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Defects in Synapse Structure and Function Precede Motor Neuron Degeneration in Drosophila Models of FUS-Related ALS

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Amyotrophic lateral sclerosis (ALS) is an adult-onset neurodegenerative disease that leads invariably to fatal paralysis associated with motor neuron degeneration and muscular atrophy. One gene associated with ALS encodes the DNA/RNA-binding protein Fused in Sarcoma (FUS). There now exist two Drosophila models of ALS. In one, human FUS with ALS-causing mutations is expressed in fly motor neurons; in the other, the gene cabeça (caz), the fly homolog of FUS, is ablated. These FUS-ALS flies exhibit larval locomotor defects indicative of neuromuscular dysfunction and early death. The locus and site of initiation of this neuromuscular dysfunction remain unclear. We show here that in FUS-ALS flies, motor neuron cell bodies fire action potentials that propagate along the axon and voltage-dependent inward and outward currents in the cell bodies are indistinguishable in wild-type and FUS-ALS motor neurons. In marked contrast, the amplitude of synaptic currents evoked in the postsynaptic muscle cell is decreased by >80% in FUS-ALS larvae. Furthermore, the frequency but not unitary amplitude of spontaneous miniature synaptic currents is decreased dramatically in FUS-ALS flies, consistent with a change in quantal content but not quantal size. Although standard confocal microscopic analysis of the larval neuromuscular junction reveals no gross abnormalities, superresolution stimulated emission depletion (STED) microscopy demonstrates that the presynaptic active zone protein bruchpilot is aberrantly organized in FUS-ALS larvae. The results are consistent with the idea that defects in presynaptic terminal structure and function precede, and may contribute to, the later motor neuron degeneration that is characteristic of ALS.

Introduction

Communication between motor neurons and their target muscles is compromised in amyotrophic lateral sclerosis (ALS). Although overt symptoms in humans and mouse models are associated with muscle atrophy after muscle denervation (Ropper and Brown, 2005), the primary defects that precipitate this loss of neuromuscular connectivity remain poorly understood. Most of what we know about the degradation of the neuromuscular synapse in ALS comes from pathological studies in mouse models of superoxide dismutase 1 (SOD1)-mediated ALS. Analysis of disease progression in these mice suggests that ALS starts in the terminal axon, leading to muscle dysfunction and denervation, and proceeds in a retrograde pattern, with degeneration of the proximal motor axon and eventual loss of neuronal cell bodies (Fischer et al., 2004; Dupuis and Loeffler, 2009). This implies that a local defect in the distal portion of the axon and/or the presynaptic terminal is a primary pathogenic event in disease progression. However, little is known about intrinsic changes in mammalian motor neuron excitability and synaptic transmission in ALS, in part because of the challenges of performing electrophysiological recordings in vivo.

Mutations in the DNA/RNA-binding protein Fused in Sarcoma (FUS), as well as in other DNA/RNA-binding proteins such as TDP43, have been implicated in both familial and sporadic cases of ALS (Sreedharan et al., 2008; Kwiatkowski et al., 2009; Vance et al., 2009). Recently, models of both FUS-related and TDP43-related ALS have been established both in the fruit fly Drosophila (Lanson et al., 2011; Wang et al., 2011; Miguel et al., 2012; Sasayama et al., 2012; Xia et al., 2012) and in zebrafish (Kabashi et al., 2010; Armstrong and Drapeau, 2013a; Armstrong and Drapeau, 2013b). Expression of human disease-causing mutants of either FUS or TDP43 in flies or fish leads to impaired locomotor activity and early death (Kabashi et al., 2010; Lanson et al., 2011). Flies that lack the gene cabeça (caz), the fly homolog of FUS, exhibit similar phenotypes that can be rescued by the expression of wild-type but not mutant human FUS (Wang et al., 2011), which is consistent with the idea that FUS-mediated ALS involves both loss of nuclear function and gain of toxic function.
from accumulation of mislocalized cytoplasmic mutant FUS. In the fly, because cell-type-specific drivers are used to restrict the expression of mutant human FUS to motor neurons, it is evident that the locomotor and life span phenotypes must arise from defects in the presynaptic cell. Morphological and electrophysiological analysis of the fish neuromuscular junction also suggests that presynaptic changes accompany the disease phenotype (Armstrong and Drapeau, 2013a).

