A Nonsense Mutation in Mouse Tardbp Affects TDP43 Alternative Splicing Activity and Causes Limb-Clasping and Body Tone Defects

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Abstract

Mutations in TARDBP, encoding Tar DNA binding protein-43 (TDP43), cause amyotrophic lateral sclerosis (ALS) and frontotemporal dementia (FTD). Attempts to model TDP43 dysfunction in mice have used knockouts or transgenic overexpressors, which have revealed the difficulties of manipulating TDP43, whose level is tightly controlled by auto-regulation. In a complementary approach, to create useful mouse models for the dissection of TDP43 function and pathology, we have identified a nonsense mutation in the endogenous mouse Tardbp gene through screening an N-ethyl-N-nitrosourea (ENU) mutant mouse archive. The mutation is predicted to cause a Q101X truncation in TDP43. We have characterised TardbpQ101X mice to investigate this mutation in perturbing TDP43 biology at endogenous expression levels. We found the TardbpQ101X mutation is homozygous embryonic lethal, highlighting the importance of TDP43 in early development. Heterozygotes (Tardbp+/Q101X) have abnormal levels of mutant transcript, but we find no evidence of the truncated protein and mice have similar full-length TDP43 protein levels as wild-type littermates. Nevertheless, Tardbp+/Q101X mice have abnormal alternative splicing of downstream gene targets, and limb-clasp and body tone phenotypes. Thus the nonsense mutation in Tardbp causes a mild loss-of-function phenotype and behavioural assessment suggests underlying neurological abnormalities. Due to the role of TDP43 in ALS, we investigated potential interactions with another known causative gene, mutant superoxide dismutase 1 (SOD1). Tardbp+/Q101X mice were crossed with the SOD1G93A/+ transgenic mouse model of ALS. Behavioural and physiological assessment did not reveal modifying effects on the progression of ALS-like symptoms in the double mutant progeny from this cross. In summary, the TardbpQ101X mutant mice are a useful tool for the dissection of TDP43 protein regulation, effects on splicing, embryonic development and neuromuscular phenotypes. These mice are freely available to the community.

Introduction

Amyotrophic lateral sclerosis is a progressive neurodegenerative disease characterised by the degeneration of upper and lower motor neurons, resulting in denervation and atrophy of skeletal muscles, leading to paralysis. Disease course is rapid, with death typically occurring within 3 to 5 years of diagnosis, usually due to paralysis and respiratory insufficiency [1]. Although the majority of ALS cases occur sporadically, approximately 10% of cases are familial (fALS) and our understanding of the genetic causes has expanded dramatically over the past few years [2,3].

Genetic and pathological findings suggest ALS lies within a spectrum of diseases including frontotemporal dementia, a neurodegenerative disorder characterised by degeneration and atrophy of specific cortical areas [1,3]. The association between ALS and FTD was strengthened following identification of TDP43, a nuclear RNA-binding protein, as a major constituent of ubiquitinated cytoplasmic aggregates in degenerating neurons in post mortem tissue from both sporadic and fALS and FTD [4–6]. Subsequently, mutations in TARDBP, the gene encoding TDP43, were identified in ALS and FTD [7–11]. Mutations in TARDBP account for approximately 4% of fALS, a smaller proportion than those caused by mutations in SOD1 (10–20%) or the repeat expansion in C9orf72 (40%), although these figures vary within different populations [12–17]. However, the observation of
TDP43 pathology in both sporadic ALS and C9orf72 patients [5], and other neurodegenerative disorders [18,19], suggests that TDP43 dysfunction may be widely relevant to ALS pathogenesis and neurodegeneration generally. TDP43 pathology is not observed in SOD1-ALS, which has led to the suggestion that divergent disease processes may underlie these forms of ALS. Currently there is evidence for and against an interaction between TDP43 and SOD1 [20–24].

TDP43 is a nuclear protein that plays crucial roles in gene expression and alternative splicing [25,26], embryogenesis [27–30], the stress response [31–36], neuronal functions such as neurite outgrowth [37–39] and many other cellular processes [30,40,41]. It is currently unknown whether mutant TDP43 causes neuronal degeneration through loss of one of these functions or via a novel gain-of-function mechanism, or through both [1,42,43].

To investigate the toxic effect of mutant TDP43, both overexpressing and knockout rodent models have been developed [44–46]. These models show overexpression of wildtype or mutant TDP43 is dose-dependently toxic and results in early mortality, recapitulating some FTD- and ALS-like phenotypes and pathological features. To address whether mutant TDP43 acts through a loss-of-function mechanism, knockout models have been developed by genetically disrupting mouse Tardbp [27–29]. Homozygous Tardbp null animals all die between embryonic day (E) 3.5 and 8.5 [27–29]. In an attempt to avoid the developmental lethality of TDP43 deficiency, a mouse with a tamoxifen-inducible null allele of Tardbp has also been produced [47]. However, disruption of both Tardbp alleles in adult mice led to rapid mortality within nine days, associated with drastic metabolic alterations [47].

Ablation of one allele of Tardbp does not affect TDP43 protein levels, leading to the finding that TDP43 autoregulates its protein levels in vivo [25,27–29,48,49]. Disruption of autoregulation is a potential pathomechanism of mutant TDP43 [50,51]. Intriguingly, despite the absence of overt pathology or perturbation of TDP43 protein levels, one study noted impaired motor performance in aged Tardbp+/− mice [28].

Although the severe effects of systemic TDP43 deficiency are a barrier to investigating loss-of-function, conditional deletion of Tardbp from motor neurons (using loxP flanked Tardbp mice crossed to Hb9-Cre recombinase expressing animals) resulted in delayed weight-gain, impaired rotarod performance, loss of spinal cord motor neurons and ALS-like pathology [52,53]. Similarly, selective disruption of Tardbp in cells (including lower motor neurons) expressing the vesicular acetylcholine transporter, led to progressive decreases in bodyweight and rotarod performance, albeit with no effect on lifespan [53]. Progressive motor impairments in these mice were associated with denervation and atrophy of skeletal muscles, but not loss of spinal cord motor neurons [53].

Loss of TDP43 function in rodents can have widespread effects on alternative splicing. Striatal depletion of TDP43 in adult mice, using antisense oligonucleotides, alters the cassette exon inclusion/exclusion of gene targets, many of which are associated with neurodegeneration [25]. Furthermore, in transgenic mice which overexpress mutant TDP43, alternative splicing patterns of target genes suggests both a loss- and gain-of-function effects [54]. Although transgenic rodents overexpressing human wildtype or mutant TDP43 model some features of ALS and FTD, and TDP43 dysfunction, it is important to remember that autoregulation of TDP43 can cause downregulation of endogenous mouse TDP43 [55], and leading to potential loss-of-function effects.

Given the dose-dependent toxicity of TDP43 in transgenic and knockout rodents, we screened an ENU archive to identify point mutations in the mouse Tardbp gene. ENU is a powerful mutagen that creates inheritable point mutations throughout the genome [56–61]. Male mice are injected with ENU, which ultimately produces a unique array of random point mutations in their sperm, in a dose-dependent manner [62]. Injected mice are mated and sperm from their male offspring harvested, with matching DNA samples, which can be screened for mutations. From screening the ENU archive at the Medical Research Council Mammalian Genetics Unit (Harwell, UK) [56], we identified a mouse Tardbp mutation predicted to cause a Q101X truncation in the N-terminus of TDP43.

The new TDP43 Q101X strain was assessed for TDP43 expression and function, and mouse survival, behaviour and motor function. We also carried out a cross to transgenic mice carrying a mutant SOD1 gene, which models ALS, to determine if there were any interaction effects between the two mutant genes. We found the TardbpQ101X mutation affected alternative splicing of selected target genes and caused behavioural phenotypes, in the absence of motor dysfunction or deleterious effects of mouse survival. Progeny of the novel cross to SOD1G93A transgenic mice did not reveal interaction effects between the novel mutation in Tardbp and the SOD1G93A transgene.

