RESEARCH ARTICLE

A new zebrafish model produced by TILLING of SOD1-related amyotrophic lateral sclerosis replicates key features of the disease and represents a tool for in vivo therapeutic screening

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ABSTRACT
Mutations in the superoxide dismutase gene (SOD1) are one cause of familial amyotrophic lateral sclerosis (ALS) in humans. ALS is a relentlessly progressive neurodegenerative disease and, to date, there are no neuroprotective therapies with significant impact on the disease course. Current transgenic murine models of the disease, which overexpress mutant SOD1, have so far been ineffective in the identification of new therapies beneficial in the human disease. Because the human and the zebrafish (Danio rerio) SOD1 protein share 76% identity, TILLING (‘targeting induced local lesions in genomes’) was carried out in collaboration with the Sanger Institute in order to identify mutations in the zebrafish sod1 gene. A T70I mutant zebrafish line was characterised using oxidative stress assays, neuromuscular junction (NMJ) analysis and motor function studies. The T70I sod1 zebrafish model offers the advantage over current murine models of expressing the mutant Sod1 protein at a physiological level, as occurs in humans with ALS. The T70I sod1 zebrafish demonstrates key features of ALS: an early NMJ phenotype, susceptibility to oxidative stress and an adult-onset motor neuron disease phenotype. We have demonstrated that the susceptibility of T70I sod1 embryos to oxidative stress can be used in a drug screening assay, to identify compounds that merit further investigation as potential therapies for ALS.

KEY WORDS: MND, ALS, SOD1, Zebrafish

INTRODUCTION
Amyotrophic lateral sclerosis (ALS) is a devastating disease that is currently incurable. Typical features include progressive muscle weakness and wasting, caused by the death of motor neurons in the motor cortex, brainstem and spinal cord. The majority of patients survive between 2 and 5 years following the onset of symptoms (Ferraiuolo et al., 2011). The current approach to drug identification for ALS has not delivered an effective intervention. Riluzole is the only disease-modifying therapy available and its effect is modest, with survival extension of only ~3 months (Bensimon et al., 1994). Although the majority of ALS cases are sporadic, 5-10% are familial (fALS). Mutations in the superoxide dismutase 1 gene (SOD1) account for 20% of fALS cases; until the recent discovery of a large hexanucleotide repeat expansion in the first intron of C9ORF72 (DeJesus-Hernandez et al., 2011; Renton et al., 2011), such mutations were thought to be the leading cause of fALS. Mutations in SOD1 have been implicated in ALS since 1993, and mutant SOD1 mice and in vitro models have been the dominant models employed to investigate pathogenesis and potential therapies for ALS (Rosen et al., 1993). Several pathogenic mechanisms have been proposed for ALS that is caused by SOD1 mutation, including glutamate excitotoxicity, oxidative stress, mitochondrial dysfunction and axonal defects, including the loss of the neuromuscular junction (NMJ) (Ferraiuolo et al., 2011). Oxidative stress is a mechanism of particular interest owing to the normal role that SOD1 plays in the cell as a cytoplasmic free-radical scavenger (Barber and Shaw, 2010). Current evidence suggests that mutation in SOD1 confers a toxic gain of function in ALS (Dal Canto and Gurney, 1994), rather than a loss of function (Reaume et al., 1996), and a component of this toxicity disrupts the normal handling of free radicals by the cell, generating oxidative stress.

Current in vivo models of ALS rely on the overexpression of mutant SOD1. The most widely used transgenic mutant SOD1 mouse model develops a very early and aggressive phenotype and, although the mutant mice develop progressive hind limb weakness leading to paralysis and death, with very predictable disease progression (Tu et al., 1996), the accelerated time course of the disease does not accurately reflect the human disease. Furthermore, many of the therapies that have seemed to be neuroprotective in the overexpressing transgenic mouse model have failed to translate into beneficial effects in human trials (Benatar, 2007; Gordon et al., 2007). There are several potential reasons for this poor translation of benefit into the human disease. Many of the murine trials have been under-powered and with inadequate attention to the potential confounding effects of gender, litter, heterogeneity of genetic background and pre-symptomatic treatment administration. Furthermore, the relevance of mouse SOD1 models to ALS more broadly has been questioned, given the presence of TDP-43-positive inclusions in motor neurons from most cases of ALS, which are absent in the SOD1-related disease subtype (Mackenzie et al., 2007). However, despite these shortcomings, the transgenic SOD1 mouse model of ALS has been extensively used for preclinical testing (Turner and Talbot, 2008; Knippenberg et al., 2010). New approaches to therapy development are urgently required, and an in vivo system allowing rapid and efficient drug screening would be a useful addition to current models.

