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# Riluzole does not improve lifespan or motor function in three ALS mouse models.

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1 **Riluzole does not improve lifespan or motor function in three ALS mouse models**

2

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35 **Key Words**

36 Riluzole, ALS, SOD1, transgenic animals

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40 **Abbreviations**

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42 ALS Amyotrophic Lateral Sclerosis

43 DMSO Dimethyl Sulfoxide

44 FUS Fused in Sarcoma

45 Ril Riluzole (2-amino-6-trifluoromethoxy benzothiazole)

46 Non-tg Non-transgenic

47 PND Post Natal Day

48 Ril Riluzole

49 SOD1 Superoxide Dismutase 1

50 TDP-43 Transactivation Domain Protein of 43 kDa

51 Tg Transgenic

52 Veh Vehicle

53

54 **Abstract**

55 **Background:** Riluzole is the most widespread therapeutic for treatment of the  
56 progressive degenerative disease amyotrophic lateral sclerosis (ALS). Riluzole gained  
57 FDA approval in 1995 before the development of ALS mouse models. We assessed  
58 Riluzole in three transgenic ALS mouse models: the SOD1<sup>G93A</sup> model (1), the TDP-  
59 43<sup>A315T</sup> model (2), and the recently developed FUS (1-359) model (3).

60

61 **Methods:** Age, sex and litter-matched mice were treated with Riluzole (22 mg/kg) in  
62 drinking water or vehicle (DMSO) from symptom onset. Lifespan was assessed and  
63 motor function tests were carried out twice weekly to determine whether Riluzole  
64 slowed disease progression.

65

66 **Results:** Riluzole treatment had no significant benefit on lifespan in any of the ALS  
67 mouse models tested. Riluzole had no significant impact on decline in motor  
68 performance in the FUS (1-359) and SOD1<sup>G93A</sup> transgenic mice as assessed by Rotarod  
69 and stride length analysis.

70

71 **Conclusions:** Riluzole is widely prescribed for ALS patients despite questions  
72 surrounding its efficacy. Our data suggests that if Riluzole was identified as a  
73 therapeutic candidate today it would not progress past pre-clinical assessment. This  
74 raises questions about the standards used in pre-clinical assessment of therapeutic  
75 candidates for the treatment of ALS.

76

77 **Introduction**

78 ALS is a progressive debilitating disease characterised by a progressive loss of motor  
79 neurons which leads to muscle wasting, paralysis and death. Despite investigation of  
80 over 60 molecules as potential therapeutics for ALS there are currently only two FDA-  
81 approved treatments, recent clinical trials are summarised in (4).

82

83 Riluzole was initially investigated as a therapeutic for ALS following a report showing  
84 that it can inhibit synaptic release of glutamate in hippocampal slices (5). It has  
85 subsequently been linked with additional pharmacological activities including  
86 modulation of AMPA (6), and GABA receptors (7), inhibition of persistent sodium and  
87 calcium currents (8, 9), and activation of AMP-activated protein kinase (10) in neurons,  
88 and stimulation of NGF and BDNF in astrocytes (11), reviewed in (12).

89

90 A phase I clinical trial revealed a 200 mg daily dose of Riluzole was well tolerated in  
91 healthy people however no ALS mouse models were available at the time for preclinical  
92 testing. Therefore it was accelerated into a randomized control trial where it showed  
93 beneficial effects which were more pronounced in patients with bulbar onset than those  
94 with limb onset (13). Following this trial Riluzole was approved by the FDA in  
95 December 1995 as patients treated with Riluzole showed significantly increased  
96 lifespan along with significantly less deterioration in muscle strength when compared  
97 to the placebo group. A larger follow up study investigated Riluzole doses of 50 mg,  
98 100 mg, and 200 mg daily, and this showed that the 100 mg dose gave the best benefit-  
99 to-risk ratio due to increased serum alanine transferase levels at the 200 mg dose (14).  
100 A third clinical trial, which included older patients treated at a later disease stage,  
101 showed Riluzole gave no benefit (15). A Cochrane Review of clinical trials published  
102 in 2012 concluded that 100 mg Riluzole daily is reasonably safe and probably prolongs  
103 lifespan for around 2-3 months in ALS patients (16). However reports persist that only  
104 a subset of patients benefit from taking Riluzole which was confirmed in a small study  
105 (Sojka 1997).