We have examined motor neuron cell body excitability, axonal conduction, synaptic transmission, and synapse structure in mutant FUS-expressing and cabeza-null flies, and report here that neuronal excitability and action potential propagation are essentially normal at the same time that neurotransmitter release is severely impaired and presynaptic active zone structure is aberrant. The results suggest that defects in presynaptic structure and function are early and are perhaps precipitating events in the pathogenesis of FUS-related ALS.
**Materials and Methods**

*Fly genetics.* Flies expressing either wild-type or R521C mutant human FUS as UAS transgenes (Lanson et al., 2011) were crossed with the motor-neuron-specific driver fly line OK371-GAL4 to drive expression of FUS exclusively in motor neurons. *caz*] flies that are null for the gene *caz* (Wang et al., 2011) were generously provided by Dr. Brian McCabe of Columbia University. For some experiments, FUS-expressing flies that express UAS-GFP under the control of the OK371-GAL4 driver were used to identify fluorescent motor neuron cell bodies for electrophysiological recordings. The flies were raised on standard fly food and genetic crosses were performed at 25°C.

*Electrophysiological recording from larval motor neuron cell bodies.* All experiments were performed on wandering late-third-instar larvae of either sex. Larvae were pinned dorsal side up and bathed in Ca²⁺-free solution containing the following (in mM): 70 NaCl, 5 KCl, 4 MgCl₂, 10 NaHCO₃, 5 trehalose, 115 sucrose, and 1 HEPES, pH 7.2. The preparation was visualized using an upright Olympus BX51WI microscope. To access motor neuron cell bodies in the ventral ganglion, Protease 14 (2 mg/ml extracellular solution; Sigma-Aldrich) was focally applied to the ganglionic sheath by applying positive pressure to a recording electrode with the tip broken to a diameter of ~10 μm. Treatment of the sheath was performed with constant laminar perfusion and the debris was removed by applying negative pressure to the electrode. Motor neuron cell bodies were visualized though the expression of a UAS-GFP transgene under the control of the OK371-GAL4 driver. For consistency, we restricted our recordings to the aCC subset of motor neurons identified separately with the RRA-GAL4 driver (Fujioka et al., 2003). The extracellular recording solution contained the following (in mM): 101 NaCl, 3 KCl, 1.8 CaCl₂, 2 MgCl₂, 1.25 Na₂HPO₄, 5 glucose, and 20.7 NaHCO₃, pH 7.2. Thin-walled borosilicate electrodes were pulled to a resistance of 4–6 MΩ when filled with intracellular solution containing the following (in mM): 102 K-gluconate, 17 NaCl, 2 CaCl₂, 5 EGTA, 0.5 MgCl₂, and 10 HEPES, pH 7.2. Standard techniques were used to record macroscopic currents in the whole-cell voltage-clamp mode and voltage in current-clamp mode with an Axopatch 200A amplifier (Molecular Devices). Data were digitized with a Digidata 1322A interface (Molecular Devices) and
stored on a PC hard drive for further analysis with pClamp10 software (Molecular Devices).

**Synaptic transmission at the larval neuromuscular junction.** Larvae were dissected and recorded in hemolymph-like saline HL3.1 containing the following (in mM): 70 NaCl, 5 KCl, 1 CaCl₂, 4 MgCl₂, 10 NaHCO₃, 5 trehalose, 115 sucrose, and 5 HEPES, pH 7.2, as described previously (Broadie and Bate, 1993; Feng et al., 2004; Ueda and Wu, 2006). The segmental nerves were severed from the ventral ganglion and stimulated with a suction electrode. Recordings were performed on ventral longitudinal muscle 6 in abdominal segments A3–A5 of third instar larvae as described previously (Broadie and Bate, 1993). All cells selected for recording had resting membrane potentials between −50 and −70 mV. Both spontaneous and evoked postsynaptic currents were recorded while the muscle cell was voltage clamped at −60 mV using an AxoClamp 2A (Molecular Devices) amplifier in single-electrode voltage-clamp mode (switching frequency 10 kHz) and sharp microelectrodes (catalog #TW150F-3; WPI) filled with 3 M KCl (5–10 M)