The zebrafish represents an alternative model for studying human disease. Zebrafish are capable of producing hundreds of transparent...
Embryos per week, which are externally fertilized and hence easily manipulated using genetic and pharmacological approaches. There are many transgenic models available, some with fluorescent reporters for easy identification of specific cell types and protein expression. Embryos and adults can also be used for behavioural studies and motor function tests. The high fecundity and relatively low maintenance costs mean that high-throughput drug screening. Compounds that have significant effects on survival in the zebrafish could be prioritised for mammalian studies and subsequently in human clinical trials, which could lead to the generation of therapies for the treatment of ALS and related motor neuron diseases. The study also demonstrates the power of TILLING for the rapid development of zebrafish mutants that accurately and reproducibly recapitulate human disease.

Fig. 1. Normal expression and function of WT and T70I Sod1. (A) Western blot with Sod1 expression in 72-hpf wild-type (WT), heterozygous T70I (Het) and homozygous T70I (Hom) zebrafish larvae. (B) Native acrylamide gel of NSC34 and 5-dpf zebrafish sonicates. The gel was stained for SOD1 activity using NBT/riboflavin. (C) Native acrylamide gel of NSC34 and 5-dpf zebrafish sonicates. The gel was stained with Coomassie blue to control for whole protein loading. (B,C) Lane 1 contains the sonicates of NSC34 cells expressing WT human SOD1. Lane 2 contains WT sod1 5-dpf zebrafish sonicates. Lane 3 contains homozygous T70I sod1 5-dpf zebrafish sonicates.

**Clinical issue**
Mutation of the superoxide dismutase 1 (SOD1) gene is the second most common cause of familial amyotrophic lateral sclerosis (ALS), a form of motor neuron disease. SOD1 encodes a protein that is responsible for reducing free oxygen radicals in the cytoplasm, and expression of mutant SOD1 can lead to oxidative stress, which could form the basis of motor neuron death in ALS. The disease, which has devastating effects on muscle function throughout the body, is currently incurable, and the only drug available has a very modest effect on disease course. Further research into identifying potential therapies for ALS is needed, particularly because efforts based on transgenic mouse models have thus far been ineffective. Zebrafish have huge potential for high-throughput drug screening because of their small size and genetic syntenic with humans. This study employs a zebrafish model to screen for potential neuroprotective therapies to treat ALS.

**Results**
In collaboration with the Sanger Institute, the authors applied ENU mutagenesis and targeting induced local lesions in genomes (TILLING) to develop a new zebrafish sod1 mutant. The novel T70I sod1 model displays late-onset motor symptoms and motor neuron loss, as seen in individuals with ALS. The authors also report that the model has a toxic gain-of-function effect, consistent with previous data on other SOD1 mutations. They also show that homozygous T70I sod1 mutant embryos have a marked susceptibility to oxidative stress compared with wild-type controls, and demonstrate that this feature can be exploited in a survival assay that could be used for drug screening. Compounds known for their antioxidant properties were tested on the zebrafish embryos in proof-of-principle assays, in which treatment with apomorphine-S provided the biggest increase in survival (66%).

**Implications and future directions**
This work provides a new animal model of ALS and a robust assay with a clear readout that together have the potential for use in high-throughput drug screening. Compounds that have significant effects on survival in the zebrafish could be prioritised for mammalian studies and subsequently in human clinical trials, which could lead to the generation of therapies for the treatment of ALS and related motor neuron diseases. The study also demonstrates the power of TILLING for the rapid development of zebrafish mutants that accurately and reproducibly recapitulate human disease.

