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Article

Upregulation of β-catenin due to loss of miR-139 contributes to motor neuron death in amyotrophic lateral sclerosis

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SUMMARY

Amyotrophic lateral sclerosis (ALS) is a fatal neurodegenerative disease characterized by the loss of motor neurons (MNs). There are no effective treatments and patients usually die within 2–5 years of diagnosis. Emerging commonalities between familial and sporadic cases of this complex multifactorial disorder include disruption to RNA processing and cytoplasmic inclusion bodies containing TDP-43 and/or FUS protein aggregates. Both TDP-43 and FUS have been implicated in RNA processing functions, including microRNA biogenesis, transcription, and splicing. In this study, we explore the misexpression of microRNAs in an iPSC-based disease model of FUS ALS. We identify the downregulation of miR-139, an MN-enriched microRNA, in FUS and sporadic ALS MN. We discover that miR-139 downregulation leads to the activation of canonical WNT signaling and demonstrate that the WNT transcriptional mediator β-catenin is a major driver of MN degeneration in ALS. Our results highlight the importance of homeostatic RNA networks in ALS.

INTRODUCTION

Amyotrophic lateral sclerosis (ALS) is a fatal neurodegenerative disease, characterized by the loss of upper and lower motor neurons (MNs) that affects 1–2/100,000 individuals worldwide (Xu et al., 2020). From clinical diagnosis, the median survival rate is 2–5 years (Brown and Al-Chalabi, 2017), with death commonly due to respiratory distress. To date, there are only two approved treatments: riluzole and edaravone, which add a few months to life (Jaiswal, 2019). Developing effective treatments has proved difficult due to the inherent complexity of ALS and unclear disease mechanisms.

Most ALS cases are not familial (fALS) and are termed sporadic (sALS). Many genes linked to fALS, including C9ORF72, TARDBP, and FUS, are involved in RNA processing (Butti and Patten, 2018), with the disruption of RNA processing seen in both fALS and sALS (Tank et al., 2018). Molecular hallmarks of fALS and sALS include the presence of TDP-43 and/or FUS cytoplasmic inclusion bodies within MN (Ikenaka et al., 2020; Deng et al., 2010). TDP-43 and FUS primarily function in RNA processing events, such as transcription, alternative splicing, and microRNA (miRNA) biogenesis (Butti and Patten, 2018). Significantly, RNA processing defects, such as intron retention (Luisier et al., 2018) and mRNA instability (Tank et al., 2018), have been observed in FUS or TDP-43 fALS as well as sALS MN, indicating that ALS models harboring these mutations may provide insights into the molecular mechanisms of both fALS and sALS.

Patient-derived induced pluripotent stem cells (iPSCs) offer an excellent opportunity to create human in vitro models of disorders like ALS that are otherwise difficult to study due to diseased cell type inaccessibility. These models provide a powerful platform to study pathogenesis in the context of the genetic background of a patient, without the need to overexpress the mutant proteins (Hawrot et al., 2020).

In this study, we deployed iPSC-based disease modeling with transcriptomic and phenotypic analysis to investigate molecular drivers of FUS ALS MN degeneration. Our results indicate that the downregulation of the MN-specific miRNA miR-139 contributes to the activation of canonical WNT signaling that drives MN degeneration. Importantly, these molecular defects are also observed in sALS MN, highlighting the shared dysregulation of RNA metabolism in fALS and sALS.

RESULTS

Patient-derived FUS MN display disease-associated phenotypes

We used iPSCs carrying the homozygous H517Q mutation in the FUS gene (H517Q/H517Q) and CRISPR-Cas9-edited
Figure 1. MN derived from FUS-ALS iPSC display disease-related phenotypes

80a, 79a indicate healthy iPSCs. H517Q/H517Q, H517Q/+ indicate ALS FUS iPSCs with a homozygous mutation or heterozygous correction of the mutation, respectively.

(A) Schematic depicting differentiation to MN from human iPSCs. See supplemental experimental procedures for further details. CHIR: CHIR09921, SB: SB431542, LDN: LDN193189.

(B) Representative image: 80a MN immunostained with ISL1, MAP2, and TUJ1 at day 34. Scale bar indicates 300 μm.

(C) 80a day 34 MN captured during ratiometric Fura-2 Ca²⁺ imaging. C1: Baseline level before stimulation. C2: Elevated Ca²⁺ during stimulation with 50 μM glutamate. Scale bar indicates 200 μm.