Methods

Ethics Statement

Mice were bred and maintained by MRC Harwell and UCL Institute of Neurology. Experiments were performed under licence from the UK Home Office and following approval from the local ethical review panels: Ethical Review Panel of MRC Harwell and Ethical Review Panel of the MRC Prion Unit, UCL Institute of Neurology. Surgery was performed under terminal anaesthesia and all efforts were made to minimise suffering.

Identification of an ENU-induced Mutation in Tardbp

DNA from the MRC Harwell ENU archive [http://www.har.mrc.ac.uk/services/dna_archive/] was screened with the LightScanner platform (Idaho Technology Inc., USA). All exons of Tardbp were screened in DNA from ~10,000 F1 ENU mutagenised animals and potential mutations confirmed with Sanger sequencing (GATC, Germany). This led to the identification of a G to T base mutation within exon 3 which is predicted to cause a Q101X nonsense (glutamine to stop codon) change in TDP43 protein.

Genetic Background

Male C57BL/6J mice treated with ENU and were crossed to C3H/HeH females. F1 progeny (C3H/HeH.C57BL/6J) were rederived and male F1 animals had sperm and DNA samples taken for archiving. F1 DNA was screened for mutations in Tardbp, and the TardbpQ101X strain was rederived from F1 sperm, used for in vitro fertilisation of C57BL/6J oocytes. All mice used for phenotyping had been back-crossed 3–4 generations onto a C57BL/6J background. Third generation (N3) back-cross C57BL/6J females were crossed with congenic C57BL/6J SOD1G93A male mice [63] to generate wildtype, single and double mutant progeny.

Genotyping

The Tardbp Q101X allele was genotyped using a Qiagen pyro-sequence. Extracted DNA (∼5 µg/µl) was amplified and sequenced following manufacturer’s instructions at 55°C annealing temperature. Primers used were forward: CAAAAGGAAAATGGAT- GAGAC, reverse: AGTTGTTTTCCAGGGGAGAC, sequenc-
ing forward: GAAAGTGAAAAGAGCAGTC, which flank the site of mutation and bind within exon 3. The pyro-sequencer converts luciferin to oxyluciferin to produce light resulting in a quantitative read-out. Thus, to measure the relative abundance of the Q101X \textit{Tardbp} transcript compared to the wild-type transcript, cDNA was generated from brain and analysed on the pyro-sequencer with the quantitative readout for the oxyluciferin generated from the wildtype (C) and mutant (T) bases used to determine relative levels of transcripts containing each base.

\textit{SOD1G93Adl} mice were maintained as hemizygotes and genotyped with transgene and control specific primers, as described on the Jackson Laboratory Mice Database (http://jaxmice.jax.org/strain/002300). Transgene primers, amplifying across exon 4: forward: CATCAGCCCGTAACTCATCTGA, which binds within intron 3, and reverse: CGCCGACTAACACTGAAATGGA, which binds within intron 4. In order to control for the reaction, the mouse gene \textit{Scna} was also amplified as the control gene to validate the mouse DNA sample. \textit{Scna} primers, forward: ATCTGGTCCTTCTTGACAAAGC, reverse: AGAAGACAATGACA, which both bind within exon 4. These produced amplicons of 236 bp and 130 bp respectively, with presence or absence of the \textit{SOD1G93A} fragment used to determine genotype.

**Figure 1.** \textit{Tardbp}^{+/Q101X} mice have reduced mutant transcript but normal TDP43 protein levels. (A) Location of the predicted Q101X truncation in TDP43. All coding exons of \textit{Tardbp} were screened for mutations using denaturing high performance liquid chromatography, with a C to T mutation identified in exon 3 which is predicted to encode a Q101X truncation, occurring before the RNA recognition motifs (RRM1/2) and the glycine-rich domain (GRD). Numbers denote amino acids in the full-length 414 residue protein. (B) Genomic DNA and transcript levels were assessed in brain tissue from male \textit{Tardbp}^{+/+} (n = 2) and \textit{Tardbp}^{+/Q101X} (n = 5) mice at 18 months of age. Data are mean ± standard deviation. (Left panel) Genomic DNA: in \textit{Tardbp}^{+/Q101X} mice, the wildtype and mutant alleles in genomic DNA are detected at a ratio of 1:1. (Right panel) cDNA: levels wildtype and mutant transcripts were quantified by pyro-sequencing; wildtype base is C and mutant base is T. RNA converted to cDNA shows a ratio of 5.25:1 for wildtype:mutant levels. The pyro-sequencer measures the relative areas under each curve, allowing comparison of relative levels of the two transcripts p < 0.001 with Fisher's test. (C) \textit{Tardbp} transcript levels, assessed using primers spanning exons 2 to 3 were not altered in brain tissue from male mice at 18 months of age. Mean \(\Delta A_CT\) of \textit{Tardbp}^{+/+} = 1.7 ± 0.3 (n = 5), \textit{Tardbp}^{+/Q101X} = 1.9 ± 0.4 (n = 5), p = 0.71 (students two-tailed t-test). Data are mean ± standard deviation. (D) No difference in full-length TDP43 protein levels (relative to loading control) between \textit{Tardbp}^{+/+} (73.0 ± 5.2) and \textit{Tardbp}^{+/Q101X} (67.8 ± 3.0) mice, using antibody directed against C-terminus of TDP43 (Proteintech 12892-1-AP). TDP43 levels (upper panel, green) were assessed from brain tissue with 5 male mice per genotype at 18 months of age. Actin (upper panel, red) was used as a loading control. Data are mean ± SEM.

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Molecular Analysis of TDP43 Alternative Splicing

RNA was extracted from tissue using TRIzol (Life Technologies, Paisley, UK) and cDNA was produced using Applied Biosystems cDNA generation kit (Life Technologies, Paisley, UK) following the manufacturer’s protocol. Oligo dT primers were used to generate the cDNA. To assess splicing reactions, 1 μl cDNA at ~ 50 ng/μl was used for polymerase chain reaction (PCR) with a Qiagen master mix (201445; Qiagen, Netherlands), and 34 thermo cycles. The PCR product was electrophoresed on a 3% agarose gel and visualised using a Bio-Rad ChemiDoc (Bio-Rad, California, USA). The forward and reverse primers for each reaction were located in exons either side of the known alternatively spliced exon for each target gene. Sort1 – forward (exon 17): CAGGAGACAAATGCGAACGT, reverse (exon 19): TGCCCCAGATAATGGGACA, Pdp1 – forward (5’UTR): GTGCTGTAGTGAGGGAAGGAC, reverse (exon 2): TGCCAGCTAGATGACCGCTCA, Knd1 – forward (exon 1): CGGCTCTCTATGGACGACCTTA, reverse (exon 4): GAGATTGTCCGAGACCCCTCA, Kcnq2 – forward (exon 3): GCGAAGACCCCTCCTCAT, reverse (exon 6): AGTGGTGCTGGAGGAAAGGAGA, CACNIP2 – forward (exon 4): CTCTATGTGGCGGAGCAGTT, reverse (exon 5): GGAGTTGTTCCAGACCCTCA, DnajC5 – forward (exon 6): TGGCCAGGATAATAGGGACA, reverse (exon 3): CAGGAGACAAATGCCAAGGT, reverse (exon 19): CAAACCCTTTCGAGT.

Assessment of TDP43 Protein Levels

Protein was harvested from tissue and cells by homogenisation in RIPA buffer with protease inhibitor cocktail (Roche). Centrifugation (13,200 x g at 4°C for 30 minutes) was used to purify the soluble fraction. SDS-page gels were Invitrogen Novex BisTris 10% gels (Life Technologies, Paisley, UK) and transferred onto immobilon low fluorescence PVDF membrane (Millipore, Massachusetts, USA). Membranes were blocked in PBS with 0.2% tween and 5% milk. Anti TDP43 antibodies were from: ProteinTech (12892-1-AP) and N-terminus antibodies from Cosmo Bio (CAG-TIP-TD-P07, Cosmo Bio, California, USA) and Abcam (ab50930, Abcam, Cambridge, UK). Control actin was labelled using Abcam primary (AC-15). Visualisation with the Li-Cor Odyssey system (LI-COR Biosciences, Nebraska, USA) used mouse and rabbit specific red and green fluorescently conjugated secondary antibodies (Li-Cor IRDye 680RD, 926-32200, 926-32211).