**Normal Sod1 expression level, but reduced enzymatic activity, in T70I sod1 zebrafish**
The presence of mutant Sod1, both in heterozygous T70I sod1 and in homozygous T70I sod1 embryos at 72 hours post-fertilisation (hpf) does not alter the total expression levels of Sod1 in comparison with WT clutch mates (Fig. 1A). The native gel in Fig. 1B demonstrates that there is SOD1 enzymatic activity in all three of the samples. Despite more protein being loaded into lane 3 (homozygous) than lane 2 (WT), there is an apparent reduction in Sod1 enzymatic activity at 65 kDa and no detectable novel activity elsewhere on the gel (Fig. 1B). This indicates that the T70I mutation causes at least a partial reduction of dismutase function of the Sod1 protein. Western blotting of native gels failed to detect any bands through the zebrafish model of sod1 ALS that replicates key features of the disease, which allows this model to be used for the screening of potential neuroprotective therapies.
smaller than 65 kDa, despite using two different polyclonal anti-SOD1 antibodies (data not shown), and hence we cannot categorically say that the band at 65 kDa on the native gel with dismutase activity consists solely of Sod1. Instead, the presence of dismutase activity at a higher molecular weight than a monomer indicates that the native gel might include components of Sod1 complexed with itself or with other proteins (Pedrini et al., 2010).

Zebrafish T70I Sod1 leads to increased motor neuronal oxidative stress in vitro

Oxidative stress represents one element of the toxicity of mutant SOD1 (Barber and Shaw, 2010). We predicted that expression of zebrafish (zf)T70I Sod1, would lead to similar susceptibility to oxidative stress as observed with the expression of human (h)G93A SOD1, in the NSC34 cellular model (Barber et al., 2006). The molecular weights of zfSod1 and mouse (m)SOD1 proteins are very similar at around 16 kDa, and the bands that they produce on a western blot overlap (Fig. 2A). The zfSod1 protein is present in the NSC34 cell lysates despite the cell culture conditions, with a temperature of 37°C rather than the 28°C temperature normally used for zebrafish maintenance. The densitometric analysis (Fig. 2B) shows that there is almost double the expression of Sod1 in the zfWT sod1 and zfT70I sod1 NSC34 cell lines, compared with the vector control NSC34 cell line. This is comparable with expression levels of transfected mutant and hWT SOD1 demonstrated in previous reports (Cookson et al., 2002; Menzies et al., 2002; Barber et al., 2009).

Serum withdrawal proved to be a very robust and consistent method for generating oxidative stress, which differentiated mSod1-expressing NSC34 cells from control cell lines. Upon serum withdrawal for 6 hours, NSC34 cells stably transfected with zfT70I sod1 or hG93A SOD1 demonstrated approximately double the oxidative stress levels compared with transfection with zfWT sod1 or a vector control (P<0.001) (Fig. 2C).

zFT70I Sod1 causes altered NMJ morphology

NMJ staining was carried out on the 11 days post-fertilisation (dpf) progeny resulting from heterozygous T70I mutant sod1 incrosses. These incrosses gave rise to a normal Mendelian ratio of WT, heterozygous and homozygous zebrafish (1:2:1). No morphological differences were noted between clutch mates.

Confocal microscopic analysis of the immunostained zebrafish showed that homozygous T70I zebrafish larvae stained for both α-bungarotoxin (post-synaptic) and SV2 (pre-synaptic) with less intensity than their WT and heterozygous clutch mates (Fig. 3). A significant decrease of ~10% (P<0.001) in colocalisation of SV2 and α-bungarotoxin in the interseptal region was observed in 11-dpf homozygous T70I larvae, as compared with their WT clutch mates (Fig. 3E). Heterozygous T70I mutant sod1 larvae displayed the largest variability in r-value (overlap coefficient, where 1 is complete overlap and 0 is no overlap), and there was no significant difference between the r-values of WT and heterozygous larvae. The decrease in r-value corresponds to a defect of the NMJ.

zFT70I Sod1 causes a reduction in spinal cord motor neurons in adult zebrafish

Choline acetyltransferase (ChAT) staining is a hallmark feature of cholinergic motor neurons. ChAT immunostaining was performed on the spinal cord sections obtained from 3-year-old zebrafish, resulting from heterozygous T70I mutant sod1 incrosses. Motor neurons in the zebrafish spinal cord are small or large depending on the maturation stage of the neurons (Becker et al., 2004). A reduction was observed in both the total number of all ChAT-
positive motor neurons and a subset of mature ChAT-positive motor neurons in the T70I mutants. Motor neurons were classified as mature if they were over 200 μm².

