1 The myopathic transcription factor DUX4 induces the production of truncated RNA-

- 2 binding proteins in human muscle cells
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25 ABSTRACT

- 26 DUX4 is an embryonic transcription factor whose misexpression in skeletal muscle causes
- 27 facioscapulohumeral muscular dystrophy (FSHD). DUX4 induces the transcription of thousands
- of RNAs and dysregulates multiple pathways that could contribute to FSHD pathophysiology.
- 29 However, lack of temporal data and the knowledge of which RNAs are actively translated
- 30 following DUX4 expression has hindered our understanding of the cascade of events that lead
- 31 to muscle cell death. Here, we interrogate the DUX4 transcriptome and translatome over time
- 32 and find dysregulation of most key pathways as early as 4 hours after DUX4 induction,
- 33 demonstrating the potent effect of DUX4 in disrupting muscle biology. We also observe
- 34 extensive transcript downregulation as well as induction, and a high concordance between
- 35 mRNA abundance and translation status. Significantly, DUX4 triggers widespread production of
- truncated protein products derived from aberrant RNAs that are degraded in normal muscle
- 37 cells. One such protein, truncated serine/arginine-rich splicing factor 3 (SRSF3-TR), is present
- in FSHD muscle cells and disrupts splicing autoregulation when ectopically expressed in
- 39 myoblasts. Taken together, the temporal dynamics of DUX4 induction show how the pathologic
- 40 presence of an embryonic transcription factor in muscle cells alters gene expression to
- 41 ultimately perturb RNA homeostasis.

42 INTRODUCTION

43 Facioscapulohumeral muscular dystrophy (FSHD) is a prevalent progressive myopathy caused 44 by misexpression of an early embryonic transcription factor, DUX4, in skeletal muscle (Hamel & 45 Tawil, 2018; Tawil, van der Maarel, & Tapscott, 2014). Sustained DUX4 expression is toxic to 46 somatic cells and induces apoptotic death, leading to skeletal muscle atrophy in individuals with 47 FSHD (Bosnakovski et al., 2008; Kowaljow et al., 2007; Rickard, Petek, & Miller, 2015; Wallace 48 et al., 2011). In the decade since the elucidation of a unifying genetic model for FSHD 49 (Lemmers et al., 2010), much work investigating the molecular consequences of DUX4 50 expression has led to the discovery of myriad altered genes and pathways that are associated 51 with FSHD pathophysiology (Campbell, Belleville, Resnick, Shadle, & Tapscott, 2018; Lim, 52 Nguyen, & Yokota, 2020). Efforts to determine which genes or pathways downstream of DUX4 53 misexpression cause FSHD are just beginning. Such understanding is critical for the 54 development of effective therapeutics for FSHD. 55 56 Previous genome-wide studies of DUX4-mediated gene expression have been performed as 57 endpoint assays in cell populations that have seen long periods of DUX4 misexpression and are 58 likely already undergoing apoptosis (Bosnakovski et al., 2019; Geng et al., 2012; Jagannathan, 59 Ogata, Gafken, Tapscott, & Bradley, 2019; Jagannathan et al., 2016; Lek et al., 2020; Resnick 60 et al., 2019: Shadle et al., 2019: Shadle et al., 2017: Sharma, Harafuji, Belavew, & Chen, 2013: 61 Whiddon, Langford, Wong, Zhong, & Tapscott, 2017). Although mechanisms underlying DUX4-62 induced cytotoxicity have been uncovered this way, it has also been shown that important 63 pathologic processes occur in distinct temporal order. For example, inhibition of the RNA quality 64 control pathway nonsense-mediated decay (NMD) by DUX4 was shown to occur early in the 65 course of DUX4 expression while proteotoxic stress was a later event (Feng et al., 2015;

66 Jagannathan et al., 2019). Therefore, genome-wide time course studies are needed to identify

67 early changes in gene expression or cellular pathways that lead to skeletal muscle cell death.68

69 While DUX4 induces widespread transcriptomic changes, proteomics shows that many genes 70 display discordant transcript and protein levels (Jagannathan et al., 2019). Given the sparse

71 nature of proteomics data, sequencing-based measurement of active translation is necessary to

72 determine if the observed discordance between RNA and protein levels in DUX4-expressing

cells is the result of altered translation regulation or protein stability. Such a measure would also

reveal whether the aberrant RNAs stabilized by DUX4-mediated NMD inhibition (Feng et al.,

75 2015) produce truncated proteins.

76 To identify early transcript- and translation-level changes induced by DUX4, we performed 77 paired RNA-sequencing (RNA-seq) and ribosome profiling (Ribo-seq) at 0, 4, 8, and 14 h 78 following the expression of DUX4 in MB135-iDUX4 human skeletal muscle myoblasts. MB135-79 iDUX4 cells are a well-characterized model of FSHD that robustly recapitulates the 80 consequences of endogenous DUX4 expression in FSHD muscle (Jagannathan et al., 2016; 81 Yao et al., 2014). While RNA-seg measures transcript abundance, Ribo-seg measures 82 ribosome-protected RNA fragments, allowing guantification of ribosome density along an mRNA 83 that serves as a proxy for active translation (Ingolia, 2014). Ribo-seg also enables precise 84 delineation of translation start and end sites to characterize the protein products made from 85 aberrant RNAs.

86

87 We found that ~1600 genes show a significant change at the transcript level after 4 h of DUX4 88 induction, including many that are repressed; most pathways known to be misregulated by 89 DUX4 are altered at this early time point. We also found a high concordance of changes in 90 mRNA abundance and translation status, suggesting that post-transcriptional modulation by 91 DUX4 (Jagannathan et al., 2019) occurs primarily at the level of protein stability. Notably, the 92 hundreds of aberrant RNAs stabilized by DUX4-mediated inhibition of NMD (Feng et al., 2015) 93 are actively translated to produce truncated proteins, including truncated RNA-binding proteins 94 (RBPs) and splicing factors. We show that one such truncated splicing factor, truncated 95 serine/arginine-rich splicing factor 3 (SRSF3-TR), is expressed in FSHD muscle cell cultures 96 and perturbs splicing homeostasis when ectopically expressed in myoblasts. Together, our 97 results illustrate the importance of misregulated RNA guality control in DUX4-induced pathology.

98 RESULTS

99

100 Identification of time points for paired RNA-seg and Ribo-seg in DUX4-expressing cells 101 Misexpression of DUX4 in skeletal muscle cells is cytotoxic (Geng et al., 2012; Jagannathan et 102 al., 2016; Rickard et al., 2015). To identify time points at which to measure transcript- and 103 translation-level changes induced by DUX4 before the onset of overt cytotoxicity, we utilized a 104 well-characterized doxycycline-inducible DUX4 human myoblast line, MB135-iDUX4 105 (Jagannathan et al., 2016), harboring a DUX4-responsive fluorescent reporter (Figure 1A). We 106 live imaged these cells every 15 min for 28 h following doxycycline treatment to induce DUX4 107 (Figure 1B, Video 1). In the absence of doxycycline, a very low level of 'leaky' DUX4 108 expression was observed, which is consistent with studies in other doxycycline-driven DUX4 109 induction systems (Bosnakovski, Chan, et al., 2017; Dandapat et al., 2014). In the presence of 110 doxycycline, expression of the DUX4-responsive fluorescent reporter was rapid and nearly 111 synchronous, with fluorescence detectable after 2 h. Most cells fluoresced after 4 h, with 112 fluorescence intensity increasing until 6 h at which point it plateaued. Cytotoxicity was first 113 observed 9 h following DUX4 induction (Figure 1B). Myoblasts continued to round up and 114 detach, resulting in a culture where most cells were dead or dying by 18 h. Given these data, we 115 limited our study to time points \leq 14 h to identify early and direct gene expression changes 116 induced by DUX4.

117

118 To study the temporal trajectory of known DUX4-induced gene expression changes in important 119 cellular pathways, we carried out quantitative reverse transcription PCR (RT-gPCR) using RNA 120 extracted from the parental MB135-iDUX4 myoblast line. At 4 h we observed robust induction of 121 the DUX4 target gene ZSCAN4 (Figure 1C, top left) and repression of the myogenic program 122 (Figure 1C, top right), which is exquisitely sensitive to DUX4 misexpression (Bosnakovski et al., 123 2018; Bosnakovski et al., 2008). In contrast, other consequences of DUX4 expression, such as 124 upregulation of pericentric human satellite II repeats (Shadle et al., 2019), activation of the 125 unfolded protein response (Jagannathan et al., 2019), perturbed Wnt signaling (C. R. Banerij et 126 al., 2015), and downregulation of oxidative stress response genes (Bosnakovski et al., 2008) 127 are more prominent at later time points (Figure 1C, remaining panels). Based on our measures 128 of DUX4 activity, perturbation of downstream pathways, and myoblast cell death, 4, 8, and 14 h 129 were chosen as informative early, mid, and late time points to measure transcript- and 130 translation-level changes induced by misexpression of DUX4, with the 0 h time point serving as

- a control. Therefore, we performed standard RNA-seq paired with Ribo-seq in parental MB135-
- 132 iDUX4 myoblasts induced to express DUX4 for 0, 4, 8, or 14 h (**Figure 1D**).

