# 1 Knockout of zebrafish desmin genes does not cause skeletal muscle

# 2 degeneration but alters calcium flux.

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# 20 Abstract

Desmin is a muscle-specific intermediate filament protein that has fundamental role in muscle 21 22 structure and force transmission. Whereas human desmin protein is encoded by a single gene, two 23 desmin paralogs (desma and desmb) exist in zebrafish. Desma and desmb show differential 24 spatiotemporal expression during zebrafish embryonic and larval development, being similarly expressed in skeletal muscle until hatching, after which expression of *desmb* shifts to gut smooth 25 muscle. We generated knockout (KO) mutant lines carrying loss-of-function mutations for each 26 gene by using CRISPR/Cas9. Desma: desmb double mutants are viable and fertile, and lack obvious 27 28 skeletal muscle, heart or intestinal defects. In contrast to morphants, knockout of each gene did

not cause any overt muscular phenotype, but did alter calcium flux in myofibres. These results
point to a possible compensation mechanism in these mutant lines generated by targeting nonsense
mutations to the first coding exon.

# 32 Introduction

Desmin is a type III intermediate filament protein that is specifically expressed in skeletal, cardiac 33 and smooth muscles. In addition to their fundamental role in maintaining the structural integrity 34 35 of the sarcomere, desmin intermediate filaments are involved in mechanotransduction and organelle positioning. Desminopathies belonging to myofibrillar myopathies are primarily 36 characterized by abnormal protein aggregates in muscle and present as progressive skeletal 37 myopathy and/or cardiomyopathy<sup>1</sup>. Although the involvement of smooth muscles is not widely 38 39 reported, some patients suffer from smooth muscle dysfunction such as swallowing difficulties, intestinal pseudo-obstruction and respiratory insufficiency<sup>2–4</sup>. The majority of desmin mutations 40 are associated with desmin aggregates; however, some mutations have been reported to not alter 41 filament assembly and network integrity in vivo or in vitro<sup>5,6</sup>. KO of desmin in mice does not affect 42 43 the viability and the development of muscles; however, muscle degeneration and cardiomyopathy are observed<sup>7,8</sup>. 44

Zebrafish is a widely used model for neuromuscular disorders<sup>9,10</sup>. Besides the advantage 45 of being the most abundant tissue in zebrafish; the gene profile, and the structural and histological 46 features of mammalian skeletal muscle are highly preserved in zebrafish<sup>11</sup>. To date, two 47 48 approaches have been used to study the effect of the loss of desmin in zebrafish. On one hand, it was shown that morpholino-mediated knockdown of desmin causes skeletal and cardiac muscle 49 myopathy<sup>12,13</sup>. On the other hand, Ramspacher et al. studied a desmin mutant line (desma<sup>sa5</sup>), 50 generated by ENU mutagenesis, which causes a truncation mutation approximately half way along 51 the molecule, that also presented with skeletal and cardiac muscle phenotype<sup>14</sup>. 52

Gene duplications are common in zebrafish and most duplicated genes have similar functions to their human orthologs. Alternatively, paralogs can be expressed in different tissues or stages of development and have distinct functions to their human orthologs. Humans carry a single copy of the *DES* gene whereas two desmin genes (*desma* and *desmb*) are found in zebrafish<sup>13</sup>. Previous work has not established the tissue specific distribution and role of these two desmin paralogs.

Here, we report the spatiotemporal expression pattern of the two desmin paralogs in 59 zebrafish, as well as the generation using CRISPR/Cas9 and characterization of putative null 60 61 zebrafish mutant lines for each of the desmin paralogs. We show that desma and desmb are expressed in skeletal muscle until 72-hour post-fertilization (hpf). After 72 hpf, expression of 62 desmb shifts to gut smooth muscle. Putative null mutation of each gene does not affect viability 63 and adults do not develop any overt phenotype. However, altered calcium flux was observed in 64 desma-KO myofibres. Surprisingly, desma; desmb double mutants also survive to adulthood with 65 no sign of muscle defect. These results show that loss-of-function of desmin genes by mutations 66 in the first coding exon result in a mild phenotype with no visible muscle degeneration but altered 67 calcium flux, in contrast to morpholino-mediated knockdown models and the desma<sup>sa5</sup> allele. 68

# 69 **RESULTS**

#### 70 Zebrafish desmin genes

Zebrafish have two desmin orthologs, desma (desmin a) and desmb (desmin b), which are located 71 on chromosome 9 and 6, respectively. Desma and desmb paralogs share 81% and 83% identity 72 with human *DES* gene. Although a single transcript is known to be encoded by human *DES* gene, 73 in ENSEMBL two isoforms are predicted to be transcribed from *desma*, *desma-1* mRNA coding 74 for a 488 amino acid protein (predicted molecular weight of 55.7 kDa) and desma-2 mRNA coding 75 for a 473 amino acid protein (predicted molecular weight of 54.1 kDa). In desma-2 transcript, exon 76 9 is skipped, corresponding to 15 amino acid located in the tail domain of Desma-1 protein 77 (Supplementary Fig. S1). The 473 amino acid Desmb protein is predicted to have a molecular 78 79 weight of 54.2 kDa. At the amino acid level, Desma-1, Desma-2 and Desmb show 80%, 82% and 80 83% similarity with human desmin protein.

### 81 Differential expression of desmin transcripts during zebrafish development

We analyzed the spatiotemporal expression patterns of *desma* and *desmb* genes during zebrafish development at several stages beginning from somitogenesis until 5 days post fertilization (dpf) by whole mount *in situ* hybridization (ISH). For detecting both *desma* transcripts (*desma-2* lacks exon 9), we synthesized an antisense probe by using a forward primer binding to exon 1 and a reverse primer binding to exon 10 (Supplementary Fig. S1). No specific staining was observed in embryos treated with sense probes for each gene (Supplementary Fig. S2). *Desma* was first

detected at 11 hpf in adaxial slow muscle precursor cells when the first somites form and remained 88 strongly expressed in somites at all examined stages (Fig. 1a). Desma was also expressed in the 89 developing heart from 35 hpf (Fig. 1b). Cryosections of 72 hpf embryos showed that desma was 90 distributed in the entire somite, including epaxial and hypaxial muscles (Fig. 1c). By 72 hpf, desma 91 was expressed in cranial muscles including external ocular muscles, opercular muscles and 92 mandibular muscles (Fig. 1d). At 96 hpf, transversal sections revealed desma expression at the 93 anterior intestine (Fig. 1d, upper panel). At 72, 96 and 120 hpf, expression of desma in pectoral 94 fin muscles was clearly visible (Fig. 1d). Desmb had an overlapping expression pattern to desma 95 from the beginning of somitogenesis until 72 hpf, including staining in trunk somites and heart 96 (Fig. 1a, b). At 72 hpf, somitic expression of desmb dramatically decreased with a residual 97 expression in the lateral edges of the myotome. By contrast, strong signal of *desmb* transcripts was 98 99 detected around the gut throughout the length of the intestine (Fig. 1a, c, e). Similar to desma, desmb was also expressed in pectoral fin muscles, operculum muscles and mandibular muscles at 100 72 hpf (Fig. 1e). These results indicate that desma and desmb expression partially overlap during 101 zebrafish muscle development. As the gut develops, *desmb* expression shifts from somitic muscle 102 103 to gut smooth muscle.