Confocal microscopic analysis of the immunostained zebrafish spinal cord sections showed a significant reduction \((P<0.01)\) by ~50% of large motor neurons with a cell body area of >200 μm² in homozygous T70I zebrafish compared with their WT clutch mates (Fig. 4).

**zfT70I causes motor impairment in adult fish**

Motor phenotypes were observed in 20-month-old homozygous T70I sod1 zebrafish resulting from a heterozygous incross. These motor phenotypes were quantified using the ViewPoint Zebralab system. The ViewPoint Zebralab ‘Tracking System’ readouts demonstrated that the homozygous T70I sod1 zebrafish spent significantly more time (20%) in the bottom third of the tank, compared with their WT clutch mates \((P<0.05; \text{Fig. } 5\text{A})\).

The ViewPoint Zebralab ‘Quantitization Program’ was used to quantify the duration of time that the zebrafish spend swimming at fast, moderate and slow speeds. Using this system it was observed that there was a 44% increase in the ratio of time (resting time/total time) \((P<0.01)\) that the homozygous T70I sod1 zebrafish spent in the ‘resting’ phase of swimming activity (Fig. 5B). The ratio of time (burst time/total time) \((P<0.01)\) that homozygous T70I sod1 zebrafish spent in the ‘burst’ phase of swimming activity (Fig. 5C) as compared with their WT clutch mates was decreased by 50%.

**Oxidative stress survival assay as proof of principle for further drug screening**

Mutant Sod1 confers significant susceptibility to oxidative stress in embryonic zebrafish, as demonstrated by a large reduction in survival of homozygous T70I sod1 embryos upon incubation with 5 mM hydrogen peroxide (H₂O₂) and 1% DMSO (mean survival 45%, s.d.=15.9) compared with WT embryos (mean survival 80% s.d.=12.3) (Fig. 6). Compounds known for their antioxidant properties were tested on the zebrafish in proof-of-principle assays in order to determine the suitability of the assay for a more high-throughput drug screen in the future.
future. All of the drugs used in the survival assay were trialled at a range of concentrations to construct dose-response curves, from which the most successful (activity versus toxicity) concentrations were taken forward (0.5 μM to 3 μM).

Olesoxime (optimal concentration 0.5 μM) and apomorphine-S (optimal concentration 3 μM) were demonstrated to protect both the WT (by 22% and 19%, respectively) and the T70I homozygous sod1 (by 62% and 66%, respectively) embryos from oxidative stress (homozygous recovery $P<0.001$, WT recovery $P<0.01$) (Fig. 6). The robust difference observed between WT and homozygous T70I sod1 zebrafish embryos in their ability to cope with oxidative stress provides a useful readout for a drug screen of compounds to ameliorate oxidative stress.

**DISCUSSION**

We have identified through TILLING a new stable zebrafish model of SOD1 ALS. Our model, which has normal expression levels of the mutant sod1 gene and protein, is different to previously published zebrafish sod1 models, which used either human mutant SOD1 mRNA injected into single-cell-stage embryos, producing a dose-dependent axonopathy in the developing zebrafish embryos (Lemmens et al., 2007), or transgenic G93R sod1 zebrafish, which show NMJ alteration and swimming defects (Ramesh et al., 2010) similar to those shown in this study.