**Fig. 5.** Kinetic analysis of the evoked synaptic currents. A. Five consecutive evoked EJCs from wild-type-FUS (top), R521C-FUS (second), and caz⁻¹ (third) larvae are shown. Bottom: EJCs are averaged, normalized to the peak amplitudes, and superimposed. B. Averaged single exponential time constants fit to the decay phase of the EJC in WT-FUS, R521C-FUS, and caz⁻¹ flies. Both R521C-FUS and caz⁻¹ decay time constants are significantly smaller than in wild-type FUS larvae (p < 0.0001). C. Averaged integrated synaptic currents in WT-FUS, R521C-FUS, and caz⁻¹ flies. Integrated synaptic currents were measured by calculating the total area under the EJC. Error bars indicate SEM. Both R521C-FUS and caz⁻¹ integrated synaptic currents are significantly smaller than in wild-type FUS larvae (p < 0.0001).

Results

**Electrical excitability of larval motor neuron cell bodies and axons is not impaired in R521C-FUS-expressing flies**

To determine the effects of the expression of mutant human FUS on larval motor neuron cell body excitability, we recorded action potential waveforms under current-clamp and voltage-dependent inward and outward currents under voltage clamp. Overshooting action potentials can be evoked in neurons expressing either wild-type (Fig. 1A) or R521C mutant human FUS (Fig. 1B) by the injection of depolarizing current via the recording electrode (Fig. 1B, inset). The neurons of both genotypes fire repetitively in response to a sustained stimulus, with no apparent accommodation. Interestingly, in the R521C-FUS-expressing motor neurons (Fig. 1B), more action potentials are evoked by the same depolarizing stimulus at all but the lowest stimulus level examined (Fig. 1C). This increase in evoked action potential frequency is not accompanied by a significant change in membrane input resistance or time constant (data not shown). A similar increase in excitability with no change in input resistance has been reported in zebrafish motor neurons that express mutant human TDP43 (Armstrong and Drapeau, 2013a) or FUS (Armstrong and Drapeau,
similar decrease is seen in the amplitude of the EJC, and a dramatic decrease in the motor neurons leads to a severe impairment in locomotor activity in R521C-FUS-expressing larvae (Lanson et al., 2011; Wang et al., 2011) does not result from an inability of motor neuron cell bodies to generate inward and outward currents or fire action potentials robustly.

To determine whether the action potentials generated in the motor neuron somata can propagate along the axon, we recorded with a suction electrode from the severed end of the segmental nerve that remains attached to the central ganglion. Spontaneous action potentials can be recorded (Fig. 3A), which presumably arise in and propagate from the cell bodies. In contrast to the evoked cell body action potentials in Figure 1, there is no statistically significant difference in the frequency of the spontaneous orthodromic action potentials recorded from the axons of wild-type FUS and R521C-FUS-expressing larvae (Fig. 3B).

**Figure 1.** Representative current traces from WT-FUS (top, black), R521C-FUS (middle, red), and caz1 (bottom, blue) flies. The muscle cell was voltage clamped at −60 mV as for Figure 4, but no stimulus was delivered to the segmental nerve. A, Representative current traces from WT-FUS (top, black), R521C-FUS (middle, red), and caz1 (bottom, blue) flies. B, Individual mEJC identified by the arrows in A are shown at higher time and amplitude resolution. n = 20 for each genotype.

**Figure 6.** Spontaneous mEJCs in WT-FUS, R521C-FUS, and caz1 flies. The muscle cell was voltage clamped at −60 mV as for Figure 4, but no stimulus was delivered to the segmental nerve. A, Representative current traces from WT-FUS (top, black), R521C-FUS (middle, red), and caz1 (bottom, blue) flies. B, Individual mEJC identified by the arrows in A are shown at higher time and amplitude resolution. n = 20 for each genotype.

**Figure 7.** Distributions of mEJC frequencies (left) and amplitudes (right) in WT-FUS (A), R521C-FUS (B), and caz1 (C) flies. Data are combined from six experiments for each genotype. The instantaneous frequency was calculated as the reciprocal of the interval between individual mEJCs (Benton and Dahanukar, 2011). Bin sizes are 0.2 Hz and 20 pA, respectively. The first 15 bins in each amplitude histogram were fit with a Gaussian function (solid lines).