### 104 Generation of *desma* and *desmb* knockout zebrafish lines

Desma and desmb mutant zebrafish lines were generated by using CRISPR/Cas9 genome editing. 105 The  $desma^{kg97}$  line exhibited 2 bp deletion and 4 bp insertion in the first exon 106 (g.198 99delGAinsTGAT, NC 007120.7) leading to a frameshift at amino acid 45 and a 107 premature stop codon after four amino acids. In the *desmb*<sup>kg156</sup> mutant line, a 5 bp deletion in exon 108 1 (c.245 249delCTTAT, NM 001077452.1) resulted in a frameshift at amino acid 82 and 109 introduced a premature stop codon after three amino acids (Fig. 2a). Heterozygous mutants showed 110 111 no defect and homozygous mutant embryos and larvae developed relatively normally, at least for the first few days (Fig. 2b-f). 112

### 113 Expression of *desma* and *desmb* in knockout lines

114 Mutations were predicted to trigger nonsense-mediated decay of mRNA or synthesis of short 115 truncated polypeptides. As expected, qRT-PCR revealed a significant decrease of *desma* 116 transcripts in *desma*<sup>kg97</sup> homozygous embryos (P=0.0119, Mann-Whitney U), while *desmb*  transcripts were significantly decreased in  $desmb^{kg156}$  homozygous embryos at 96 hpf compared to wild type (WT) (*P*=0.0079, Mann-Whitney U) (Fig. 2b). Consistent with El-Brolosy et al.<sup>15</sup>, weak *desmb* or *desma* mRNA up-regulation may occur in the respective mutant embryos (Fig. 2b).

In order to investigate the potential compensatory effect of *desmin* paralog upregulation in 120 mutant embryos, expression patterns of *desmb* and *desma* transcripts were determined in *desma*<sup>kg97</sup> 121 or  $desmb^{kg156}$  homozygous mutant embryos, respectively. In  $desma^{kg97/kg97}$  embryos, the expression 122 pattern of desmb transcripts (Fig. 2c) was similar to that of wild-type embryos (Fig. 1a). Somitic 123 expression of *desmb* was observed in *desma<sup>kg97/kg97</sup>* at 48 hpf, and a shift towards gut expression 124 after 72 hpf. Beyond 72 hpf, no *desmb* staining was detected in somites of *desma<sup>kg97/kg97</sup>* embryos. 125 In desmb<sup>kg156/kg156</sup> embryos, desma was still strongly expressed in somites and visible in the 126 anterior intestine at 96 hpf (Fig. 2c). In order to confirm the loss-of-function of desmin in mutants 127 at the protein level, expression of Desma and Desmb in WT, *desma<sup>kg97</sup>* or *desmb<sup>kg156</sup>* homozygous 128 embryos at 96 hpf was determined by immunofluorescent staining using a desmin antibody 129 recognizing both proteins (Fig. 2d-f). In WT embryos, desmin staining was detected in both 130 somites and gut (Fig. 2d). As expected from ISH results, somitic expression was lost and gut was 131 preferentially stained in desmakeg97/kg97 mutants (Fig. 2f). In desmbkg156/kg156 mutants, expression in 132 somites and anterior intestine were preserved while staining of the middle and posterior intestine 133 was lost (Fig. 2e). We conclude that mutation of each gene reduces the cognate mRNA and protein 134 at all stages examined and that compensation by up-regulation of the unmutated paralogous gene 135 136 is at best very weak.

## 137 General larval characteristics of desmin mutants

In order to evaluate the effects of the absence of *desma* or *desmb* on development, WT, *desma<sup>kg97</sup>* 138 and *desmbkg156* homozygous embryos/larvae were compared. No significant difference was 139 observed in the number of viable eggs between  $desma^{kg97/kg97}$  and WT (P=0.6842, Mann-Whitney 140 U) or desmb<sup>kg156/kg156</sup> and WT (P=0.9654, Mann-Whitney U) (Fig. 3a). Hatching period (48-72) 141 hpf<sup>16</sup>) is a critical process in embryonic development and reduction in the hatching rate could be 142 associated with reduced muscle function<sup>17</sup>. The time course showed that although different 143 hatching times were observed between groups, no significant difference was found between 144 hatching rates of mutant and WT embryos at 72 hpf with over 94% of embryos hatched 145

146 ( $desma^{kg97/kg97}$  vs. WT, P=0.9902;  $desmb^{kg156/kg156}$  vs WT, P=0.3186, repeated measures two-way 147 ANOVA, Bonferroni post *hoc* test) (Fig. 3b).

Similar mortality rates were observed with no significant difference between homozygous mutants of  $desma^{kg97}$  or  $desmb^{kg156}$  and WT embryos ( $desma^{kg97/kg97}$  vs. WT, P=0.1287;  $desmb^{kg156/kg156}$  vs. WT, P=0.6239, repeated measures two-way ANOVA) (Fig. 3d). Finally, among surviving larvae, no statistically significant difference in body length was observed between WT and mutant groups (P=0.2036 for WT vs.  $desma^{kg97/kg97}$ ; P=0.1893 for WT vs.  $desmb^{kg156/kg156}$ , Mann-Whitney U) (Fig. 3c).

#### 154 Mutant larvae show no neuromuscular defect

Muscle fibre integrity and somite morphology length were investigated by staining muscle actin with rhodamine phalloidin in 96 hpf embryos (Fig. 3e). At this stage, mainly *desma* is expressed in somites. No muscle lesion or detachment of fibres was observed in mutants and no significant difference was found in somitic length between mutants and WT (*P*=0.7912 for WT vs. *desma*<sup>kg97/kg97</sup>; *P*=0.4923 for WT vs. *desmb*<sup>kg156/kg156</sup>, Mann-Whitney U) (Fig. 3f).