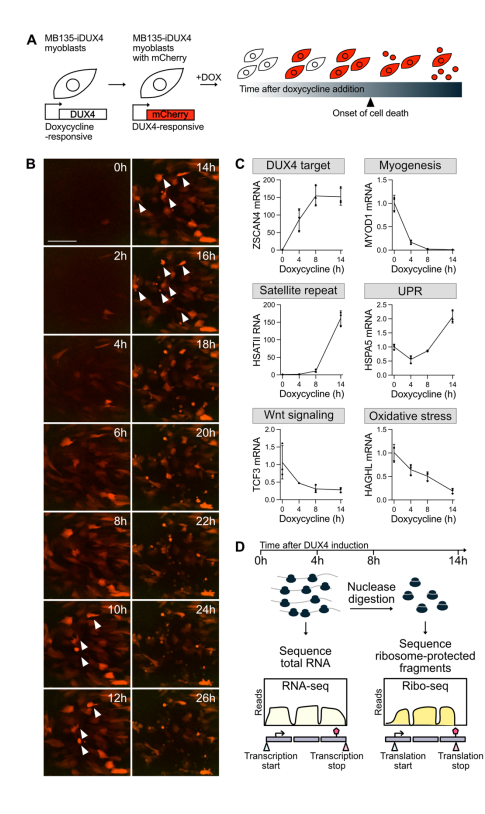


Figure 1. Synchronous expression of DUX4 in MB135-iDUX4 myoblasts enables time course analyses of downstream gene expression changes. (A) Schematic representation of the inducible DUX4 expression systems used in this study and the experimental outline to identify informative time points prior to the onset of cell death. (B) Still images from live cell fluorescence microscopy of MB135iDUX4/ZSCAN4mCherry myoblasts every 2 h following treatment with doxycycline to induce DUX4. Arrowheads indicate overtly dying cells. Scale bar, 150 µm. (C) Relative RNA levels of ZSCAN4, MYOD1, HSATII, HSPA5, TCF3, and HAGHL determined by RT-gPCR following treatment with doxycycline to induce DUX4 in MB135-iDUX4 myoblasts. Error bars denote the standard deviation from the mean of three biological replicates, which are shown as individual data points. (D) A schematic representation of the paired RNA-seq and Ribo-seq experiment. RNA was harvested from MB135-iDUX4 myoblasts treated with doxycycline to induce DUX4 for 0, 4, 8, or 14 h and either directly sequenced (RNA-seq) or digested with nuclease to degrade non-ribosome-protected RNA and the protected fragments sequenced (Ribo-seg).

133 DUX4 triggers early-onset disruption of key pathways involved in FSHD

134 We examined DUX4-induced transcriptome changes revealed by our RNA-seg dataset 135 (Supplementary Table 1). As expected, the housekeeping gene RPL27 had constant, robust 136 RNA expression throughout the time course (Figure 2A, top). Transcripts of a DUX4 target 137 gene, ZSCAN4, were absent in uninduced cells but highly expressed at 4 h and increased with 138 time, while transcripts from a myogenic gene, MYOD1, displayed the opposite trend (Figure 2A, 139 middle and bottom). Genome-wide, DUX4 altered the expression of thousands of transcripts, 140 with known DUX4 targets (Yao et al., 2014) showing increasing upregulation throughout the 141 time course (Figure 2B). Notably, there were 1674 genes whose expression significantly 142 changed after only 4 h of DUX4 induction with similar numbers activated and repressed. Genes 143 underlying pathways known to be involved in FSHD pathology are already significantly altered 144 at the 4 h time point (Figure 2 – figure supplement 1A), suggesting that DUX4 sets up 145 cascades of misregulation very early after its expression in muscle cells.

146

147 To identify novel pathways altered by DUX4, we used k-means clustering to group genes 148 significantly altered (defined as absolute log2 fold change > 1 and adjusted p-value < 0.01) at 149 any point during the time course into five clusters (Figure 2 – figure supplement 1B) and 150 carried out gene ontology (GO) analysis on each cluster (Figure 2C-E, Supplementary Table 151 2). We observed that Cluster 1, which is comprised of 181 genes rapidly induced upon DUX4 152 expression, returns GO categories that could underlie DUX4's normal role in establishing an 153 early embryonic program - negative regulation of cell differentiation, positive regulation of cell 154 proliferation, and DNA-templated transcription. The 692 genes that are rapidly silenced (Cluster 155 5) are enriched for GO categories related to myogenesis, positive regulation of cell 156 differentiation, and cytoskeleton organization. Together, this is suggestive of a general switch 157 away from a differentiated muscle program and towards a proliferative phenotype. Cluster 2 158 (583 genes) that is less robustly induced and Cluster 4 (2462 genes) that is less robustly 159 repressed are enriched for broad GO categories illustrative of the many fundamental cellular 160 processes that are altered upon DUX4 misexpression. Cluster 3 (2243 genes), which is induced 161 only at the late 14 h time point is enriched for GO categories mRNA splicing, ribonucleoprotein 162 transport, and ubiquitin-dependent process, which have previously been reported as some of 163 the major signatures of DUX4-induced gene expression (Geng et al., 2012; Jagannathan et al., 164 2016; Rickard et al., 2015). Together, these time course RNA-seg data show that each pathway 165 misregulated by DUX4 has a unique temporal signature following DUX4 induction and suggest 166 that a combination of perturbed cellular systems underlies DUX4 toxicity.

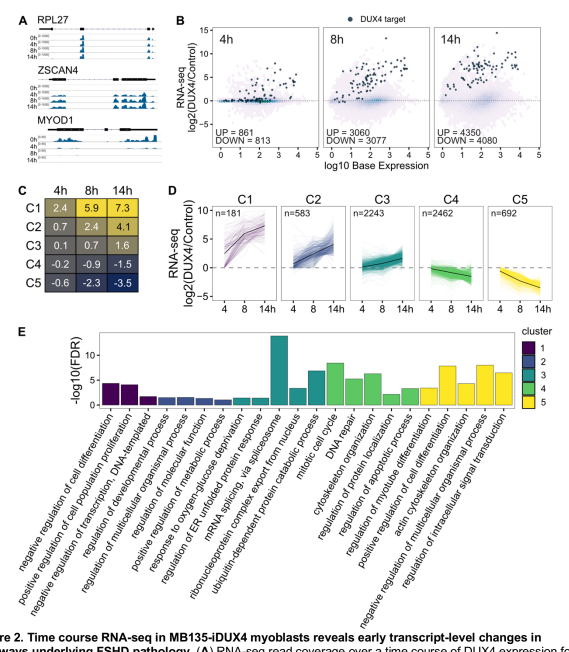


Figure 2. Time course RNA-seq in MB135-iDUX4 myoblasts reveals early transcript-level changes in pathways underlying FSHD pathology. (A) RNA-seq read coverage over a time course of DUX4 expression for housekeeping gene RPL27 (top), DUX4 target gene ZSCAN4 (middle), and myogenic factor MYOD1 (bottom). (B) M-A plots for RNA-seq data after 4, 8, and 14 h of DUX4 induction compared to the 0 h control. DUX4 target status defined as in (Yao et al., 2014). (C) Heatmap depicting the clustering of all genes significantly differentially expressed upon DUX4 induction. The numbers within the heatmap represent the mean log2 fold change for the genes within that cluster. C, cluster. (D) Log2 fold change in RNA expression from the 0 h time point is shown for each gene in each cluster defined in (C). C, cluster. The thick black line represents the cluster mean. (E) GO analysis results of selected gene sets (biological process complete) that are significantly enriched in each cluster defined in (C).

167 Most DUX4-induced coding transcripts are robustly translated

- 168 A previous study that used stable isotope labeling by amino acids in cell culture (SILAC) mass
- 169 spectrometry to measure the DUX4 proteome revealed extensive post-transcriptional gene
- 170 regulation including discordant changes at the RNA versus protein level, particularly for genes

171 involved in RNA quality control (Jagannathan et al., 2019). While powerful, this study could not 172 provide a complete index of all expressed proteins due to inherent limitations in mass 173 spectrometry technology. Ribo-seq, by measuring which mRNAs are actively translated, is 174 capable of identifying the proteins being produced with greater depth than proteomics. 175 Additionally, DUX4 induces eIF2 α phosphorylation at later time points in MB135 myoblasts 176 (Jagannathan et al., 2019; Shadle et al., 2017), which is thought to generally inhibit cap-177 dependent translation (Proud, 2005), but the effect of this on protein expression in DUX4-178 expressing cells is unknown. Therefore, we asked whether transcript level changes driven by 179 DUX4 misexpression were echoed at the level of translation by comparing the RNA-seg and 180 Ribo-seg datasets. To verify the guality of our Ribo-seg data, we confirmed that reads exhibited 181 the characteristic 3 nucleotide periodicity indicative of ribosome-protected RNA fragments 182 (Figure 3 – figure supplement 1). Representative Ribo-seg read coverage plots of the 183 housekeeping gene RPL27 showed constant, robust translation throughout the time course 184 (Figure 3A, top). In contrast, a DUX4 target gene, ZSCAN4, showed no coverage in uninduced 185 cells, low ribosome density beginning at 4 h, and active translation at 8 and 14 h (Figure 3A, 186 middle). MYOD1 was robustly translated at 0 h but rapidly downregulated (Figure 3A, bottom). 187 The changes in translation status at these specific genes mirrored the differences seen in their 188 mRNA levels (Figure 2A).

