# **Neuron**

# ALS/FTD Mutation-Induced Phase Transition of FUS **Liquid Droplets and Reversible Hydrogels into Irreversible Hydrogels Impairs RNP Granule Function**

### **Highlights**

- FUS phase transitions between monomer, liquid droplet, and hydrogel states
- FUS mutants induce further phase transition into irreversible fibrillar hydrogels
- Irreversible hydrogels sequester RNP cargo and impair RNP granule function
- Formation of non-amyloid fibrillar hydrogels provides a compelling causative mechanism for neurodegeneration

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#### In Brief

Murakami et al. show that FUS transitions between monomer, liquid droplet, and hydrogel states during uptake and release of RNP granule cargo. FUS mutations accelerate transition into fibrillar hydrogels that trap RNP cargo, impair RNP granule function, and cause neurodegeneration.







# ALS/FTD Mutation-Induced Phase Transition of FUS Liquid Droplets and Reversible Hydrogels into Irreversible Hydrogels Impairs RNP Granule Function

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#### **SUMMARY**

The mechanisms by which mutations in FUS and other RNA binding proteins cause ALS and FTD remain controversial. We propose a model in which low-complexity (LC) domains of FUS drive its physiologically reversible assembly into membrane-free, liquid droplet and hydrogel-like structures. ALS/ FTD mutations in LC or non-LC domains induce further phase transition into poorly soluble fibrillar hydrogels distinct from conventional amyloids. These assemblies are necessary and sufficient for neurotoxicity in a C. elegans model of FUS-dependent neurodegeneration. They trap other ribonucleoprotein (RNP) granule components and disrupt RNP granule function. One consequence is impairment of new protein synthesis by cytoplasmic RNP granules in axon terminals, where RNP granules regulate local RNA metabolism and translation. Nuclear FUS granules may be similarly affected. Inhibiting formation of these fibrillar hydrogel assemblies mitigates neurotoxicity and suggests a potential therapeutic strategy that may also be applicable to ALS/FTD associated with mutations in other RNA binding proteins.

#### INTRODUCTION

Mutations in a growing list of RNA binding proteins (RBPs) including TDP-43, FUS, ataxin2, hnRNPA1, and B2, are associated with sporadic and inherited forms of amyotrophic lateral sclerosis and frontotemporal dementia (hereafter "ALS/FTD") (van der Zee and Van Broeckhoven, 2014). A neuropathological hallmark of these disorders is the deposition of poorly soluble assemblies of the mutant RBP in the nucleus and cytoplasm of neurons in the brain and spinal cord of ALS/FTD patients.

The pathogenic significance of these intraneuronal assemblies has been the subject of much speculation. Hypotheses have included that they might (i) be irrelevant secondary phenomena, (ii) form conventional neurotoxic amyloids, or (iii) bind and trap crucial RNA sequences (Couthouis et al., 2011; Gendron et al., 2010; Neumann et al., 2007; Rademakers et al., 2012; Ticozzi et al., 2010). Although there are data to support the notion that these aggregates may be amyloid like (Fang et al., 2014; Robinson et al., 2013), other evidence suggests that they are distinct from classical amyloids (Cairns et al., 2007; Dormann et al., 2009; Murakami et al., 2012). Specifically, we and others have shown that in contrast to classical amyloids, pathological RBP deposits are (1) soluble in urea, (2) have low  $\beta$  sheet content, (3) have a mixed granular/fibrillar appearance under EM, (4) do not bind amyloidophyllic dyes (e.g., Thioflavin T), and (5) when



fluorescently labeled, do not display the marked reductions in in vivo fluorescent lifetimes that are typical of conventional amyloids (Kaminski Schierle et al., 2011) (Figures S1–S3).

Recently, attention has refocused on the possibility that mutant ALS/FTD-associated RBPs might be incorporated into and alter the function of ribonucleoprotein (RNP) granules (Li et al., 2013; Mitchell and Parker, 2014; Ramaswami et al., 2013; Wolozin, 2012). However, these hypotheses have not been directly proven.