Touch-evoked response assay is a widely used method to assess neuromuscular function 160 in 48 hpf zebrafish embryos<sup>18</sup>. We performed motility experiments in a blinded manner on siblings 161 from three groups of heterozygous mutant in-crosses and genotyped them after. First, the average 162 escape time of siblings from *desma<sup>kg97/+</sup>* in-crosses was assessed and no significant difference was 163 found in desmakeg97 heterozygous (P=0.122, Mann-Whitney U) or homozygous (P=0.5529, Mann-164 Whitney U) mutants compared to WT (Fig. 3g). Similarly, no significant difference in the average 165 escape time was found in *desmb*<sup>kg156</sup> heterozygous (P=0.3968, Mann-Whitney U) or homozygous 166 (P=0.6131, Mann-Whitney U) mutants compared to WT (Fig. 3h). Since both desma and desmb 167 are expressed in somites up to this stage (48 hpf) and might potentially compensate for the absence 168 of each other, we finally evaluated the effect of the absence of both proteins in siblings from 169 desma<sup>kg97/+</sup>; desmb<sup>kg156/kg156</sup> double mutants. No significant difference in the average escape time 170 was found in homozygous double mutants (desmakeg97/kg97; desmbkg156/kg156) compared to 171  $desma^{+/+}$ :  $desmb^{kg156/kg156}$  (P=0.3165, Mann-Whitney U) and in  $desma^{kg97/+}$ :  $desmb^{kg156/kg156}$ 172 embryos compared to  $desma^{+/+}$ ;  $desmb^{kg156/kg156}$  (P = 0.2334, Mann-Whitney U) (Fig. 3i). 173

#### 175 Mutant adult skeletal muscle shows no muscle degeneration

Because no skeletal muscle phenotype was observed in mutant larvae, surviving one-year-old 176 adults were investigated for muscle defects. Some homozygous single and double mutants 177 178 survived until adulthood with no sign of behavioral defect. Western blot was performed using myotomal skeletal muscle protein extracts isolated from adult fish (WT, homozgous and 179 heterozygous desmakg97 mutants, homozygous and heterozygous desmbkg156 mutants and 180 desma<sup>kg97/kg97</sup>: desmb<sup>kg156/kg156</sup> double mutants). As desma was shown to be predominantly 181 expressed in somites from 72 to 120 hpf, the two bands detected in WT and *desmb*<sup>kg156</sup> mutants 182 were predicted to correspond to Desma-1 (predicted molecular weight 55.7 kDa) and Desma-2 183 184 (predicted molecular weight 54.1 kDa) isoforms. In agreement with that, both bands were absent in *desma<sup>kg97</sup>* homozygous mutant and double mutants while still expressed in homozygous 185 desmb<sup>kg156</sup> mutant (Fig. 4a). Nevertheless, histopathological examination of hematoxylin-eosin 186 stained skeletal muscle tissue sections revealed no pathological changes in homozygous single or 187 double mutants (Fig. 4b). General morphology of skeletal fibre shape and size, integrity of 188 sarcoplasm was normal. There was an absence of cytoplasmic aggregates, signs of degeneration 189 or regeneration, internal nuclei, inflammatory cell infiltration, increase in connective or adipose 190 191 tissue in interstitium (Fig. 4b). We conclude that fish entirely lacking wild-type desmin generate and maintain functional skeletal muscle in the context of a zebrafish aquarium. 192

### 193 Altered calcium flux in *desma* mutant fibres

Although anatomically apparently wild-type, adult *desma*<sup>kg97/kg97</sup> mutants were physiologically 194 defective in calcium handling. To investigate amplitude and time course of calcium signals 195 196 released in response to a depolarizing voltage stimulus, individual fibres were dissected from WT (N=8), desma<sup>kg97</sup> (N=11) or desmb<sup>kg156</sup> (N=8) homozygous mutant 1-year-old fish. Calcium flux 197 along fibres was monitored in the isolated fibres by Fluo-4 AM after four consecutive depolarizing 198 stimuli with the amplitude of the current pulse kept constant at 100 nA and the duration increased 199 200 by 10 ms at each pulse (Fig. 5a-b). Amplitudes of the calcium emission signals in the first (10 ms) stimulus and fourth (40 ms) stimulus were compared between mutants and WT. To eliminate 201 variance induced by amplitude, offset and fibre size differences, amplitude values were baseline 202 corrected and divided by the diameter of each fibre. Amplitudes of the first or the fourth calcium 203 transient were significantly lower in desmakeg97/kg97 fibres compared to WT fibres (P=0.0008 for 204

first transient; P=0.0025 for fourth transient, Mann-Whitney U) (Fig. 5c-d). In contrast, no significant difference was found in the amplitude of calcium flux between  $desmb^{kg156/kg156}$  fibres and WT fibres (P=0.0881 for first transient; P=0.6657 for fourth transient, Mann-Whitney U) (Fig. 5c-d).

Time course of the longest calcium transient (40 ms, fourth transient) has been analyzed 209 for comparing the waveform of the responses and dissecting each phase of the transient. Signals 210 have been baseline corrected and then normalized. Mean amplitude values as a function of time 211 were compared between desmakg97/kg97 and WT or desmbkg156/kg156 and WT. No significant 212 difference was found in the rising and the plateau phases of the recorded transients between the 213 experimental groups (Figure 5e). However, the decay phase was significantly faster in 214 desmakg97/kg97 mutant fibres compared to WT (\* indicates P<0.05, repeated measures ANOVA, 215 Bonferroni post hoc test). No significant difference was found in desmbkg156/kg156 mutant fibres 216 compared to WT (Fig. 5e). 217

# 218 **DISCUSSION**

We demonstrated differential expression of desma and desmb paralogs during zebrafish 219 development. Although previous studies showed that *desma* is the main gene expressed in somites, 220 no study did clearly distinguish the spatial and temporal differences in the expression of the two 221 genes. We showed that until 72 hpf desma and desmb expression overlap but later in development, 222 their expression diverges. While desmb expression shifts from somites to gut, desma remains as 223 the predominantly expressed paralog in somites. Note that the residual desmb expression detected 224 in the lateral edges of the myotome might reflect a transient expression in new muscle fibres<sup>19</sup>. 225 226 Reduction of Desmin protein in adult *desma* mutants confirms that Desma is the major desmin protein in adult zebrafish skeletal muscle. Desmb appears the major desmin in smooth muscle, at 227 least in the gut. So zebrafish *desmb* mutants may be particularly useful for understanding the role 228 of desmin in smooth muscle without interference by additional phenotypes from skeletal muscle. 229 230 In future studies, it will be necessary to take into account this divergent expression when studying the role of desmin in zebrafish. 231

