SOD1 targeted to the mitochondrial intermembrane space prevents motor neuropathy in the Sod1 knockout mouse

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Motor axon degeneration is a critical but poorly understood event leading to weakness and muscle atrophy in motor neuron diseases. Here, we investigated oxidative stress-mediated axonal degeneration in mice lacking the antioxidant enzyme, Cu,Zn superoxide dismutase (SOD1). We demonstrate a progressive motor axonopathy in these mice and show that Sod1−/− primary motor neurons extend short axons in vitro with reduced mitochondrial density. Sod1−/− neurons also show oxidation of mitochondrial—but not cytosolic—thioredoxin, suggesting that loss of SOD1 causes preferential oxidative stress in mitochondria, a primary source of superoxide in cells. SOD1 is widely regarded as the cytosolic isoform of superoxide dismutase, but is also found in the mitochondrial intermembrane space. The functional significance of SOD1 in the intermembrane space is unknown. We used a transgenic approach to express SOD1 exclusively in the intermembrane space and found that mitochondrial SOD1 is sufficient to prevent biochemical and morphological defects in the Sod1−/− model, and to rescue the motor phenotype of these mice when followed to 12 months of age. These results suggest that SOD1 in the mitochondrial intermembrane space is fundamental for motor axon maintenance, and implicate oxidative damage initiated at mitochondrial sites in the pathogenesis of motor axon degeneration.

Keywords: SOD; axon; neuromuscular junction; motor neuron disease; mitochondria

Abbreviations: O2− = superoxide; SOD1 = Cu,Zn superoxide dismutase

Introduction

Motor axons are the anatomical and functional link between spinal motor neurons and skeletal muscles. Degeneration of motor axons at the neuromuscular junction is an early feature of motor neuron disease in animal models (reviewed in Fischer and Glass, 2007) and in humans (Bjornskov et al., 1984; Maselli et al., 1993). Moreover, axonal degeneration is sufficient to cause weakness...
and muscle atrophy, even in the absence of motor neuron cell death (Gould et al., 2006; Rouaux et al., 2007; Suzuki et al., 2007). Axon degeneration is therefore a key pathologic event and therapeutic target in motor neuron disease, although the pathogenic mechanisms that initiate axon degeneration are poorly understood.

Evidence from Cu,Zn superoxide dismutase (SOD1) knockout mice suggests that this antioxidant enzyme is essential for motor axon maintenance. Sod1−/− mice develop normally and lack behavioral deficits up to 6 months of age (Reaume et al., 1996), but ageing Sod1−/− mice perform poorly on the Rotarod and exhibit accelerated skeletal muscle atrophy (Muller et al., 2006). Spinal motor neuron and ventral root axon numbers are normal at 17 and 19 months, respectively (Flood et al., 1999; Shefner et al., 1999), but muscle fibre atrophy and fibre-type grouping, suggestive of chronic denervation and reinnervation, are detectable by 6 months (Flood et al., 1999). EMG recordings also show spontaneous activity and progressive loss of motor units (Shefner et al., 1999). Thus, genetic deletion of SOD1 spares motor neurons and proximal axons, but may be detrimental to distal motor axons.

SOD1 is one of three superoxide dismutases in mammalian cells that catalyse the conversion of superoxide (O2−) to H2O2 (McCord and Fridovich, 1969). SOD1 is traditionally considered to be the cytosplasmic isoform, SOD2 the mitochondrial isoform and SOD3 the secreted form (Zelko et al., 2007). However, SOD1 is also found in the mitochondrial intermembrane space (Weisiger et al., 2001; Iñárrrea, 2002; Vijayvergiya et al., 2005). The functional significance of SOD1 in the intermembrane space remains to be defined, although a protective role is likely. Mitochondria isolated from Sod1−/− mouse muscle (Muller et al., 2007; Jang et al., 2010) and Sod1−/− Caenorhabditis elegans (Yanase et al., 2009) exhibit increased generation of reactive oxygen species including O2− and H2O2. Similarly, antisense knockdown of SOD1 in vitro causes preferential oxidation of mitochondrial—not cytosolic—proteins, loss of mitochondrial membrane potential and decreased ATP production (Aquilano et al., 2006). Thus, loss of SOD1 may result in an accumulation of mitochondrial reactive oxygen species, leading to oxidative damage and mitochondrial dysfunction.

Here, we demonstrate that targeted replacement of SOD1 only in the mitochondrial intermembrane space rescues motor axon outgrowth and normalizes the mitochondrial redox state in Sod1−/− neurons in vitro, and prevents weakness and neuromuscular junction denervation in Sod1−/− mice followed up to 12 months of age. These data suggest that mitochondrial oxidative stress is an underlying cause of distal motor axonopathy, and demonstrate that localization of SOD1 in the intermembrane space is sufficient for the survival of motor axons.

Materials and methods

Animals

Sod1−/− mice, generated by Huang and colleagues (1997), were obtained from Marie Csete (Emory University) (Muller et al., 2006). Sod1−/− males were crossed with thy1-YFP16 females (Feng et al., 2000) to obtain Sod1−/−, thy1-YFP16 breeders, expressing yellow fluorescent protein (YFP) in all motor axons. Both transgenic lines were in a C57BL/6 background. For ease of description, thy1-YFP16 is omitted from the genotype and expression should be assumed. Mice were housed in microisolator cages on a 12 h light–dark cycle and given free access to food and water. Genotyping was by standard polymerase chain reaction analysis on tail-snip DNA. YFP status was determined by fluorescent examination of epidural nerve fibres in ear punch biopsies.