189

190 On a genome scale, DUX4 altered the translation status for thousands of transcripts, with later 191 time points showing larger differences and known DUX4 targets being translated at increasing 192 levels throughout the time course (Figure 3B, Supplementary Table 3). Strikingly, most genes 193 were concordantly up or downregulated at the level of transcript and translation status (Figure 194 **3C**. Materials and Methods) at all three time points with only a small number of genes showing 195 some discordance. GO analysis of the discordantly regulated genes returned significant results 196 only for the gene set (n = 137) that showed a mild translation downregulation at the 14 h time 197 point with pathways such as protein targeting to ER and viral transcription being enriched 198 (Supplementary Table 3). This likely represents a small subset of genes that are affected by 199 the inhibition of cap-dependent translation due to $eIF2\alpha$ phosphorylation. In contrast, a pairwise 200 correlation analysis of the RNA-seq. Ribo-seq and our previous SILAC proteomics 201 (Jagannathan et al., 2019) data demonstrated that both RNA and translation status are 202 discordant with protein levels (Figure 3D). Overall, these data show a high concordance of 203 transcript level and ribosome occupancy in DUX4-expressing cells, demonstrating that post-

204 transcriptional modulation by DUX4 likely occurs at the level of protein stability rather than

205 protein synthesis.

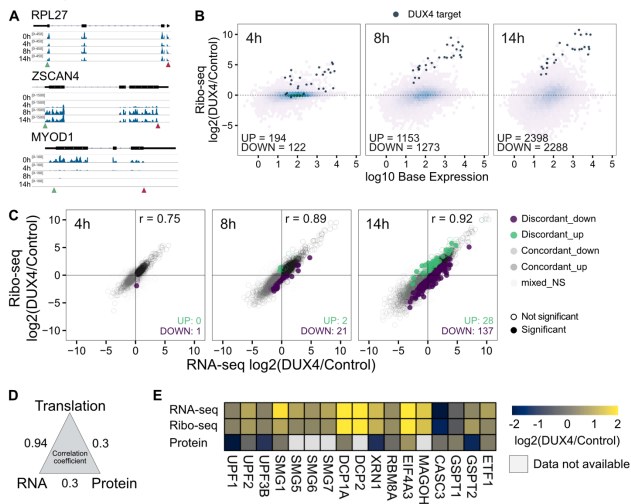


Figure 3. Ribo-seq shows high concordance between transcript levels and translation status. (**A**) Ribo-seq read coverage over a time course of DUX4 expression for housekeeping gene RPL27 (top), DUX4 target gene ZSCAN4 (middle), and myogenic factor MYOD1 (bottom). Green triangle, translation start. Red triangle, translation stop. (**B**) M-A plots for Ribo-seq data after 4, 8, and 14 h of DUX4 induction compared to the 0 h control. (**C**) Scatter plot of RNA-seq versus Ribo-seq log2 fold change after 4, 8, and 14 h of DUX4 expression. Significance defined as adjusted p-value < 0.01 for Ribo-seq fold change. (**D**) Pearson's correlation coefficient for the log2 fold change values computed for genes with quantification in all datasets (RNA-seq, Ribo-seq, and SILAC proteomics; n = 4854 genes). (**E**) Heatmap showing RNA-seq, Ribo-seq, and proteomics log2 fold change values of NMD factors.

- 206 We have previously shown that DUX4 induction leads to detectably lower protein abundance of
- 207 key NMD factors such as UPF1, SMG6, and XRN1, and that UPF1 is proteolytically degraded
- 208 (Feng et al., 2015; Jagannathan et al., 2019). We sought to determine if other NMD factors
- follow a mechanism similar to UPF1 or if their downregulation could be due to lowered
- 210 translation of these proteins. Comparing RNA abundance from our RNA-seq data, translation
- 211 status from our Ribo-seq data, and protein level from our previous proteomics data
- (Jagannathan et al., 2019) for all NMD factors showed that transcript level and translation is

- highly concordant, while protein level is not (**Figure 3E**). This confirms that multiple key NMD
- factors are indeed downregulated at the protein level without a corresponding change in their
- 215 RNA level or translation status, pointing to protein stability as the mode of regulation by DUX4.
- 216

217 DUX4 causes widespread truncated protein production

218 Having established that most transcripts induced by DUX4 are robustly translated, we wanted to 219 determine if truncated proteins are produced from RNAs containing premature translation 220 termination codons (PTCs) stabilized as a result of DUX4-mediated NMD inhibition (Feng et al., 221 2015). Western blot analysis showed that levels of the key NMD factor UPF1 were reduced by 222 65% after only 4 h of DUX4 induction (Figure 4A). UPF1 levels fall further to 13% and 1% of 223 baseline at 8 and 14 h, respectively. This confirmed our previous observation that NMD 224 inhibition is an early event in the course of DUX4 expression (Feng et al., 2015), and suggested 225 that proteins resulting from the translation of NMD-targeted, stabilized aberrant RNAs should

- appear in our dataset.
- 227

228 To ask if and when aberrant RNAs are translated, we used ORFquant (Calviello, Hirsekorn, & 229 Ohler, 2020), a new pipeline that identifies isoform-specific translation events from Ribo-seq 230 data. We then used DEXSeg (Anders, Reves, & Huber, 2012) to conduct exon-level differential 231 analysis on the set of ORFquant-derived open reading frames, using Ribo-seg data. This 232 analysis identifies changes in relative exon usage to measure differences in the expression of 233 individual exons that are not simply the consequence of changes in overall transcript level. After 234 4 h of DUX4 induction 397 genes showed differential expression of specific exons, of which only 235 24 are predicted NMD targets (Figure 4B), whereas later time points showed a greater fraction 236 of exons that are unique to NMD targets as differentially expressed (**Supplementary Table 4**). 237 We grouped exons based on their NMD target status and calculated their fold change in 238 ribosome footprints at 4, 8, and 14 h of DUX4 expression compared to the 0 h time point 239 (Figure 4C). We observed a progressive and significant increase in the translation status of 240 NMD-targeted exons at 8 and 14 h, confirming the translation of stabilized aberrant RNAs in 241 DUX4-expressing cells. 242

Translation of an NMD target typically generates a prematurely truncated protein. To ask how
 the specific truncated proteins being produced in DUX4-expressing myoblasts might functionally
 impact cell homeostasis, we conducted GO analysis of the 74 truncated proteins being actively
 translated at 14 h of DUX4 induction (Figure 4D, Supplementary Table 4). Strikingly, the

- truncated proteins are enriched for genes encoding RBPs involved in mRNA metabolism and
- specifically, splicing (Supplementary Table 4). Examples include genes such as IVNS1ABP,
- 249 SRSF3, SRSF6, and SRSF7 (Figure 4E). Thus, not only are NMD targets stabilized by DUX4
- 250 expression but they also produce truncated versions of many RBPs, which could have
- significant downstream consequences to mRNA processing in DUX4-expressing cells.

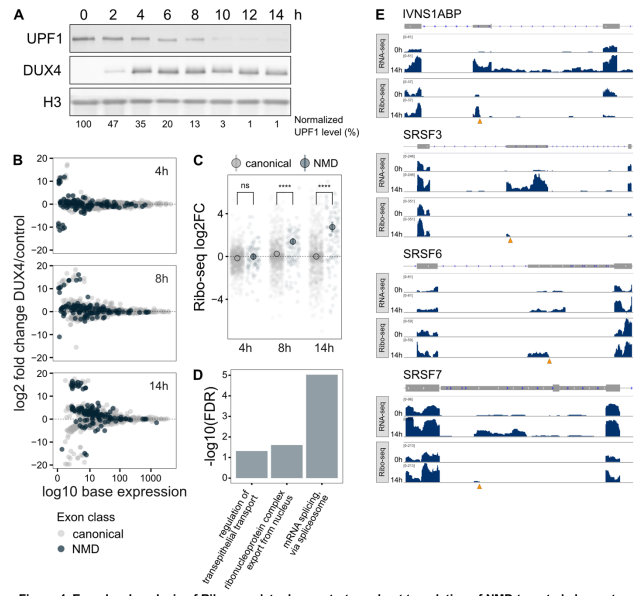


Figure 4. Exon-level analysis of Ribo-seq data demonstrates robust translation of NMD-targeted aberrant RNAs. (A) Western blot analysis for UPF1, DUX4, and Histone H3 (loading control) over a time course of DUX4 expression following doxycycline induction in MB135-iDUX4 myoblasts. (B) M-A plot of exon-level analysis of Ribo-seq data from 4, 8, and 14 h of DUX4 induction. The x-axis represents mean expression calculated at the level of each exon within a gene. (C) Exon-level log2 fold change Ribo-seq values at 4, 8, or 14 h of DUX4 expression for canonical and NMD exons. Statistical testing was performed using a two-sided Wilcoxon test. (D) GO analysis results of selected gene sets (biological process complete) for all NMD targets that are translated at 14 h. (E) RNA-seq and Ribo-seq coverage over splicing-related genes IVNS1ABP, SRSF3, SRSF6, and SRSF7. The orange triangles denote PTCs.