Table 1. TardbpQ101X/Q101X embryos die in development before E6.5.

| Genotype       | Timed mating | n  | Tardbp+/+ | Tardbp+Q101X | TardbpQ101X/Q101X | Resorptions |
|----------------|--------------|----|-----------|---------------|-------------------|-------------|
| E7.5–11.5      | 47           | 12 | 21        | 0             | 14                |             |
| E6.5           | 25           | 9  | 11        | 0             | 5                 |             |

Timed matings were set-up to establish survival of TardbpQ101X/Q101X mice. No homozygous embryos were identified at time-points assessed. We were unable to genotype resorptions. At E6.5 χ² = 8.3, p = 0.0158 versus an expected ratio of 1:2:1 – Tardbp+/+, Tardbp+Q101X, TardbpQ101X/Q101X.

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Figure 2. The Q101X mutation in Tardbp alters splicing of TDP43 targets Sort1 and Pdp1. (A) Exon inclusion of Sort1, determined from cDNA generated from brain from 18-month old mice, is significantly increased in Tardbp+Q101X (n = 5) mice compared to Tardbp+/+ mice (n = 5). Bar charts show abundance of exon-included transcript relative to total transcript (included and excluded). Sort1 exon inclusion: Tardbp+/+ 24.4%±1.3% versus Tardbp+Q101X 31.3%±2.6%, p = 0.044 (student’s two-tailed t-test). Data are mean±SE. (B) Exon inclusion of Pdp1 is also significantly increased in brain tissue from Tardbp+Q101X mice at 18 months of age. Exon inclusion: Tardbp+/+ 9.0%±2.0% (n = 5), Tardbp+Q101X 50.3%±1.3% (n = 5), p = 0.0013 (student’s two-tailed t-test). Data are mean±SE.

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Embryonic Lethality of TardbpQ101X/Q101X Mutant Mice

Embryonic lethality was assessed by culling female mice at selected days post conception using UK Home Office defined Schedule 1 methods. Embryos were harvested and tissue collected for genotyping.
Lifespan
All mice were maintained and assessed in accordance with UK Home Office project licence regulations. Mice were weighed every two weeks until the endpoint of the assessment was reached, or until a humane endpoint, defined as when mice showed either lunched posture, 20% bodyweight loss or limb paralysis. Only females were used for the lifespan analysis of Tardbp\(^{+/+}\) mice, as aged males were used for pathological analysis at scheduled time points.

Behavioural Analysis of Mice

Grip-strength was longitudinally assessed (BioSeb, France) by measurements using all four limbs and the two front paws alone. Recordings were taken the week following the SHIRPA assessment, from 8 weeks of age to 104 weeks of age.

Motor performance on an accelerating rotarod was assessed with three trials per day, on three days within the testing week. The average value of all nine runs was recorded (Ugo Basile, Varese, Italy). The rotarod accelerated from 4 to 40 revolutions per minute over a maximum of 300 seconds. Rotarod was assessed from 8 weeks of age to 56 weeks of age and was measured in the same week as the grip strength test.

Behavioural analysis of mice was performed using the SHIRPA (SmithKline Beecham, Harwell, Imperial College, Royal London Hospital, phenotype assessment) protocol as previously described [64]. Phenotyping was carried out blind to mouse genotype.

SHIRPA was undertaken every two months from 7 weeks of age for the Tardbp\(^{+/-}\) Q101X phenotyping cohort; female Tardbp\(^{+/-}\) n = 12, female Tardbp\(^{+/-}\) Q101X n = 15, male Tardbp\(^{+/-}\) n = 16, male Tardbp\(^{+/-}\) Q101X n = 16. Beyond one year, only the female cohort was phenotyped up to end stage or two years of age. For the double mutant cross between Tardbp\(^{+/-}\) Q101X and SOD1G93Adl, progeny were phenotyped monthly from 7 weeks of age up to endstage between 31–38 weeks. This cohort contained, females: Tardbp\(^{+/-}\) Q101X n = 10, Tardbp\(^{+/-}\) Q101X n = 5, Tardbp\(^{+/-}\), SOD1G93Adl n = 14, Tardbp\(^{+/-}\) Q101X, SOD1G93Adl n = 14. Males: Tardbp\(^{+/-}\) n = 12, Tardbp\(^{+/-}\) Q101X n = 10, Tardbp\(^{+/-}\), SOD1G93Adl n = 13, Tardbp\(^{+/-}\) Q101X, SOD1G93Adl n = 14.

Within the SHIRPA, limb-clasping was assessed by suspending the mice by the tail, with a score of 0 given for no clasping and 1 given for front or hind-paw clasping. For analysis, only mice which showed clasping over two successive assessments and maintained this phenotype in all remaining assessments were scored as limb-clasping. Body tone was assessed by holding mice near the base of the tail; front paws were placed to grip a grid above the SHIRPA arena and the hindlimbs were suspended slightly above the grid. With the free hand, the thumb and index finger were placed around the pelvic region and lower thorax and then rounds of lateral compression, while continuously keeping the thumb and finger against the mouse, were used to feel muscular tone with a reflexive elicited response. If mice were anxious and elicited a jerk/jump response they were then re-tested. Mice were scored 2 for normal body tone, 1 for soft tone, and 0 for flaccid tone. Mice
were only considered to have body tone phenotype if they showed flaccid tone over two consecutive assessments and maintained this phenotype.

Startle response and pre-pulse inhibition were tested as described on the Empress database (http://empress.har.mrc.ac.uk). The Tardbp+/Q101X cohort was assessed at 10 and 22 weeks of age and the Tardbp+/Q101X, SOD1G93Adl cohort assessed at 10, 18 and 22 weeks of age.

The Open Field test was undertaken to assess both motor- and anxiety-based behaviour, and analysed with software from Noldus (Wageningen, Netherlands). The test was completed as described on the Empress database but was modified to increase the recording time from 20 to 30 minutes. The Tardbp+/Q101X cohort was assessed at 14 and 30 weeks of age. Progeny from the Tardbp+/Q101X, SOD1G93Adl cross were assessed at 14 weeks of age.