To generate mitoSOD1 transgenic mice, human SOD1 complementary DNA was inserted in a prion promoter vector (MoPPrXho) (Borchelt et al., 1996) at the Xhol site, between PrP exons 2 and 3. The start codon of human SOD1 was removed and substituted by an in-frame DNA linker of 6 nt, containing a BamHi site and coding for Gly–Ser. We then inserted a 561 nt complementary DNA encoding the first 187 amino acids of mouse mitofilin (GenBank accession code: NM_029673) (John et al., 2005) in frame with the 5′-end of the human SOD1 plus linker. The mitofilin complementary DNA was obtained by polymerase chain reaction amplification of a mouse complementary DNA library using primer sequences derived from the mouse genome database.

The plasmid was microinjected into fertilized eggs of B6CBF1 mice. Offspring harbouring the transgene were identified by polymerase chain reaction that identified the following primers: CCGCTCGAGATGCTGCGGGCTGTTCAG (sense) and CCGCTCGAGTTATTGGGCGATCCCAAT (antisense) to generate a 1027 bp product. Five lines of mitoSOD1 transgenic mice were identified. Male founders were crossed with B6SJLF1/J females (Jackson).

A two-step mating scheme was used to generate mice expressing only mitoSOD1. Sod1−/− males were crossed with mitoSOD1 females to generate F1 heterozygotes (mitoSOD1,Sod1+/−). Sod1−/− and mitoSOD1,Sod1+/− mice were then crossed to generate the target genotype (mitoSOD1,Sod1−/−) and littermate controls.

Neuromuscular junction morphology

Tibialis anterior muscle was dissected, pinned in mild stretch and immersion fixed for 30 min in 4% paraformaldehyde at room temperature. Fixed muscles were cryoprotected in 30% sucrose/phosphate-buffered saline (48–72 h at 4 °C), flash frozen in supercooled isopentane and 35 μm frozen sections were cut longitudinally through the entire muscle. Acetylcholine receptors at the motor endplate were labelled with Alexa Fluor 555-conjugated α-bungarotoxin (Invitrogen), 1:5000 in phosphate-buffered saline (30 min, room temperature). Motor axon terminals were identified by YFP fluorescence. Innervated, intermediate and denervated endplates were defined by complete, partial or absent overlap between nerve terminal and endplate, respectively. All endplates were evaluated in every fourth section. No difference was seen between innervation in male and female mice, therefore the data were pooled.

Mitochondrial isolation from mouse tissues

Tissues (brain and spinal cord) were homogenized in a Dounce homogenizer in MS-EGTA buffer (225 mM mannitol, 75 mM sucrose, 5 mM HEPES, 1 mM EGTA, pH 7.4), and centrifuged at 2000 g for 5 min (4 °C). The supernatant was centrifuged at 15 000 g for 20 min to generate the cytosolic fraction (supernatant) and crude mitochondrial fraction (pellet). The cytosolic fraction was centrifuged twice (22 000 g for 20 min) to eliminate membrane contamination. The crude
mitochondrial fraction was resuspended and washed twice in MS-EGTA. To prepare purified mitochondria, the crude mitochondrial pellet was layered onto 9 ml of 23% Percoll in MS-EGTA and centrifuged at 25 000g for 11 min. The pellet was resuspended in MS-EGTA and centrifuged three times at 14 000g for 14 min. The final purified mitochondrial pellet was resuspended in MS-EGTA at ~10 mg/ml. All reagents were from Sigma.

Western blot analyses

The expression and mitochondrial localization of mitoSOD1 were tested by western blot of crude mitochondria prepared from spinal cord. Proteins in whole homogenates (50 μg), cytosol (20 μg) and mitochondria (20 μg) were separated on a 12% sodium dodecyl sulphate polyacrylamide gel, transferred to polyvinylidene difluoride membranes (BioRad) and immunoblotted with sheep anti-SOD1 (1:5000, Calbiochem), mouse anti-Tim23 (1:5000, Stressgen), goat anti-Akt1 (1:5000, Santa Cruz) or goat anti-Hsp60 (1:5000, Stressgen).

Blue native gel electrophoresis

Spinal cord mitochondrial proteins (50 μg) were separated on a 10–16% gradient blue native gel as previously described (Schagger and von Jagow, 1991). After electrophoresis, proteins were transferred to polyvinylidene difluoride and immunoblotted for SOD1 and core II subunit of complex III (mouse anti-CIII, 1:2500, Molecular Probes). A high molecular weight ladder (Amersham) was used to estimate protein size.

Alkaline extraction of mitochondrial proteins

Alkaline extraction of mitochondrial proteins was performed as described (Vijayvergiya et al., 2005). Briefly, mitochondria (50 μg) were incubated in the presence or absence of 0.1 M Na₂CO₃ (pH 11.5), with or without Triton X-100 (1%), for 30 min at 4°C, then centrifuged at 91000g for 25 min. The pellet was saved. Supernatant proteins were precipitated with ice-cold 12% trichloroacetic acid and centrifuged at 18000g for 15 min, followed by an ice-cold acetone wash. Pellet and precipitated supernatant proteins were analysed by western blot as described above.

Mitoplast preparation

Mitoplasts were prepared as previously described (Acin-Perez et al., 2009). Briefly, purified mitochondria (300 μg) were suspended in MS-EGTA, water (1/10 volume) and digitonin (1 mg digitonin/5 mg mitochondrial protein), and the mixture was incubated on ice for 45 min. KCl (150 mM) was then added, followed by incubation for 2 min on ice and centrifugation at 18000g for 20 min. The mitoplast pellet was resuspended to 0.5 mg/ml in MS-EGTA. The supernatant containing the post-mitoplast fraction was subjected to trichloroacetic acid precipitation as described above.