106

107 In this study we aimed to assess the therapeutic effect of Riluzole on lifespan and motor  
108 function in three separate ALS mouse models. Surprisingly we found that systemic  
109 dosing of Riluzole in drinking water from symptom onset had no effect on lifespan or

110 motor function in any of the preclinical ALS mouse models tested, emphasizing the  
111 difficulties regarding the use of transgenic ALS mouse models in the pre-clinical  
112 assessment of therapeutic candidates for the treatment of ALS.

113

114

## 115 **Methods**

116

### 117 **Animal strains**

118 SOD1<sup>G93A</sup> mice (C57B6.Cg-Tg (SOD1<sup>G93A</sup>) 1Gur/J mice were purchased from The  
119 Jackson Laboratory (Bar Harbor, Maine) and originally generated in the laboratory of  
120 Professor Siddique (1). The SOD1<sup>G93A</sup> transgene copy number was verified in breeding  
121 males from our colony, see Supplementary Figure 1. TDP-43<sup>A315T</sup> mice on a congenic  
122 C57Bl/6 background (B6.Cg-Tg(Prnp-TARDBP\*A315T)95Balo/J) were  
123 purchased from The Jackson Laboratory (Bar Harbour, Maine, USA) and originally  
124 generated in the laboratory of Dr Baloh (2). FUS (1-359) mice generated in the  
125 laboratory of Professor Buchman (3) were re-derived at the Institute of Molecular  
126 Genetics ASCR, Prague, Czech Republic, they are congenic on the C57Bl/6  
127 background.

128

### 129 **Animal Maintenance**

130 Mice were housed at constant temperature (22 °C) on a 12 h light/dark cycle (07:00 h  
131 on, 19:00 h off), with *ad libitum* food and water. Experimental mice from the TDP-  
132 43<sup>A315T</sup> colony were weaned at post-natal day (PND) 21 at which time they were  
133 switched on to a high fat jelly diet (DietGel Boost, Clear H20 Maine, USA). Pups from  
134 litters of the same generation were housed in groups of 3-5 per cage. Genotyping was  
135 performed using primers and conditions for SOD1<sup>G93A</sup> and TDP-43<sup>A315T</sup> available at  
136 [www.jax.org](http://www.jax.org), and for FUS (1-359) in (3). Ethical approval was received for this project  
137 from the RCSI Research Ethics Committee (REC447 & REC1122) and licences were  
138 obtained from the Health Products Regulatory Authority (HPRA: AE19127/P003 and  
139 AE19127/P004).

140

### 141 **Assessment of lifespan and disease progression**

142 Animals in the study were age-, sex- and litter-matched according to ALS community  
143 preclinical guidelines (17) and sample size power calculations are provided in  
144 Supplementary Data. End stage of disease in all strains was determined by loss of  
145 righting reflex when mice were placed on their back for 15 seconds according to the  
146 ALS guidelines (17). Non-transgenic littermates were culled when no transgenic mice  
147 remained in the cage. Motor function performance was assessed by Rotarod (Stoelting,  
148 IL, USA) and stride length measurements and weight was monitored. Mice were trained  
149 on motor function equipment for 2 weeks prior to the start of recording. Observers were  
150 blinded to the treatment groups when motor function data was being recorded.

151

### 152 **Statistical analysis**

153 Motor function data are presented as mean +/- SEM and statistical significance was  
154 assessed by one way ANOVA with post-hoc Tukey's test. Survival data were analysed  
155 by Kaplan-Meier curves with significance determined by Mantel-Cox test. Statistical  
156 analyses were performed in SPSS statistics software (IBM).

157

### 158 **Drug preparation and dosing**

159 Riluzole was purchased in powdered form from AKScientific (California, USA) and  
160 reconstituted in DMSO at 22 mg/ml. The stock was diluted to a final concentration of  
161 137.5 µg/ml in drinking water. Based on the assumptions that an adult mouse weighs  
162 approximately 25g and drinks 4 ml liquid per day, this sums up to an approximate dose  
163 of 22 mg/kg/day. This dose was chosen based on previous studies as it has been shown  
164 that plasma Riluzole concentrations reach similar levels in mice as those in ALS  
165 patients treated with 50 mg Riluzole twice daily (18-21). Vehicle solutions were made  
166 using DMSO at the same dilution.