Pathology

Brains from Tardbp+/+ and Tardbp+/Q101X male mice (both n = 3) were harvested for histopathological analysis at one year of age. Mice were perfused transcardially with 0.9% saline followed by 4% paraformaldehyde (PFA). Following dissection, brains were transferred to formalin and subsequently embedded in paraffin wax. Sections were stained with haematoxylin and eosin, and antibodies against GFAP (3 ug/mL, DAKO, Z0334), IBA-1 (1.7 ug/mL, WAKO Chemicals, 019-19741), p62 (5.6 ug/mL, Progen Biotechnik, GP62-C), MBP (2 ug/mL, Covance, SMI-94R) and neurofilament (0.5 ug/mL, Sigma, N3389) using a Ventana automated immunohistochemical staining machine (Ventana Medical Systems, Tuscon, USA). Once incubated with appropriate secondary antibodies, immunoreactivity was developed with 3,3′-diaminobenzidine and counterstained with haematoxylin.
Figure 5. No interactive effects between Tardbp<sup>Q101X</sup> and SOD1<sup>G93A</sup> mice. (A) Grip strength of SOD1<sup>G93A</sup> mice (both sexes) is not affected by the Q101X mutation in Tardbp. SOD1<sup>G93A</sup> mice show progressive loss of grip strength (versus Tardbp<sup>+/+</sup> p = 0.002, LSD test from one-way ANOVA), however there is no difference between Tardbp<sup>+/+</sup>, SOD1<sup>G93A</sup> and Tardbp<sup>Q101X</sup>, SOD1<sup>G93A</sup> mice (p > 0.05). Data are mean ± SEM. (B) Progressive bodyweight loss in SOD1<sup>G93A</sup> mice is not affected by the Q101X mutation in Tardbp. Data are mean ± SEM. (C) Survival plots for female (left panel) and male (right panel) mice show no effect of the Q101X mutation on survival of SOD1<sup>G93A</sup> mice. Endstage was defined as humane end-point or greater than 20% weight loss. Tardbp<sup>+/+</sup>, SOD1<sup>G93A</sup> 256 ± 4 days n = 22, Tardbp<sup>Q101X</sup>, SOD1<sup>G93A</sup> 263 ± 8 days n = 19 (total number including both sexes, with equal ratio), p = 0.227 (Kaplan Meier log rank statistic) (D) Body tone phenotype of Tardbp<sup>Q101X</sup> is replicated but not significantly different in Tardbp<sup>Q101X</sup>, SOD1<sup>G93A</sup> mice (female mice: left panel; male mice: right panel). Mean age at which body tone phenotype was shown for both sexes combined: Tardbp<sup>+/+</sup> = absent, Tardbp<sup>Q101X</sup> = 32 weeks of age (more than 50% of females), Tardbp<sup>+/+</sup>, SOD1<sup>G93A</sup> = absent, Tardbp<sup>Q101X</sup>, SOD1<sup>G93A</sup> n = 26 weeks (50% for both sexes), Tardbp<sup>+/+</sup>, SOD1<sup>G93A</sup> versus Tardbp<sup>Q101X</sup>, SOD1<sup>G93A</sup> p = 0.042, Tardbp<sup>+/+</sup>, SOD1<sup>G93A</sup> versus Tardbp<sup>Q101X</sup>, SOD1<sup>G93A</sup> p = 0.982, Tardbp<sup>+/+</sup> versus Tardbp<sup>Q101X</sup> p = 0.006 (Kaplan Meier log rank statistic). Overall p = 0.008, with no significant difference between sexes within genotypes. Female mice: Tardbp<sup>+/+</sup> n = 10, Tardbp<sup>Q101X</sup> n = 10, Tardbp<sup>+/+</sup>, SOD1<sup>G93A</sup> n = 15, Tardbp<sup>Q101X</sup>, SOD1<sup>G93A</sup> n = 14. Male mice: Tardbp<sup>+/+</sup> n = 12, Tardbp<sup>Q101X</sup> n = 10, Tardbp<sup>+/+</sup>, SOD1<sup>G93A</sup> n = 14, Tardbp<sup>Q101X</sup>, SOD1<sup>G93A</sup> n = 14.

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**Primary Embryonic Motor Neuron Culture and Quantification of Neurite Outgrowth**

Mixed ventral horn cultures were prepared as previously described [65]. Briefly, primary embryonic motor neurons were isolated on E13.5, following cervical dislocation of pregnant females and hysterectomy. Ventral horns were dissected from individual embryos and dissociated by 10 minute incubation with trypsin (final concentration 0.025%, Type XII-S, Sigma Aldrich, Paisley, UK) and three trituration steps in 400 μl L-15, 50 μl 4% bovine serum albumin (BSA, Sigma Aldrich,) in L-15 and 50 μl DNase (1 μg/ml, Sigma Aldrich.). 1 ml 4% BSA was added to the cell suspension to form a cushion and cells were centrifuged at 239xg, room temperature for 5 minutes. Cell pellets were resuspended and then plated onto glass coverslips (precoated with polyornithine and laminin for at least 2 hours each) and maintained in neurobasal medium, supplemented with 1% penicillin-streptomycin (50 units/ml penicillin, 50 μg/ml streptomycin), 2% B27 supplement, 2% horse serum, 0.05% 50 mM β-mercaptoethanol, 0.5 mM L-glutamine (all from Invitrogen, Paisley, UK), 0.1 ng/ml brain-derived neurotrophic factor, 0.1 ng/ml glial-derived neurotrophic factor and 0.5 ng/ml ciliary neurotrophic factor (all from Peprotech, London, UK).

After 18 hours in vitro, coverslips were immunostained with β-III-tubulin (Covance, Harrogate, UK) to identify neuronal cells and fluorescent images were captured at x20 magnification with a Leica DFC colour camera (Leica, Wetzlar, Germany) running Leica Application Suite Version 2.8.1 (Leica, Wetzlar, Germany). From 3 independent cultures, 10 random fields were imaged from 3 coverslips per genotype. To assess neurite outgrowth in primary embryonic motor neuron cultures, only cells identified with β-III-tubulin, with at least 3 neurites whose full length could be seen, were selected. The length of every selected neurite and branch was measured with Metamorph (Molecular Devices, Berkshire, UK) and data transferred into Microsoft Excel (Microsoft Corporation, Redmond, Virginia, US), where mutant values were expressed as a percentage of wildtype values. Data were plotted for the mean neurite length, and the mean longest neurite length.

**In vivo Assessment of Neuromuscular Function**

Neuromuscular function was assessed in vivo in anaesthetised mice (4.5% chloral hydrate, i.p. 1 ml/100g bodyweight; Sigma Aldrich), as previously described [66,67]. Anaesthesia was checked regularly throughout the procedure and supplemented when necessary with a quarter of the initial dose of anaesthetic.

The distal tendons of the tibialis anterior (TA) and extensor digitorum longus (EDL) muscles were exposed, cut and attached to isometric force transducers (Dynamometer UFI Devices, Welwyn Garden City, UK), which were connected to a MultitracRecorder (Lectromed Multitrace 2, Letchworth Garden City, UK), and a Picoscope recorder (Pico Technology, Cambridgeshire, UK) which captured the data.

The sciatic nerve was exposed in the mid-thigh region and sectioned proximally before being placed over a platinum stimulating electrode and kept moist with 0.9% saline. Muscle length was adjusted for maximum twitch force. The sciatic nerve was then stimulated with 0.02 ms square wave pulses at a supramaximal voltage (10V) to record maximum single twitch force. Maximum tetanic force was determined by stimulating the sciatic nerve with trains of stimuli at 40, 80 and 100 Hz for 450 ms. From the maximum twitch force recorded, muscle contraction characteristics were determined by measurement of the time taken to reach peak contraction (time-to-peak; TTP) and the time to reach half relaxation (½RT). Muscle fatigue characteristics of the EDL were assessed by a fatigue test, in which the muscle was stimulated at 40 Hz for 250 ms per second, for 180 seconds, and the resulting muscle contractions recorded on a pen electrode (Lectromed Multitrace 2). The fatigue index (FI), a measure of fatiguability, was determined from the trace by expressing the final force (F180) as a ratio of initial force (F0).

To determine the number of surviving motor units innervating the EDL, the sciatic nerve was stimulated with voltage of increasing intensity, from a sub-threshold voltage to maximum, resulting in the gradual recruitment of motor units, reflected as stepwise increments in twitch tension, which were counted to estimate the number of functional motor units.

**Morphological Assessment of Motor Neuron Survival**

Following the acute physiological experiment, the mice were terminally anaesthetised with an overdose of 4.5% chloral hydrate (i.p.) and transcardially perfused with 0.9% saline followed with 4% PFA. Spinal cords were dissected from the spinal column and left overnight in 4% PFA at 4°C before cryoprotection in 30% sucrose in PBS at 4°C. Cross-sections of the lumbar region (L2-L6) of the spinal cord were cut at 20 μm thickness on a cryostat and Nissl stained with gallocyanin [68]. Motor neuron survival was established by counting the number of large polygonal neurons with a minimum diameter of 20 μm within the sciatic motor pool [69]. Only those cells with a clear nucleolus and distinct Nissl-dense cytoplasm were counted in every 3rd section to prevent recounting the same neurons twice in consecutive sections. Counting was performed blind to genotype using a Leica DMR microscope and using a minimum for 40 sections.