**Synaptic transmission at the larval neuromuscular junction is severely impaired in R521C-FUS-expressing and caz1-null flies**

We next examined synaptic transmission at the larval neuromuscular junction by voltage clamping the postsynaptic muscle cell and stimulating the presynaptic segmental nerve. In larvae expressing only the GAL4 driver but no FUS transgene, robust excitatory junctional currents (EJCs) can be evoked by nerve stimulation, and presynaptic expression of wild-type FUS does not influence the amplitude of the EJC (Fig. 4A,B). In contrast, expression of R521C-FUS in the motor neurons leads to a dramatic decrease in the amplitude of the EJC, and a similar decrease is seen in caz1-null flies (Fig. 4A,B). No such decrease in EJC amplitude is observed in R521C-FUS transgenic larvae in the absence of the GAL4 driver (data not shown), indicating that the synaptic defect is associated with the expression of the mutant protein.

Interestingly, the decay of the EJC is much faster in R521C-expressing and in caz1 larvae (Fig. 5A), with decay time constants that are ∼35% of those in the wild-type FUS larvae (Fig. 5B). This substantial change in the decay kinetics, in combination with the decrease in EJC amplitude, results in an even more profound decrease in the integrated synaptic current (Fig. 5C).

To further extend the analysis of synaptic transmission, we recorded spontaneous miniature EJCs (mEJCs) from the different fly genotypes. mEJC frequency is used widely as a reliable reporter of presynaptic neurotransmitter release, which often is
Figure 6. Imaging of presynaptic boutons and active zones at the neuromuscular junction in larvae of the indicated genotypes. A, Confocal imaging of NMJ boutons stained with an antibody that recognizes the active zone marker bruchpilot indicates comparable morphology in wild-type and R521C-FUS-expressing flies. Scale bar, 5 μm. B, Analysis of bruchpilot staining at active zones by STED microscopy reveals that the proportion of active zones (AZs) exhibiting a central hole is reduced in R521C-FUS-expressing flies. Error bars indicate SEM. Statistical significance was evaluated by unpaired Student’s t-test (p < 0.01). C–D, Representative STED images of bruchpilot staining at active zones in wild-type and R521C-FUS-expressing flies, respectively. Insets 1, Higher-magnification images of “doughnut-shaped” active zones. Insets 2, Higher-magnification images of active zones without a distinguishable central hole. Scale bars, 0.5 μm.

difficult to measure directly. As shown in the traces in Figure 6A, fewer spontaneous mEJCs are observed in R521C-FUS-expressing and caz null larvae compared with larvae that express wild-type FUS. However, an examination of individual mEJCs at higher time and amplitude resolution (Fig. 6B) suggests that there is no difference among genotypes in the amplitude or kinetics of the unitary mEJC. These conclusions are reinforced by the frequency and amplitude histograms shown in Figure 7. The frequency histograms (Fig. 7, left) demonstrate that the distribution of mEJCs is shifted markedly to the lower-frequency range in both R521C-FUS (Fig. 7B) and caz null (Fig. 7C) flies. Interestingly, although an inspection of the low-time-resolution current traces in Figure 6A suggests that mEJC amplitudes are affected by FUS genotype as well, the amplitude histograms (Fig. 7, right) show clearly that the unitary mEJC size is the same in all three genotypes (Fig. 6B). Therefore, the apparent difference in mEJC amplitudes (Fig. 6A) reflects the higher frequency of mEJCs and simultaneous release of several synaptic vesicles in the wild-type larvae (note the higher amplitude peaks in Fig. 7A, right). The Gaussian fits to the first peak in each amplitude histogram yield amplitude values of 100 ± 41 pA (mean ± width at half-peak amplitude) in wild-type FUS larvae, 97 ± 59 pA in R521C-FUS larvae, and 93 ± 45 pA in caz null larvae. Therefore, quantal size is unchanged by R521C-FUS expression or caz ablation, but the number of synaptic vesicles undergoing evoked release (quantal content, defined as evoked EJC amplitude/unitary mEJC amplitude) decreases from 200 in WT-FUS flies to 37 and 39 in R521C-FUS and caz null flies, respectively.

Structure of presynaptic active zones is aberrant in R521C-FUS-expressing flies

As yet, there is no consensus as to whether fly models of FUS- or TDP43-related ALS exhibit morphological defects at the larval neuromuscular junction, as measured by confocal microscopy. As shown previously (Lanson et al., 2011), expression of mutant bruchpilot-positive active zones, but the pattern of bruchpilot staining is different (Fig. 8D). First, there is a 50% reduction in the number of active zones with discernible doughnut holes in R521C-FUS flies (Fig. 8B). In addition, the bruchpilot-labeled structures appear disorganized, as they are in flies lacking RIM-binding protein, a key protein component of active zones (Liu et al., 2011).