(D) Time course of MN response to glutamate. X axis indicates relative time in seconds. Y axis indicates the whole field average (WFA) 340:380 emission ratio for Fura-2, which is proportional to intracellular Ca²⁺ concentration.

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heterozygous iPSC (H517Q/+) derived from this line (Bhinge et al., 2017), as well as two healthy iPSC lines to evaluate disease-associated phenotypes in the FUS ALS MN. We deployed a robust protocol to differentiate all iPSCs into spinal MN (Figure 1A). Day 30 neurons expressed the mature MN markers ISL1, MAP2, NF-H, and TUJ1 (Figures 1B and 1E). In addition, our iPSC-derived MN demonstrated an increase in intracellular Ca2+ concentration in response to glutamate stimulation, indicating expression of glutamate receptors, a key feature of functionally mature MNs (Figures 1C and 1D). Across five independent differentiations, all four iPSC lines could be differentiated to ISL1+ MN at an efficiency of ~80% (Figures 1E and 1F), validating the reproducibility of our protocol.

We assessed MN survival by counting ISL1+ nuclei at days 30 and 45 for all of the genotypes. FUS MNs displayed >50% loss in vitro, while the healthy MNs displayed <15% loss (Figures 1G and S1A). FUS MN also displayed higher levels of apoptotic cells compared to healthy MN (Figure 1H). Heterozygous correction of the mutation rescued the survival deficit in FUS MN (Figures 1G, 1H, and S1A), indicating that the neuronal loss was due to the FUS H517Q mutation. However, the survival deficits in the H517Q/+ MN were significantly higher than in the healthy MNs (Figures 1G, 1H, and S1A). Thus, a single mutant FUS allele was enough to drive MN degeneration. Finally, morphometric analysis of our neuronal cultures revealed that, on average, FUS H517Q/H517Q MN had reduced neurite length compared to the healthy controls and H517Q/+ MN (Figures 1I and S1A). Overall, our in vitro iPSC model recapitulated essential characteristics of MN degeneration observed in ALS.

**FUS MN display widespread dysregulation of miRNA expression**

miRNAs are post-transcriptional regulators of gene expression that suppress the translation of their target mRNAs and can fine-tune signaling pathways (Ha and Kim, 2014). Dysregulation of miRNA expression has been observed in fALS and sALS (Emde et al., 2015; Figueroa-Romero et al., 2016; De Santis et al., 2017; Reichenstein et al., 2019; Tung et al., 2019; Kurita et al., 2020). Hence, we used small RNA sequencing (RNA-seq) to profile miRNA expression levels in our disease model. Principal-component analysis (PCA) and hierarchical clustering indicated that the major source of variation in our data was the underlying sample genotype (Figures 2A and S1B). Sample 79A R1 was an outlier and was excluded from further analysis. DESeq2 analysis identified 81 miRNAs in the FUS H517Q/H517Q MN and 38 miRNAs in the H517Q/+ MN as differentially regulated (adjusted p < 0.05, absolute log2 fold change >0.5) (Figure 2B).

Almost equal numbers of miRNAs were upregulated and downregulated (41 up versus 39 down, H517Q/H517Q versus healthy and 15 up versus 23 down, H517Q/+ versus healthy) in FUS ALS MN. These results contrast with miRNA profiling data obtained in sALS MNs, where most miRNAs were observed as downregulated (Emde et al., 2015). One reason for this discrepancy could be that the profiling was performed at different stages. We profiled miRNA expression levels at day 30 before the onset of degenerative phenotypes, while postmortem analysis represents the end stage of the disease. Likely, in the end stages of the disease, miRNA biogenesis is severely impaired, resulting in a global miRNA downregulation.

**MiR-139 is downregulated in FUS and sALS MN**

We compared our expression data with three miRNA datasets generated from sALS tissue, and our previous study in which we profiled miRNAs dysregulated in FUS P525L MNs (Figure 2C). To identify high-confidence miRNA candidates that are dysregulated in FUS and sALS, we only retained miRNAs that showed a significant expression change in at least three out of the six datasets analyzed. The overlap of differentially expressed miRNAs between ALS datasets was low, highlighting the well-known polygenic architecture of ALS (Figure 2C). MiR-139-5p, hereafter referred to as miR-139, was the only miRNA that was commonly downregulated across FALS and sALS datasets. Furthermore, miR-139 is downregulated in patient serum and postmortem spinal cord tissue (Raheja et al., 2018; Kurita et al., 2020). qRT-PCR analysis of our iPSC-derived MNs confirmed the downregulation of miR-139 in FUS
Figure 2. MiR-139-5p is downregulated in ALS MN

(A) Principal-component analysis (PCA) of 844 miRNAs in FUS H517Q/H517Q, FUS H517Q/+ and healthy MN harvested at day 30. N = 2 independent differentiations.