**Neuromuscular Junction Immunofluorescence**

Mice were killed by cervical dislocation. Abdominal muscles were dissected, isolated in PBS, stained for 40–60 minutes in 4% paraformaldehyde. Axons were visualised by staining with anti-150 kDa Neurofilament M rabbit and SV2 mouse antibodies overnight at 4°C at dilutions of 1:250 and 1:200, respectively. After washing for 1 hour in PBS, samples were incubated with secondary antibody (150 kDa Neurofilament M: goat anti-rabbit IgG FITC-conjugated antibody, 1:100 dilution; SV2: goat anti-mouse IgG FITC-conjugated antibody, 1:100 dilution) and subsequently washed with PBS. Whole muscles were mounted in Mowiol and imaged using a BioRad Radiance 2000 confocal microscope mounted on a Nikon Eclipse E600FN platform, using 20x and 40x water immersion objectives.

**Statistical Analysis**

Kaplan-Meier Log rank statistic was used to compare phenotypes in the SHIRPA to define age of phenotype onset. For other tests including grip strength, rotarod, startle response, open field, RNA and protein levels; comparisons were made using a two-tailed ANOVA test with Least Squared Difference (LSD) and/or Bonferroni post-hoc tests or two tailed T-test. Mean neurite length and axon length were analysed using a multi-level model in Stata 12 (StataCorp, Texas, USA), to compare neurite and axon lengths between wildtype and mutant motor neurons from independent experiments. For in vivo assessment of neuromuscular function a Mann-Whitney U-test was used to compare between Tardbp+/+ and Tardbp+/+D20X mice at 18 months of age. A one-way ANOVA, with Tukey’s test was used to compare groups at 32-33 weeks of age.
Results

Identification of an ENU-induced Point Mutation in Mouse Tardbp

The MRC Harwell ENU archive of over 10,000 DNAs was screened and a C to T mutation in exon 3 of Tardbp was identified. The mouse strain was rederived onto a C57BL/6 background using standard in vitro fertilisation techniques. The mutation is predicted to cause a Q101X truncation in the N-terminus of TDP43 whose longest isoform in mouse is 414 amino acids. The truncation occurs before any predicted functional domains of the protein (Figure 1A).

The Q101X Mutation in Tardbp is Homozygous Embryonic Lethal

To investigate embryonic lethality, heterozygous (Tardbp+/Q101X) mice were intercrossed to produce homozygous animals. No Tardbp+/Q101X/Q101X progeny were detected at birth, or between E7.5-11.5 (n = 47 embryos) or earlier at E6.5 (n = 25 embryos; Table 1). Therefore the Tardbp+/Q101X mice die in early development prior to E6.5.

Reduced Level of Mutant Transcript but Normal Level of Wildtype Transcript in Tardbp+/Q101X Brain

As expected, genomic DNA from Tardbp+/Q101X mice showed equal levels of wildtype (53.0%±5.2%) and mutant alleles (47.0%±5.2%) (Figure 1B). We then investigated whether the Q101X mutation had effects on Tardbp transcript abundance. Using a Qiagen pyro-sequencer with primers flanking the mutation, we assessed transcript levels from brain cDNA in Tardbp+/Q101X mice and found a significant difference in the relative abundance of wildtype to mutant transcripts at 18 months of age (86%±1%: 16%±1%, p<0.001; Figure 1B) and 3 months of age (86%±1%: 14%±1%, p<0.001) – A ratio of 5.25:1 instead of the anticipated 1:1 ratio. Using this assay we can assess whether transcripts are wildtype or mutant, but we cannot distinguish different wildtype transcripts, however it demonstrates a greatly reduced level of mutant compared to wildtype transcript in Tardbp+/Q101X mice.

We went on to assess transcript levels by rPCR. As the Tardbp+/Q101X mutation occurs within exon 3, we quantified Tardbp transcript levels using primers for regions spanning exons 2 to 3 and exons 3 to 4 (data not shown) in brain tissue from 18 month old mice. Both sets of primers amplify regions either side of the Tardbp+/Q101X mutation and do not overlap the mutation. Using these primers, transcript abundance did not differ between Tardbp+/+ and Tardbp+/Q101X mice (1.7±0.3 versus 1.9±0.4 respectively; Figure 1C), demonstrating equivalent levels of wildtype transcripts.

Thus in heterozygous mouse brain (1) the mutant transcript is detectable, but at significantly lower levels than the wildtype transcript, and (2) levels of Tardbp transcripts including exon 2,3 and 4 are the same as in wildtype littermates.

No Detectable Truncated but Normal Full-length TDP43 Protein Levels in Tardbp+/Q101X Mice

We assayed for truncated TDP43 Q101X protein by probing western blots with antibodies (Cosmo Bio: CAC-TIP-TD-P07, Abcam: ab50930) directed against the C-terminus of TDP43, and no truncated protein was detected in brain tissue from heterozygotes at 18 months of age (Figure S1A, B).

As previous reports have demonstrated that TDP43 autoregulates its protein levels [25,28,48,49], and because transcript levels are the same in wildtype and mutant mouse brain, we assessed TDP43 protein levels using the Li-Cor Odyssey system and normalising to alpha actin content. Using an antibody directed against the C-terminus of TDP43 (ProteinTech: 12892-1-AP), a region beyond the predicted point of truncation, on brain from male mice at 18 months of age, no difference was seen in TDP43 protein levels (relative to loading control) between Tardbp+/+ (73.0±5.2) and Tardbp+/Q101X (67.8±3.0) mice (n = 6 per genotype) (Figure 1D). Similar results were obtained with this antibody in spinal cords at 18 months of age for both soluble and insoluble fractions (Figure S1C-D) as well as at 3 or 12 months of age in brain or in spinal cord at 3 months, or in mouse embryonic fibroblasts (data not shown).

Thus Tardbp+/Q101X mice have equivalent, wildtype Tardbp transcript and TDP43 protein levels as wildtype littermates, but also have low levels of mutant transcripts, although no mutant protein that we could detect in brain tissue.

Loss of Splicing Function of Selected Target Genes in Tardbp+/Q101X Mice

Although Tardbp+/Q101X mice showed overall normal wildtype TDP43 protein levels, we examined whether these mice retained complete TDP43 function. Depletion of TDP43, therefore a loss-of-function, has been previously shown to disrupt genome-wide cassette exon inclusion/exclusion in target genes [25]. Using splicing patterns of TDP43 target genes as a paradigm to assess TDP43 function, we characterised the exon inclusion/exclusion patterns of five established target genes: Sort1, Pdp1, Dnaj5, Knd5 and Kcnd2 for which TDP43 promotes exon skipping, and Kcnd2 for which TDP43 promotes exon inclusion.

With RNA extracted from brain tissue at 18 months of age, we found significant differences in the splicing patterns of Sort1 (exon 18 inclusion: Tardbp+/+24.4%±1.3%, Tardbp+/Q101X 31.3%±2.6%); Figure 2A) and Pdp1 (exon 1 inclusion: Tardbp+/+39.0%±2.0%, Tardbp+/Q101X 50.3%±1.3%; Figure 2B) but no difference in the splicing patterns of Dnaj5, Knd5 and Kcnd2 (data not shown). This shows that in two target genes in the brain, the splicing function of TDP43 in Tardbp+/Q101X mice is altered in a manner similar to that following TDP43 depletion [25], indicating possible partial loss-of-function. The altered splicing pattern for Sort1 was also confirmed with RNA extracted from mouse embryonic fibroblasts (data not shown).