Although the recent genetic discoveries in the field of ALS particularly relating to alterations in TARDBP (Kabashi et al., 2008), FUS/TLS (Kwiatkowski et al., 2009) and C9ORF72 (DeJesus-Hernandez et al., 2011; Renton et al., 2011) do identify previously unknown disease mechanisms requiring investigation, clearly SOD1 remains of key importance for understanding motor neuron degeneration and there is a need for better models to complement the repertoire of models currently available. Our T70I mutant zebrafish model replicates several of the features of human ALS, expresses the mutant protein at physiological levels and has properties that facilitate high-throughput drug screening.

**T70I mutation located in zinc-binding region of sod1**

The T70I sod1 mutation occurs immediately adjacent to the conserved region in the zinc-binding loop of the Sod1 protein (Nordlund et al., 2009). The zinc ion is crucial for Sod1 enzyme stability, and the pathogenic mechanism of the T70I mutation might be due to the alteration of the zinc-binding properties of T70I Sod1. Mutations in the zinc-binding loop and the zinc-binding sites, such as G72C (Stewart et al., 2006), D76Y (Andersen et al., 1997) and H80A (Alexander et al., 2002), are associated with classical ALS.
phenotypes. Cytoplasmic inclusions positive for both ubiquitin and SOD1 were identified in lower motor neurons and anterior horn cells at post mortem in G72C (Stewart et al., 2006) and H80A (Alexander et al., 2002) ALS patients, providing evidence that mutations affecting the region of the zinc-binding loop of SOD1 cause protein instability, incorrect dimerisation of SOD1 and aggregation of the SOD1 protein.

**T70I Sod1 is stable and has a partial reduction in dismutase activity**

Mutation in SOD1 causes ALS by a toxic gain-of-function mechanism (Reaume et al., 1996). A wide range of enzymatic activity of SOD1 has been described, and the different levels are associated with different mutations within the SOD1 gene. Despite this observation, all SOD1 mutations cause fALS, implying that the normal or impaired enzymatic function of SOD1 is irrelevant to the pathogenesis and progression of ALS. Our data are consistent with these earlier observations. We have demonstrated that T70I Sod1 is stable and that this mutation is associated with a partial reduction in dismutase activity. The native zebrafish Sod1 was observed to have an apparent weight of ~65 kDa, double that of the dimerised human and mouse SOD1. Although an unexpected finding, a tetrameric form of Sod1 has previously been described in teleost skin (Nakano et al., 1995), and our findings suggest the possibility that zebrafish Sod1 also exists in a tetrameric form.

**T70I sod1 mutation is associated with susceptibility to oxidative stress**

In mammalian ALS models, SOD1 mutation confers susceptibility to oxidative stress and this mechanism has been identified as one of the key pathways in ALS pathogenesis (Barber and Shaw, 2010). We have demonstrated that NSC34 cells transfected with zfT70I sod1 were viable when grown under normal conditions and that, upon serum withdrawal for 6 hours, the zfT70I-sod1-transfected cells demonstrated increased oxidative stress as measured by reactive oxygen species (ROS) production in a dichlorofluorescein (DCF) assay. The response to oxidative stress of the zfT70I-sod1-transfected NSC34 cell line was comparable to that observed in the established hG93A-SOD1-transfected NSC34 cell line (Barber et al., 2009). These data support the pathogenicity of the zfT70I mutation via a toxic gain of function mediated in part through an increased susceptibility to oxidative stress.

**T70I zebrafish display an early NMJ phenotype and a late motor phenotype**

Since 2004, there has been an increasing body of evidence that points to defects in the NMJ as one of the primary pathological events in ALS (Fischer et al., 2004; Dupuis et al., 2009). The T70I model exhibits early abnormalities at the NMJ. At 11 dpf there is a significant reduction in colocalisation of α-bungarotoxin and SV2 in the interseptal region, between homozygous T70I sod1 zebrafish and their WT and heterozygous clutch mates. This observation might be due to either a failure of the NMJs to form correctly during embryogenesis, or due to a later disruption of the NMJ, followed by the dying back of the axon. Further investigation will be required to determine which of these mechanisms is operating. Our data are consistent with evidence from other studies that point to NMJ alteration as an early pathogenic event and support the validity of the T70I zebrafish model (Fischer et al., 2004; Dupuis et al., 2009; Dupuis and Loeffler, 2009; Ramesh et al., 2010).