(B) Volcano plot of differentially expressed miRs in H517Q/H517Q versus healthy MNs (left panel) and H517Q/+ versus healthy MNs (right panel). MiR-139 is shown as a triangle (arrow).

(C) Heatmap of miRs identified as differentially expressed in at least 3 ALS datasets out of 6 analyzed. Red indicates upregulation, while blue indicates downregulation. Only miR-139 was commonly downregulated across all datasets (orange rectangle). sALS_spinalcord: postmortem cervical and thoracic spinal cord tissue from ALS patients versus healthy controls (Figueroa-Romero et al., 2016), sALS_LCM_MN: MNs enriched by laser capture microdissection of lumbar spinal cord tissue from ALS patients versus healthy controls (Emde et al., 2015). sALS_ventralhorn: lumbar ventral horn tissue of sporadic ALS versus non-degeneration controls (Reichenstein et al., 2019).

(D) Quantitative RT-PCR (qRT-PCR) confirmation of miR-139-5p downregulation in FUS H517Q MNs. N = 3–5 independent differentiations.

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MiR-139 is a regulator of WNT signaling

Downstream targets of a miRNA can reveal its functional role within a particular cell type. To determine the downstream targets of miR-139 in MNs, we overexpressed miR-139 in the H517Q/H517Q line and analyzed downstream changes in genome-wide gene expression using RNA-seq. PCA confirmed well-separated experimental conditions, indicating that miR-139 overexpression (OE) induced significant changes in gene expression (Figure 3A). We observed an enrichment of predicted miR-139 targets in the set of genes downregulated by miR-139 OE (Figure 3B). Furthermore, we observed a significant enrichment of miR-139 7mer seed sequences in the downregulated gene set, indicating that our analysis could detect direct miRNA targets (Figure 3C). Gene set enrichment analysis (GSEA) (Subramanian et al., 2005; Liberzon et al., 2011) identified targets of the WNT mediator β-catenin (CTNNB1) in the top 10 gene sets that were enriched in genes downregulated by miR-139 OE (Figure 3D). Analysis of miR-139 predicted targets revealed that several WNT mediators, including β-catenin, are potential direct targets of miR-139 (Figure 3E).

β-Catenin is the major effector/transcriptional mediator of the canonical WNT signaling pathway (Figure S2A). Previous studies have revealed that miR-139 inhibits WNT/β-catenin signaling in immortalized/cancer cells (Anton et al., 2011; Long et al., 2017; Gu et al., 2014). Hence, we explored β-catenin protein abundance and activity after miR-139 modulation in MNs. OE of miR-139 in FUS H517Q/H517Q MN resulted in a 40% reduction in β-catenin protein levels, while mRNA levels were reduced ~20% (Figures 3F, S2B, and S2C). In healthy 80a MNs, miR-139 OE resulted in >50% reduction in β-catenin protein levels (Figures 3F and S2B), indicating that miR-139 can inhibit canonical WNT signaling in MN. These data also reflect that miRNAs may only inhibit translation and not necessarily cause transcript degradation. To confirm that miR-139 targets β-catenin directly, we conducted a 3' UTR luciferase reporter assay in healthy MNs. Mutation of the predicted miR-139 binding site in the CTNNB1 3' UTR (identified by Targetscan) resulted in a >3-fold increase in luciferase signal compared to the non-mutated 3' UTR, indicating that β-catenin is a direct target of miR-139 in human MNs (Figure 3G).

Since miR-139 is downregulated in sALS, we hypothesized that WNT pathway genes targeted by miR-139 (miR-139-WNT gene set) would be upregulated in sALS datasets. We observed a significant enrichment of the miR-139-WNT gene set in genes upregulated in sALS MN (Figure 3K), suggesting that disruption of the miR-139-WNT axis may be a common theme across fALS and sALS.