Loss of TDP43 affects neurite outgrowth in a neuroblastoma cell line, primary mouse cortical neurons and Drosophila neurons [37,70,71]. Following on from the potential loss of splicing function of at least some TDP43 target genes in the Tardbp+/Q101X mice, we assessed neuronal development in these animals by establishing primary embryonic motor neuron cultures and measuring neurite outgrowth after 18 hours in vitro. We observed no difference in mean longest neurite length (Tardbp+/+93.2%±4.4% of Tardbp+/+ value) or mean neurite length (Tardbp+/+93.2%±4.4% of Tardbp+/+ value, n = 117 neurons, of Tardbp+/+ value, n = 175 neurons) compared to Tardbp+/+ motor neurons (Figure S2).

Behavioural Characterisation of Tardbp+/Q101X Mice

Grip-strength, rotarod performance and SHIRPA were assessed in male up to one year of age and in female mice up to two years of age. Longitudinal assessment of motor behaviour using grip-strength and rotarod tests revealed no abnormalities in Tardbp+/Q101X mice compared with wildtype controls, in both sexes up to 52 weeks of age and in females to 104 weeks (endpoint of the assessment) (Figure 3A & 3B). There were no significant differences in bodyweight, measured every two weeks, over the lifespan.
of Tardbp+/+ and Tardbp+/Q101X littermates (Figure 3C) and survival did not significantly differ between the two groups (Tardbp+/+ survival = 709±266 days, n=9; Tardbp+/Q101X survival = 634±49 days, n=15; p = 0.235; Figure 3D). Known causes of premature death, such as cancer, did not differ between wildtype and Tardbp+/Q101X mice.

Additionally, as part of the behavioural testing, startle response and pre-pulse inhibition were assessed at 10 and 22 weeks, as well as open field at 14 and 30 weeks of age, but yielded no significant differences between wildtype and heterozygous mutant littermates (data not shown).

Abnormal Limb-clasping and Body Tone in Tardbp+/Q101X Mice

Although motor behaviour and survival were not affected in Tardbp+/Q101X mice, during the SHIRPA assessment significant differences from wildtype littermates were seen with two phenotypes: hindlimb-clasping and body tone (Figure 4). Hindlimb-clasping occurred in 60% of Tardbp+/+ females by two years of age, n = 15, with an average onset of 61 weeks (Figure 4A). Male Tardbp+/Q101X mice did not show this phenotype to a significant extent; however they were not assessed beyond 52 weeks of age.

As well as limb-clasping, Tardbp+/Q101X mice also showed abnormal, softer, abdominal body tone, a component of the SHIRPA which is poorly reported in the literature. This is a highly penetrant phenotype, which manifested with age, and was seen in more than 80% of male and female Tardbp+/Q101X mice, with a mean age of onset in female Tardbp+/Q101X mice of 32 weeks of age (Figure 4B). In comparison, less than 50% of littermate control mice showed a soft body tone (3 of 16 male mice and 4 of 12 female mice at 52 weeks of age: overall 5 of 28). To determine whether the body tone phenotype was associated with reduced neuronal innervation, we examined neuromuscular junctions of abdominal muscles, including the external oblique muscle. No denervation or abnormal neuromuscular junctions were observed (Figure S3).

No Evidence of Hindlimb Neuromuscular Dysfunction in Tardbp+/Q101X Mice at 18 Months of Age

Although no signs of motor dysfunction were evident in grip strength or rotarod performance of Tardbp+/Q101X mice, we conducted an in vivo assessment of hindlimb neuromuscular function, a more sensitive measure of functional decline than grip strength or rotarod tests, in a cohort of male mice at 18 months of age (Tardbp+/+ n=12, Tardbp+/Q101X n=4).

Examples of isometric force recordings are shown in Figure S4. The mean maximum tetanic force of tibialis anterior (TA) and extensor digitorum longus (EDL) muscles are summarised in Figure S4, and muscle contraction, relaxation and fatigue characteristics, as well as EDL motor unit survival are summarised in Figure S5. No differences were seen between Tardbp+/+ and Tardbp+/Q101X male mice in any parameter assessed (data in Table S1).

Alongside neuromuscular assessment, brains from Tardbp+/+ and Tardbp+/Q101X mice were examined for pathology at one year of age. No overt structural abnormalities were observed using a panel of markers against GFAP, IBA-1, p62 and MBP (data not shown).

Does the Tardbp+/Q101X Mutation Affect Transgenic SOD1-ALS Mouse Phenotypes?

Since there is evidence to suggest that SOD1 and TDP43 may interact [20–22], and in order to establish whether motor dysfunction may manifest in Tardbp+/Q101X mice in the presence of an ALS-related stress, Tardbp+/Q101X mice were crossed with SOD1+/Q101X transgenic mice which have been previously been shown to model ALS, including deterioration of hindlimb neuromuscular function and a shortened lifespan ranging between 31–38 weeks [64].

Progeny from this cross were assessed from 7 weeks of age by SHIRPA, grip-strength and rotarod tests, all of which become progressively more abnormal in SOD1+/Q101X mice [63]. Although phenotypes previously seen in Tardbp+/+ and SOD1+/Q101X mice were replicated in the equivalent progeny, no genetic interaction effects were observed in Tardbp+/Q101X, SOD1+/Q101X (double mutant offspring Figure 5), i.e. the mutant offspring were not significantly more badly affected than their SOD1+/- littermates, other than for too relative minor effects on (1) TA muscle relaxation time, and (2) EDL muscle weight, as below. Deficits in grip-strength, bodyweight and survival induced by the SOD1 transgene were not affected by the Q101X mutation (Figure 5A, B and C). The body tone phenotype observed in Tardbp+/Q101X mice was replicated in Tardbp+/Q101X, SOD1+/Q101X mice, which had a mean age of onset of 26 weeks, and this was not exacerbated by the SOD1+/Q101X transgene (Figure 5D).

Physiological assessment of isometric muscle force in TA and EDL hindlimb muscles of 32-33-week old male mice (Tardbp+/+ n=6, Tardbp+/Q101X n=6, Tardbp+/+ n=7, Tardbp+/Q101X, SOD1+/Q101X n=7), a symptomatic stage of disease in SOD1+/Q101X animals, showed that expression of Tardbp+/Q101X had no effect on mean tetanic muscle force (Figure S6A, B). Assessment of TA contraction and relaxation characteristics did not reveal any changes in the TTP of TA muscles (Figure S6C). However, relaxation time of TA muscles was significantly slower in Tardbp+/Q101X, SOD1+/Q101X mice compared to Tardbp+/+, SOD1+/Q101X mice (20.1 ms±3.5 ms n = 10, and 30.4 ms±2.8 ms n = 10, respectively, p = 0.007; Figure S6D) but did not differ between Tardbp+/+ and Tardbp+/Q101X mice. Contraction times of EDL muscles did not differ between groups (Figure 5E). Although relaxation time of the EDL was not significantly different between Tardbp+/+, SOD1+/Q101X and Tardbp+/Q101X, SOD1+/Q101X littersmates, a significant difference was evident between Tardbp+/+ and Tardbp+/Q101X mice (21.4 ms±1.3 ms n = 7, and, 15.9 ms±0.7 ms n = 9, p<0.05, albeit not in a manner which would suggest dysfunction - slower relaxation times are associated with neuromuscular dysfunction (Figure S6F). We note we did not see this difference in previous studies comparing just Tardbp+/+ and Tardbp+/Q101X littersmates. No differences were observed in the fatigue characteristics between groups (Figure S6G). The number of surviving EDL motor units was also assessed and did not reveal significant differences between any groups (Figure S6H; all data also presented in Table S2).

TA and EDL weights were also recorded. TA weight did not differ between Tardbp+/+ and Tardbp+/Q101X mice, but was reduced in Tardbp+/+ and Tardbp+/Q101X mice compared to Tardbp+/+, SOD1+/Q101X littersmates, although this difference was not statistically significant (41.7±2.1 mg, n = 12 and 46.3 mg±1.9 mg, n = 9, respectively; p = 0.051; Figure S7A). EDL muscle weight did not differ between Tardbp+/+ and Tardbp+/Q101X mice, however a significant reduction was seen in Tardbp+/+ and Tardbp+/Q101X mice compared to Tardbp+/+, SOD1+/Q101X mice (10.1 mg±0.7 mg, n = 8 and 12.6 mg±0.9 mg, n = 8, respectively; p = 0.035; Figure 5B).

