314; Il1b2, *P* = 0.0421; Itgb2, *P* = 0.0215; Il1r1, *P* = 0.0092). Data are presented as mean ± s.e.m. **P* < 0.05, ***P* < 0.01, ****P* < 0.001.

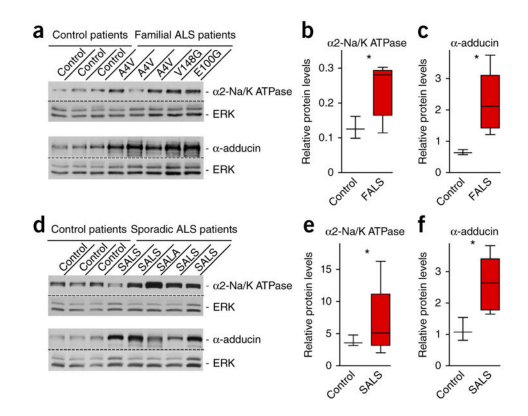


Figure 6.

The α 2-Na/K ATPase/ α -adducin complex is upregulated in spinal cord in familial and sporadic ALS patients. (a) Immunoblots from patients with familial ALS (n = 5) revealed elevated protein levels of α 2-Na/K ATPase (top) and α -adducin (bottom) in spinal cord lysates as compared to control patients (n = 3). ERK served as a loading control. (b,c) Quantification of the relative densitometry protein levels for α 2-Na/K ATPase and α adducin relative to the internal control ERK; two-tailed unpaired *t* test: α 2-Na/K ATPase, *P* = 0.0302; α -adducin, *P* = 0.0336. Blots shown in **a** and **b** are cropped. Full-length blots are presented in Supplementary Figure 11. (d) Immunoblots from patients with sporadic ALS (n = 5) showed elevated protein levels of α 2-Na/K ATPase (top) and α -adducin (bottom) in spinal cord lysates as compared with control patients (n = 3). (e,f) Quantification of the relative densitometry protein levels for α 2-Na/K ATPase, P = 0.0467; α -adducin, P = 0.0181. Data are presented as mean \pm s.e.m. *P < 0.05.