252 Truncated SRSF3 is present in FSHD myotubes and perturbs RNA homeostasis

253 To explore the role truncated proteins play in DUX4-induced cellular phenotypes, we chose 254 SRSF3 for further characterization. SRSF3 is an SR family protein that possesses an N-terminal 255 RNA-binding RNA recognition motif (RRM) and a C-terminal arginine/serine (RS)-rich domain 256 responsible for protein-protein and protein-RNA interactions. SRSF3 is a multifunctional splicing 257 factor involved in transcriptional, co-transcriptional, and post-transcriptional regulation and has 258 been implicated in a variety of human pathologies including heart disease, Alzheimer's, and 259 cancer (More & Kumar, 2020; Zhou et al., 2020). Examination of our paired RNA-seg and Ribo-260 seq data revealed robust expression of SRSF3 NMD-targeted exon 4 after 8 h of DUX4 261 induction, followed by robust translation of this exon that ends at the site of the PTC (Figure 262 5A).

263

264 To confirm translation of the aberrant SRSF3 RNA stabilized by DUX4, we carried out polysome 265 profiling using sucrose density gradient separation. As expected, the polysome profile after 14 h 266 of DUX4 expression showed a higher fraction of 80S monosomes compared to polysomes 267 (Figure 5B, top left), consistent with our observation of eIF2a phosphorylation and general 268 downregulation of translation at this and later time points of DUX4 expression (Jagannathan et 269 al., 2019; Shadle et al., 2017). We extracted RNA from various fractions and found that RPL27 270 mRNA was ribosome-bound in both control and DUX4-expressing cells, as evidenced by its 271 association with monosomes and polysomes and as expected for a housekeeping gene (Figure 272 5B, top right). In contrast, DUX4 mRNA and mRNA of the DUX4 target gene ZSCAN4 are highly 273 translated only in DUX4-expressing myoblasts, as indicated by their enrichment in heavy 274 polysomes (Figure 5B, middle). The normal SRSF3 isoform (SRSF3-Excl) is enriched in heavy 275 polysomes in both control and DUX4-expressing myoblasts but is less abundant in the latter 276 (Figure 5B, bottom left). In contrast, the NMD-targeted isoform of SRSF3 (SRSF3-Incl), also 277 enriched in heavy polysomes in both conditions, was significantly more abundant in DUX4-278 expressing cells (Figure 5B, bottom right). These data confirm that aberrant SRSF3 mRNA is 279 being actively translated into truncated protein in DUX4-expressing myoblasts. 280

To determine if we could detect truncated SRSF3 protein, we generated a custom antibody
 recognizing a 10 amino acid C-terminal neo-peptide unique to SRSF3-TR. This custom SRSF3 TR antibody was able to recognize FLAG-tagged SRSF3-TR exogenously expressed in 293T
 cells and endogenous SRSF3-TR immunoprecipitated from DUX4-expressing MB135-iDUX4
 myoblasts (Figure 5C). We also used a commercial SRSF3 antibody that recognizes an N-

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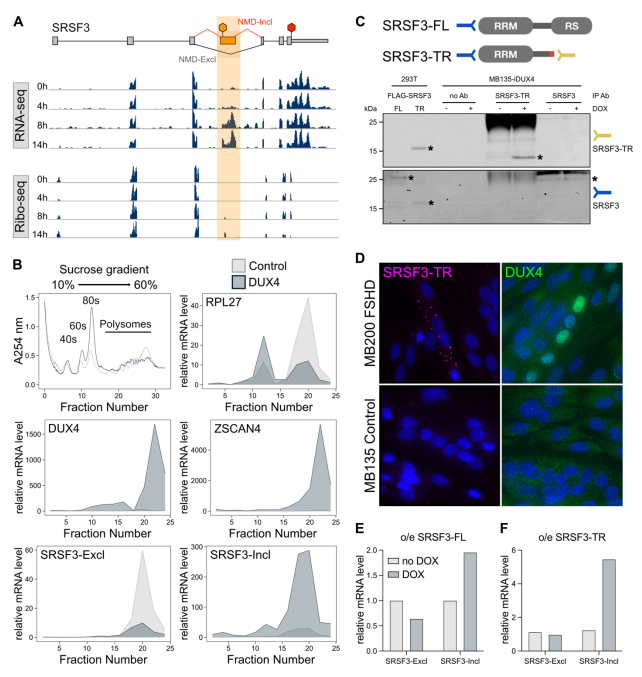


Figure 5. Truncated SRSF3 protein could disrupt RNA processing in FSHD myotubes. (**A**) RNA-seq and Riboseq coverage over SRSF3. The PTC-containing exon 4 is highlighted. The red hexagon indicates the normal stop codon while the orange hexagon denotes the PTC. (**B**) Absorbance at 254 nm across a sucrose density gradient of lysates from control MB135-iDUX4 myoblasts and MB135-iDUX4 myoblasts expressing DUX4 for 14 h (top left). RTqPCR measurement of RPL27, DUX4, ZSCAN4, SRSF3-Excl, and SRSF3-Incl mRNA levels in DUX4-expressing myoblasts relative to control myoblasts from collected fractions (remaining panels). (**C**) Detection of SRSF3 and SRSF3-TR in whole cell extracts from 293T cells exogenously expressing FLAG-tagged full-length (FL) or truncated (TR) SRSF3 as compared to protein lysates from MB135-iDUX4 myoblasts treated with (+) or without (-) doxycycline (DOX) to induce DUX4 and immunoprecipitated with a custom anti-SRSF3-TR antibody, no antibody (Ab), or a commercial SRSF3 antibody. IP, immunoprecipitation. Asterisks denote proteins of interest. (**D**) Immunofluorescence staining in MB135 control and MB200 FSHD myotubes differentiated for 72 h and stained with DAPI (blue) and rabbit anti-DUX4 (green) or custom rabbit anti-SRSF3-TR (pink) antibody. (**E**) SRSF3-Excl and -Incl mRNA isoform levels as determined by RT-qPCR following induction of full-length (FL) or truncated (TR) SRSF3 in MB135-iFLAG-SRSF3-FL or -TR myoblasts by treatment with doxycycline for 72 h.

286 terminal epitope common to both the full-length and truncated SRSF3. This antibody detected 287 both exogenously expressed full-length and truncated FLAG-SRSF3, and endogenous full-288 length SRSF3, but was insufficient to visualize endogenous SRSF3-TR (Figure 5C), possibly 289 due to lower affinity for this protein isoform in an immunoprecipitation assay. To determine if 290 SRSF3-TR was present in FSHD myotubes expressing endogenous levels of DUX4, we carried 291 out immunofluorescence for SRSF3-TR or DUX4 in differentiated FSHD and control muscle 292 cells. While there was no detectable SRSF3-TR staining in control cells, in DUX4-expressing 293 FSHD cultures SRSF3-TR appeared in cytoplasmic puncta (Figure 5D). Together, these data 294 show that endogenous SRSF3-TR is present at detectable levels in DUX4-expressing MB135-295 iDUX4 and FSHD cells.

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297 To explore the functional consequences of SRSF3-TR expression, we exogenously expressed 298 FLAG-tagged full-length or truncated SRSF3 in healthy muscle cells. As previously described, 299 full-length SRSF3 decreased the level of the normal SRSF3 mRNA isoform and increased the 300 aberrant isoform ((Jumaa & Nielsen, 1997); Figure 5E). Strikingly, SRSF3-TR led to a 5-fold 301 upregulation of the endogenous SRSF3 aberrant mRNA isoform (Figure 5F). These data 302 suggest that DUX4-induced translation of SRSF3-TR may feedback to create more aberrant 303 RNA and therefore more truncated protein, highlighting the importance of misregulated post-304 transcriptional processes in DUX4-induced pathology.

305 DISCUSSION

306 We paired RNA-seg and Ribo-seg across a time course of DUX4 expression in MB135-iDUX4 307 human skeletal muscle myoblasts, providing an integrative temporal view of the transcriptome 308 and translatome in a well-accepted cellular model of FSHD. Our choice to examine early time 309 points, before overt DUX4-induced cytotoxicity, provides a view into early DUX4-driven 310 regulation that builds on prior work at later time points (Bosnakovski et al., 2019; Chew et al., 311 2019; DeSimone, Leszyk, Wagner, & Emerson, 2019; Geng et al., 2012; Jagannathan et al., 312 2019; Jagannathan et al., 2016; Lek et al., 2020; Resnick et al., 2019; Shadle et al., 2019; 313 Shadle et al., 2017; Sharma et al., 2013; Whiddon et al., 2017). Taken together, our results 314 demonstrate that critical pathways underlying FSHD pathophysiology are perturbed earlier than 315 previously understood and provide datasets that should serve as an informative resource for the 316 FSHD community as it works to develop effective FSHD therapeutics. 317

318 Our time course RNA-seq data revealed a large number of both up and downregulated genes 319 after only 4 h of DUX4 expression. While the former result is expected given DUX4's role as a 320 transcriptional activator (Geng et al., 2012), this degree of early gene silencing is remarkable 321 and would require rapid cessation of transcription. There is no evidence that DUX4 acts as a 322 direct repressor of transcription (Bosnakovski et al., 2018); therefore, our results suggest that 323 there exists one or more unstudied DUX4-induced transcriptional repressors responsible for 324 early gene downregulation. Indeed, of the 883 genes significantly upregulated by DUX4 at 4 h, 325 28 are DNA-binding transcription factors known to effect gene repression and at least 5 more 326 are DNA-binding proteins with an as-yet-undetermined role in gene regulation. This novel 327 hypothesis provides a mechanism by which DUX4 inhibits gene expression that is unique from 328 previously suggested models invoking competition of DUX4 with the homeodomain transcription 329 factors PAX3 and PAX7 (C. R. S. Banerji et al., 2017; Bosnakovski, Daughters, Xu, Slack, & 330 Kyba, 2009; Bosnakovski, Toso, et al., 2017; Bosnakovski et al., 2008) or transcriptional 331 interference from DUX4-induced non-coding transcripts (Bosnakovski et al., 2018). Future work 332 could determine which, if any, of the DNA-binding repressors induced by DUX4 cause early 333 DUX4-mediated gene downregulation.