It should be noted, however, that the NMJ of zebrafish are not distinctly organized, unlike the human and mouse NMJ, which form pretzel-shaped structures. The NMJs in zebrafish are present all along the axonal tracts. Thus, characterising denervation in zebrafish is more difficult than in mammalian systems. Additionally, owing to the polyneuronal innervation in zebrafish, individual muscle fibres that are totally devoid of NMJs are rarely present until the terminal stages of disease (Ramesh et al., 2010). Thus, reduced colocalisation and/or reduced pre/post-synaptic (SV2 and α-bungarotoxin) labelling is used as an indicator of NMJ abnormality in larval zebrafish (Boon et al., 2009; Ramesh et al., 2010).

The early NMJ phenotype does not lead to a measurable impairment of motor function until the adult zebrafish reach ~20 months of age. It is possible that the polyneuronal innervation of zebrafish muscle or the regenerative capabilities of zebrafish neurons prevent the onset of motor symptoms at a younger age (Fleisch et al., 2011). At 20 months, a motor defect is clearly observed and the adult homozygous T70I sod1 zebrafish spend a greater proportion of time at the bottom of the tank and in the ‘resting’ phase of motor activity, compared with their WT clutch mates. This increased time spent in the resting phase might correlate with a progressive loss of motor function that is developing in the homozygous T70I sod1 zebrafish.

The slow degenerative process in T70I zebrafish coupled with high regenerative capacity of the zebrafish makes it difficult to determine exactly when motor neuron loss starts in these mutants. Hence, adults were chosen for quantification of motor neuron loss because we expected to see the maximum difference between the mutants and WT at this stage. Although we cannot rule out neuronal loss earlier than 2 years, we clearly demonstrate that older zebrafish show the characteristic hallmark features of ALS, i.e. age-related loss of motor neurons, thus validating the model.

ChAT staining is the gold standard measure of cholinergic motor neurons in all animal species. Becker et al. have shown different sizes of motor neurons in the zebrafish spinal cord (Becker et al., 2004). Small motor neurons are generally thought to represent young motor neurons, which become larger as they mature. The total number of motor neurons in the homozygous zebrafish spinal cord was significantly reduced as compared with the WT zebrafish (Fig. 4). To ensure that only large mature motor neurons were counted, a size threshold was set and only motor neurons with an area greater than 200 μm² were counted. As expected, even the large mature motor neurons were reduced in the mutants as compared with the WT spinal cord. Thus, this reduction in ChAT motor neurons truly reflects the death of motor neurons in the mutants. The loss of motor neurons in the homozygous T70I sod1 mutant adult zebrafish replicates a key pathological feature of the disease in humans. This, along with the development of a later adult phenotype, resembles human ALS more closely than the aggressive murine models and this might in part be due to the more physiological level at which the mutant protein is expressed.

**T70I zebrafish can be used to identify potential therapeutic targets for ALS**

Susceptibility to oxidative stress at 24 hpf was chosen as the preferred readout for drug screening experiments because it represents a clear and easily measurable outcome that could be performed simultaneously in large numbers of zebrafish embryos. The other potential readouts, the NMJ phenotype and motor phenotype, were considered too time consuming to be practical within a future high-throughput screen. We identified two antioxidant compounds, apomorphine-S and olesoxime, to assess the feasibility of utilising the T70I sod1 embryos in drug screening assays. Both apomorphine-S and olesoxime give rise to significant
protection to the WT and homozygous T70I sod1 zebrafish embryos against oxidative stress. These data indicate that the susceptibility of T70I sod1 embryos to oxidative stress can be used as a readout in a future large-scale drug screening programme.

Through TILLING we have developed a novel zebrafish model of ALS caused by T70I sod1 mutation. The model demonstrates key features of ALS: an early NMJ phenotype, susceptibility to oxidative stress and an adult-onset motor phenotype. We have demonstrated that the susceptibility of T70I sod1 embryos to oxidative stress can be used in a drug screening assay to identify compounds that merit further investigation as potential therapies for ALS.