Proteinase K treatment of mitoplasts

Proteinase K treatment of mitoplasts was performed as previously described (Vijayvergiya et al., 2005). Briefly, mitoplasts (25 μg) were incubated in the presence or absence of proteinase K (20 μg/ml), with or without Triton X-100 (0.1%), for 1 h on ice. Proteolysis was stopped by adding 2 mM phenylmethylsulphonyl fluoride.

SOD1 activity: spectrophotometric assay

SOD1 activity was assessed as previously described (Vives-Bauza et al., 2007) with minor modifications. The assay measures the reduction of acetylated cytochrome c by O₂⁻ generated by xanthine/xanthine oxidase. All reagents were from Sigma except for xanthine oxidase (Calbiochem). Spinal cord mitochondria (50 μg) and cytosolic fractions (10 μg) were incubated in 1 ml of reaction buffer (50 mM phosphate buffer, pH 7.8, 0.1 mM EDTA, 1 mM NaN₃, 100 mM xanthine, 2 U xanthine oxidase and 25 mM partially acetylated cytochrome c). Cytochrome c reduction was followed spectrophotometrically at 550 nm, at 25°C for 3 min. The activity of SOD1 (KCN-insensitive) was determined by adding 2 mM KCN, and subtracting residual activity from total SOD activity. One unit of SOD activity was defined as the amount of enzyme required to inhibit the rate of reduction of cytochrome c by 50%. Note that in measuring SOD1 activity in mitochondria it is necessary to minimize the interference associated with the interaction of cytochrome c with cytochrome c oxidase and cytochrome c reductases, using partially acetylated cytochrome c as described (Azzi et al., 1975; Kuthan et al., 1986). Nevertheless, we found some residual cytochrome c reduction in the mitochondria of Sod1−/− samples (Fig. 5E). This can be explained by the fact that the preparation of acetylated cytochrome c actually contains up to 40% non-acetylated cytochrome c, which can react with mitochondrial oxidases and reductases.

Primary motor neuron culture

Spinal motor neurons were enriched from E12.5 mouse embryos by density centrifugation (Zhang et al., 2006). Spinal cords were isolated, incubated in 0.05% trypsin (Worthington, 37°C, 10 min) and dissociated by pipetting up and down in neurobasal medium (Invitrogen) containing 0.1% trypsin inhibitor (Sigma), 100 μM Dnase (Worthington) and 0.4% bovine serum albumin. Cells were centrifuged through 4% bovine serum albumin (400g, 5 min), resuspended and centrifuged through 10% (v/v) Optiprep (Axis Shield, 700g, 10 min). The interface was aspirated, centrifuged through a second bovine serum albumin cushion, and resuspended in growth medium consisting of neurobasal with 2% B27 supplement minus antioxidants (Invitrogen), 2% horse serum (Invitrogen), 0.5 mM Glutamax (Invitrogen), BDNF, CNTF, GDNF and NT-3 (10ng/ml, Peprotech). Cells were plated on coverslips coated with polyornithine (10 μg/ml, Sigma) and Matrigel (1:25, BD Bioscience). By this method, 84.7 ± 4.8% of cells at 24 h were immunoreactive for the motor neuron marker, HB9 (1:1000, Abcam). Individual spinal cords were kept separate during cell isolation and genotype subsequently determined by polymerase chain reaction.

Axon length was determined by systematic random sampling of cells along a pre-marked grid. Motor neurons were identified morphologically under phase contrast and photographed at ×20 on an Olympus IX51 inverted microscope using an Olympus Qcolor3 digital camera. Axons were manually traced and measured using ImageJ software (http://rsb.info.nih.gov/ij/). In cells with multiple processes, the axon was considered to be the longest process. The average number of neurons analysed in each of four trials was 130 ± 41 (Sod1+/−), 158 ± 34 (Sod1−/−) and 163 ± 19 (Sod1−/−), or 500–600 neurons per group (n = 4 was used for statistical analysis).
Mitochondrial density

Mitochondrial density in axons was evaluated using Mitotracker Red CM-H$_2$XROS (Invitrogen). Dye was added to cells at 250 nM in serum-free medium (30 min, 37°C). Cells were returned to growth medium for 10 min and fixed in pre-warmed 4% paraformaldehyde (15 min, room temperature). Coverslips were inverted onto slides using anti-fade mounting medium (Vectorshield+DAPI; Vector Labs), and cells were examined under standard fluorescence microscopy. In some experiments, Mitotracker staining was followed by immuno-labeling with an antibody specific for human SOD1 (Sigma, 1:500). Twenty-five neurons per coverslip were selected and photographed at $\times40$ magnification by unbiased coverslip scanning. Mitochondrial density was evaluated morphologically using ImageJ software. Lengths of individual mitochondria were measured, combined and divided by the length of the axon. At least three replicates per genotype were obtained over the course of multiple motor neuron preparations.