334

335 The profound consequences of DUX4 misexpression on cell identity were also revealed in our 336 time course RNA-seq results. GO analyses showed that DUX4-expressing myoblasts begin to 337 turn on genes that promote cell proliferation and inhibit cell differentiation, and silence genes 338 that promote myogenesis, early after DUX4 induction. Thus, skeletal muscle – a post-mitotic,

differentiated cell type – is pushed towards a proliferative, naïve embryonic state. This conflicted
cell identity may contribute to apoptosis, as has been proposed before (Ashoti, Alemany, Sage,
& Geijsen, 2021; Geng et al., 2012). Interestingly, a few hours after DUX4 begins forcing
expression of an embryonic gene expression program, and just as overt apoptosis appears,
cells respond by altering core regulatory processes such as RNA splicing and localization, and
protein ubiquitination. Whether these changes underlie cellular "coping" mechanisms that delay
death or underpin cytotoxicity remains to be determined.

347 We recently described the DUX4-induced proteome using MB135-iDUX4 cells (Jagannathan et 348 al., 2019). Although this catalog of expressed proteins remains incomplete due to partial 349 coverage and restricted dynamic range inherent to mass spectrometry, our proteomics data 350 demonstrated that hundreds of genes are post-transcriptionally modulated by DUX4. But, these 351 experiments could not determine if this regulation was mediated at the level of protein synthesis 352 or degradation. The time course Ribo-seq presented here complements and extends this earlier 353 work, and suggests that most DUX4 post-transcriptional regulation, including of multiple key 354 NMD factors, occurs via proteolysis. We did uncover a small number of genes regulated at the 355 level of protein synthesis at the 14 h time point of DUX4 expression, likely representing the 356 subset of transcripts impacted by eIF2 α phosphorylation induced by DUX4, as previously 357 reported (Jagannathan et al., 2019; Shadle et al., 2017).

358

359 Loss of NMD leads to the stabilization of aberrant RNAs (Kurosaki & Maguat, 2016). However, 360 few studies have looked at whether aberrant RNAs are translated and what proteins they might 361 produce. Here, we demonstrate that DUX4-induced NMD inhibition causes truncated protein 362 production in muscle cells. The confirmed existence of truncated proteins in DUX4-expressing 363 cells has implications for how we understand cytotoxicity. Protein truncation could result in a 364 dominant negative function that inhibits the activity of the remaining, cell critical full-length 365 protein. Truncated proteins might misfold and facilitate the formation of protein aggregates, such 366 as those observed in FSHD myotubes (Homma, Beermann, Boyce, & Miller, 2015; Homma, 367 Beermann, Yu, Boyce, & Miller, 2016; Shadle et al., 2017). Additionally, some DUX4-induced 368 truncated proteins contain unique C-terminal extensions, or neo-peptides, that could serve as 369 novel antigens and might induce inflammation in FSHD muscle. Importantly, we were able to 370 detect truncated proteins in FSHD muscle cell cultures, thereby validating our findings from the 371 inducible DUX4 system in a more physiological setting and raising the possibility that these 372 molecules could be used clinically as functionally relevant and FSHD-specific biomarkers.

373 Future work is required to establish whether truncated proteins, either individually or in 374 combination, contribute to cell death.

375

376 Many of the identified DUX4-induced truncated proteins are RBPs and splicing factors. It is well-377 established that DUX4 alters RNA splicing (Geng et al., 2012; Jagannathan et al., 2019; 378 Jagannathan et al., 2016; Rickard et al., 2015) and therefore interesting to speculate that 379 truncated RBPs and splicing proteins might be responsible for inducing global RNA processing 380 defects. Such misprocessing would generate aberrant RNAs that could act to further overwhelm 381 the already inhibited NMD pathway. As an example of this phenomenon, we observed that 382 exogenous expression of the truncated splicing factor SRSF3-TR causes a significant increase 383 in the level of the corresponding aberrant SRSF3 RNA. Splicing factors are known to regulate 384 and coordinate their expression via various forms of autoregulation (Konigs et al., 2020; Leclair 385 et al., 2020; Muller-McNicoll, Rossbach, Hui, & Medenbach, 2019). Overexpression of full-length 386 SRSF3 leads to a reduction in the level of normal SRSF3 transcript while at the same time 387 activating the production of aberrant SRSF3 RNA that gets degraded by NMD in a balancing act 388 mechanism that prevents SRSF3 accumulation (Jumaa & Nielsen, 1997). Our results suggest 389 that SRSF3-TR facilitates its own production and disrupts normal autoregulatory processes. 390 leading to a new model in which DUX4-expressing cells hijack the splicing protein 391 autoregulatory network and flip it – instead of acting to limit the production of full-length protein 392 the creation of truncated protein is amplified. 393 394 In summary, our study provides a critical missing piece towards a comprehensive

395 characterization of the major steps in DUX4-induced gene expression. We demonstrate that

396 DUX4 induces cascading changes at all levels of gene regulation that are invisible at steady

397 state, offering a glimpse into how developmentally regulated transcription factors expressed at

398 the wrong time in the wrong context can wreak havoc leading to human disease.

399	MATERIALS AND METHODS
400	
401	RESOURCE AVAILABILITY
402	
403	Lead contact
404	Further information and requests for resources and reagents should be directed to and will be
405	fulfilled by the lead contact, Sujatha Jagannathan (sujatha.jagannathan@cuanschutz.edu).
406	Materiala availability
407	Materials availability
408	The cell lines and antibody generated in this study are available upon request. Plasmids
409 410	generated in this study have been deposited to Addgene (plasmid #171951, #171952, #172345, and #172346).
411	
412	Data and code availability
413	The RNA-seq and Ribo-seq data generated during this study are available at GEO (accession
414	number GSE178761). The code generated during this study are available at GitHub
415	(https://github.com/sjaganna/2021-campbell_dyle_calviello_et_al).
416	
417	EXPERIMENTAL MODEL AND SUBJECT DETAILS
417 418	EXPERIMENTAL MODEL AND SUBJECT DETAILS
	EXPERIMENTAL MODEL AND SUBJECT DETAILS Cell lines and culture conditions
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- 433 dihydrochloride (VWR). MB135-iFLAG-SRSF3-FL and -TR myoblasts were additionally
- 434 maintained in 10 µg/mL blasticidin S HCI (Thermo Fisher Scientific). Induction of DUX4 and
- 435 SRSF3 transgenes was achieved by culturing cells in 1 µg/mL doxycycline hyclate (Sigma-
- 436 Aldrich). Differentiation of myoblasts into myotubes was achieved by switching the fully
- 437 confluent myoblast monolayer into DMEM containing 1% horse serum (Thermo Fisher
- 438 Scientific) and Insulin-Transferrin-Selenium (Thermo Fisher Scientific). All cells were incubated
- 439 at 37 °C with 5% CO₂.
- 440

441 METHOD DETAILS

- 442
- 443 Cloning
- 444 pTwist-FLAG-SRSF3_Full.Length_Codon.Optimized and pTwist-FLAG-
- 445 SRSF3_Truncated_Codon.Optimized plasmids were synthesized by Twist Bioscience. To
- 446 construct pCW57.1-FLAG-SRSF3_Full.Length_Codon.Optimized-Blast and pCW57.1-FLAG-
- 447 SRSF3_Truncated_Codon.Optimized-Blast plasmids, the SRSF3 open reading frames were
- subcloned into pCW57-MCS1-P2A-MCS2 (Blast) (a gift from Adam Karpf, Addgene plasmid
- 449 #80921) (Barger, Branick, Chee, & Karpf, 2019) by restriction enzyme digest using EcoRI and
- 450 BamHI (New England Biolabs).
- 451

452 Antibody generation

- 453 Purified SRSF3-TR peptide (Cys-PRRRVTIMSLLTTL) was used as an immunogen and
- 454 polyclonal rabbit anti-SRSF3-TR antibody production was done in collaboration with Pacific
- 455 Immunology (Ramona, CA). The antisera from all animals were screened for reactivity by ELISA
- 456 against the immunogen and with western blots and immunofluorescence against transfected
- 457 SRSF3-TR.
- 458

459 Transgenic cell line generation

- 460 Lentiviral particles expressing doxycycline-inducible FLAG-SRSF3-FL or -TR transgenes were
- 461 generated by co-transfecting 293T cells with the appropriate lentivector, pMD2.G (a gift from
- 462 Didier Trono, Addgene plasmid #12259), and psPAX2 (a gift from Didier Trono, Addgene
- 463 plasmid #12260) using Lipofectamine 2000 Transfection Reagent (Thermo Fisher Scientific). To
- 464 generate polyclonal SRSF3 transgenic cell lines, MB135 myoblasts were transduced with
- 465 Ientivirus in the presence of 8 μg/mL polybrene (Sigma-Aldrich) and selected using 10 μg/mL
- 466 blasticidin S HCl.