167

### 168 **Results**

169 We investigated the potential therapeutic effect of Riluzole in three genetically distinct  
170 mouse models of ALS: the SOD1<sup>G93A</sup> mice (1); the newer TDP-43<sup>A315T</sup> (2) and FUS  
171 (1-359) mouse models (3), Figure 1 shows a comparison of symptom onset and disease  
172 progression in these models. Mutations in SOD1 contribute to approximately 20% of  
173 familial and 2% of sporadic ALS cases (22). A mouse model containing multiple copies  
174 of a human mutant SOD1<sup>G93A</sup> transgene is the most established model for preclinical



175 testing of therapeutics (1). SOD1<sup>G93A</sup> mice show uniform disease progression with  
176 transgenic (Tg) animals showing symptoms from PND 90, therefore treatment was  
177 started at PND 90. Mice develop a slow progressing hind limb weakness which  
178 eventually leads to paralysis and the humane end point in our colony occurs at PND  
179 160-180.

180

181 The TDP-43<sup>A315T</sup> mouse congenic on a C57Bl/6 background develops severe  
182 gastrointestinal problems which leads to premature death at around PND 100 due to  
183 neuronal degeneration within the solar plexus, which leads to loss of innervation in the  
184 gut and severely decreased gastrointestinal motility (23, 24). Feeding a high calorie  
185 jellified diet alleviates the gastrointestinal defect and allows the mice to live long  
186 enough to develop a motor neuron disease phenotype (24, 25). This TDP-43<sup>A315T</sup> model  
187 shows variation between male and female mice, with female mice living almost twice  
188 as long as male mice and showing more variable disease penetrance (23, 26), therefore  
189 we used male TDP-43<sup>A315T</sup> mice raised on a high fat jelly diet. Treatment began at PND  
190 60 due to the observation that the gastrointestinal defects manifest from PND 60  
191 onwards and are not completely alleviated in this model (25).

192

193 The FUS (1-359) mice express a truncated fragment of the human FUS gene under the  
194 Thy-1 promoter, which leads to cytoplasmic mis-localisation and aggregation of FUS  
195 protein in neurons (3). Tg mice develop a severe motor phenotype which displays  
196 considerable variation in symptom onset and a rapid disease course (time between  
197 symptom onset and death is less than 2 weeks). This mirrors the fast disease progression  
198 often seen in FUS mutant fALS patients (27).

199

200 To determine whether Riluzole could extend lifespan we performed a lifespan study in  
201 FUS (1-359) mice. Age, sex, and litter-matched groups were treated with Riluzole (22  
202 mg/kg; Tg n=8 male and 8 female, Non-Tg n=4 male and 4 female) in drinking water  
203 or vehicle (DMSO; Tg n=12 male and 12 female, Non-tg n=6 male and 6 female). The  
204 earliest death in our FUS (1-359) colony occurred at PND 64; therefore treatment began  
205 at PND 50. Kaplan-Meier survival analysis for Tg mice treated with Riluzole and  
206 vehicle show that there was no significant difference between treatment groups (Fig  
207 2A, p=0.271). Analysis of the genders separately revealed that female Tg mice, but not  
208 male Tg mice, showed an increase in lifespan with Riluzole treatment compared to

209 vehicle-treated mice (Fig 2B and C), however this was not statistically significant  
210 ( $p=0.265$ ).

211

212 To assess whether Riluzole treatment could delay symptom onset or reduce the rapid  
213 decline in motor function in FUS (1-359) mice we assessed motor performance and  
214 monitored weight throughout treatment. Age-, sex- and litter-matched groups of mice  
215 were treated with Riluzole (22 mg/kg) in drinking water (Tg n=16, Non-transgenic  
216 (Non-Tg) n = 8) or vehicle (DMSO; Tg n=24, Non-Tg n=12). No significant difference  
217 in motor function ability could be detected between Tg mice treated with Riluzole or  
218 vehicle by Rotarod (Fig 3A) or stride length analysis (Fig 3B). No differences were  
219 observed when analysis was performed on gender separated groups (data not shown).  
220 Interestingly, the motor neuron degeneration that occurred in the FUS (1-359) mice was  
221 not accompanied by a change in weight as seen in other ALS mouse models (Fig 3C).  
222 Non-Tg mice treated with Riluzole (22 mg/kg) or vehicle (DMSO) showed no  
223 difference in viability or behaviour and motor function was not affected (Fig 3).