Primary cortical neuron cultures

Primary cortical neurons were isolated from E15.5 mouse embryos. Brains were removed in ice-cold dissecting buffer (2.5 mM HEPES, 4 mM NaHCO$_3$, and 35 mM glucose in Hanks’ balanced salt solution) and cortices were dissected free of subcortical structures and meninges and incubated in 0.25% Trypsin/EDTA (Sigma), for 15 min at 37°C. Tissue was dissociated in dissection buffer containing 0.1 mg/ml trypsin inhibitor (Sigma) and 200 U/ml DNase (Worthington). Cells were pelleted by centrifugation at 218g for 5 min and resuspended in growth medium consisting of neurobasal (Invitrogen), 5% foetal bovine serum (Atlanta Biologicals), 200 mM GlutaMAX (Invitrogen) and 2% B27 minus antioxidants (Invitrogen). Cells were plated in poly-L-lysine coated six-well plates at a density of 20000 cells/cm$^2$. Cells were maintained for 3 days in serum-free medium (30 min, 37°C). Twenty-five neurons per coverslip were selected and photographed using the Nernst equation:

$$ E_0 = E_0^\circ + (RT/NF) \times \log([o]/[red]). $$

where $E_0^\circ = -256$ mV (thioredoxin-1) or $-330$ mV (thioredoxin-2). Note that protein levels were not equalized between samples. Only relative changes between reduced and oxidized bands are of quantitative significance.

Confocal imaging

Images were captured on a Zeiss LSM 510 NLO META system, coupled to a Zeiss Axiovert 100M inverted microscope. Neuromuscular junction z-stacks were obtained with a Plan-Neofluar ×40 (NA 1.3) oil objective with optical slice thickness of 1 μm. Motor neurons were imaged with a Plan-Neofluar ×20 (NA 0.3) objective or Plan-Neofluar ×100 (NA 1.3) oil objective. Z-stacks were compressed and images exported using LSM Image Examiner software (Zeiss).

Statistical analysis

Results are expressed as mean ± SD, and comparisons were made by ANOVA with Tukey post hoc analysis ($p=0.05$), unless otherwise specified, using Prism software (GraphPad).

Results

Loss of SOD1 causes a progressive motor neuropathy

Previous studies of the Sod1$^{+/−}$ mouse provide compelling, although indirect, evidence for a motor axonopathy (Flood et al., 1999; Shefner et al., 1999). To facilitate morphological analysis of neuromuscular junctions, we first crossed Sod1$^{−/−}$ mice ( Muller et al., 2006) with thy1-YFP16 mice (Feng et al., 2000) to generate mice expressing YFP in all motor axons. Absence of SOD1 protein and enzymatic activity was verified in Sod1$^{−/−}$, thy1-YFP16 offspring by western blot and SOD1 zymography (Supplementary Fig. 1).

We observed progressive hind-limb weakness in Sod1$^{−/−}$ mice, evidenced by significant loss of grip strength by 12 months (Fig. 1C). Whereas most Sod1$^{+/+}$ and Sod1$^{+/−}$ mice could remain suspended from a wire grid for at least 300 s, the mean latency to fall for Sod1$^{−/−}$ animals was 134 s ($p < 0.001$). Sod1$^{−/−}$ mice had obvious difficulty gripping the wire with their hind limbs, and hind limb grip was typically lost first, followed by forelimb grip (Supplementary Videos 1–4).

To determine how loss of SOD1 affects distal motor axons, we examined neuromuscular junction morphology in the tibialis anterior muscle, which undergoes significant (~50%) atrophy in 20-month-old Sod1$^{−/−}$ mice ( Muller et al., 2006). Neuromuscular junction innervation at 1, 4, 12 and 18 months of age was evaluated by the overlap between YFP-positive motor axon terminals and motor endplates labelled with Alexa Fluor 555-conjugated α-bungarotoxin (Fig. 1A and B). At 1 month, the tibialis anterior muscle was fully innervated in Sod1$^{−/−}$ mice. At 4 months, 69.5 ± 6.2% of endplates were innervated in Sod1$^{−/−}$ mice, compared to 97.1 ± 0.9% and 99.6 ± 0.3% in Sod1$^{+/+}$ and Sod1$^{+/−}$ mice, respectively ($p < 0.001$). By 18 months, only 33.9 ± 8.5% of Sod1$^{−/−}$ endplates were innervated compared to 92.2 ± 2.8% in Sod1$^{+/+}$ mice and 79.9 ± 15.5% in Sod1$^{+/−}$ mice ($p < 0.001$). This demonstrates that SOD1 is required for maintenance of motor neuron function.
axons in vivo, and suggests that distal motor axons are sensitive to oxidative stress-mediated injury.

**Axon defects in primary motor neuron cultures**

To examine axonal defects intrinsic to Sod1−/− motor neurons, we cultured primary motor neurons from E12.5 mice in medium free of standard antioxidant supplements (Fig. 2A). Sod1−/− motor neurons were short-lived compared with controls. No viable Sod1−/− cells remained at 72 h in culture. At 24 h, Sod1−/− motor neurons had markedly shorter axons compared with controls (Fig. 2C). Mean axon length in Sod1−/− cells (±SEM) was 50±3 μm at 24 h, compared with 93±5 and 83±3 μm for Sod1+/+ and Sod1+/− cells, respectively (P < 0.001).

Given a previous report of mitochondrial damage in Sod1−/− cells (Aquilano et al., 2006), we tested whether mitochondrial density was altered in Sod1−/− axons (Fig. 2B). At 24 h in culture, mitochondria were labelled with Mitotracker Red-CM-H2XRos, the motor neurons were fixed and individual mitochondria along the length of the axon were measured. Mitochondrial density was expressed as the cumulative length of all mitochondria in the axon, divided by the total length of the axon (Fig. 2D).
By this approach, a 40% decrease in mitochondrial density was observed in Sod1−/− axons compared to Sod1+/+ axons (P < 0.01).