MATERIALS AND METHODS

Materials
All general chemicals and reagents, unless otherwise stated, were obtained from Sigma-Aldrich (Poole, Dorset, UK). Polymerase chain reaction (PCR) primers were purchased from Sigma-Genosys (Haverhill, UK) or Integrated DNA Technologies (IDT; Coralville, USA). Tissue culture consumables were from Gibco Life Technologies (Paisley, Scotland) and tissue culture plastics were from Greiner (Gloucestershire, UK).

Zebrafish husbandry
Adult and larval zebrafish stocks were kept at 28.5°C in the University of Sheffield zebrafish facility and bred according to established protocols (Westerfield, 2000). All zebrafish husbandry and experimental procedures were performed in accordance with the UK Animals (Scientific Procedures) Act 1986.

TILLING
Male Tübingen Longtail (TL) zebrafish were treated with a 6-week course of 3.0 mM ENU in order to mutagenise their gametes. These fish were subsequently bred with female TL zebrafish, and the resulting F1 progeny were allowed to reach adulthood. This F1 generation formed the TILLING library (Wienholds et al., 2002).

Nested primers were designed for amplics of interest using the Laboratory Information Management System for Identification of Mutations by Sequencing and TILLING (LIMSTILL) website (http://limstill.niob.knaw.nl), with the internal set of primers being M13-tagged for ease of sequencing by the Sanger Institute.

Cell culture
Neuroblastoma × Spinal Cord hybrid 34 (NSC34) cells (Cashman et al., 1998) were cultured using standard techniques (Cookson et al., 1998). ZfWT and zfT70I sod1 inserts were cloned into the pRRESneo vector and used to transfect NSC34 cells. The selection pressure G418 was used throughout to ensure selection at a concentration of 250 μg/ml.

Cellular oxidative stress assays
NSC34 cells were grown in 96-well tissue culture plates in phenol-red-free DMEM containing 10% FBS until 30–40% confluent. Oxidative stress was induced by 3-hour serum withdrawal. Cytosolic ROS levels were measured using DCF fluorescence (Barber et al., 2009). Carboxy-H2DCFDA (Molecular Probes, Paisley, UK) was added to NSC34 cells to a final concentration of 5 μM, and DCF fluorescence was read at Ex485 nm/Em530 nm after 1 hour. Cell death was simultaneously measured by adding ethidium homodimer-1 (EthD1, 0.3 μM, Molecular Probes) to the culture medium, and the fluorescence was measured at Ex530 nm/Em645 nm. When different cell lines were compared, the DCF results were normalised to the cell number, which was determined by measuring EthD1 fluorescence after the cells had been freeze-thawed.

Western blotting
Western blotting was carried out according to standard protocols. Owing to co-migration of zSod1 and native mSOD1 in the NSC34 cells, densitometry was required to determine the level of overexpression. The film blots were placed inside an Alphalnotech Multimage Light Cabinet (Flowgen, Berks, UK), photographed and densitometry carried out using Alphalnotech software (Alphalmager 1220 v5.1). Ratios of protein of interest to an actin loading control were calculated and, from these ratios, the level of overexpression of Sod1 could be determined.

Native gel for enzymatic activity
Samples were placed into a non-reducing lysis buffer [1× PBS, 1 Proteinase Inhibitor Cocktail (PIC) tablet (Roche, Basel, Switzerland) per litre] and sonicated [Soniprep 150, MSE (UK) Ltd, London] for 30 seconds. Prior to running the samples on the polyacrylamide gel, the samples were added to a 2× loading buffer (160 mM Tris-HCl pH 7.0, 20% glycerol, 0.0025% Bromophenol blue) and incubated at room temperature (RT) for 10 minutes. Protein samples were run on 8% acrylamide gels. Molecular weight standards were added to determine protein size, along with the 25 μg protein samples. An identical gel was run to Coomassie stain to ensure equal loading was achieved.

The gel was first soaked in 1.23 mM NBT for 15 minutes at RT on a shaker, briefly washed in H2O2 before being soaked again in the dark in 100 mM potassium phosphate buffer (pH 7.0) containing 28 mM TEMED and 28 μM riboflavin for another 15 minutes at RT on a shaker. The gel was briefly washed again in H2O2 before being placed onto a transilluminator to develop (Beauchamp and Fridovich, 1971).