Finally, no differences were seen in the number of motor neurons in the sciatric motor pool of Tardbp+/+, SOD1+/Q101X and Tardbp+/Q101X, SOD1+/Q101X mice (Figure S7C).
Taken together, these findings show that the Q101X TDP43 mutation does not significantly affect the hindlimb neuromuscular phenotype of SOD1<sup>Q51<sub>106</sub></sup> mice.

**Discussion**

We have carried out a comprehensive assessment of a novel mutant Tardbp mouse with a nonsense mutation predicted to cause a truncation in the N-terminus of TDP43. Characterisation of the Tardbp<sup>Q101X</sup> strain has revealed novel molecular and behavioural phenotypic changes that will help to shed more light on the normal role and pathophysiology of TDP43.

We found Q101X mutation was homozygous embryonic lethal and Tardbp<sup>Q101X/Q101X</sup> embryos died in utero before E6.5. This result indicates a likely loss of function and adds further support for a crucial role for TDP43 during embryonic development; complete deficiency of TDP43 is embryonic lethal [27–29].

Heterozygous Tardbp<sup>+/Q101X</sup> mice had a decrease in the relative abundance of mutant compared to wildtype transcripts. This might arise from either non-sense mediated decay of mutant mRNA due to the presence of an early stop codon or downregulation of expression from the mutant allele. Further investigation may shed more light on Tardbp autoregulation mechanisms [25,48,49] as wildtype transcripts (likely to be full-length) were at similar levels between wildtype and heterozygous mutant littermates.

Full-length TDP43 protein levels are equivalent between Tardbp<sup>+/+</sup> and Tardbp<sup>+/Q101X</sup> littersmates, and we could not detect truncated Q101X protein in brain from aged heterozygous mice. Thus, although we did not look during development, or in all tissues, we might expect these heterozygotes to be normal. Curiously, however, assessment of TDP43 alternative splicing function in Tardbp<sup>+/Q101X</sup> mice showed aberrant exon inclusion in two of five gene targets tested, and Tardbp<sup>+/Q101X</sup> mice developed hindlimb-clasping and an intriguing body tone phenotype. A number of possibilities could explain these phenotypes, including: (1) The upregulation of the wildtype transcripts in response to the null allele (Q101X) may not reach totally normal levels as in Tardbp<sup>+/+</sup> mice and small, undetectable differences, perhaps in particular cellular populations, may give rise to these phenotypes and/or (2) the mutant transcript or very low levels of truncated protein may interfere with normal TDP43 function.

Exon inclusion of two targets, Sort1 and Pdp1, was altered in the same manner as that following TDP43 depletion [25], therefore suggesting a partial loss-of-function effect of the TDP43 Q101X mutation. However, cassette exon inclusion/exclusion of 3 other target genes: Dmunc3, Kcnj5 and Kcnj2, did not differ between Tardbp<sup>+/+</sup> and Tardbp<sup>+/Q101X</sup> mice. Moreover, neurite outgrowth, a process also known to be affected by loss of TDP43 function, was not affected in primary embryonic motor neurons from Tardbp<sup>+/Q101X</sup> mice. It is unclear why cassette exon inclusion/exclusion of Sort1 and Pdp1 are affected yet other targets are not, although this could perhaps indicate differential dependence on TDP43 for regulation of alternative splicing.

Hindlimb claspimg is often observed in rodent models of neuromuscular disease including ALS and has been associated with lower motor neuron degeneration, although it can also be caused by muscular dystrophy [72] and loss of cortical neurons [73]. It has also been noted following motor neuron-specific depletion of TDP43 [52]. Pathological analysis of brains from Tardbp<sup>+/Q101X</sup> mice did not reveal overt signs of degeneration and in vivo assessment of hindlimb neuromuscular function did not suggest lower motor neuron degeneration at 18 months of age.

When body tone of wildtype mice is examined, a basal muscular tone with a reflexive elicited response is normally felt by the assessor [74]. Loss of body tone may therefore be due to muscular alterations or dysfunction of the neuronal reflexive control. Body tone differences have been reported in rat [73] and mouse studies [76,77]. Reduced body tone was detected in Diazepam treated mice [77], which also affected mouse posture by reducing muscle tone in the abdomen and limbs. Mice treated with the acetylcholine esterase inhibitor donepezil also showed a body tone reduction alongside multiple toxic effects ranging from motor deficits through to decreased arousal and pilorecursion [76]. In these reports, body tone reduction is part of a wider array of phenotypes which are the result of muscular and neurological effects. Clearly this phenotype warrants further investigation to identify the cellular pathology in Tardbp<sup>+/Q101X</sup> mice.

In the Tardbp<sup>+/Q101X</sup> mice, the softer body tone and limb-clasping are the only phenotypes detected, and mice may have subtle slowly-progressing neuromuscular deficits that remain mild throughout life. Thus, possible loss-of-function effects in Tardbp<sup>+/Q101X</sup> mice may manifest as subtle neurological dysfunction rather than neuronal loss. Consistent with this possibility, neuronal atrophy, but crucially not loss, has been observed in adult mice with conditional knockout of Tardbp in motor neurons [53]. Furthermore, heterozygous knockout (Tardbp<sup>+/−</sup>) mice displayed motor deficits on an inverted grid test but without evidence of pathological changes in motor neurons and, as here, TDP43 protein levels auto-regulated to match those of wildtype littersmates [27]. Thus potentially, Tardbp<sup>+/−</sup> and Tardbp<sup>+/Q101X</sup> mice may share similar perturbations resulting in muscular and neurological alterations, although no motor deficits were detected in Tardbp<sup>+/Q101X</sup> mice showing that they likely model differing aspects of TDP43 biology.

As TDP43 and SOD1 have been proposed to interact [20–22], we investigated progeny of crossing Tardbp<sup>+/Q101X</sup> mice with the SOD1<sup>Q51<sub>106</sub></sup> transgenic model of ALS. This SOD1-ALS model was chosen in preference to the more commonly used SOD1<sup>Q51<sub>106</sub></sup> mouse, in which ALS-like symptoms are seen much earlier and progress more rapidly [68], so we could identify subtle effects. The TDP43 mutation did not affect survival, bodyweight loss or disease course in Tardbp<sup>+/Q101X</sup>, SOD1<sup>Q51<sub>106</sub></sup> progeny compared to SOD1<sup>Q51<sub>106</sub></sup> littermates. However, TA relaxation time was slower in the double mutants and more extensive atrophy of the EDL muscle was found, albeit in the absence of significant changes in muscle force or motor unit survival, compared to SOD1<sup>Q51<sub>106</sub></sup> mice. Taken together, these results show that although the Q101X mutation partially disrupts TDP43 function, it does not notably modify the ALS-phenotype of SOD1<sup>Q51<sub>106</sub></sup> mice. These findings support published work which was unable to detect any interaction between SOD1 and TDP43 in a C.elegans model of ALS [24].