232 Reduced *desma* or *desmb* mRNAs in their respective mutants suggest the mRNAs are 233 degraded by nonsense-mediated decay, such that only tiny amounts of the severely truncated 234 proteins would be produced. Such fragments lack the highly conserved  $\alpha$ -helical central rod

domain responsible for filament formation and all known protein-protein interaction domains of 235 Desmin<sup>20</sup>. Intriguingly, no overt phenotype was observed in some double homozygous mutants. 236 237 In contrast to desmin morphants where both *desma* and *desmb* were targeted with morpholinos <sup>13</sup>, we found that *desma<sup>kg97</sup>* or *desmb<sup>kg156</sup>* alleles have no effect on hatching of eggs, mortality rate and 238 length of larvae. While motility is altered in 48 hpf desmin morphants, no significant alteration 239 was detected in *desma<sup>sa5</sup>* embryos<sup>13,21</sup>. By reasoning that this could be due to a compensation by 240 desmb, which is still expressed in somites at 48 hpf, we performed motility assays in both single 241 mutant and double mutant embryos but found no additional neuromuscular defect in larvae mutant 242 in single or both desmin genes. At the histological level, no sign of muscle degeneration or 243 disorganisation of fibres was observed in *desma<sup>kg97</sup>* or *desm*b<sup>kg156</sup> mutant skeletal muscles in either 244 embryos and adults. In contrast, the desmin morphants and *desma*<sup>sa5</sup> mutant muscles of embryos 245 were disorganized, with disruption of sarcomeres<sup>13,14</sup>. Our findings raise the possibility that a 246 haploinsufficient dominant effect in *desma<sup>sa5</sup>* mutants may disrupt muscle; such an explanation 247 would no account for the morphant phenotype. We conclude that at least two further alleles of 248 desma may be needed to resolve these differences: firstly, a deletion allele that would 249 250 simultaneously prevent any Desma polypeptide synthesis and any RNA-triggered compensation and, secondly, an allele lacking morpholino binding ability but nevertheless encoding wild-type 251 252 protein to prove that the morphant phenotype is due to *desma* binding.

We found that the amplitude of calcium signals released in response to a depolarizing 253 current stimulus applied to fibres was significantly lower in desmakeg97 compared to WT. In 254 addition, while rising and plateau phases of calcium signals were similar between all groups, the 255 decay phase was shorter in *desmakg97* mutants compared to WT. These results indicate that 256 mechanisms regulating cytosolic calcium are altered and fibres are not efficient in generating a 257 258 long lasting calcium increase in the absence of desmin. These results are concordant with calcium flux defects observed in desmin mutant mice and hearts of desma<sup>sa5</sup> embryos<sup>14</sup>. Our data support 259 the idea that desmin has a similar role in calcium conduction in skeletal muscle fibres and suggest 260 that some aspects of the *desma<sup>sa5</sup>* phenotype may reflect loss of function of Desma. 261

Although not the focus of this study, it should be noted that heart and gastroinstestinal development in mutant lines have not been extensively analyzed. No heart oedema such as observed in desmin morphant or overt morphological abnormality in the gut have been observed in  $desma^{kg97}$ ,  $desmb^{kg156}$  or double mutant embryos and adults.

Many loss-of-function mutants not exhibiting an overt phenotype and failing to phenocopy 266 morphants have been previously reported<sup>22-24</sup>. El-Brolosy et al. proposed transcriptional 267 268 adaptation triggered by mutant RNA degradation as a mechanism to explain this discrepancy<sup>15,25</sup>. According to this model, decay of the mutant mRNA results in upregulation of related genes, 269 alleviating or suppressing the phenotype observed in morphants. Such adaptation could explain 270 the milder phenotype observed in mutants created in our study compared to desmin morphants. 271 However, it has been shown that *desma<sup>sa5</sup>* allele results in a muscle phenotype similar to that of 272 morphants, even though the desmasa5 allele predicts an in-frame stop codon and no desma mRNA 273 was detected in mutants<sup>14,21</sup>. We also observe a severe reduction in *desma* mRNA in our *desma<sup>kg97</sup>* 274 mutant. The specific mechanism leading to the absence of mRNA was not investigated either in 275 desma<sup>sa5</sup> mutants or in the present study. It is possible that the desma<sup>sa5</sup> allele is not as efficient as 276 desmakg97 allele in triggering an adaptation mechanism for yet unknown reasons. However, our 277 data suggest that the compensating gene in fish carrying our *desma* allele is not *desmb*, the only 278 gene known to have high and extensive homology to *desma*. It will be interesting to compare these 279 models in the contexts of mRNA decay and transcriptional adaptation. Alternatively, short 280 281 truncated polypeptides not detected in Western blot might play a role in this process before being rapidly degraded. In conclusion, unraveling modifier genes and proteins that compensate for the 282 283 loss of desmin in our models will bring further insights in the role of desmin in muscle.

# 284 METHODS

#### 285 Ethical approval and zebrafish maintenance

All animal procedures were approved by the Hacettepe University Animal Experimentations Local 286 287 Ethics Board (2014/07-08). Adult zebrafish (AB) (Danio rerio) were kept in 14/10 hr light-dark cycle at 28.5°C. Adults are fed twice a day, with dry feed in morning and Artemia spp. in evening. 288 For spawning, male and female fish were placed in a breeding tank with a separator. Next morning 289 the separator was removed, viable eggs were collected and rinsed in E3 medium (5 mM NaCl, 0.17 290 291 mM KCl, 0.33 mM CaCl<sub>2</sub>, 0.33 mM MgSO<sub>4</sub>, 10<sup>-5</sup>% Methylene Blue). Embryos were kept in E3 medium for five days and placed into adult tanks containing system water. Embryos were fed five 292 293 times a day with dry feed and Artemia.