To investigate this possibility, we utilized FUS as a tractable example of the RBP-proteinopathies. We describe here the results of an extensive array of biophysical, biochemical, cellular, and animal model experiments conducted over the past 4 years. In partial agreement with earlier reports (Han et al., 2012; Kato et al., 2012), we show that in a process driven by the LC domain at the N terminus of FUS, FUS undergoes phase transitions, reversibly shifting between dispersed, liquid droplets, and hydrogel-like phases. Crucially, we demonstrate that pathogenic FUS mutants severely limit this ability to reiteratively shift between these phases. In fact, FUS mutants increase the propensity of FUS to condense into poorly soluble, stable, fibrillar hydrogel-like assemblies. We show that these stable "irreversible" FUS assemblies: selectively entrap other ribonucleoproteins, impair local RNP granule function, and attenuate new protein synthesis in the axon terminals of cultured neurons. We do not exclude the possibility that formation of fibrillar hydrogels may have similar disruptive effects on nuclear RNP granules. Importantly, we show that modulation of these protein phase transitions may be a tractable therapeutic strategy.

#### **RESULTS**

To rigorously test the idea that mutations in FUS alter its assembly into RNP granules, we first explored the biology of FUS in a C. elegans model of FUS proteinopathy (Murakami et al., 2012). In this model, equivalent levels of wild-type or mutant human FUS protein are expressed under the neuron-specific Prgef-1 promoter with a GFP tag, which does not alter the activity or location of FUS (Murakami et al., 2012). We focused on animals (n ≥ 3 lines per mutant) expressing wild-type FUS (FUS(WT)) or one of five ALS/FTD mutants: FUS (R495X), FUS(R522G), FUS(R524S), FUS(P525L), and FUS501(which models several ALS/FTD-associated C-terminal truncating mutations) (Figure 1A). These mutations span a wide range of predicted aggregation propensities as calculated using the Zyggregator method (Tartaglia and Vendruscolo, 2008): FUS(R495X): Zagg = 0.44; FUS(R522G): Zagg = 0.09; FUS(R524S): Zagg = 0.05; FUS(P525L): Zagg = 0.12; FUS501: Zagg = 0.52 (Figure S4). The aggregation propensities of these mutants correlate with their phenotypic severities in C. elegans and in humans, ranging from mild (R522G) to severe (FUS (P525L), FUS (R495X), and FUS501) (Murakami et al., 2012) (Figure S5).

This *C. elegans* model displays several features observed in human FUS-related disorders (Murakami et al., 2012). Thus, in animals expressing wild-type FUS, FUS is expressed physiologically in the nucleus, and causes no detectable neurotoxicity (Figures S6 and S7). In contrast, animals expressing mutant

FUS display age-dependent neurotoxicity manifested by impaired motor function and reduced lifespan (Figures S7 and S8). These phenotypes are accompanied by nuclear and cytoplasmic deposits of FUS that are insoluble in high-salt and RIPA buffers but are soluble in 8 M urea (Figure 1B). The ex vivo abundance of these 8 M urea-soluble species correlates with the in vivo abundance of visible cytoplasmic FUS assemblies in neurons ( $r^2 = 0.96$ ) (Figure S8). As is the case in human brain expressing mutant FUS, these pathological FUS assemblies are biochemically and biophysically distinct from conventional neurotoxic amyloids (Figures S1–S3).

#### Non-Amyloid 8 M Urea-Soluble FUS Assemblies Cause Neurotoxicity

Two lines of evidence indicate that FUS assemblies drive neurotoxicity. First, in animals expressing equivalent quantities of total FUS protein, the abundance of 8 M urea-soluble FUS assemblies correlates directly with neurotoxicity as measured by motor activity ( $r^2 = 0.69$ ) and lifespan ( $r^2 = 0.88$ ) (Figure S7). Second, within each line, neurotoxicity can be increased or decreased by manipulating the abundance of the 8 M urea-soluble assemblies. In lines expressing ALS/FTD-associated mutant FUS, increasing the abundance of these 8 M urea-soluble assemblies (e.g., by heat-stress-induced depletion of molecular chaperones) exacerbated motor deficits and accelerated mortality (Figure S9; p < 0.01; n = 16 animals/ strain). Conversely, reducing the abundance of 8 M urea-soluble assemblies, either by treatment with protein aggregation inhibitors such as scyllo-inositol, or by overexpression of hsp-70, rescued both motor function and mortality of mutant FUS strains but not in non-transgenic N2 or FUS(WT) animals (Figure S10: scyllo-inositol: p < 0.01: n = 20 animals/ strain; hsp-70: p < 0.001, n = 20 animals/strain).