224

225 We then went on to assess Riluzole (22 mg/kg) in drinking water in SOD1<sup>G93A</sup> mice.  
226 These mice show a more uniform onset of degeneration and a more gradual decline in  
227 motor function (see Fig 1). Age and litter-matched groups were treated from symptom  
228 onset (PND 90) with Riluzole (22 mg/kg, Tg n=4 male and 4 female, non-Tg n=2 male  
229 and 2 female) in drinking water or vehicle (DMSO, Tg n=5 male and 8 female, Non-  
230 Tg n=4 male and 2 female). Riluzole had no significant effect on lifespan in transgenic  
231 mice from the SOD1<sup>G93A</sup> colony when compared to vehicle (Fig. 4;  $p=0.427$ ). No  
232 differences were observed in gender separated groups (data not shown).

233

234 To determine whether Riluzole could delay symptom onset or improve motor function  
235 in SOD1<sup>G93A</sup> mice treated from symptom onset (PND 90) were assessed by weekly  
236 motor function testing (Fig 5). Rotarod analysis revealed Tg mice showed a slow  
237 decline in motor function over 10 weeks and there was no significant difference  
238 between Tg mice treated with Riluzole (22 mg/kg) in drinking water or vehicle (DMSO,  
239 Fig 5A). Similarly Riluzole treatment had no effect on stride length or weight which  
240 both declined gradually across disease progression in Tg SOD1<sup>G93A</sup> mice (Fig 5B & C).

241

242 Finally we assessed the efficacy of Riluzole from PND 60 in age- and litter-matched  
243 male TDP-43<sup>A315T</sup> mice raised on a high calorie jellified diet. Kaplan-Meier analysis of  
244 survival revealed no significant difference between Riluzole (average lifespan 170.8  
245 days +/- 10.9, n=6) and vehicle (average lifespan 166.7 days +/- 10.7, n=6 p=0.79 two  
246 tailed t test) (Fig 6). Despite the high calorie jelly diet the intestinal phenotype in the  
247 TDP-43<sup>A315T</sup> mice is not completely corrected (see (25)) making assessment of motor  
248 performance in these mice difficult, therefore only lifespan data was recorded.

249

250

## 251 **Discussion**

252 Despite widespread use of the SOD1<sup>G93A</sup> ALS mouse model in preclinical trials many  
253 therapeutics that show promising results have failed to show positive effects in patients.  
254 This was highlighted as an issue impacting development of novel therapeutics in the  
255 recently published ALS community guidelines which recommended development of  
256 new ALS mouse models to improve the translatability of preclinical research in ALS  
257 (17). Riluzole received FDA approval in 1995 before ALS mouse models were widely  
258 available; however two later studies showed that Riluzole (in drinking water or food)  
259 can extend lifespan in SOD1<sup>G93A</sup> mice (18, 28). Subsequent studies into the efficacy of  
260 Riluzole have been performed: in low copy number SOD1<sup>G93A</sup> mice Riluzole in  
261 drinking water (22 mg/kg) from PND 40 delayed symptom onset (29), in high copy  
262 number SOD1<sup>G93A</sup> mice Riluzole (30 mg/kg) in drinking water from PND 60 had no  
263 significant effect on lifespan in one study (30) but treatment with Riluzole (16 mg/kg)  
264 in drinking water from PND 30 showed significant lifespan extension in another (31).  
265 The different outcomes of these studies has been attributed to different treatment  
266 paradigms, low animal numbers, and lack of gender balanced groups. In 2008 the ALS  
267 Therapy Development Institute (TDI) systematically reviewed compounds which had  
268 been published as significantly increasing lifespan in SOD1<sup>G93A</sup> mice (19).  
269 Unfortunately they could not replicate the published beneficial effects, including those  
270 for Riluzole (22 mg/kg in drinking water) which had no significant effect on lifespan  
271 (19). An ALS TDI update re-assessing nine compounds found that none of the initial  
272 preclinical trial results could be replicated (32).