#### 294 Whole mount *in situ* hybridization

Embryos were fixed in 4% PFA in PBS (w/v) overnight at 4°C, dehydrated in 50% and %100 295 methanol washes and stored at -20°C at least overnight before use. Probe templates were amplified 296 297 from cDNA isolated from 96 hpf wild-type embryos. Reverse primers for antisense probe contained T3 promoter sequence while forward primers for sense probes contained T7 promoter 298 sequence. Primers sequences can be found as Supplementary Table S1 online. Digoxigenin-299 labelled probes were generated using T3 or T7 RNA polymerases. Whole mount in situ mRNA 300 hybridization was performed as described<sup>26</sup>. Embryos were photographed as on Zeiss Axiophot 301 302 with Axiocam (Carl Zeiss, Oberkochen, Germany) using Openlab software (Agilent, Santa Clara, CA, USA). 303

### 304 Generation of mutant zebrafish lines

CRISPR-Cas9 genome editing was performed as described in Fin et al<sup>24</sup>. CRISPR-Cas9 sgRNAs 305 showing minimal off-target sites were designed by using the online software ZiFiT Targeter. 306 Oligos encoding sgRNAs targeting the first exon of desma (GRCz11, Chr9: 7539113-7539132, 307 GGTCACCTCGTAAACTCTGG) and desmb (GRCz11, Chr6: 308 13891417-13891437, GGCTATACTCGCTCTTATGG) were cloned into pDR274 plasmid (Addgene). sgRNAs were 309 synthesized using T7 RiboMAX Large Scale RNA production system (Promega) following the 310 manufacturer's protocol. Transcribed sgRNAs were purified using sodium acetate and ethanol 311 312 precipitation and quantified on Qubit fluorometer. Cas9 mRNA was transcribed from pCS2-Cas9 plasmid using mMessage mMachine SP6 Kit (Ambion) and purified by lithium chloride 313 precipitation. One-cell stage embryos were injected with 2 nl containing 80-200 pg of sgRNA and 314 100 pg of Cas9 mRNA. At 48 hpf, ten embryos were analysed for mosaicism at targeted loci using 315 316 high resolution melt analysis. Injected embryos were raised to adulthood and back-crossed to identify transmitted mutations in F1 progeny. Homozygotes for each gene were generated by in-317 318 cross of F2 heterozygotes. Primers sequences used for genotyping by Sanger sequencing can be found as Supplementary Table S1 online. 319

## 320 Quantitative real-time PCR

Total RNA was isolated from a pool of 20 mechanically homogenized embryos at 96 hpf for each sample (n=8) using TRItidy G (AppliChem) and 500 ng of cDNA was synthesized using QuantiTect Reverse Transcription Kit (Qiagen) according to manufacturer's protocol. Primer

sequences for *desma* (covering both *desma-1* and *desma-2*), *desmb* and *actb1* can be found as Supplementary Table S1 online. qPCR was performed in triplicates using SensiFAST SYBR No-ROX Kit (Bioline) in Rotor-Gene 6000 (Corbett Life Science). *Desma* and *desmb* expression levels were normalized to *actb1* expression, mutant expression levels were calculated relative to wild-type using  $\Delta\Delta$ Ct method<sup>27</sup>.

#### 329 **Protein Extraction and Immunoblotting**

For protein extraction, 1-year-old fish were euthanized using overdose of tricaine (MS222, 300 330 mg/l). Skeletal muscle tissue was immediately dissected and fresh-frozen in liquid nitrogen. Tissue 331 was pulverized and homogenized in blending buffer (16 M Tris HCl, 200 M EDTA, 20% SDS, 332 1X protease inhibitor cocktail, volume adjusted to 5 ml with dH<sub>2</sub>O) by sonication on ice. Lysates 333 were then centrifuged and supernatant transferred in a fresh tube. Total protein concentrations were 334 determined by bicinchoninic acid Protein Assay (Pierce, 23225). Equal amounts of protein (40 µg) 335 were loaded on a 13% polyacrylamide gel and transferred onto a nitrocellulose membrane using 336 semi-dry transfer, blocked for one hour at room temperature (%5 nonfat dried milk in %0,2 TBS-337 T) and probed with anti-desmin antibody (1:1000; Sigma, D8281) overnight at 4°C or anti-lamin 338 B1 antibody (Abcam, ab90169) followed by appropriate HRP-conjugated secondary antibody. 339 Chemiluminescence detection was done by using SuperSignal<sup>TM</sup> West Femto Maximum Sensivity 340 341 substrate (Thermo Scientific, 34095) in GeneGnome device.

#### 342 Whole-mount Immunofluorescence

96 hpf embryos were fixed in %4 PFA overnight at +4°C and washed with PBTx (%0.01 Triton
X-100 in PBS). Embryos were permeabilized in 50 µg/ml Proteinase K for 2 hours and re-fixed in
2% PFA for 20 minutes at room temperature. Embryos were blocked in %5 BSA for 2 h at room
temperature and incubated with anti-desmin antibody (1:20; Sigma, D8281) for two overnights.
After two overnight incubations with Alexa-Fluor 488 conjugated secondary antibody, embryos
were mounted with DAPI onto glass bottom dishes for imaging with Zeiss LSM Pascal laser
scanning confocal microscope.

# 350 Larval Phenotype and Morphology

After successful mating, embryos were collected, counted and recorded for the following rates.
Hatching rate was reported as cumulative percentage of hatched embryos at 24 hpf, 48 hpf and 72

hpf. Mortality rate was expressed as the percentage of death embryos for 5 dpf. For body length

measurements, at least 16 embryos at 96 hpf were mounted, photographed (The Imaging Source,

355 DFK 41AU02) and measured from the mouth tip to the tail base using ImageJ (Version 1.49u).

- 356 Values for body length were presented as mean body length in cm. All the experiments except
- body length measurements (three times) were performed at least eight times.

#### 358 Phalloidin Staining

Phalloidin staining was performed on 96 hpf embryos (N=8) as described<sup>28</sup>. Following fixation in 4% PFA and permeabilization with 50  $\mu$ g/ml Proteinase K, embryos were incubated in Alexa-Fluor 488 conjugated Phalloidin (Invitrogen, A12379) overnight at +4°C. Embryos were then mounted onto glass bottom dishes and photographed with Zeiss LSM Pascal laser scanning confocal microscope. Somite lengths were measured by using ImageJ (Version 1.49u).

#### 364 Hematoxylin Eosin Staining

Adult zebrafish were freshly frozen in methyl-butanol cooled down in liquid nitrogen and then taken into O.C.T compound for sectioning. Tissue sections were incubated in hematoxylin for 5 minutes, 1% acid-alcohol solution for 1 minute, 1% ammoniac for 1 minute, eosin solution for 3 minutes respectively by washing with distilled water in between steps. Finally, sections were washed with alcohol and xylene, and mounted. Sections were scanned with Nikon Eclipse E-400 microscope and photographed by DXM 1200F digital camera.