Embryonic survival assay
Dechorinated 24 hpf T70I homozygous and WT embryos were placed into Petri dishes and H2O2 was added to a final concentration of 5 mM to half of the Petri dishes and incubated at 28°C for 24 hours before counting for survival as judged by the presence of a heartbeat at 48 hpf.

Drug treatment
Apomorphine-S was identified through an in-house screening cascade of CNS-penetrant Nrf2 activators (Barber et al., 2009), and has been observed to delay the onset of ALS symptoms in transgenic G93A SOD1 mice (Mead et al., 2013). Oleosoxime has been demonstrated to prolong motor neuron survival via modulation of the mitochondrial permeability transition pore and was recently evaluated in a phase 3 trial in human ALS (Bordet et al., 2007). Dose-response curves were conducted to establish an optimal drug concentration. These were done by assessing the toxicity of the drug on three plates of 60 embryos at various doses and represented as percent survival. Optimal doses were then added at gastrulation (final concentration of 1% DMSO), and exposure continued for the duration of the assay. 5 mM H2O2 was added as the oxidative stressor at 24 hpf and survival was measured at 48 hpf.

NMJ staining
NMJ staining was carried out using the protocol described by Ramesh et al. (Ramesh et al., 2010). WT and T70I homozygous embryos were stained in the same tube to accurately compare the NMJ staining. Heads were removed prior to mounting for genotyping. The interseptal region is defined as the region in between the myoseptal regions. This is the region where the longer axons terminate (Meyers et al., 1986). The ventral half of the interseptal region was used for analysis.

ChAT immunohistochemistry
ChAT immunohistochemistry was performed as described (Clemente et al., 2004). Briefly, fresh frozen sections (20 μm) of spinal cord were collected from 3-year-old zebrafish that had been fixed in 4% PFA in 0.1 M phosphate buffer (pH 7.4) for 10 minutes. The sections were washed twice in 0.1 M PB (pH 7.4) for 5 minutes per wash. To decrease background staining, the sections were incubated in 2% H2O2/PBS for 20 minutes. Sections were washed three times in PBDT with 0.2% Tween 20 for 5 minutes, incubated in PBDT with 0.2% Tween and 10% normal donkey serum (NDS) for 2 hours at RT, followed by incubation in goat polyclonal anti-ChAT antibody (1:125 in PBDT with 0.2% Tween and 10% NDS, Chemicon International) for 2 days at 4°C. Sections were washed four times with PBS with 0.2% Tween 20 for 15 minutes each and incubated in biotinylated donkey anti-
goat secondary antibody (1:250 in PBDT with 5% NDS, Santa Cruz Biotechnology) for 2 hours at RT. Sections were washed four times with PBST with 0.2% Tween 20 for 15 minutes each and washed twice with 0.01 M PB (pH 7.4) for 15 minutes. Sections were incubated in Vectastain RTU reagent (Vector Labs) for 90 minutes and were washed four times with 0.01 M PB (pH 7.4) for 15 minutes each. The reaction product was visualized with 0.025% 3,3-diaminobenzidine (DAB) and 0.0033% H2O2 in 0.01 M PB (pH 7.4) or 0.2 M Tris (pH 7.6). The course of the reaction was monitored and stopped by washing in PBST. Sections were serially dehydrated for 10 minutes each in 70% and 100% EtOH, cleared in xylene for 15 minutes, and mounted with Permount (Fisher Scientific). Sections from the dorsal fin region were blinded to genotype and each section was counted for ChAT-80.

Supplementary material available online at this publication site.

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Supplementary material
Supplementary material available online at http://dmm.biologists.org/lookup/suppl/doi:10.1242/dmm.012013/-DC1

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Supplementary Fig. 1. A Blast P of zebrafish and human SOD1, annotated to show the location of the T70I mutation in zebrafish Sod1 (in blue), and also the location of disease causing mutations in the zinc binding loop of human SOD1 (in red).