273

274 Here we utilised Riluzole as a benchmark to assess the suitability of other preclinical  
275 mouse models. Initially we used the FUS (1-359) mouse model (3) but found no  
276 significant effect of Riluzole (22 mg/kg in drinking water) on lifespan when compared  
277 to vehicle, irrespective of gender, and no significant effect on motor performance. We  
278 next trialled Riluzole in SOD1<sup>G93A</sup> mice and our data support the results from the ALS  
279 TDI in that we saw no significant effect of Riluzole on lifespan or motor performance  
280 (19, 32). Finally we assessed Riluzole in TDP-43<sup>A315T</sup> mice on a high calorie jellified  
281 diet; Riluzole had no significant effect on lifespan compared to vehicle. Hence we  
282 conclude that Riluzole does not extend lifespan or improve motor performance in three  
283 preclinical ALS mouse models.

284

285 During the writing of this manuscript Edaravone was granted FDA approval as a  
286 therapeutic for ALS. Interestingly, however this drug also failed to show consistent,  
287 beneficial effects on lifespan in ALS rodent models. Edaravone is a free-radical  
288 scavenger which was originally investigated for its neuroprotective effects following  
289 cerebral ischaemia (reviewed in (33)). Interestingly Edaravone (15 mg/kg daily i.p.)  
290 showed improved motor function and preserved motor neurons in the spinal cord in  
291 SOD1<sup>G93A</sup> female mice, however it had no significant effect on survival (34). In a  
292 further study in the SOD1<sup>H46R</sup> rat model Edaravone had no significant effect on survival  
293 (35). Edaravone was licenced for treatment of cerebral ischaemia in Japan in 2001,  
294 therefore it was fast-tracked into clinical trials for ALS despite the negative results in  
295 preclinical models (36-39). This highlights the challenges facing novel therapeutics for  
296 ALS where many drugs fail at preclinical trials including the only two currently  
297 licenced therapeutics.

298

299 Given the pleiotropic nature of ALS there is potential that the mouse models used here  
300 do not fully recapitulate the pathophysiology of ALS, or the precise defect targeted by  
301 Riluzole. The transgenic models used here recapitulate several important aspects of  
302 ALS pathogenesis but we have to assume that no single model generated to date  
303 captures them all. The SOD1<sup>G93A</sup> model recapitulates many aspects of familial ALS  
304 caused by mutations in the SOD1 gene, including aggregation of mutant SOD1 protein  
305 and impaired proteasome function, however SOD1 mutations account for

306 approximately 20% of fALS and only 2% of sALS cases suggesting the wider relevance  
307 of this model may be limited (22). Aberrant RNA processing has been identified as a  
308 pathological mechanism in ALS with a majority of sporadic ALS patients showing  
309 cytoplasmic TDP-43 positive inclusions in neurons (40). We did not observe a  
310 beneficial effect of Riluzole in the TDP-43<sup>A315T</sup> model, however this model does not  
311 develop cytoplasmic TDP-43 inclusions (2) and despite the high calorie jelly diet the  
312 intestinal phenotype is not completely corrected (25), hence this model has limitations.  
313 Therefore we utilised the newer FUS (1-359) mice (3), which develop FUS positive  
314 neuronal inclusions that are distinct from stress granules (41) but may affect RNA  
315 metabolism via sequestration of endogenous FUS, recapitulating proteinopathy and  
316 RNA metabolism defects. The FUS model is the most wide-ranging of our three  
317 models; yet treatment with Riluzole did not extend lifespan or improve motor function.

318

319 ALS mouse models provide valuable tools to investigate the pathogenic mechanism of  
320 ALS-associated mutations and can provide important information on common  
321 pathways involved in pathogenesis which may reveal therapeutic targets. However, our  
322 study raises important questions surrounding the use of transgenic ALS mouse models  
323 in preclinical studies and the stringency by which we assess success. If Riluzole were  
324 investigated today as a novel therapeutic it would not proceed on to clinical trials yet it  
325 has documented beneficial effects in a subset of ALS patients. Conversely, many  
326 therapeutics for ALS that show significant benefit in ALS mouse models fail in clinical  
327 trials, highlighting the need for development of additional platforms (such as patient-  
328 derived, induced pluripotent stem cells) for the pre-clinical testing of novel ALS  
329 therapeutics.