### 371 Motility Assay

For motility assay 48 hpf embryos from in-crosses of desmakeg97/+, desmbkg156/+ 372 or desmakg97/+;desmbkg156/kg156 adults were used. Motility assays were performed by previously 373 described protocol<sup>18</sup>. A petri dish was placed on a transparent sheet with concentric circles 5 mm 374 apart. One embryo at a time was positioned in the centre of the dish and stimulated using a 375 376 dissection needle. The escape response of the embryo was recorded using a high-speed camera 377 (Huawei, Mate 10 Lite, 16 MP, 120 fps). The movements of the embryo-midpoint between the two eyes-were tracked using a custom software that works based on template matching 378 379 algorithm. The elapsed time between the last frame before touch stimulus and the first frame after the body of the fish contacts the 10 mm circle was computed from the digitized embryo movements 380

with a resolution of 8.3 ms (120 fps). Experiments were performed blindly and after video
accusation, DNA was extracted from each embryo for genotyping.

#### 383 Calcium Flux

384 1-year-old fish were euthanized using overdose of tricaine (MS222, 300 mg/l). After removal of the skin, tissue samples were incubated into 5 mg/ml collagenase solution in PBS (without Ca<sup>2+</sup> 385 and Mg<sup>2+</sup>) at 28.5 °C for 20 minutes on a shaker at 50 rpm. Samples have been triturated gently 386 under dissecting microscope in order to dissociate muscle fibres from the tissue components until 387 388 rode-shaped muscle fibres were obtained. Dissociated fibres were incubated in DMEM containing 10% FBS and 1% antibiotic-antimycotic overnight at 28.5 °C and 5% CO<sub>2</sub> environment. On the 389 390 following day, fibres were incubated in 2.5 µM Fluo-4 AM for 20 minutes in 5% CO<sub>2</sub>. Under a laser scanning confocal microscope (Zeiss LSM Pascal), glass microelectrodes (filled with 3M 391 KCl) were inserted into muscle fibres in a current clamp mode<sup>29</sup>. A depolarizing current stimulus 392 was delivered through the microelectrode to excite the fibre and evoke a calcium signal. The 393 evoked calcium-specific fluorescent emission signal was recorded in "Line Scan" mode of 394 confocal microscope, supplying information about a line cross sectioning the muscle fibre at a high 395 frequency in tandem with the electrical recordings. Data were analyzed in MATLAB (The 396 397 MathWorks, Inc). The stimulus waveform, which consisted of four consecutive pulses increasing in duration, was used in all the experiments. The amplitude of the current pulse was kept constant 398 at 100 nA while the durations were 10 ms, 20 ms, 30 ms, 40 ms respectively. Experiments were 399 done by using three WT, three  $desma^{kg97/kg97}$  and two  $desmb^{kg156/kg156}$  fish and at least eight fibres 400 401 from each group were recorded.

# 402 **Statistics**

403 Data were statistically analyzed by GraphPad Prism 8 (GraphPad Software Inc.) by nonparametric 404 Mann-Whitney U test (Two-tailed) when two groups were compared. The results were considered 405 significant when P<0.05. For analysis of time course experiments (hatching rate, mortality rate 406 and time course calcium flux), repeated measures two-way ANOVA was used and Bonferroni post 407 *hoc* test was used for multiple comparisons. All error bars were presented as mean±SEM and bars 408 were represented as median.

409

# 410 Acknowledgements

- 411 This work was supported by The Scientific and Technological Research Council of Turkey, Project
- 412 no. 214S174 to P.R.D.

# 413 **Competing interests**

414 The author(s) declare no competing interests.

# 415 Author Contributions

- 416 P.R.D. designed, P.R.D. and S.M.H. supervised the study. G.K.K. wrote the paper, E.K.M. made
- 417 contribution to writing the paper and assembled the figures. G.K.K., E.K.M, C.K., S.U., B.S., B.E.,
- 418 I.U., N.D., B.T. and N.P. performed the experiments and/or analyzed the results. All authors
- 419 interpreted the data and edited the manuscript.

# 420 Data Availability

- 421 Supporting information is available in Supplementary files and further information is available
- 422 from the corresponding author upon request.

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# 500 Figure Legends

Fig. 1. Whole mount in situ mRNA hybridisation of embryos at the indicated stages for 501 antisense probes to desma and desmb. (a) Lateral views are anterior to top, dorsal to left for 13-502 22 hpf; anterior to left except 11 hpf which is a dorsal view. 48-120 hpf whole mounts are anterior 503 to left, dorsal to top. Scale bar: 250 µm. (b) Arrows indicate heart in frontal view for left panel 504 (desma) and ventral view for right panel (desmb) of 35 hpf embryos. (c) Transversal sections where 505 506 dorsal is top. (d) Upper left panel is a lateral view at 96 hpf, scale bar: 250 µm. Dashed line represents the position of the transversal section in upper right panel where dorsal is top, scale bar: 507 508 100 µm. Lower panels are lateral views of zebrafish heads at 72 and 120 hpf treated with desma probe. Scale bar: 100 µm. (e) Left panel is a ventral view with anterior at top. Right panels are 509 510 lateral views. Ai, anterior intestine; eom, extraocular muscles; fast, fast muscles; fm, pectoral fin muscles; hm, hypaxial muscles; mdm, mandibular muscles; om, opercular muscles; slow, slow 511 muscles; som, somites. Scale bar: 250 µm. 512

Fig. 2. Generation of desma and desmb knockout lines. (a) Alignments and chromatograms of 513 wild-type DNA sequences with mutant alleles, and predicted mutant polypeptide sequences. In 514 DNA sequences, yellow highlights gRNA target sequence, hyphens show deleted bases, inserted 515 516 bases are indicated in red font. In protein sequences, the first residue affected by the frameshift is indicated in red font, asterisks represent early stop codons. (b) Quantitative real-time PCR results 517 showing the expression of desma mRNA (upper graphic) and desmb mRNA (lower graphic) in 518 wild-type and homozygous mutant 96 hpf embryos (N=5). A significant decrease of desma 519 transcripts in *desma<sup>kg97</sup>* homozygous embryos (P=0.0119, Mann-Whitney U) was observed while 520 desmb transcripts were significantly decreased in desmb<sup>kg156</sup> homozygous embryos at 96 hpf 521 compared to WT (P=0.0079, Mann-Whitney U). (c) Left panel shows whole mount in situ mRNA 522 hybridisation of *desma<sup>kg97</sup>* homozygous embryos at the indicated stages for antisense probes to 523 desmb. Right panel shows whole mount in situ mRNA hybridisation of desmb<sup>kg156</sup> homozvgous 524 embryos at the indicated stages for antisense probes to desma. Scale bar: 250 µm. Ai, anterior 525 intestine; som, somites. (d-f) Whole mount immunofluorescence staining of desmin with an anti-526 527 desmin polyclonal antibody (Sigma, D8281) recognizing both Desma and Desmb, in wild-type and mutant 96 hpf embryos. Scale bar: 500 µm. Boxes with dashed lines delineate areas that were 528