WNT signaling is activated in FUS ALS MNs

As miR-139 is downregulated in ALS MNs and inhibits WNT signaling, we expected WNT signaling to be overactivated in our FUS H517Q MN lines. Accordingly, we found upregulation of several WNT ligands and WNT target genes in the FUS MN by qRT-PCR (Figure 4A). In addition, SuperTopFlash luciferase assays showed an increase in β-catenin activity (Figure 4B), while traditional western blot assays identified an increase in β-catenin protein levels (Figure 4C) in FUS MN compared to healthy MNs. Quantification of β-catenin protein levels, using the highly sensitive and quantitative in-cell western assay, confirmed a greater than 3-fold increase in β-catenin levels in the H517Q/H517Q MNs relative to healthy control MNs (Figures 4D and S3C).

To expand our observations beyond the H517Q FUS mutation, we evaluated WNT activity in our previously...
characterized FUS P525L iPSC MN (Figure S3A), alongside the appropriate isogenic control (De Santis et al., 2017). In-cell western assays confirmed significant upregulation of β-catenin protein in FUS P525L MNs (Figures 4D and S3C), similar to observations in the FUS HS17Q ALS MNs.

Inhibition of canonical WNT signaling improves MN survival

We hypothesized that inhibiting WNT signaling should improve FUS ALS MN survival. We also targeted pathways known to synergistically interact with WNT signaling and activated in ALS, including mitogen-activated protein (MAP) kinase family members p38 (MAPK14), ERK1 and -2, and JNK1, -2, and -3, as well as TP53 (Bhinge et al., 2017; Sama et al., 2017; Gibbs et al., 2018; Cervenka et al., 2011; Damalas et al., 1999). Furthermore, we treated cells with a pan-cell-cycle-dependent kinase (CDK) inhibitor as MN death could be due to WNT-driven activation of the cell cycle (Bryja et al., 2017; Frade and Ovejero-Benito, 2015). We used FUS HS17Q/HS17Q MN as these neurons displayed greater survival deficits than the FUS HS17Q/+ MN. FUS ALS MN were treated with small molecule inhibitors of the selected pathways at the indicated concentrations at day 30 and treatment was continued until day 45 (Figures 4E and S4A).

Figure 3. MiR-139-5p targets WNT signaling in FUS ALS MNs

(A) PCA using the top 500 variable genes from miR-139 overexpression (OE) in FUS HS17Q MNs. RNA collected at day 35.

(B) Cumulative distribution function (CDF) plot comparing the fold changes of predicted miR-139 targets (blue curve) relative to all genes (black curve) in response to miR-139 OE. p values were estimated using the Kolmogorov-Smirnov test.

(C) Enrichment of 7mers starting at positions 1–15 of miR-139-5p in the 3' UTR of genes downregulated upon miR-139-5p OE. The miRNA sequence is shown at top for reference. p values were estimated using a hypergeometric distribution and corrected for multiple hypothesis testing using the Benjamini-Hochberg method. A p value threshold of 0.01 is indicated by the vertical dotted line.

(D) Left panel: GSEA analysis of miR-139 OE using curated gene sets from MSigDB. X axis displays the log transformed p values. Genes downregulated upon CTNNB1 deletion in intestinal epithelial cells (FEVR_CTNNB1_TARGETS_DN) were identified in the top 10 gene sets with a significant negative enrichment score (i.e., these gene sets were downregulated after miR-139-5p OE). Right panel: GSEA enrichment plot for the gene set FEVR_CTNNB1_TARGETS_DN. X axis shows genes sorted from most upregulated (left-hand side) to most downregulated (right-hand side). Enrichment p values for the gene sets were estimated using 1,000 random simulations and adjusted using the Benjamini-Hochberg method.

(E) MiR-139-5p is predicted to directly target several WNT-associated genes, including the WNT transcriptional mediator CTNNB1. Predicted targets were obtained from 3 databases: miRWalk, Targetscan, and miRDB (Slichter et al., 2018; Agarwal et al., 2015; Wong and Wang, 2015). Score indicates the number of databases supporting each interaction.

(F) In-cell western blot analysis to measure β-catenin protein levels in response to miR-139-5p overexpression (OE) in FUS HS17Q/HS17Q ALS (N = 5) and healthy 80A MNs (N = 3).

(G) Relative luminescence of the Renilla luciferase gene, carrying the CTNNB1 3' UTR with the predicted miR-139 binding site (wild type [WT]) or with the mutated site (mutant). Mutation of the miRNA binding site rescues luciferase activity when transfected in human MNs. N = 5.

(H) SuperTopFlash dual luciferase assay to measure β-catenin activity upon inhibition of miR-139 in SH-SYSY cells. N = 3.