# Neurotoxicity Is Driven by the FUS Low-Complexity Domain

The N-terminal low-complexity (LC) domain of FUS (residues 2–214) has been reported to induce the assembly of wild-type FUS into hydrogel-like assemblies (Han et al., 2012; Kato et al., 2012). Consistent with this observation, calculation of the biophysical aggregation propensity of FUS revealed that the LC domain is highly prone to aggregate. However this tendency is offset by the lower aggregation propensity of the C-terminal domains (Figure S4). Taking these observations together, we hypothesized that (1) ALS/FTD mutations in the FUS LC domain (e.g., S96del and G156E) might directly increase the propensity of the LC domain to induce FUS assembly into higher order structures; while (2) ALS/FTD mutants in the C terminus would indirectly have the same effect by reducing the anti-aggregation effect of the C terminus (Figure S4).

In agreement with these predictions, animals expressing mutant FUS constructs lacking the LC domain had significantly less 8 M urea-soluble assemblies and less neurotoxicity (Figure 1C; n = 20 animals/strain; p < 0.01 for 8 M urea FUS and lifespan; p < 0.001 for thrashing score; Figure S11). Conversely, expression of FUS(LC) alone induced robust 8 M urea-soluble assemblies accompanied by increased motor deficits and mortality (Figure 1D, p < 0.01, Figure S12; p < 0.001; n = 20 animals/line). As predicted, the severity of these FUS(LC)-induced phenotypes was further

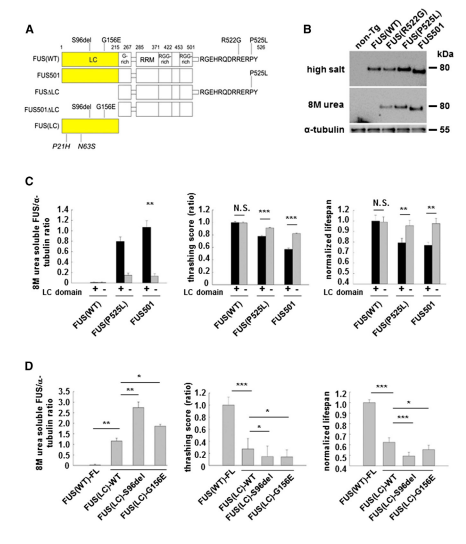


Figure 1. FUS Constructs, Cellular Localization, and Neurotoxicity of Mutant FUS

(A) FUS constructs and mutations. Full-length FUS constructs include wild-type FUS(WT) and ALS/FTD mutations (R522G, R524S, P525L, and R495X), or FUS (FUS501), which mimics several ALS/FTD truncation mutants. LC motif at residues 2-214 was deleted in FUS(WT) \( \Delta LC, \text{ FUS(P525L)} \) ΔLC, and FUS501ΔLC. FUS(LC)-only constructs include wild-type FUS(LC), pathogenic S96del or G156E mutants, or clinically benign P21H and N63S sequences. LC, low-complexity motif; G-Rich, glycine-rich region; RRM, RNA recognition motif; RGG-rich, arginine-glycine-glycine-rich

(B) FUS(WT) is located in neuronal nuclei: mutant FUS accumulates as 8 M urea-soluble assemblies in neuronal cytoplasm. Western blot of high-salt buffer extracted whole-worm lysates (top), 8 M urea extracted lysates (middle), and α-tubulin (bottom).