The mitochondrial redox state in neurons depends on SOD1 expression

We then investigated how deletion of SOD1 in neurons influences the cytosolic versus mitochondrial redox state. To avoid introducing oxidation by subcellular fractionation, we took advantage of the fact that mitochondria have a distinct thioredoxin system from the cytosol, and these can be resolved by using different antibodies on the same sample. Thioredoxins are small, multi-functional proteins with a highly conserved active site dithiol motif (Arné r and Holmgren, 2000). They serve as protein disulphide reductases and participate in reactive oxygen species removal via peroxiredoxins. Redox western analysis of thioredoxin-1 (cytosolic) versus thioredoxin-2 is a sensitive measure of compartmental changes that can occur independent of overall cellular oxidation (Halvey et al., 2005; Hansen et al., 2006a, b).

Primary cortical neurons were isolated from Sod1−/− mice and littermate controls, maintained in culture for 3 days, and harvested for redox western visualization of oxidized and reduced thioredoxin (Fig. 3). Sod1+/+ neurons showed a thioredoxin-1 redox potential of −273 ± 2 mV and a thioredoxin-2 redox potential of −365 ± 2 mV. These values are consistent with previous data from other cell types, showing that thioredoxin-1 is relatively oxidized at baseline compared with thioredoxin-2 (Halvey et al., 2005; Hansen et al., 2006a). The thioredoxin-1 redox potential in Sod1−/− neurons (−271 ± 1 mV) was identical to wild-type. However, Sod1−/− neurons showed significant oxidation of thioredoxin-2, with a redox potential of −342 ± 7 mV, representing a +23 mV shift toward a more oxidizing potential as compared to controls (P < 0.01). This demonstrates oxidative stress occurring...
within mitochondria of $\text{Sod}1^{-/-}$ neurons, at a time when the cytosolic redox state is unchanged.

## Targeting of wild-type human SOD1 to the mitochondrial intermembrane space

To test the hypothesis that SOD1 in the mitochondrial intermembrane space plays a role in motor axon maintenance, transgenic mice were generated that express wild-type human SOD1 targeted to the mitochondrial intermembrane space (Fig. 4A). We fused the N-terminal of SOD1 to the mitochondrial targeting sequence of mouse mitofilin, a mitochondrial inner membrane protein (John et al., 2005). The 34 amino acid mitofilin targeting sequence, cleavable by matrix metalloproteinases, was followed by 153 amino acids of the mature protein, including 26 amino acids of the transmembrane domain and a 2 amino acid linker between mitofilin and the N-terminus of human SOD1. The resulting construct coded for a 38 kDa chimeric protein (mitoSOD1). This targeting strategy was aimed at anchoring human SOD1 to the outer side of the mitochondrial inner membrane, facing the intermembrane space, while conferring enough flexibility for SOD1 to fold and mature into the active enzyme. Tethering to the inner membrane was designed to prevent unwanted escape of mitoSOD1 into the cytosol.

### Generation and characterization of mice expressing only mitoSOD1

Next we crossed mitoSOD1 and $\text{Sod}1^{-/-}$ mice in a two-step breeding process, to generate $\text{mitoSOD1, Sod}1^{-/-}$ mice expressing SOD1 only in the intermembrane space. Littermate $\text{mitoSOD1, Sod}1^{+/+}$ and generation of transgenic mice, the construct was cloned downstream of the mouse prion promoter, which drives high expression in brain, spinal cord, skeletal muscle, heart and kidney, low expression in lung and spleen and virtually no expression in liver (Wang et al., 2005).

Five lines of transgenic mice with different levels of expression were obtained. For this study, we utilized the line with the highest level of mitoSOD1 expression. The pattern of mitoSOD1 expression in different tissues was investigated by western blot of whole tissue homogenates (Fig. 4B). The 38 kDa mitoSOD1 protein was detected in brain and spinal cord of transgenic mice and not in non-transgenic controls (Fig. 4B). MitoSOD1 was also detected in skeletal muscle, heart and kidney, while lung, spleen and liver expressed low to undetectable levels (Supplementary Fig. 2).
Sod1+/+ mice were used as controls. Fractionation of spinal cord homogenates confirmed that the mitoSOD1 was detected only in the mitochondrial fraction from mitoSOD1,Sod1+/+ and mitoSOD1,Sod1−/− mice, and was undetectable in the cytosol (Fig. 5A). As expected, no mouse SOD1 was detected in the mitochondria or cytosol of Sod1−/− and mitoSOD1, Sod1−/− spinal cords.

Protein alkaline extraction of brain mitochondria from mitoSOD1 mice showed the transgenic protein almost exclusively in the insoluble pellet, whereas cytochrome c, a soluble protein of the intermembrane space, was found in the supernatant (Fig. 5B). Most of the mitoSOD1 transgenic protein was released in the soluble fraction after treatment of the mitochondrial membranes with Triton X-100. This indicates that mitoSOD1 is anchored to the mitochondrial inner membrane through its transmembrane domain.

To study the intra-mitochondrial localization of mitoSOD1, mitoplasts were prepared by removing the mitochondrial outer membrane from mitoSOD1,Sod1+/+ brain mitochondria. MitoSOD1 was associated with mitoplasts and absent from the post-mitoplast supernatant containing the outer membrane and intermembrane space fractions (Fig. 5C). Following digestion of mitoplast surface proteins with proteinase K, most of the mitoSOD1 was degraded to a smaller ~19 kDa fragment. This fragment displayed gel migration properties roughly corresponding to those of human SOD1, which is proteinase K resistant (Vijayvergiya et al., 2005), indicating that proteinase K digested the portion of the protein anchoring SOD1 to the inner membrane. This result was reproduced by solubilization of mitoplasts with Triton X-100 followed by proteinase K digestion. Taken together, these results confirm that mitoSOD1 is bound to the mitochondrial inner membrane facing the intermembrane space.