(I) qRT-PCR analysis of miR-139-5p expression upon activation of WNT signaling using CHIR09921 in SH-SYSY cells. N = 3.

(J) Schematic of proposed negative feedback loop between miR-139 and the WNT pathway.

(K) CDF plot of miR-139-5p WNT targets (blue) and all expressed genes (black) in differentially expressed genes in laser capture microdissected sALS MN relative to healthy controls. GSE18920: MNs were derived from lumbar spinal cords (12 ALS and 10 controls) and gene expression analyzed using microarrays. GSE76220: MN were derived from lumbar spinal cords (13 ALS and 8 controls) and gene expression analyzed by RNA-seq. The p value was calculated using the Kolmogorov-Smirnov test. N: number of independent differentiations. Error bars indicate SEMs. *p < 0.05, **p < 0.01, ***p < 0.001 by Student's t test, unless otherwise stated.
A

B

C

D

E

F

G

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the upregulation of WNT alone is enough to drive degeneration in iPSC-derived MNs.

To investigate the relevance of the identified pathways to ALS, we performed GSEA on the sALS MN datasets using gene sets related to WNT/β-catenin or p38/MAPK perturbation. Enrichment analysis showed significant upregulation of several WNT and p38/MAPK gene sets in sALS MNs (Figure 4G). Overall, our results indicate that the hyperactivation of WNT signaling, at least partly due to the downregulation of miR-139, drives MN degeneration in both fALS and sALS MN. Interestingly, miR-139 inhibition has been shown to activate p38/MAPK signaling in colon tumor tissue (Zou et al., 2016). It is likely that the downregulation of miR-139-5p also leads to activation of p38/MAPK in MNs.

**Mutant FUS downregulates miR-139 potentially by a gain-of-function mechanism**

FUS and TDP-43 are involved in different stages of the miRNA biogenesis pathway (Butti and Patten, 2018). To assess whether FUS influences miR-139 through a loss of function, we depleted FUS in SH-SYSY cells and measured miR-139 expression by qRT-PCR (Figures 5A and 5B). Although we could deplete FUS by 85% using small interfering RNA (siRNA) (Figure 5A), the observed downregulation of miR-139 was less than 20% (Figure 5B). Depletion of FUS in healthy MN progenitors for 3 days failed to produce any change in miR-139 levels (Figure 5B). This suggested that miR-139 downregulation was not due to a loss of function of FUS, supporting recent work indicating that FUS-associated ALS is due to toxic gain of function (Korobeynikov et al., 2022; Sharma et al., 2016). As FUS has been shown to regulate RNA polymerase II (RNAPII)-mediated transcription (Schwartz et al., 2012), we hypothesized that mutant FUS may affect miR-139 at the transcriptional level.

We quantified primary miRNA (pri-miRNA) levels from RNA-seq data by measuring intronic reads mapping to host genes (supplemental experimental procedures) (Gaidatzis et al., 2015). We tested the feasibility of this approach using our RNA-seq data for the FUS P525L MN. Here, we had previously analyzed expression levels of protein-coding genes and miRNAs in parallel (De Santis et al., 2017). We used the MIRIAD database (Hinske et al., 2017) to identify intronic miRNAs and their corresponding host genes expressed in the FUS P525L or isogenic MN (Figure 5C). Next, we plotted fold changes in the mature miRNAs expression against the fold changes of the intronic read counts of their host genes. We identified a modest but significant correlation between the miRNAs and their host gene intron expression (Figure 5D), suggesting that the expression level of mature miRNAs was largely due to transcriptional control. Example exceptions to this statement are indicated in Figure 5D by a blue asterisk.

Our analysis showed expression changes in the mature as well as the pri-miRNA transcript of miR-139 (Figure 5D, arrow). MiR-139 is located within an intronic region of the PDE2A gene (Kent et al., 2002). Analysis of intronic reads mapping to PDE2A showed that pri-miR-139 transcript levels were reduced by 30% in the FUS P525L MN compared to the isogenic controls (Figure 5E). This was similar to the 35% reduction seen for the mature miR-139 in the FUS P525L MN (Figure 5D). Intrinsic miRNAs 185, 22, and 26a, which were identified as unchanged in our iPSC MN model did not display any changes in their pri-miRNA levels, as expected (Figure 5E).