(C) FUS LC domain is necessary and sufficient for FUS assembly and neurotoxicity. Deletion of LC domain from FUS(P525L) and FUS 501 reduces formation of 8 M urea-soluble FUS (left panel), improves motor function (middle panel), and lifespan in mutant FUS lines (right panel), but has no impact on FUS(WT) animals. N.S., not significant. \*\*p < 0.01; \*\*\*p < 0.001.

(D) FUS LC domain is necessary and sufficient for FUS assembly and neurotoxicity. Overexpression of wild-type FUS(LC) alone is associated with increased 8 M urea-soluble assemblies (left panel), impaired motor function (middle panel), and lifespan (right panel). The ALS/FTD FUS(LC)-mutations (S96del and G156E) further augment neurotoxicity. \*p < 0.05, \*\*p < 0.01, and \*\*\*p < 0.001.

Error bars are SEM.

enhanced in animals expressing FUS(LC) domain containing the ALS/FTD associated mutants: S96del or G156E (Figure 1D; p < 0.05; n = 20 animals /line, 3 replications; Figure S12).

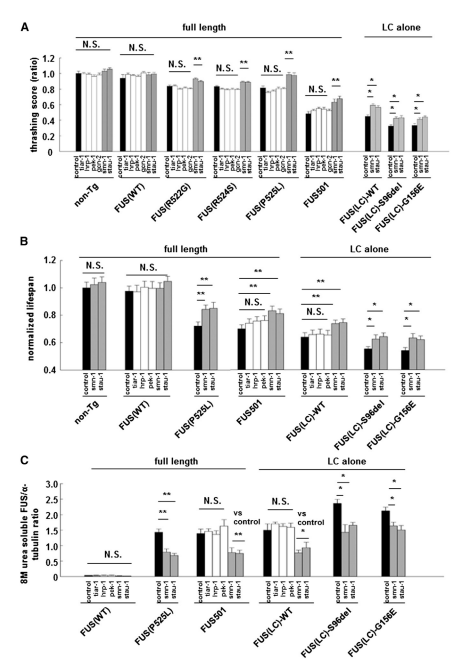
#### **Manipulation of Other Components of FUS-Containing RNP Granules Rescues Toxicity**

Pathological FUS assemblies contain other components of RNP granules including other RNA-binding proteins (Li et al., 2013; Mitchell and Parker, 2014; Ramaswami et al., 2013; Wolozin, 2012). We wondered whether some of these other components may stabilize FUS assemblies, as is the case with many other heteromeric complexes. If so, then knockdown of a subset of these RNP granule components might destabilize the pathological complexes and ameliorate neurotoxicity. We selected seven RNPs that (1) are components of RNP granules, (2) have human and C. elegans homologs, and (3) have validated neuron-specialized RNAi clones available to knock down the C. elegans homolog. This list comprised CBP (cbp-1), EIF2AK (pek-1), EIF2AK3 (gcn-2), STAU-1 (stau-1), SMN (smn-1), hnRNP1 (hrp-1), and TIA-1 (tiar-1) (upper case = human; lower case = C. elegans ortho-

log). RNAi of cbp-1 was uninterpretable because it caused nonspecific effects in non-transgenic animals. RNAi of tia-1, hrp-1, gen-2, and pek-1 did not alter thrashing scores, lifespan, or abundance of 8 M soluble FUS levels in animals expressing FUS(WT), mutant FUS, or FUS(LC) (white bars in Figures 2A-2C; p = N.S). In contrast, RNAi knockdown of stau-1 or smn-1 partially rescued neurotoxicity, reduced 8 M urea-soluble FUS assemblies, and reduced visible cytoplasmic FUS deposits in all mutant FUS strains tested (gray bars in Figures 2A-2C and S13; p < 0.05, n = 20 animals/strain, three replications).

To determine whether the rescue effect was mediated at the level of FUS granules, we co-expressed RFP-tagged STAU-1 or SMN in animals expressing various FUS cDNAs. STAU-1 and SMN strongly colocalized ( $\geq$ 72% ± 5.3% overlap) with cytoplasmic assemblies comprised of mutant full-length FUS or aggregation-prone FUS(LC) domain. STAU-1 and SMN did not colocalize with FUS(WT) (6.0% ± 3.1%; Figures 3D and S14; n = 20 neurons; p < 0.001).