MitoSOD1 enzyme activity was assessed in whole spinal cord lysates by standard SOD1 zymography (Supplementary Fig. 3). An additional band of KCN-sensitive SOD1 activity was identified in mitoSOD1,Sod1+/+ and mitoSOD1,Sod1−/− mice, demonstrating that mitoSOD1 protein is enzymatically active. To address subcellular localization and avoid disruption of mitoSOD1
membrane tethering, SOD1 activity was also measured spectrophotometrically in spinal cord cytosol and intact mitochondrial fractions (Fig. 5D). We found 3- and 3.5-fold higher SOD1 activity in mitoSOD1,Sod1−/− and mitoSOD1,Sod1+/− mitochondria, respectively, compared to Sod1+/+. Note that these values were obtained after subtraction of non-specific cytochrome c reductase activity present in mitochondria (indicated by the dashed line in Fig. 5D). KCN-sensitive SOD activity was undetectable in the cytosolic fraction of mitoSOD1,Sod1−/− spinal cord. These data confirm that mitoSOD1 is enzymatically active and that its activity is exclusively localized to mitochondria. We also found a moderate but significant reduction of SOD1 activity in mitoSOD,Sod1+/− cytosol, as compared to Sod1+/+. Since the active form of SOD1 is predominantly a dimer (Tainer et al., 1982; Hornberg et al., 2007), we looked for mitoSOD1 oligomerization by blue native gel electrophoresis and western blot of spinal cord mitochondria. In this experiment, proteins are maintained in their native state and separated in a non-denaturing gel, which allows the preservation of protein complexes that are detected with specific antibodies. In mitoSOD1,Sod1+/− and mitoSOD1,Sod1−/− mitochondria, we observed a major band with an estimated size of 100–120 kDa (Fig. 5E) and a smear of higher molecular weight protein complexes, which indicate higher orders of oligomerization than a simple dimer. Since the monomer of mitoSOD1 is 38 kDa, this result suggests that the transgenic protein oligomerizes on the mitochondrial IM. In addition, we observed that mitochondria of mitoSOD1,Sod1−/− contained a larger amount of oligomeric SOD1 than mitoSOD1,Sod1−/−, suggesting that endogenous SOD1 in mitochondria participates to the formation of multiple species of oligomeric complexes.

The proposed scheme for import and maturation of mitoSOD1 in mitochondria is shown in Fig. 5F. We suggest that mitoSOD1 enters the intermembrane space through the translocator of the outer membrane, where the mitochondrial targeting sequence is engaged in the translocator of the inner membrane, exposed to the matrix and cleaved by matrix metalloproteinases. Upon processing, the transmembrane domain allows for the insertion of the protein in the inner membrane with the SOD1 portion facing the intermembrane space. We propose that mitoSOD1 folds and acquires metals in the intermembrane space to reach its mature and enzymatically active state. Active dimers/oligomers of SOD1 may result from the juxtaposition of adjacent mitoSOD1 molecules on the inner membrane.

**MitoSOD1 rescues axon outgrowth and mitochondrial defects in vitro**

Primary motor neurons were cultured from mitoSOD1,Sod1−/− mice, and mitochondrial localization of human SOD1 was verified by immunocytochemistry (Fig. 6A). Staining with a monoclonal antibody specific for human SOD1 revealed a punctate distribution. This staining was absent in cells negative for the mitoSOD1 transgene (not shown). Double labelling with Mitotracker Red CM-H2XRos revealed precise colocalization of human SOD1 staining with mitochondria.

Axon outgrowth was measured in primary motor neurons at 24 and 48 h (Fig. 6B). MitoSOD1 showed a robust protective effect in mitoSOD1,Sod1−/− motor neurons, restoring growth characteristics and axon length to wild-type levels. Axon length...
Mitochondrial SOD1 protects motor axons

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Mitochondrial SOD1 preserves neuromuscular junction integrity and grip strength in vivo

Cohorts of Sod1−/− and mitoSOD1,Sod1−/− mice (and littermate controls) were followed to 12 months of age for assessment of neuromuscular junction innervation and grip strength. Morphological analysis of neuromuscular junction in tibialis anterior muscle showed a robust protective effect of mitoSOD1 at 4, 8 and 12 months of age (Fig. 7A and B). At 4 months, 92.5 ± 0.3% of endplates were fully innervated in mitoSOD1,Sod1−/− mice, which were indistinguishable from controls (92.9 ± 1%). As the animals aged to 8 and 12 months, mitoSOD1 Sod1−/− neuromuscular junctions remained fully innervated.

Similarly, on grip strength analysis, mitoSOD1,Sod1−/− mice performed as well as controls, compared to Sod1−/− littermates that showed a significantly reduced latency to fall (Fig. 7C). These data demonstrate that mitoSOD1 is sufficient to prevent both the pathological and phenotypical hallmarks of motor axonopathy in Sod1−/− mice.