467 Plasmid transfections

- 468 293T cells were transfected with pTwist-FLAG-SRSF3_Full.Length_Codon.Optimized and
- 469 pTwist-FLAG-SRSF3_Truncated_Codon.Optimized plasmids using Lipofectamine 2000
- 470 Transfection Reagent following the manufacturer's instructions.
- 471

472 Live cell imaging

- 473 MB135-iDUX4/ZSCAN4-mCherry myoblasts were induced with doxycycline hyclate to turn on
- 474 DUX4 expression and subjected to time lapse imaging using the IncuCyte S3 incubator
- 475 microscope system (Sartorius). Images were collected every 15 min from the time of
- 476 doxycycline addition (t = 0 h) to 28 h.
- 477

478 RNA extraction and RT-qPCR

- 479 Total RNA was extracted from whole cells using TRIzol Reagent (Thermo Fisher Scientific)
- 480 following the manufacturer's instructions. Isolated RNA was treated with DNase I (Thermo
- 481 Fisher Scientific) and reverse transcribed to cDNA using SuperScript III reverse transcriptase
- 482 (Thermo Fisher Scientific) and random hexamers (Thermo Fisher Scientific) according to the
- 483 manufacturer's protocol. Quantitative PCR was carried out on a CFX384 Touch Real-Time PCR
- 484 Detection System (Bio-Rad) using primers specific to each gene of interest and iTaq Universal
- 485 SYBR Green Supermix (Bio-Rad). The expression levels of target genes were normalized to
- that of the reference gene *RPL27* using the delta-delta-Ct method (Livak & Schmittgen, 2001).
- 487 The primers used in this study are listed in the Key Resources Table.
- 488

489 RNA-seq library preparation and sequencing

- Total RNA was extracted from whole cells using TRIzol Reagent following the manufacturer's
 instructions. Isolated RNA was subjected to ribosomal RNA depletion using the Ribo-Zero rRNA
 Removal Kit (Illumina). RNA-seq libraries were prepared using the NEXTflex Rapid Directional
 qRNA-Seq Kit (Bioo Scientific) following the manufacturer's instructions and sequenced using
 75 bp single-end sequencing on the Illumina NextSeq 500 platform by the BioFrontiers Institute
- 495 Next-Gen Sequencing Core Facility.
- 496

497 Ribosome footprinting

498 Ribo-seq was performed as described previously (Calviello et al., 2020) using six 70% confluent

- 499 10 cm dishes of MB135-iDUX4 cells per condition. Briefly, cells were washed with ice-cold
- 500 phosphate-buffered saline (PBS) supplemented with 100 µg/mL cycloheximide (Sigma-Aldrich),

501 flash frozen on liquid nitrogen, and lysed in Lysis Buffer (PBS containing 1% (v/v) Triton X-100 502 and 25 U/mL TurboDNase (Ambion)). Cells were harvested by scraping and further lysed by 503 trituration ten times through a 26-gauge needle. The lysate was clarified by centrifugation at 504 20.000 g for 10 min at 4 °C. The supernatants were flash frozen in liquid nitrogen and stored at -505 80 °C. Thawed lysates were treated with RNase I (Ambion) at 2.5 U/µL for 45 min at room 506 temperature with gentle mixing. Further RNase activity was stopped by addition of SUPERaseIn 507 RNase Inhibitor (Thermo Fisher Scientific). Next, ribosome complexes were enriched using 508 MicroSpin S-400 HR Columns (GE Healthcare) and RNA extracted using the Direct-zol RNA 509 Miniprep Kit (Zymo Research). Ribo-Zero rRNA Removal Kit was used to deplete rRNAs and 510 the ribosome-protected fragments were recovered by running them in a 17% Urea gel, staining 511 with SYBR Gold (Invitrogen), and extracting nucleic acids that are 27 to 30 nucleotides long 512 from gel slices by constant agitation in 0.3 M NaCl at 4 °C overnight. The recovered nucleic 513 acids were precipitated with isopropanol using GlycoBlue Coprecipitant (Ambion) as carrier and 514 treated with T4 polynucleotide kinase (Thermo Fisher Scientific). Libraries were prepared using 515 the NEXTflex Small RNA-Seg Kit v3 (Bioo Scientific) following the manufacturer's instructions 516 and sequenced using 75 bp single-end reads on an Illumina NextSeq 500 by the BioFrontiers 517 Institute Next-Gen Sequencing Core Facility.

518

519 RNA-seq and Ribo-seq data analysis

520 Fastq files were stripped of the adapter sequences using cutadapt. UMI sequences were 521 removed, and reads were collapsed to fasta format. Reads were first aligned against rRNA 522 (accession number U13369.1), and to a collection of snoRNAs, tRNAs, and miRNA (retrieved 523 using the UCSC table browser) using bowtie2 (Langmead & Salzberg, 2012). Remaining reads 524 were mapped to the hq38 version of the genome (without scaffolds) using STAR 2.6.0a (Dobin 525 et al., 2013) supplied with the GENCODE 25 .gtf file. A maximum of two mismatches and 526 mapping to a minimum of 50 positions was allowed. De-novo splice junction discovery was 527 disabled for all datasets. Only the best alignment per each read was retained. Quality control 528 and read counting of the Ribo-seg data was performed with Ribo-seQC (Calviello, Sydow, 529 Harnett, & Ohler, 2019).

530

531 Differential gene expression analysis of the RNA-seq data was conducted using DESeq2 (Love,

Huber, & Anders, 2014). Briefly, featureCounts from the subread R package (Liao, Smyth, &

533 Shi, 2014) was used to assign aligned reads (in BAM format) to genomic features supplied with

the GENCODE 25. gtf file. The featureCounts output was then supplied to DESeq2 and

differential expression analysis was conducted with the 0 h time point serving as the reference
sample. Genes with very low read count were filtered out by requiring at least a total of 10 reads

- 537 across the 12 samples (3 replicates each of the 0, 4, 8, and 14 h samples). Log2 fold change
- shrinkage was done using the apeglm function (Zhu, Ibrahim, & Love, 2019).
- 539

540 Differential analysis of the RNA-seg and Ribo-seg data was performed using DESeg2, as 541 previously described (Calviello et al., 2021; Chothani et al., 2019), using an interaction model 542 between the tested condition and RNA-seq – Ribo-seq counts. Only reads mapping uniquely to 543 coding sequence regions were used. In addition, ORFquant (Calviello et al., 2020) was used to 544 derive de-novo isoform-specific translation events, by pooling the Ribo-seQC output from all 545 Ribo-seq samples, using uniquely mapping reads. DEXSeq (Anders et al., 2012) was used to 546 perform differential exon usage along the DUX4 time course data, using Ribo-seg counts on 547 exonic bins and junctions belonging to different ORF guant-derived translated regions. NMD 548 candidates were defined by ORFquant as open reading frames ending with a stop codon 549 upstream of an exon-exon junction.

550

551 GO category analysis

552 Gene Ontology (GO) analysis was conducted using the web tool http://geneontology.org, 553 powered by pantherdb.org. Briefly, statistical overrepresentation test using the complete GO 554 biological process annotation dataset was conducted and p-values were calculated using the 555 Fisher's exact test and False Discovery Rate was calculated by the Benjamini-Hochberg 556 procedure.

557

558 Polysome profiling

559 Polysome profiling was performed as previously described (Merrick & Hensold, 2001; Miura, 560 Andrews, Holcik, & Jasmin, 2008) with the following modifications. Four 70% confluent 15 cm 561 dishes of MB135-iDUX4 cells per condition were treated with 100 µg/mL cycloheximide for 10 562 min. transferred to wet ice, washed with ice-cold PBS containing 100 µg/mL cvcloheximide, and 563 then lysed in 400 µL Lysis Buffer (20 mM HEPES pH 7.4, 15 mM MgCl₂, 200 mM NaCl, 1% 564 Triton X-100, 100 µg/mL cycloheximide, 2 mM DTT, and 100 U/mL SUPERaseIn RNase 565 Inhibitor) per 15 cm dish. The cells and buffer were scraped off the dish and centrifuged at 566 13,000 rpm for 10 min at 4 °C. Lysates were fractionated on a 10% to 60% sucrose gradient 567 using the SW 41 Ti Swinging-Bucket Rotor (Beckman Coulter) at 36,000 rpm for 3 h and 10 568 min. Twenty-four fractions were collected using a Gradient Station ip (BioComp) and an FC

- 569 203B Fraction Collector (Gilson) with continuous monitoring of absorbance at 254 nm. RNA
- 570 from each fraction was extracted using TRIzol LS Reagent (Thermo Fisher Scientific) following
- 571 the manufacturer's instructions. RT-qPCR was carried out as described above.
- 572

573 **Protein extraction**

Total protein was extracted from whole cells using TRIzol Reagent following the manufacturer's
instructions, excepting that protein pellets were dissolved in Protein Resuspension Buffer (0.5 M
Tris base, 5% SDS). Isolated protein was quantified using the Pierce BCA Protein Assay Kit
(Thermo Fisher Scientific) according to the manufacturer's protocol. Protein was mixed with 4X
NuPAGE LDS Sample Buffer (Thermo Fisher Scientific) containing 50 mM DTT and heated to

- 579 70 °C before immunoblotting.
- 580

581 Immunoprecipitation

582 MB135-iDUX4 myoblasts were treated with or without doxycycline for 14 h and then trypsinized 583 prior to lysis on ice in 1 mL of Lysis Buffer (50 mM Tris-HCl pH 7.5, 150 mM NaCl, 1% NP-40) 584 containing protease inhibitors (Sigma Aldrich). Lysates were precleared using Protein G 585 Sepharose (Thermo Fisher Scientific) for 1 h prior to an overnight incubation at 4 °C with either 586 anti-SRSF3 or anti-SRSF3-TR antibody. Protein G Sepharose was added the following morning 587 for 5 h to bind the antibody, and beads were subsequently washed 5 times with 1 mL cold Lysis 588 Buffer. After the final wash, 4X NuPAGE LDS Sample Buffer containing 50 mM DTT was added 589 directly to the beads and samples heated to 70 °C for protein elution before immunoblotting.