200 zoomed and focused on intestine. Pictures showing DAPI staining of each embryo are included at their bottom right. (d) Upper panel is lateral view of a WT embryo with anterior to left, dorsal to top. Lower panel is ventral view with anterior to left, arrowheads indicate the gut. (e) Upper panels are lateral views of  $desmb^{kg156/kg156}$  embryo with anterior to left, dorsal to top. Lower left panel shows anterior intestine (orange arrowheads), lower right panel shows middle and posterior intestine (white arrowheads). (f) Lateral view of  $desma^{kg97/kg97}$  with anterior to left, dorsal to top.

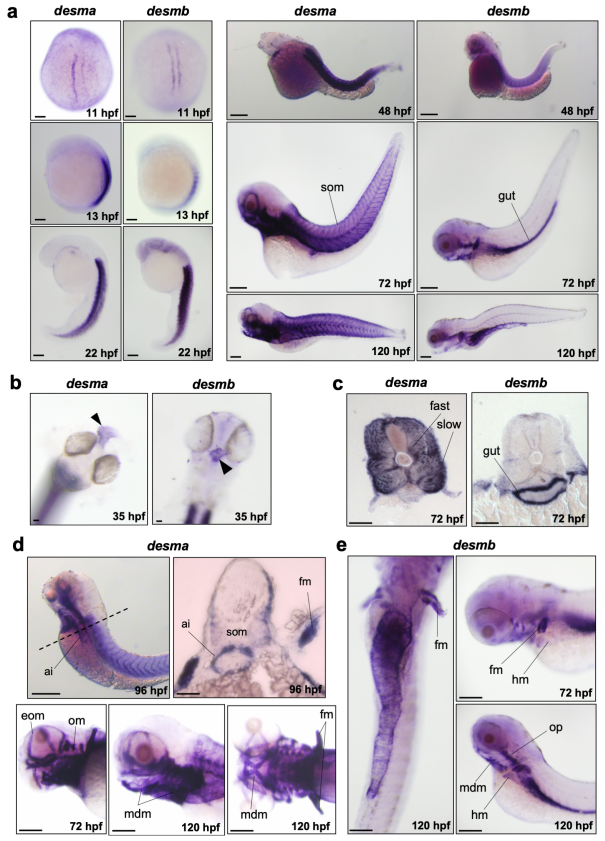
Fig. 3. Evaluation of neuromuscular defect in mutant larvae. (a) The number of viable embryos 535 after successful mating of WT fish (n=10) compared to homozygous  $desma^{kg97}$  (n=10) (P=0.6842, 536 Mann-Whitney U) or homozygous desmb<sup>kg156</sup> (n=8) (P=0.9654, Mann-Whitney U) fish. (b) 537 538 Comparison of hatching rate between homozygous mutants and WT 24, 48 and 72 hpf embryos At 72 hpf, no significant difference in the hatching rate was found between mutants and WT 539 ( $desma^{kg97/kg97}$  vs. WT, P=0.9902;  $desmb^{kg156/kg156}$  vs WT, P=0.3186, repeated measures two-way 540 ANOVA, Bonferroni post hoc test) (c) Body length of 96 hpf WT (N=18) and homozygous 541 desma<sup>kg97</sup> (N=25) or homozygous desmb<sup>kg156</sup> (N=16) mutant embryos (P=0.2036 for WT vs. 542 desma<sup>kg97/kg97</sup>; P=0.1893 for WT vs. desmb<sup>kg156/kg156</sup>, Mann-Whitney U). (d) Cumulative mortality 543 rate from 1 to 5 dpf homozygous desmakeg97 mutants (N=1247) compared to WT (N=2460) 544 (P=0.1287, repeated measures two-way ANOVA, Bonferroni post hoc test) and homozygous 545 desmbkg156 mutants (N=2615) compared to WT (P=0.6239, repeated measures two-way ANOVA, 546 Bonferroni post hoc test). e. Optical sections of the mid-trunk region of WT and homozygous 547 mutants 96 hpf embryos stained with phalloidin (N=8). Scale bars: 50  $\mu$ m. (f) Somite lengths ( $\mu$ m, 548 N=8) were measured from optical sections of WT and mutant embryos stained with phalloidin 549 (P=0.7912 for WT vs. desma<sup>kg97/kg97</sup>; P=0.4923 for WT vs. desmb<sup>kg156/kg156</sup>, Mann-Whitney U). (g-550 i) Touch-evoked escape time (ms) of 48 hpf embryos. Motility experiments were performed blind 551 on siblings from (g)  $desma^{kg97/+}$ , (h)  $desmb^{kg156/+}$  or (i)  $desma^{kg97/+}$ ;  $desmb^{kg156/kg156}$  in-crosses 552 followed by *post hoc* genotyping. g. Heterozygous *desma<sup>kg97</sup>* embryos (N=23) were compared to 553 WT (N=9) (P=0.122, Mann-Whitney U) and  $desma^{kg97/kg97}$  (N=15) compared to WT (P=0.5529, 554 Mann-Whitney U). h. Heterozygous desmb<sup>kg156</sup> embryos (N=6) were compared to WT (N=38) 555 (P=0.3968, Mann-Whitney U) and desmbkg156/kg156 (N=6) compared to WT (P=0.6131, Mann-556 Whitney U). i. Homozygous double mutants (N=8) were compared to  $desma^{+/+}$ :  $desmb^{kg156/kg156}$ 557 (n=15) (P=0.3165, Mann-Whitney U) and desmakeg97/+; desmbkg156/kg156 embryos (n=34) compared 558 to  $desma^{+/+}$ :  $desmb^{kg156/kg156}$  (P=0.2334, Mann-Whitney U). 559

560 Figure 4. Desmin protein expression and histological examination of adult skeletal muscle.

(a) Immunoblotting of skeletal muscle protein extracts of WT,  $desma^{kg97/kg97}$ ,  $desma^{kg97/+}$ , *desmb^{kg156/kg156*,  $desmb^{kg156/+}$  and  $desma^{kg97/kg97}$ ;  $desmb^{kg156/kg156}$  double mutant fish, showing expression of putative Desma-1 (55.7 kDa) and Desma-2 (54.1 kDa) isoforms. Lamin B1 was labeled as loading control. (b) Transversal cryosections of mid-trunk somites from WT and homozygous single and double mutant adults stained with hematoxylin eosin showing no pathological changes. Scale bar: 250 µm.