Discussion

Altered mitochondrial function and dynamics have emerged as a major cause of axonal degeneration in peripheral neuropathy and motor neuron disease (Baloh, 2008; Magrane and Manfredi, 2009). Here, we investigated how SOD1 expression in the mitochondrial intermembrane space influences motor axon survival. A protective function for SOD1 in the intermembrane space has been suggested by reports of mitochondrial oxidative stress and...
oxidative damage due to deletion of SOD1 (Aquilano et al., 2006; Muller et al., 2007; Jang et al., 2010), although one group reported that SOD1 potentiates mitochondrial reactive oxygen species generation due to respiratory chain inhibition (Goldsteins et al., 2008). None of these studies addressed the functional effect of intermembrane space-localized SOD1 on oxidative stress-mediated pathology.

**Mitochondrial superoxide production and intermembrane space SOD1**

The protective effect of mitoSOD1 suggests that SOD1 in the intermembrane space is important for regulating physiologic $\text{O}_2^-$ levels. The major source of intracellular $\text{O}_2^-$ production is the electron transport chain, located along the mitochondrial inner membrane. Complex I releases $\text{O}_2^-$ into the mitochondrial matrix (the site of SOD2 expression), and complex III releases $\text{O}_2^-$ bidirectionally into the matrix and intermembrane space (Han et al., 2001; Muller et al., 2004). SOD1 may, therefore, play a protective role by neutralizing complex III-derived $\text{O}_2^-$ in the intermembrane space. This is supported by our finding of thioredoxin-2 oxidation in the setting of a normal thioredoxin-1 redox potential, and by a previous report of preferential protein oxidation in mitochondria, rather than the cytosol, following SOD1 knockdown (Aquilano et al., 2006).

Our data suggest that loss of SOD1, at least acutely, causes a redox imbalance within mitochondria that is not transmitted to the cytosol. Previous reports of increased mitochondrial reactive oxygen species release in $\text{Sod1}^{-/-}$ mice and *C. elegans* did not address the question of the cytosolic redox state (Muller et al., 2007; Yanase et al., 2009; Jang et al., 2010). Eichuri and colleagues (2005) measured cytosolic and mitochondrial aconitate activity in $\text{Sod1}^{-/-}$ mouse liver, and found a significant decrease in the activity of both enzymes with age, suggesting an excess of $\text{O}_2^-$ in both compartments. Whether cytosolic aconitate inactivation in liver represents spilling of mitochondrial $\text{O}_2^-$ into the cytosol, or excess $\text{O}_2^-$ from other typically minor sources, such as cellular oxidases and cytochrome P450s, is unclear.

We cannot exclude the possibility that mitoSOD1 in the intermembrane space may be scavenging $\text{O}_2^-$ generated by other sources, such as membrane-associated NADPH oxidase. However, $\text{O}_2^-$ is unlikely to diffuse far from its site of production without undergoing reactions to produce other reactive oxygen species. Moreover, negatively charged $\text{O}_2^-$ cannot freely diffuse across lipid bilayers (Gus’kova et al., 1984), although it is transmitted by voltage-dependent anion channels (Han et al., 2003), and a small fraction exists in the membrane permeable protonated form ($\text{HO}_2^-$) at neutral pH (Halliwell and Gutteridge, 1990). $\text{O}_2^-$ may also spontaneously convert to membrane-permeable $\text{H}_2\text{O}_2$ at a rate 4-fold slower than the SOD1-catalysed reaction. However, excess cytosolic $\text{H}_2\text{O}_2$ would be detected as thioredoxin-1 oxidation in our assay, since both thioredoxin-1 and thioredoxin-2 are oxidized by $\text{H}_2\text{O}_2$ (Hansen et al., 2006b).

The contribution of SOD1 to $\text{O}_2^-$ neutralization from other sources could be tested by alternative targeting strategies, taking care to prevent SOD1 from being accessible for mitochondrial import. For example, SOD1 could be tethered to the inner face of the plasma membrane or targeted to the endoplasmic reticulum. Further analysis is also needed to define the compartmental redox changes leading to denervation *in vivo*, although subcellular fractionation analysis of neuromuscular junction contents is not feasible with current techniques. New methods for assessing mitochondrial versus cytosolic redox state at the neuromuscular junction *in vivo* are needed.

This study also raises the question of the relative roles of SOD1 and SOD2 in maintaining the mitochondrial redox state and in motor axon maintenance. Loss of SOD2 in mice causes neonatal lethality associated with CNS and cardiac pathology and extensive mitochondrial damage, demonstrating a vital role in development (Lebovitz et al., 1996). Conditional knockout of SOD2 in adult motor neurons does not cause motor neuron loss, denervation or muscle atrophy up to 9 months of age, although it is associated with accelerated axon degeneration following nerve transection (Misawa et al., 2006). We did not directly address how mitoSOD1 affected SOD2 levels or activity in the current study, but SOD2 activity was previously shown to be unaffected in $\text{Sod1}^{-/-}$ mice (Muller et al., 2006), and is therefore unlikely to account for our observation of preferential mitochondrial redox changes in this model.

**Mitochondrial SOD1 import**

Like many other intermembrane space proteins, native SOD1 lacks a canonical mitochondrial targeting sequence. Mitochondrial import of SOD1 depends on the level of copper chaperone for SOD1 in mitochondria, which in turn depends on oxygen levels and import via the Mia40/Erv1 disulphide relay system (Kawamata and Manfredi, 2008; Reddehase et al., 2009). As a consequence, mitochondrial localization of endogenous SOD1 cannot be experimentally manipulated without influencing the mitochondrial localization of other proteins.