590

591 Immunoblotting

592 Protein was run on NuPAGE Bis-Tris precast polyacrylamide gels (Thermo Fisher Scientific)

alongside PageRuler Plus Prestained Protein Ladder (Thermo Fisher Scientific) and transferred

- to Odyssey nitrocellulose membrane (LI-COR Biosciences). Membranes were blocked in
- 595 Intercept (PBS) Blocking Buffer (LI-COR Biosciences) before overnight incubation at 4 °C with
- primary antibodies diluted in Blocking Buffer containing 0.2% Tween 20. Membranes were
- 597 incubated with IRDye-conjugated secondary antibodies (LI-COR Biosciences) for 1 h and
- 598 fluorescent signal visualized using a Sapphire Biomolecular Imager (Azure Biosystems) and
- 599 Sapphire Capture software (Azure Biosystems). When appropriate, membranes were stripped
- 600 with Restore Western Blot Stripping Buffer (Thermo Fisher Scientific) before being re-probed.
- Band intensities were quantified by densitometry using ImageJ (Schneider, Rasband, & Eliceiri,
- 602 2012).

603 Immunofluorescence

604 Cells were fixed in 10% Neutral Buffered Formalin (Research Products International) for 30 min 605 and permeabilized for 10 min in PBS with 0.1% Triton X-100. Samples were then incubated 606 overnight at 4 °C with primary antibodies, followed by incubation with 488- or 594-conjugated 607 secondary antibodies for 1 h prior to counterstaining and mounting with Prolong Diamond 608 Antifade Mountant with DAPI (Thermo Fisher Scientific). Slides were imaged with a DeltaVision 609 Elite deconvolution microscope, CoolSNAP HQ² high-resolution CCD camera, and Resolve3D 610 softWoRx-Acquire v7.0 software. Image J software (Schneider et al., 2012) was used for image 611 analysis.

612

613 Antibodies

- The antibodies used in this study are anti-DUX4 (Abcam 124699), anti-Histone H3 (Abcam
- 615 1791), anti-SRSF3 (Thermo Fisher Scientific 33-4200), anti-SRSF3-TR (this paper), anti-
- 616 RENT1/hUPF1 (Abcam ab109363), Drop-n-Stain CF 488A Donkey Anti-Rabbit IgG (Biotium
- 617 20950), Drop-n-Stain CF 594 Donkey Anti-Rabbit IgG (Biotium 20951), IRDye 650 Goat anti-
- Mouse IgG Secondary Antibody (LI-COR Biosciences 926-65010), and IRDye 800CW Goat
- 619 anti-Rabbit IgG Secondary Antibody (LI-COR Biosciences 926-32211).
- 620

621 QUANTIFICATION AND STATISTICAL ANALYSIS

622

623 Data analysis, statistical tests, and visualization

All data analysis and statistical tests were performed in the R programming environment and

- relied on Bioconductor (Huber et al., 2015) and ggplot2 (Wickham, 2016). Plots were generated
- 626 using R plotting functions and/or the ggplot2 package. Bar and line graphs were generated
- 627 using GraphPad Prism software version 9.0. Biological replicates were defined as experiments
- 628 performed separately on distinct samples (i.e. cells cultured in different wells) representing
- 629 identical conditions and/or time points. No outliers were eliminated in this study. All statistical
- 630 tests were performed using R functions.

631 ACKNOWLEDGEMENTS

- 632 We thank Stephen Tapscott for the MB135-iDUX4/ZSCAN4-mCherry cell line. We thank Jeffrey 633 Kieft for his guidance in carrying out polysome profiling. We thank Neelanian Mukherjee, Olivia 634 Rissland, Srinivas Ramachandran, and all members of the Jagannathan laboratory for insightful 635 manuscript feedback. We thank the BioFrontiers Institute Next-Gen Sequencing Core Facility, 636 which performed the Illumina sequencing and library construction. This work was supported by 637 the RNA Bioscience Initiative, University of Colorado Anschutz Medical Campus (S.J.), Friends 638 of FSH Research and The Chris Carrino Foundation for FSHD AWD-194864 (S.J.), the FSHD 639 Society FSHS-82018-01 (A.E.C. and M.D.), the California Tobacco-Related Disease Research 640 Grants Program 27KT-0003 (S.N.F.), and the National Institutes of Health DP2GM132932
- 641 (S.N.F.).
- 642

643 AUTHOR CONTRIBUTIONS

A.E.C., M.C.D., and S.J. conceived and designed the study. A.E.C, M.C.D., M.A.C., and T.F.

performed experiments. A.E.C., L.C., T.M., R.F., A.E.G., S.N.F., and S.J. analyzed data. A.E.C.

- and S.J. wrote the paper with input from all authors.
- 647

648 **DECLARATION OF INTERESTS**

649 The authors declare no competing interests.

650 SUPPLEMENTAL FILES

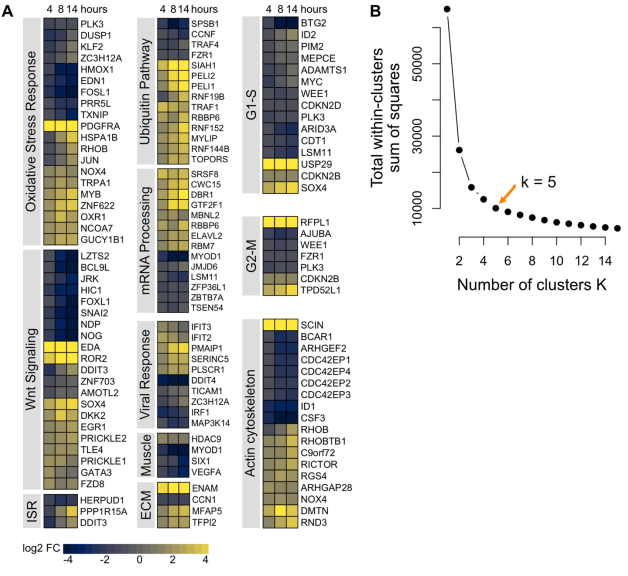


Figure 2 – figure supplement 1. Gene-level analysis of early changes in pathways impacted by DUX4. (A) Heat maps of transcript-level changes in genes that belong to various biological pathways known to be misregulated by DUX4, as defined by the Gene Ontology Biological Process classification system accessed through GSEA/MSigDB database. The subset of genes that showed significant change in expression at the 4 h time point of DUX4 induction are plotted. (**B**) Elbow method analysis was used to determine appropriate cluster number for Figure 2C-E.

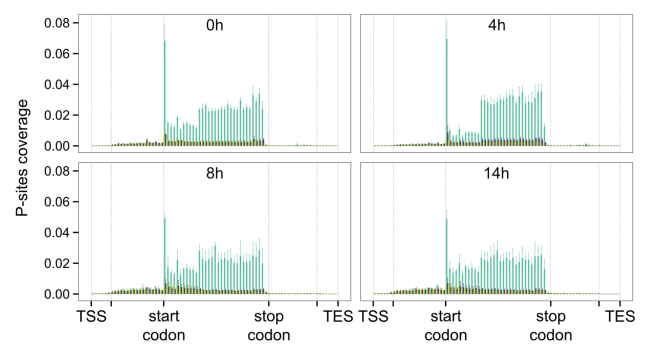


Figure 3 – figure supplement 1. Ribo-seq quality control. Aggregate profile of P-sites coverage (as calculated by Ribo-seQC) depicting single nucleotide resolution of Ribo-seq data along the time course. Each frame is shown with a different color. Error bars represent the standard deviation from three biological replicates.

651 Video 1. Time course imaging following DUX4 expression.

- 652 Live cell fluorescence microscopy recording of MB135-iDUX4/ZSCAN4-mCherry myoblasts
- treated with doxycycline to induce DUX4 expression.
- 654

655 **Supplementary Table 1.**

- 656 DESeq2 differential gene expression analysis results for DUX4 time course RNA-seq data at 4,
- 657 8, or 14 h post-induction compared to 0 h (control).
- 658

659 Supplementary Table 2.

- 660 Cluster analysis of RNA-seq log2 fold change at 4, 8, or 14 h of DUX4 induction compared to 0
- h (control); and Gene Ontology analysis of genes within each cluster.
- 662

663 Supplementary Table 3.

- 664 ORFquant analysis results for DUX4 time course RNA-seq and Ribo-seq data at 4, 8, or 14 h
- 665 post-induction compared to 0 h (control); and Gene Ontology analysis of genes with
- 666 downregulated ribosome density at 14 h.
- 667

668 Supplementary Table 4.

- 669 DEXSeq analysis results for DUX4 time course Ribo-seq data at 4, 8, or 14 h post-induction
- 670 compared to 0 h (control); and Gene Ontology analysis of NMD targets upregulated at 14 h.