To determine quantitatively whether bruchpilot-labeled active zones with discernible doughnut holes are misshapen in R521C-FUS flies compared with those in wild-type FUS transgenic flies, we examined the relationship between the size of the central hole and overall active zone area and used a fractal-based analysis to generate a quantitative measure of the complexity of the active zones. Examples of individual active zones in the two genotypes are outlined in Figure 9A and averaged active zones are shown in Figure 9B. The areas of individual active zones are similar in wild-type FUS and R521C-FUS-expressing flies (Fig. 9C). However, the overall size of the bruchpilot active zone is less well correlated with the size of the doughnut hole in the R521C-FUS flies (Fig. 9D). Furthermore, fractal analysis of the active zones indicates that they are significantly less complex (Fig. 9E) and more circular (Fig. 9F) in R521C-FUS than in wild-type FUS transgenic flies. In sum, superresolution microscopy reveals that active zones in R521C-FUS flies are aberrantly organized and misshapen compared with their wild-type counterparts.

Discussion

The demonstration that mutations in the DNA/RNA-binding proteins TDP43 and FUS are associated with both familial and sporadic ALS has profoundly influenced thinking about the pathogenesis of the disease (for reviews, see Lagier-Tourenne and Cleveland, 2009; Polymenidou et al., 2012). It is becoming evident that the impact of these findings will not be restricted to ALS, because mutations in these same proteins are implicated in at least one other neurodegenerative disorder, frontotemporal human FUS does not affect presynaptic bouton number or overall synaptic morphology, and caz null flies were reported to have normal neuromuscular junction morphology as well (Wang et al., 2011). However, aberrant synaptic morphology has been described in flies lacking caz or expressing both wild-type and mutant human FUS (Xia et al., 2012). In the present study, confocal imaging revealed no obvious differences in bouton size or number between wild-type FUS and R521C-FUS-expressing flies (Fig. 8A), but we did not investigate this in depth. Instead, we imaged the presynaptic active zone protein bruchpilot using superresolution STED microscopy. Consistent with previous work in wild-type flies (Kittel et al., 2006), STED imaging of bruchpilot in wild-type FUS transgenic flies reveals that it is localized in structures thought to be individual active zones, many of which are doughnut shaped (Fig. 8C). Presynaptic calcium channels critical for neurotransmitter release are thought to be localized within the central doughnut holes (Liu et al., 2011). Transgenic flies expressing R521C-FUS appear to have normal numbers of
Several animal models of FUS/TDP43-related ALS are becoming very useful for investigating cellular and molecular aspects of disease progression (Lanson and Pandey, 2012). Transgenic rats that express R521C but not wild-type human FUS exhibit a progressive paralysis and degeneration of motor neurons, as well as neuronal loss in the cortex and hippocampus (Huang et al., 2011). In the zebrafish, a more genetically tractable vertebrate model, expression of mutant human TDP43 or FUS causes disruption of motor neuron projections, larval motor deficits, and synaptic dysfunction (Kabashi et al., 2010; Armstrong and Drapeau, 2013a; Armstrong and Drapeau, 2013b). Finally, several groups have exploited the power of Drosophila genetics to produce flies that express either wild-type or mutant human FUS transgenes and have found that expression of the mutant protein leads to locomotor impairment and neurodegeneration (Lanson et al., 2011; Xia et al., 2012). The fly model confirms the enormous advantage of being able to restrict expression of the mutant FUS exclusively to the presynaptic motor neuron. Interestingly, expression of the wild-type protein also produces some disease-like phenotypes, but they appear to be later in onset and less severe. Knock-out of caz, the gene that encodes the fly homolog of FUS, produces behavioral and morphological defects similar to those caused by transgenic expression of mutant human FUS (Wang et al., 2011; Xia et al., 2012).