Fig. 5. Calcium flux in isolated fibres. (a) Calcium flux along fibres was monitored in vivo by 567 Fluo-4 AM after four consecutive depolarizing stimuli. Top panel, membrane potential responses 568 to current stimulus pulses with varying durations of 10, 20, 30, 40 ms with the amplitude of the 569 current pulse kept constant at 100 mV and the duration increased by 10 ms at each pulse. Middle 570 panel, evoked calcium transients in line scan mode (tx) to current stimuli. Bottom panel, integrated 571 emission signal as a function of time for each experimental group. (b) Membrane potential 572 response to the first stimulus of 10 ms (top) or to the fourth stimulus of 40 ms. (c) Baseline 573 corrected amplitude divided by fibre diameter ( $\Delta F/\mu m$ ) of the calcium emission signals during the 574 first (10 ms) stimulus and fourth (40 ms) stimulus were compared between mutants and WT. (e) 575 576 Time course analysis of the baseline corrected and normalized amplitude of the calcium transient from the longest stimulus (40 ms) was represented as mean amplitude values as a function of time. 577 Homozygous *desma<sup>kg97</sup>* fibres (N=11, red curve) or homozygous *desmb<sup>kg156</sup>* fibres (N=8, orange 578 curve) were compared to WT (N=8, blue curve). (\* indicates time points where P<0.05, repeated 579 580 measures two-way ANOVA, Bonferroni post hoc test).





# Figure 2.

#### a desma

 WT
 CCTCAGGTTCCTCTGGCTCCTCAAGACTGACCTCCAGAGTTTACGAGGTGACCAAGAG

 kg97
 CCTCAGGTTCCTCTGGCTCCTCAAGACTGACCTCCAAGACTGACCTCCAAGACTGACCA

 -2+4
 Protein caquanca

 Protein sequence

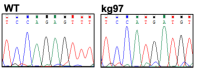
 WT
 MSTKYSASAESASSYRRTFGSGLGSSIFAGHGSSGSSGSSRLTSRVYEVTKSSASPHF

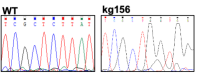
 kg97
 MSTKYSASAESASSYRRTFGSGLGSSIFAGHGSSGSSGSSGSSRLTSMMFTR\*

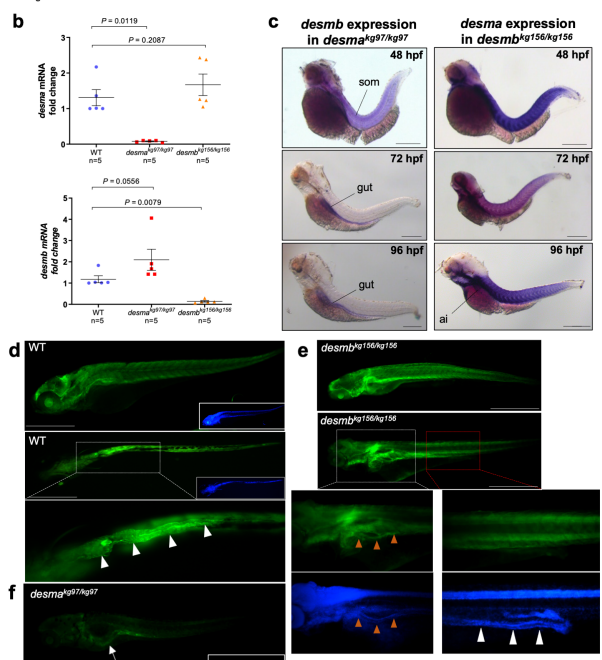
#### desmb

#### **Protein sequence**

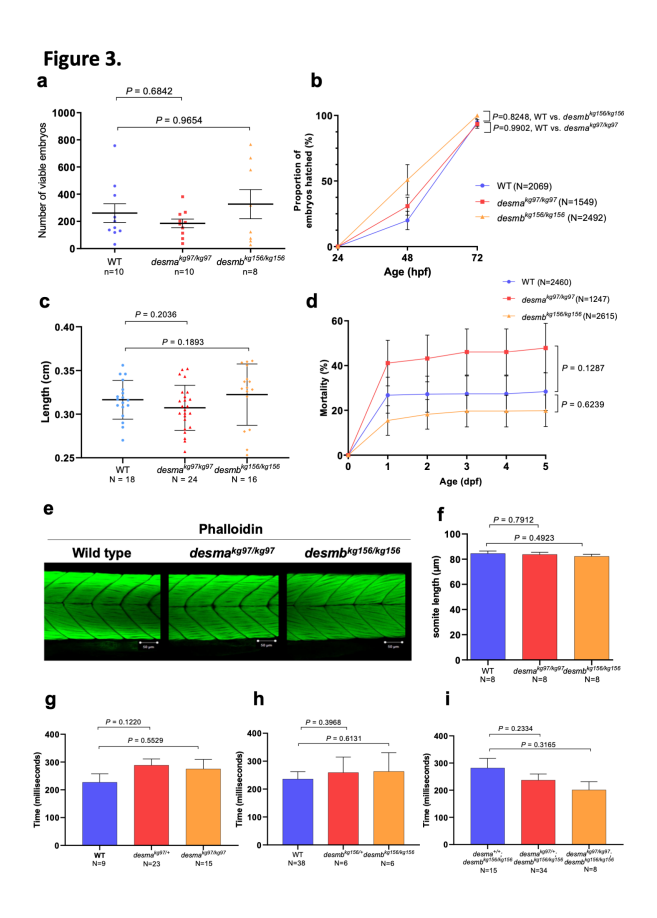
WT MSHSYATSSSSYRMFGGPGYMTP...PGFSGYRASSYSMPNLSAGYTRSYGGMGE kg156 Mshsyatssssyrmfggpgymtp...PGFsgyrassysMpnlsagytrWwdG\*







gut



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