330

331

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497

498 **Disclosure Statement**

499 The authors report no conflicts of interest.

500 **Figure legends**

501

502 **Figure 1: Cartoon comparison of disease progression in the three ALS mouse**  
503 **models used in this study.** Graphical representation of disease onset and duration in  
504 A) FUS (1-359), B) TDP-43<sup>A315T</sup> male mice on a high calorie jellified diet and C)  
505 SOD1<sup>G93A</sup> ALS mouse models.

506

507 **Figure 2: Riluzole treatment does not extend lifespan in the FUS (1-359) mouse**  
508 **model.** A) Kaplan-Meier analysis of survival in transgenic FUS (1-359) mice treated  
509 with Vehicle (blue, DMSO, n=24) or Riluzole (green, 22 mg/kg, n=16) in drinking  
510 water from PND 50. No significant effect on lifespan was recorded. B) Lifespan  
511 analysis in male transgenic FUS (1-359) mice treated with DMSO (n=12) or Riluzole  
512 (n=8) showed no significant difference in lifespan. C) Lifespan analysis in female  
513 transgenic FUS (1-359) mice showed no significant difference between DMSO (n=12)  
514 or Riluzole (n=8) treated groups.

515

516 **Figure 3: Riluzole treatment does not improve motor function in the FUS (1-359)**  
517 **mouse model.** A) Rotarod assessment of motor function was performed throughout  
518 treatment in transgenic (Tg) and non-transgenic (Non-Tg) FUS (1-359) mice treated  
519 with Riluzole (22 mg/kg) or Vehicle (DMSO) in drinking water from PND 50 onwards.  
520 No significant difference in onset or rate of decline could be detected. B) Analysis of  
521 stride lengths revealed no significant difference between Tg mice treated with vehicle  
522 (DMSO) or Riluzole (22 mg/kg). C) No significant difference in weight could be  
523 detected between Tg or Non-Tg mice, or between those treated with Riluzole (22  
524 mg/kg) or vehicle (DMSO). Data shows mean +/- SEM, statistical significance was  
525 assessed by one way ANOVA with post-hoc Tukey's.

526

527 **Figure 4: Riluzole treatment does not improve lifespan in the SOD1<sup>G93A</sup> mouse**  
528 **model.** Kaplan-Meier analysis of survival in the SOD1<sup>G93A</sup> mouse model treated with  
529 Riluzole (22 mg/kg) or vehicle (DMSO) in drinking water from PND 90 onwards.  
530 There is no significant difference in survival between Riluzole treated Tg mice (green,  
531 n=8) and vehicle (DMSO) treated mice (blue, n=13, p=0.427).

532

533 **Figure 5: Riluzole treatment does not improve motor function in the SOD1<sup>G93A</sup>**  
534 **mouse model.** A) Assessment of coordination and balance via rotarod testing revealed  
535 no significant difference between Tg mice treated with Riluzole (22 mg/kg, n=8) or  
536 vehicle (DMSO) treated mice (n=13). Both groups showed a gradual decrease in motor  
537 skills across disease progression. B) There was no significant difference between stride  
538 lengths measured across disease progression in Tg mice treated with Riluzole or vehicle  
539 (DMSO) control. C) There was no significant difference in weight between Tg mice  
540 treated with Riluzole and Tg mice treated with vehicle (DMSO). Data shows mean +/-  
541 SEM, statistical significance was assessed by one way ANOVA with post-hoc Tukey's.  
542

543 **Figure 6: Riluzole treatment does not extend lifespan in TDP-43<sup>A315T</sup> mice.** Kaplan-  
544 Meier analysis of mice treated with Riluzole (22 mg/kg) or vehicle (DMSO) in drinking  
545 water from PND 60 onwards. There was no significant difference in survival between  
546 Tg mice treated with Riluzole (green, n=6) or vehicle (DMSO, blue, n=5, p=0.975).