The colocalization of STAU-1 and SMN with mutant 8 M ureasoluble FUS granules was semi-specific. Thus hnRNP1, which



has no effect on FUS toxicity (Figures 2A–2C), poorly colocalized with mutant FUS assemblies (Figure 3; hnRNP1 colocalization: FUS(WT) and FUS501  $\leq 6.0\% \pm 3.1\%$ ; p = N.S.). We have reported similar lack of colocalization for PAB-1 (Murakami et al., 2012). Intriguingly, TIAR-1, which is a known component of stress granules, did colocalize with pathological cytoplasmic FUS assemblies, and as described below, its presence in mutant FUS granules may have functional significance (TIAR-1 colocalization: FUS(WT) = 2.0%  $\pm$  2.0%; FUS501 = 84.0%  $\pm$  5.0%; p < 0.001) (Figure 3). Taken together, these results suggest that pathological FUS assemblies selectively trap other RNPs. This conclusion led us to ask two further questions. First, could physiological FUS assemblies form reversible structures that tran-

Figure 2. Selected RNA Binding Proteins Co-Localized with FUS in Mutant FUS Assemblies and Their RNAi Knockdown Partially Rescued Neurotoxicity

(A) A directed RNAi screen of selected LC-domain containing RBPs revealed that RNAi knockdown of SMN (smn-1) and STAU-1 (stau-1) (gray bars) partially rescued motor function, whereas control RNAi knockdown (black bars) and RNAi knockdown of TIA-1 (tiar-1), hnRNP (hrp-1), EIF2AK (pek-1), and EIF2AK3 (gcn-2) (white bars) had no effect (N.S, not significant; \*p < 0.05 and \*\*p < 0.01).

(B) RNAi knockdown of SMN and STAU-1 (gray bars) rescued lifespan, whereas RNAi of the other RNPs (white bars) and control RNAi (black bars) had no effect. N.S., not significant. \*p < 0.05 and \*\*p < 0.01.

(C) RNAi knockdown of SMN and STAU-1 (gray bars) reduced 8 M urea-soluble FUS assemblies whereas control RNAi (black bars) and RNAi of the other RNPs (white bars) had no effect. N.S: not significant. \*p < 0.05; \*\*p < 0.01.

Error bars are SEM.

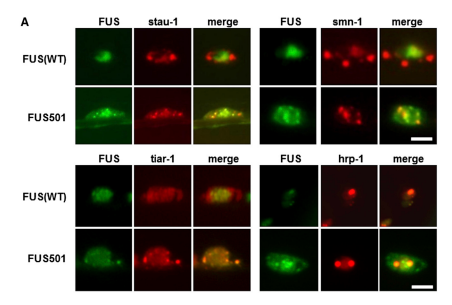
siently sequester RNP cargo upon gelformation, but then release it upon melting? Second, do irreversible FUS assemblies permanently trap their RNP cargo?

#### FUS LC Domain Induces Reversible Liquid Droplet and Hydrogel Formation

To address these questions, we began by exploring the impact of FUS mutations on RNP granule formation in SHSY5Y mammalian neuronal cells transiently transfected with YFP-tagged wild-type FUS, P525L, or FUS501 constructs. As was the case in vivo in our *C. elegans* model, in cells expressing YFP-tagged wild-type FUS, FUS was predominantly localized as spherical intranuclear droplets (Figure 4A, left panel). In cells expressing YFP-tagged mutant FUS, FUS was present as similar structures in the cyto-

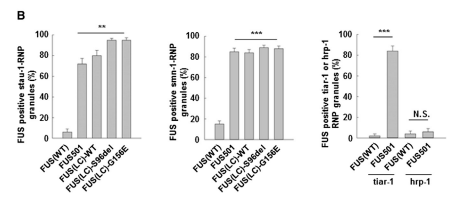
plasm and nucleus. Live cell imaging revealed that the nuclear droplets were static, but the cytoplasmic droplets were motile and on occasion fused with each other in a process previously likened to liquid droplets (Brangwynne et al., 2009) (Figure 4A, left panel).