An artificial targeting approach was therefore required to express SOD1 exclusively in the intermembrane space. The mitochondrial targeting sequence was chosen because it creates an integral membrane protein, anchoring SOD1 to the outer face of the inner mitochondrial membrane. Endogenous SOD1 is normally thought to be soluble in the intermembrane space, and can be released from mitochondria due to spontaneous increases in outer membrane permeability (Li et al., 2006). Subcellular fractionation studies in *mitoSOD1*, $\text{Sod1}^{-/-}$ tissues demonstrate that anchoring of mitoSOD1 effectively prevents unwanted release into the cytosol. SOD1 activity assays in isolated (intact) mitochondria also verify that mitoSOD1 is enzymatically active; therefore, tethering SOD1 to the inner membrane does not abolish its activity. Moreover, native gel electrophoresis demonstrates that mitoSOD1 is capable of oligomerization. Whether this accounts for measured SOD1 activity remains to be seen, but is not strictly required, as isolated SOD1 monomers retain ~10% activity compared to SOD1 dimers (Bertini et al., 1994).

The expression of mitoSOD1 in *mitoSOD1*, $\text{Sod1}^{-/-}$ spinal cord results in a modest but significant decrease of cytosolic SOD1 activity as compared to $\text{Sod1}^{-/-}$ mice (Fig. 5D). The cause for this reduction is unknown, but one hypothesis is that the increased
amount of SOD1 localized in the intermembrane space may decrease the amounts of copper or copper chaperone for SOD1 available for the maturation and function of endogenous SOD1 in the cytosol (Kawamata and Manfredi, 2008). This change was insufficient to cause a phenotypic change in MitoSOD1, Sod1+/− mice, which was indistinguishable from wild-type controls in multiple assays.

Mitochondrial defects and motor axonopathy

The spatiotemporal sequence of motor pathology has been investigated in numerous mouse models of motor neuron disease, including mutant SOD1-mediated familial amyotrophic lateral sclerosis (Fischer et al., 2004; Pun et al., 2006), spinal muscular atrophy (Murray et al., 2008), progressive motoneuropathy (Holtmann et al., 1999) and others (reviewed in Fischer and Glass, 2007). Distal axonal degeneration, or ‘dying-back’ at the neuromuscular junction is an early pathological feature in these models and appears to correlate most closely with disease progression. Here, we report progressive denervation in Sod1−/− mice reminiscent of these traditional models of motor neuron disease, occurring in the setting of chronic oxidative stress.

Maintenance of extensive axonal processes relies on proper trafficking and function of mitochondria. For example, mutations in MFN2 and GDAP1, regulators of mitochondrial fission and fusion, cause peripheral neuropathy in Charcot–Marie–Tooth disease (Niemann et al., 2005; Baloh et al., 2007). In Drosophila, inactivation of dMiro, a cargo adaptor for anterograde mitochondrial transport, causes accumulation of mitochondria in motor neuron cell bodies and depletion in axons, leading to morphological abnormalities at the neuromuscular junction (Guo et al., 2005). Exposure to the mitochondrial toxin, rotenone (a complex I inhibitor), has also been shown to cause axonal degeneration in vitro (Press and Milbrandt, 2008).

Growing evidence suggests that aberrant mitochondrial trafficking and mitochondrial dysfunction promote distal axonal degeneration in motor neuron disease, although the pathogenic mechanism(s) are less clear. Biochemical and structural defects in mitochondria are well described in mutant SOD1 models of familial amyotrophic lateral sclerosis (Kong and Xu, 1998; Mattiazzi et al., 2002; Menzies et al., 2002), in which mutant SOD1 accumulates and forms insoluble aggregates in mitochondria (Liu et al., 2004). Alterations in axonal transport of mitochondria are also seen in mutant SOD1 models, and abnormal mitochondrial clustering and a relative depletion of mitochondria in axons have been reported (De Vos et al., 2007; Magrané et al., 2009; Sotelo-Silveira et al., 2009). This parallels anecdotal observations from human amyotrophic lateral sclerosis showing abnormal mitochondrial trafficking in proximal axons and cell bodies of motor neurons (Sasaki and Iwata, 2007). Primary culture models of spinal muscular atrophy demonstrate diminished ATP production, mitochondrial depolarization and reduced mitochondrial density in axons (Acsadi et al., 2009; Wen et al., 2010). Similar findings have also been reported in models of spinal and bulbar muscular atrophy (Piccioni et al., 2002; Ranganathan et al., 2009). Thus despite having different upstream causes, mitochondrial pathology appears to lie along a final common pathway of motor axon degeneration.

In the current study, we report a 40% decrease in mitochondrial density in Sod1−/− axons, preventable by introduction of mitoSOD1. We utilized a dye that accumulates in polarized mitochondria, and therefore we cannot exclude that mitochondria were still present in Sod1−/− axons in the depolarized state. Further investigation is needed to determine the link between oxidative stress and altered mitochondrial trafficking in this model, and to investigate whether mitochondria are depleted in motor axon terminals prior to denervation in vivo. Importantly, we demonstrate that mitochondrial-targeted intervention in this model is sufficient to restore the density of polarized mitochondria in axons and to prevent denervation in vivo. It is unclear if mitoSOD1 would be protective in other models of motor neuropathy, where endogenous SOD1 expression is normal, although may be beneficial in models with a strong oxidative stress component.

Conclusion

Targeted replacement of SOD1 in the mitochondrial intermembrane space is sufficient for robust protection against motor axonopathy and mitochondrial abnormalities in Sod1−/− motor neurons. This suggests that mitochondrial damage is an underlying cause of distal motor axonopathy in Sod1−/− mice. Moreover, the functional requirement for SOD1 in motor axons may be based on a protective role in mitochondria.

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Supplementary material

Supplementary material is available at Brain online.

References