Next, we conducted deep RNA-seq on ribosomal depleted total RNA for the FUS H517Q/H517Q and healthy 80A MNs to obtain better coverage of reads mapping to intronic regions of the genome. As observed in the P525L dataset, we identified a significant correlation between changes in expression of the mature miRNAs and their corresponding

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**Figure 4. ALS MNs display increased WNT**

(A) qRT-PCR analysis of mRNA expression of WNT genes and ligands in FUS ALS MNs compared to healthy controls (79a and 80a). N = 3. (B) SuperTopFlash luciferase assay showing relative luciferase units for β-catenin activity in H517Q/H517Q and H517Q/+ MN relative to 80a and 79a healthy controls. N = 3–4. (C) Representative western blot (left panel) and quantification (right panel) of active β-catenin protein in healthy and FUS ALS MNs. β-catenin normalized to α-tubulin and shown relative to healthy controls. N = 3. (D) In-cell protein quantification of β-catenin in FUS H517Q/H517Q MN relative to healthy control 80a (N = 5) and FUS P525L MN relative to isogenic control (N = 3). (E) Quantification of ISL1+ FUS H517Q/H517Q MNs at day 45 relative to day 30 after treatment with small-molecule inhibitors of the indicated pathways. Horizontal dotted line indicates MN loss in the DMSO-only control. N = 3, p values were estimated using Student’s 2-tailed t test. (F) Caspase-3/7 activity relative to viable cells in response to overexpression of β-catenin versus GFP control in day 23 healthy (80A) MNs. N = 3. (G) GSEA on sALS MN datasets using gene sets relevant to WNT or p38/MAPK signaling. Blue bars indicate p38/MAPK gene sets. Orange bars indicate WNT/CTNNB1 gene sets. p values were estimated using 1,000 random simulations and adjusted using the Benjamini-Hochberg procedure. N: number of independent differentiations. Error bars indicate SEMs. p values were estimated by Student’s 1-tailed t test unless otherwise indicated. *p < 0.05, **p < 0.01, ***p < 0.001.
Figure 5. Mutant FUS transcriptionally downregulates miR-139 potentially through a gain-of-function mechanism

(A) Relative FUS gene expression after siRNA-mediated knockdown in SH-SY5Y cells. Fold changes were normalized to scrambled (negative control) siRNA. N = 3 independent transfections. p values were estimated using Student’s 1-tailed t test.

(B) qRT-PCR analysis of miR-139-5p expression in response to siRNA-mediated FUS knockdown in SH-SY5Y cells and MN progenitors derived from healthy (80a) iPSC. Gray bars: scrambled siRNA used as a negative control; blue bars: siRNA against FUS. N = 3. Fold changes were normalized to scrambled siRNA.

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host genes (Figure 5F). We observed a 60% reduction in intronic reads of PDE2A in FUS H517Q/H517Q MN (Figure 5G), with the reduction proportional to that seen for the mature miR (Figures 2B and 5F). Our results indicate that the observed changes in mature miR-139 levels are at least partly due to the corresponding downregulation of the primary miR-139 transcript.

These results demonstrate the potential of using deep RNA-seq data to predict transcriptional changes in pri-miRNAs and indicate that mutant FUS may impede transcription of the pri-miR-139 transcript. However, when we performed a similar analysis on RNA-seq data obtained from sALS MNs (GSE76220), we did not observe any decrease in the intronic reads of PDE2A. Our analysis suggests that downregulation of pri-miR-139 transcription could be specific to FUS mutations only.

**DISCUSSION**

It is unclear whether the widespread downregulation of miRNAs observed in ALS MN is an effect of dying neurons or if it contributes to neuronal death. In addition, the differential susceptibility of MN in ALS may be due to genes that are differentially active in MNs, but are disrupted in disease. We observed that miR-139 was activated during MN differentiation, is highly expressed and enriched in mature MN, and was downregulated in both mutant FUS and sALS MN. This suggested that the downregulation of miR-139 in MNs can lead to the disruption of homeostatic mechanisms. We found that miR-139 inhibits the canonical WNT pathway in human MNs through the direct targeting of β-catenin, and is reciprocally activated by an increase in WNT signaling. This negative feedback loop between miR-139 and WNT may be required to keep WNT activity within a desired range in MNs.