We show here that the motor neuron cell bodies of R521C-FUS-expressing larvae generate normal voltage-dependent inward and outward currents and fire action potentials in response to depolarizing stimuli. Furthermore, these action potentials can propagate normally along the axon. Indeed, if anything, the FUS-ALS motor neurons are somewhat more excitable than neurons that express wild-type FUS, again in agreement with the results in zebrafish (Armstrong and Drapeau, 2013a; Armstrong and Drapeau, 2013b) and prior findings in SOD1 mice (Kuo et al., 2004; van Zundert et al., 2008; Fritz et al., 2013). The biophysical mechanism of this widely observed increase in motor neuron excitability in ALS models remains to be determined.

Despite the fact that motor neuron cell body and axon excitability are normal or enhanced, we find that synaptic function at the larval neuromuscular junction is profoundly impaired in flies that overexpress R521C-FUS, as well as in caz-null flies. Evoked synaptic transmission is decreased by >80% and spontaneous mEJC frequency is also reduced substantially. These findings re-inforce those reported for the zebrafish larval neuromuscular junction after expression of mutant human TDP43 (Armstrong and Drapeau, 2013a) or FUS (Armstrong and Drapeau, 2013b). In the fly, although we cannot be certain whether the effect of caz knock-out is pre- or postsynaptic, the changes we observe in mEJC frequency strongly suggest a major presynaptic contribution; a similar inference has been drawn in the zebrafish. Furthermore, in the fly, there is no uncertainty about the primary location of the altered synaptic transmission; a similar inference has been drawn in the zebrafish.
presynaptic locus in the case of R521C-FUS expression because the OK-371GAL4 driver restricts expression to the presynaptic motor neuron and, again, mEJC frequency is decreased substantially. Quantal analysis reveals that the unitary mEJC amplitude is the same in R521C-FUS-expressing and caz larvae as in larvae expressing wild-type FUS. This lack of change in quantal size suggests that the size and neurotransmitter content of synaptic vesicles is not affected by FUS genotype. Rather, the decrease in evoked synaptic transmission in the FUS-ALS larvae is most readily explained by a decrease in quantal content, the number of vesicles released in response to a given stimulus. Such a decrease in evoked release probability is consistent with the decrease in spontaneous mEJC frequency observed in the FUS-ALS larvae.

It is surprising and intriguing that the decay of the evoked EJC is considerably faster in the R521C-FUS and caz larvae. Such a change in the kinetics of the EJC might be consistent with a change in the deactivation properties of the postsynaptic glutamate receptors, perhaps secondary to the initial presynaptic defect. Conversely, another possibility is that presynaptic glutamate uptake systems are among the many targets of FUS and are altered in R521C-FUS-expressing and caz motor neurons and that this accounts for the changes in EJC kinetics. A detailed analysis of glutamate receptor properties and glutamate diffusion and reuptake will be required to distinguish between these interesting possibilities and to explain how the kinetics of the evoked EJC are altered by FUS genotype whereas those of the unitary mEJC are not.

Examination of presynaptic bouton structure by confocal microscopy did not reveal any gross morphological abnormalities in the present study or in Lanson et al. (2011) (but see Xia et al., 2012). In contrast, using STED microscopy, we show that the presynaptic active zones, measured by staining the active zone protein bruchpilot, are misshapen and disorganized in R521C-FUS larvae. Although bruchpilot itself is essential for the formation of active zones and neurotransmitter release (Kittel et al., 2006), we use it here as a marker of active zone organization, as has been done previously (Liu et al., 2011). Misshapen bruchpilot-labeled active zones like those we observe in R521C-FUS-expressing larvae are associated with impaired calcium channel clustering and a decrease in calcium influx (Liu et al., 2011) and, in the zebrafish, treatment with calcium channel agonists can rescue both locomotor and synaptic transmission defects induced by expression of mutant TDP43 (Armstrong and Drapeau, 2013a). It will be interesting to use STED microscopy to examine calcium channel organization at active zones in FUS-aberrant Drosophila larvae.

Our results demonstrate clearly that structural abnormalities at the synapse and defects in neurotransmitter release precede motor neuron cell body and axonal degeneration in two fly models of FUS-related ALS. Many questions remain to be explored, including the identity of the critical FUS target(s), the mechanism by which overexpression of mutant FUS and knock-out of endogenous caz both produce the same synaptic phenotype, and the way that synaptic dysfunction leads ultimately to motor neuron degeneration. Our findings and those in the zebrafish point to the importance of early therapeutic intervention at the synapse in developing novel treatments for ALS.

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