In summary, the first ENU-induced point mutant Tardbp mouse, the Tardbp<sup>Q101X</sup> strain, provides further evidence for the important role of TDP43 in development and for autoregulation of TDP43. Tardbp<sup>+/Q101X</sup> mice have a likely partial loss-of-function phenotype of exon inclusion of selected targets, and these mice are a useful new genetic model for investigating TDP43 function in vivo. Moreover, Tardbp<sup>+/Q101X</sup> mice develop abnormal hindlimb and body tone phenotypes, in the absence of overt neurodegeneration. The underlying cause of these phenotypes is remains unknown and may indicate previously unknown functions of TDP43. We note that this new mouse strain is freely available for the community to further investigate TDP43 biology.
Supporting Information

Figure S1 TDP43 protein levels do not change in Tardbp+/Q101X mice. (A, B) N-terminal anti-TDP-43 antibodies do not show any novel truncated TDP-43 band in Tardbp+/Q101X soluble or RIPA-insoluble brain fractions from 18 months-old males. N-terminal antibodies used were: (A) Cosmo Bio (CAC-TIP-TD-P07) and (B) Abcam (ab50930). (C) No significant differences in full-length TDP43 protein levels relative to tubulin between Tardbp+/+ (0.99±0.20) and Tardbp+/Q101X (0.76±0.14) using an antibody directed against C-terminus of TDP43 (Proteintech 12892-1-AP). TDP43 levels (green) were assessed from RIPA soluble (p = 0.375) and RIPA-insoluble fractions (p = 0.123) from whole spinal cord lysates using 3 mice per genotype at 18 months of age. Tubulin (red) was used as a loading control. Data are mean±SEM. (TIF)

Figure S2 Neurite outgrowth of primary embryonic motor neurons is not affected by the TardbpQ101X mutation. (A) Representative images of primary embryonic motor neurons stained for the neuronal marker B-III tubulin (red) and DAPI (blue) from Tardbp+/+ and Tardbp+/Q101X embryos. (B) Mean longest neurite length was not significantly different between Tardbp+/+ and Tardbp+/Q101X motor neurons (n = 175 and n = 117 neurons, respectively). (C) Neither were any differences seen between mean neurite length of Tardbp+/+ and Tardbp+/Q101X motor neurons. Data are mean±SEM from 3 independent experiments. (TIF)

Figure S3 TardbpQ101X mice present a normal innervation pattern of the external abdominal oblique muscle. (A, B) Representative immunofluorescence images of whole mount abdominal oblique muscle from Tardbp+/+ (A) and Tardbp+/Q101X (B) mice at ~1 year of age. A normal innervation pattern was present for both genotypes. Postsynaptic, presynaptic and axonal regions were identified by acetylcholine receptor (red), synaptic vesicle protein (green) and neurofilament (green) staining, respectively. Scale bar represent 50 μm in both images. (TIF)

Figure S4 Examples of recordings from in vivo assessment of neuromuscular function of male Tardbp+/+ mice at 18 months of age. (A) Example recording of maximum twitch (smaller peak) and tetanic force (larger peak) from TA muscle. (B) Example trace from an EDL muscle illustrating how contraction time (TTP) and relaxation time (1/2RT) are calculated from maximum twitch force recordings. (C) Example trace demonstrating motor unit number estimation of the EDL. (D) Trace recording of fatigue characteristic of an EDL muscle, where the fatigue index (FI) is calculated as the ratio of force after 180 seconds (F180) compared to initial force (F0). (TIF)

Figure S5 No evidence of neuromuscular dysfunction in male Tardbp+/Q101X mice at 18 months of age. (A) Maximum tetanic force recorded from TA muscles was not significantly different between Tardbp+/+ (n = 12 muscles) and Tardbp+/Q101X (n = 6) mice. (B) No difference was seen in the maximum tetanic force of EDL muscles in Tardbp+/+ (n = 8) and Tardbp+/Q101X (n = 5) mice. (C&D) Assessment of TA contraction (TTP) and relaxation (1/2RT) characteristics in Tardbp+/+ and Tardbp+/Q101X mice (n = 11, and n = 6, respectively) did not reveal any significant differences. (E&F) EDL muscle contraction (TTP) and relaxation (1/2RT) characteristics did not differ between Tardbp+/+ and Tardbp+/Q101X mice (n = 8 and n = 5, respectively). (G) A fatigue index of EDL muscles was established but did not show any difference between Tardbp+/+ (n = 7) and Tardbp+/Q101X (n = 3) mice. (H) The number of motor units innervating EDL muscles was assessed but also failed to show any difference between Tardbp+/+ (n = 8) and Tardbp+/Q101X (n = 4) mice. (TIF)

Figure S6 Assessment of hindlimb neuromuscular function in male Tardbp+/Q101X, SOD1G93Adl mice. (A) Maximum tetanic force recorded from the TA was not significantly different between Tardbp+/+ and Tardbp+/Q101X mice (both n = 10), or between Tardbp+/+, SOD1G93Adl and Tardbp+/Q101X SOD1G93Adl mice (both n = 10). (B) EDL tetanic force did not differ between Tardbp+/+ (n = 8) and Tardbp+/Q101X (n = 9) mice, or between Tardbp+/+, SOD1G93Adl (n = 10) and Tardbp+/Q101X SOD1G93Adl (n = 9) mice. (C&D) Contraction (TTP) and relaxation (1/2RT) of TA muscles did not differ between Tardbp+/+ (n = 9 and n = 8) and Tardbp+/Q101X (n = 13, n = 12) mice, and although TTP did not differ between Tardbp+/+ and Tardbp+/Q101X (n = 10) and Tardbp+/Q101X SOD1G93Adl (n = 10) mice, relaxation time was significantly slower in Tardbp+/Q101X SOD1G93Adl mice compared to Tardbp+/+, SOD1G93Adl mice (both n = 10; p = 0.007). (E) Contraction of EDL muscles was not different between Tardbp+/+ (n = 8) and Tardbp+/Q101X (n = 9) or between Tardbp+/+, SOD1G93Adl (n = 10) and Tardbp+/Q101X SOD1G93Adl (n = 10) mice. (F) EDL relaxation time was significantly quicker in Tardbp+/Q101X (n = 9) mice compared to Tardbp+/+ mice (n = 7; p<0.05), however no difference was observed between Tardbp+/+, SOD1G93Adl (n = 10) and Tardbp+/Q101X SOD1G93Adl (n = 10) mice. (G) Fatigue characteristics of the EDL, defined as the FI, did not differ between Tardbp+/+ (n = 9) and Tardbp+/Q101X (n = 8) mice, or between Tardbp+/+, SOD1G93Adl (n = 7) and Tardbp+/Q101X SOD1G93Adl (n = 7) mice. (H) The number of surviving motor units of EDL muscles was not significantly different between Tardbp+/+ (n = 8) and Tardbp+/Q101X (n = 8) mice, or between Tardbp+/+, SOD1G93Adl (n = 8) and Tardbp+/Q101X, SOD1G93Adl (n = 10) mice. (TIF)

Figure S7 Muscle weights and motor neuron survival in male Tardbp+/+, SOD1G93Adl and Tardbp+/Q101X, SOD1G93Adl mice at 32-33 weeks of age. (A) No difference between TA muscle weights of Tardbp+/+ (n = 12) and Tardbp+/Q101X (n = 17) mice, or between Tardbp+/+, SOD1G93Adl (n = 9) and Tardbp+/Q101X SOD1G93Adl mice (n = 12) (p = 0.051). (B) EDL muscle weight was similar between Tardbp+/+ (n = 12) and Tardbp+/Q101X (n = 17) mice, but was showed a significant difference between Tardbp+/+, SOD1G93Adl (n = 8) and Tardbp+/Q101X SOD1G93Adl (n = 8) mice (p = 0.033). (C) The number of motor neurons of the sciatic pool (L2-L6) was counted in Tardbp+/+, SOD1G93Adl (n = 3) and Tardbp+/Q101X, SOD1G93Adl (n = 4) mice and did not reveal any differences. (TIF)

Table S1 Normal hindlimb neuromuscular function in Tardbp+/Q101X male mice at 18 months of age. (DOCX)

Table S2 The Q101X mutation in TDP43 does not affect neuromuscular function in 32-33 week old SOD1G93Adl mice. (DOCX)

Author Contributions

Performed the experiments: TR PM PF HMO GB RK VP SB AA-A. Analyzed the data: TR PM PF SB AA-A. Contributed reagents/materials/analysis tools: SB LG AA-A EF. Wrote the paper: TR PM PF LG AA-A EF.
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