To investigate the biophysics of FUS assembly into these liquid droplet assemblies, and to test the hypothesis that mutations in FUS might alter the dynamics of this process, we expressed and purified recombinant wild-type full-length FUS and full-length FUS containing one of five different mutations (FUS(R522G), FUS(R524S), FUS(P525L), FUS501, and FUS(G495X)). At 23°C in near-physiological protein and salt concentrations (1–5  $\mu$ M protein; 150 mM NaCl), all of the FUS proteins were fully dispersed



#### Figure 3. STAU-1, SMN, and TIAR-1 but Not hnRNP1 Colocalize with Mutant FUS in **Cytoplasmic RNP Granules**

STAU-1, SMN, and TIAR-1 (red) colocalized with pathological intraneuronal cytoplasmic and axonal assemblies of mutant full-length FUS501. wild-type FUC-LC domain and mutant FUS(LC) domain (green) but did not colocalize with nuclear FUS(WT) (green). \*\*\*p < 0.001. Scale bars, 2.0 μm. Additional data are shown in Figures S3B-S3D. STAU-1 colocalization: FUS(WT) = 6.0% ± 3.1%:  $FUS501 = 72.0 \pm 5.3\%$ ;  $FUS(LC) = 80.0 \pm 5.2\%$ ; FUS(LC)-S96del = 95.0 ± 1.7%; FUS(LC)-G156E =  $95.0 \pm 2.2$ ; n = 20 neurons; p < 0.001; SMN colocalization: (FUS(WT) = 15.0 ± 3.1%; FUS  $501 = 85.0 \pm 3.4\%$ ; FUS(LC) = 84.0 ± 3.1%; FUS(LC)-S96del = 89.0 ± 2.3%; FUS(LC)-G156E =  $88.0 \pm 2.5\%$ ; n = 20 neurons; p < 0.001). Scale bars, 2.0 um. Error bars are SEM.



liquid droplet and then to more stable assemblies, and to assess the consequences of these phase transitions, we adapted a previously published method (Han et al., 2012; Kato et al., 2012) that provides a simple and tractable platform to produce a working model of reversible (physiological) and irreversible (pathological) assembly of FUS into higher-order structures. Briefly, when 0.25-1 mM solutions of wild-type FUS(LC) in 200-500 mM NaCl are cooled to 4°C, they condensed into gels. Upon rewarming to 23°C, the gels re-dissolve into a clear solution

(Figure 5A). This liquid → gel → liquid cycle can be assayed by placing the protein solution into a siliconized microtube and then varying the temperature between 4°C and 23°C. Liquid assemblies readily fall under gravity when the tube is inverted. Gelled assemblies remain at the bottom of the tube (Figure 5A).

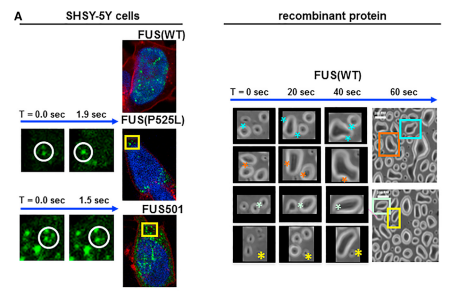
Using this assay, wild-type FUS(LC) (Figure 5B, black bar, left panel) and FUS(LC) domains containing either of the non-pathogenic variants (P2IH or N63S) all cycled four to five times before becoming irreversibly gelled (Figure 5B, white bars, left panel). In contrast, the ALS/FTD-associated mutant FUS(LC) proteins (S96del or G156E) became irreversibly gelled within two cycles (Figure 5B, gray bars, left panel, p < 0.01, n = 4 replications).