Dysregulation of WNT, either overactivation or inhibition, has been implicated in causing neurodegeneration in adults. Decrease in canonical WNT signaling has been linked to Alzheimer’s disease (AD), in which inhibition of canonical WNT and activation of the non-canonical WNT/PCP leads to synaptic disassembly (Purro et al., 2014; Sellers et al., 2018). However, the upregulation of WNT pathway components has been associated with Huntington’s disease (HD) and ALS. For example, in Drosophila neurons, expression of mutant huntingtin protein stabilizes β-catenin and leads to neuronal death, which can be reversed by the RNAi-mediated knockdown of β-catenin (Godin et al., 2010). In ALS, increased levels of WNT ligands, receptors, and downstream targets have been identified in human postmortem tissue (spinal cord, anterior horn) and SOD1G93A transgenic mouse models (spinal cord) (Yu et al., 2013; Chen et al., 2012a, 2012b; González-Fernández et al., 2019).

ALS-associated mutant Ubiquilin 4 (UBQLN4) has been linked to the upregulation of β-catenin when expressed in mouse spinal MN in vitro and zebrafish MN in vivo (Edens et al., 2017). Pharmacological inhibition of the nuclear localization of β-catenin rescued axonal defects observed in these models (Edens et al., 2017). β-Catenin was also identified as a key driver of neurodegeneration in animal models of spinal muscular atrophy (SMA), an MN disease affecting infants and children (Wishart et al., 2014). These observations indicate that the upregulation of WNT/β-catenin signaling can be detrimental to MN survival and homeostasis. WNT has been shown to be upregulated through autocrine mechanisms in cancer cell models (Bafico et al., 2004; Schlange et al., 2007), as well as neuronal cell populations (Wexler et al., 2009; Ryu et al., 2013). Our data suggest that miR-139 may also regulate WNT signaling through targeting WNT ligands, in addition to inhibiting β-catenin, to break the autocrine loop. Overall, our results indicate that WNT activity is tightly controlled in human MN to
maintain homeostasis and prevent neurodegeneration. Expression of miR-139 is one of the mechanisms involved in this regulation. It is unlikely that miR-139 is the sole miRNA responsible for regulating WNT activity in MN. Indeed, other miRNAs implicated in regulating WNT signaling have been observed to be downregulated in ALS MNs. For example, miR-375 targets the canonical WNT pathway and is downregulated in FUS P525L MN (Wang et al., 2013; De Santis et al., 2017). MiR-218 is the most significantly downregulated miRNA in sALS MNs and has been shown to inhibit as well as promote WNT signaling (Reichenstein et al., 2019; Tu et al., 2013; Anton et al., 2011; Zhao et al., 2019). Consequently, WNT signaling is under the control of multiple miRNAs, and the upregulation of WNT activity in ALS MNs is likely due to the dysregulation of multiple WNT-regulating miRNAs.

How does WNT hyperactivation lead to MN death? WNT signaling is typically implicated in cell growth, proliferation, and specification (Clevers, 2006). Previous studies have observed an upregulation in the cell cycle in ALS MNs and inhibition of CDKs improve disease phenotypes (Bhinge et al., 2017; Rojas-Prats et al., 2021; Ranganathan and Bowser, 2010). An attractive possibility is that cell-cycle reactivation through aberrant WNT signaling contributes to MN death (Trade and Ovejero-Benito, 2015; Bryja et al., 2017). However, inhibition of the cell cycle by a pan-CDK inhibitor led to a modest increase in MN survival and was toxic at higher doses. It is possible that some CDKs have a critical cell-cycle-independent role to play in MN homeostasis. In this case, targeting specific CDKs may improve ALS MN survival as observed in SMA (Hor et al., 2018; Ling et al., 2010). Hence, mislocalization of these RNA-binding proteins may cause defects in miRNA biogenesis. Our findings raise the possibility that FUS may also affect the transcription of ALS-associated miRNAs. We observed a higher absolute fold change for the mature miR-139 expression in comparison to the intronic PDE2A levels, for both H517Q and P525L datasets. Therefore, in FUS-ALS, the downregulation of miR-139 could be due to a combination of reduced pri-miR-139 transcription and downstream miRNA processing defects. However, in sALS, we could not detect significant changes in the miR-139 pri-miRNA, suggesting that defects in miRNA processing are likely to be the primary cause of miR-139 downregulation.

Overall, our results highlight the importance of RNA processing in ALS and demonstrate the knock-on effect a single miRNA can have on transcriptomic equilibrium that can disrupt MN homeostasis.

**EXPERIMENTAL PROCEDURES**

**miRNA overexpression**

Adeno-associated viruses (AAVs) expressing miR-139 or small hairpin RNA (shRNA) against OCT4 were generated using the AAV-pro Helper Free System (Takara). OCT4 is not expressed in MN; hence, this shRNA was used as a non-targeting control. The mature miR-139-5p sequence was expressed using a shRNA design to ensure the expression of miR-139-5p and not miR-139-3p (Brummelkamp et al., 2002; Qu et al., 2012) (Figure S5). MNs were transfected at day 23 and collected for RNA at day 34.