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852 APPENDIX

853

854 KEY RESOURCES TABLE

Reage nt type (specie s) or resour ce	Designation	Source or reference	Identifiers	Additional information
cell line (Homo sapiens)	293T	ATCC	Cat# CRL- 3216, RRID:CVCL_ 0063	
cell line (<i>H.</i> sapiens)	MB135	Stephen Tapscott		Female control myoblast line
cell line (<i>H.</i> sapiens)	MB135-iDUX4	(Jagannathan et al., 2016)		Harbors doxycycline- inducible DUX4 transgene
cell line (<i>H.</i> sapiens)	MB135- iDUX4/ZSCAN4- mCherry	Stephen Tapscott		Harbors doxycycline- inducible DUX4 transgene and DUX4- responsive fluorescent reporter
cell line (<i>H.</i> sapiens)	MB135-iFLAG-SRSF3- FL	This paper		Harbors doxycycline- inducible, FLAG- tagged, codon- optimized, full-length SRSF3
cell line (<i>H.</i> sapiens)	MB135-iFLAG-SRSF3- TR	This paper		Harbors doxycycline- inducible, FLAG- tagged, codon- optimized, truncated SRSF3
cell line (<i>H.</i> sapiens)	MB200	Stephen Tapscott		Male FSHD myoblast line
antibod y	anti-DUX4 (Rabbit monoclonal)	Abcam	Cat#ab12469 9, RRID:AB_109 73363	IF(1:200), WB(1:1000)
antibod y	anti-Histone H3 (Rabbit polyclonal)	Abcam	Cat# ab1791, RRID:AB_302 613	WB(1:5000)

antibod y	anti-SRSF3 (Mouse monoclonal)	Thermo Fisher Scientific	Cat# 33- 4200, RRID:AB_253 3119	WB(1:250)
antibod y	anti-SRSF3-TR (Rabbit polyclonal)	This paper		IF(1:200), WB(1:500)
antibod y	anti-RENT1/hUPF1 (Rabbit monoclonal)	Abcam	Cat# ab109363, RRID:AB_108 61979	WB(1:1000)
recomb inant DNA reagent	pCW57.1-FLAG- SRSF3_Full.Length_Co don.Optimized-Blast (plasmid)	This paper		All-in-one lentiviral vector for doxycycline- inducible expression of FLAG-tagged, codon- optimized, full-length SRSF3
recomb inant DNA reagent	pCW57.1-FLAG- SRSF3_Truncated_Cod on.Optimized-Blast (plasmid)	This paper		All-in-one lentiviral vector for doxycycline- inducible expression of FLAG-tagged, codon- optimized, truncated SRSF3
recomb inant DNA reagent	pCW57-MCS1-P2A- MCS2 (Blast) (plasmid)	Addgene	Plasmid #80921, RRID:Addgen e_80921	
recomb inant DNA reagent	pMD2.G (plasmid)	Addgene	Plasmid #12259; RRID:Addgen e 12259	
recomb inant DNA reagent	psPAX2 (plasmid)	Addgene	Plasmid #12260; RRID:Addgen e_12260	
recomb inant DNA reagent	pTwist-FLAG- SRSF3_Full.Length_Co don.Optimized (plasmid)	This paper		For expression of FLAG-tagged, codon- optimized full-length SRSF3 in mammalian cells; synthesized by Twist Bioscience
recomb inant DNA reagent	pTwist-FLAG- SRSF3_Truncated_Cod on.Optimized (plasmid)	This paper		For expression of FLAG-tagged, codon- optimized truncated SRSF3 in mammalian cells; synthesized by Twist Bioscience
sequen ce- based reagent	DUX4 transgene F	(Shadle et al., 2019)	qPCR primers	TAGGGGAAGAGGTA GACGGC

sequen ce-	DUX4 transgene R	(Shadle et al., 2019)	qPCR primers	CGGTTCCGGGATTC CGATAG
based reagent				
sequen ce- based reagent	HAGHL F	This paper	qPCR primers	AGTTTGCCCAGAAAG TGGAG
sequen ce- based reagent	HAGHL R	This paper	qPCR primers	ATCCTCATCCCTCTT CTTAGCC
sequen ce- based reagent	HSATII F	(Shadle et al., 2019)	qPCR primers	TGAATGGAATCGTCA TCGAA
sequen ce- based reagent	HSATII R	(Shadle et al., 2019)	qPCR primers	CCATTCGATAATTCC GCTTG
sequen ce- based reagent	HSPA5 F	(Lin et al., 2007)	qPCR primers	CGGGCAAAGATGTC AGGAAAG
sequen ce- based reagent	HSPA5 R	(Lin et al., 2007)	qPCR primers	TTCTGGACGGGCTT CATAGTAGAC
sequen ce- based reagent	MYOD1 F	Stephen Tapscott	qPCR primers	AGCACTACAGCGGC GACT
sequen ce- based reagent	MYOD1 R	Stephen Tapscott	qPCR primers	GCGACTCAGAAGGC ACGTC
sequen ce- based reagent	SRSF3 F	(Feng et al., 2015)	qPCR primers	TGGAACTGTCGAATG GTGAA
sequen ce- based reagent	SRSF3-Excl R	(Feng et al., 2015)	qPCR primers	CTTGGAGATCTGCG ACGAG
sequen ce- based reagent	SRSF3-Incl R	(Feng et al., 2015)	qPCR primers	GGGTGGTGAGAAGA GACATGA
sequen ce-	TCF3 F	(Zeisel, Yitzhaky,	qPCR primers	CCCCAGGAGAATGA ACCAG

based		Bossel Ben-		
reagent		Moshe, &		
		Domany,		
		2013)		
sequen	TCF3 R	(Zeisel et al.,	qPCR primers	CCTTCCCGTTGGTGA
ce-		2013)		CAG
based				
reagent				
sequen	RPL27 F	(Jagannathan	qPCR primers	GCAAGAAGAAGATC
ce-		et al., 2016)		GCCAAG
based		,,		
reagent				
sequen	RPL27 R	(Jagannathan	qPCR primers	TCCAAGGGGATATC
ce-		et al., 2016)		CACAGA
based		et al., 2010)		CACACA
reagent	ZSCAN4 F	(Geng et al.,	qPCR primers	TGGAAATCAAGTGG
sequen	ZSCAN4 F		qPCR primers	
ce-		2012)		СААААА
based				
reagent	700000			
sequen	ZSCAN4 R	(Geng et al.,	qPCR primers	CTGCATGTGGACGT
ce-		2012)		GGAC
based				
reagent				
peptide	Insulin-Transferrin-	Thermo	41400045	
,	Selenium	Fisher		
recomb		Scientific		
inant				
protein				
peptide	Protein G Sepharose	Thermo	101241	
	•	Fisher		
recomb		Scientific		
inant				
protein				
	Recombinant human	Promega	G5071	
P = P = 00	basic fibroblast growth			
, recomb	factor			
inant				
protein				
chemic	Blasticidin S HCl	Thermo	R21001	
al		Fisher		
		Scientific		
compo		Scientinic		
und,				
drug	Overlah en 1. 11		000705	
chemic	Cycloheximide	Sigma-Aldrich	239765	
al				
compo				
und,				
drug				

chemic	Doxycycline hyclate	Sigma-Aldrich	D9891	
al compo und, drug				
chemic al compo und, drug	Lipofectamine 2000 Transfection Reagent	Thermo Fisher Scientific	11668-030	
chemic al compo und, drug	Puromycin dihydrochloride	VWR	97064-280	
chemic al compo und, drug	TRIzol Reagent	Thermo Fisher Scientific	15596018	
comme rical assay or kit	BCA Protein Assay Kit	Pierce	23225	
comme rical assay or kit	Direct-zol RNA Miniprep Kit	Zymo Research	R2051	
comme rical assay or kit	NEXTflex Rapid Directional qRNA-Seq Kit	Bioo Scientific	NOVA-5130- 02D	
comme rical assay or kit	NEXTflex Small RNA- Seq Kit v3	Bioo Scientific	NOVA-5132- 06	
comme rical assay or kit	Ribo-Zero rRNA Removal Kit	Illumina	MRZH11124, discontinued	
comme rical assay or kit	SuperScript III First- Strand Synthesis System	Thermo Fisher Scientific	18080051	
softwar e, algorith m	Code used for RNA-seq and Ribo-seq figure generation	This paper		The code used for the RNA-seq and Ribo-seq figure generation can be accessed via github at https://github.com/sjag

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softwar e, algorith m	GraphPad Prism	GraphPad Prism (https://graph pad.com)	RRID:SCR_0 02798	Version 9
softwar e, algorith m	ImageJ	ImageJ (https://imagej .nih.gov)	RRID:SCR_0 03070	