**miRNA inhibition**

SH-SY5Y cells were plated in 96-well optically clear white tissue culture plates (Greiner) 24 h before transfection with a negative or miR-139-5p inhibitor (IDT) using lipofectamine STEM reagent (Thermo Fisher Scientific). Inhibitors were used at a final concentration of 20 nM. For luciferase assay, inhibitors were co-transfected with 50 ng STF19 plasmid (Psichek2, Promega) modified to contain 19 binding sites for β-catenin/TCF upstream of the firefly luciferase gene, and 5 ng Renilla luciferase plasmid (pRLcmv, Promega) as an internal control. Media was replaced 24 h post-transfection and the luciferase assay completed 72 h post-transfection as per protocol (Dual-Glo Luciferase Assay System [Promega]). Readings for Firefly and Renilla luciferase were taken on the CLARIOstar Plus microplate reader (BMG Labtech).

**SuperTopFlash luciferase assays**

Day 30 MNs were plated in 96-well optically clear white tissue culture plates (Greiner). Cells were transfected with 700 ng STF19 plasmid (Psichek2, Promega) and 100 ng Renilla luciferase plasmid (pRLcmv, Promega). Cells were transfected using a calcium-phosphate method modified from Jiang and Cheng (2006). A final concentration of 12.4 mM CaCl₂ was used per well, and transfection mixes were 1/10 of the volume covering cells. Plasmid and CaCl₂ master mixes were mixed gradually into Hank’s balanced salt solution (HBSS) to prevent the formation of large transfection complexes, and incubated for 20 min at room temperature before addition to wells. Cells were incubated with transfection mixes at 37°C, 5% CO₂ for 4.5 h, then washed once with 300 mM sodium acetate dissolved in DMEM:F12 medium for 5 min, 37°C to dissolve transfection complexes. Cultures were maintained in neuronal differentiation media supplemented with brain-derived neurotrophic factor (BDNF) and glial cell line-derived neurotrophic factor (GDNF) for a further 72 h, then lysed according to the Dual-Glo Luciferase Assay System (Promega).

**UTR luciferase assay**

The miR-139 binding site in the 3’ UTR of the CTNNB1 transcript was identified using Targetscan (Agarwal et al., 2015) and extended by 100 bp either side of the predicted site. The resultant sequence
was cloned downstream of the Renilla luciferase gene in the PSICheck2 vector (Promega C8021). To mutate the miR-139 binding site, we deleted positions 2, 4, and 6 of the predicted miRNA binding site. Day 30 healthy MNs (80A) were transfected using the same method described in MN SuperTopFlash luciferase assays. MNs were collected 72 h post-transfection and processed using the Dual-Glo Luciferase Assay System (Promega).

**β-catenin overexpression**

AAVs expressing β-catenin-EGFP or EGFP only were generated using the AAVpro Helper Free System (Takara). β-Catenin-EGFP plasmid was a gift from Alpha Yap (Addgene plasmid #71367) (Han et al., 2014). Healthy MN progenitor cells were transduced at day 9 before notch inhibition. MNs were matured for 2 weeks before harvesting at day 23 for caspase/cell titer assays. Longer cultures were not viable due to cell death caused by β-catenin overexpression.

**Data and code availability**
The accession number for the sequencing data reported in this paper is GEO: GSE203173.

**SUPPLEMENTAL INFORMATION**
Supplemental information can be found online at https://doi.org/10.1016/j.stemcr.2017.02.019.

**AUTHOR CONTRIBUTIONS**
S.H., S.C.N., A.T., C.B., and A.B. designed and performed the experiments and analyzed the data. S.H. analyzed the RNA-seq and microarray data. C.F. and N.S.D. helped with imaging. A. Randall performed the Ca²⁺ imaging assays. A. Rosa provided the FUS P525L iPSC lines. A.B. and L.W.S. funded the project. A.B. conceptualized the project. S.H. analyzed the RNA-seq and miRNA-seq data, and analyzed the data. S.H., S.C.N., A.T., C.B., and A.B. designed and performed the experiments. All of the authors contributed toward editing the manuscript.

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**CONFLICTS OF INTERESTS**
The authors declare no competing interests.

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