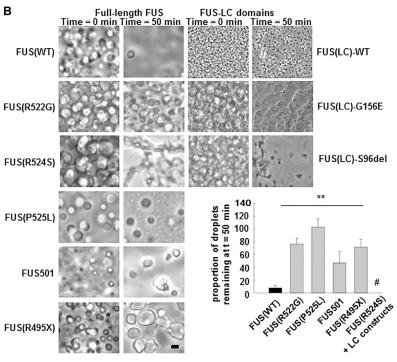
Recombinant FUS assemblies generated in this way displayed biochemical and biophysical characteristics identical to those of FUS assemblies observed in tissues from humans and from our C. elegans models. Thus, the ungelled, liquid FUS(LC) domain was fully soluble in RIPA buffer, like wild-type FUS in normal human and C. elegans tissues. In contrast, the irreversibly gelled recombinant FUS(LC) domains, regardless of whether generated by multiple cycles of wild-type FUS(LC) or fewer cycles of mutant FUS, were poorly soluble in RIPA buffer but fully soluble in 8 M

in solution. When cooled to 4°C to reduce thermal motion and thus stabilize weak interactions, they all assembled into visible liquid droplets that frequently fused with each other over timescales of 20-40 s (Figure 4A, right panel). When droplets composed of FUS(WT) were maintained at 23°C, they almost completely dissolved within 50 min (Figure 4B). In contrast, when liquid droplets composed of mutant FUS were treated in the same way, more than 50% of mutant FUS assemblies persisted at 50 min (Figure 4B, p < 0.01 n = 3 replications). Crucially, several of the full-length FUS mutants (especially FUS (R524S)) and all of the LC-domain constructs tended to produce dark, angled, or spiculated structures, which we interpreted to be solid (Figure 4B). These analyses suggest that pathogenic FUS mutations favor phase transitions of FUS from a monomeric dispersed state into a liquid droplet and then into a more stable solid form. These observations were robust but were difficult to quantify in detail because liquid droplet formation is exquisitely sensitive to changes in ionic strength and temperature of the buffer.

### **ALS/FTD-Associated Mutations in the FUS LC Domain Alter Hydrogel Properties**

To quantitatively assess the biophysical features that govern phase transition of FUS from soluble monomer to reversible





urea-again like pathological FUS deposits in human and C. elegans tissues (Figure 5C, top panel). Furthermore, negative stain electron microscopic (EM) analysis of recombinant irreversible FUS(LC) assemblies revealed a loose meshwork of fibrils, the diameter of which (14.00  $\pm$  1.0 nm, n = 20 fibrils) was indistinguishable from that of fibrils from brain tissue of a human with a FUS-P525L mutation  $(14.4 \pm 3.4 \text{ nm}, \text{n} = 56 \text{ fibrils}, \text{p} = \text{N.S.},$ Figure 5C, bottom panel) (Uchikado et al., 2006). The EM structure for reversible FUS assemblies could not be assessed because their lifetimes in the reversible state (minutes) is shorter than the time required to fix and prepare them for EM analysis. Nevertheless, these observations suggest that the recombinant

#### Figure 4. Recombinant FUS Generates Liquid Droplet-Like Assemblies, the Reversibility of which Is Reduced by ALS/FTD Mutants

(A) (Left panel) SHSY5Y cells transiently transfected with YFP-tagged wild-type FUS show static intranuclear spherical structures. (Top) In the same cell type, mutant FUS localizes in the cytoplasm as mobile spherical structures ("droplets") that move, fuse, and disappear (middle and bottom-time series of two fusion events). (Right panel) Cooling of 1-5 µM solutions of either full-length FUS or FUS(LC) domain, induces spontaneous phase transition into liquid droplets, which spontaneously fuse (\*), split over seconds, and redisperse into solute over  $\sim\!50$  min.

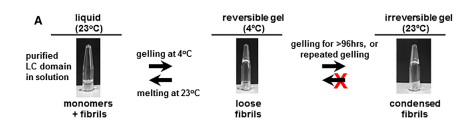
(B) Full-length FUS(WT) liquid droplets are more labile than full-length mutant FUS or FUS(LC) droplets. Both mutant and wild-type FUS proteins form multiple bright spherical liquid droplets after cooling (left column). FUS(WT) droplets almost completely disperse by 50 min at 23°C (right column). FUS proteins containing ALS/FTD mutants formed more stable droplets, many of which were still visible at 50 min at 23°C. The residual mutant assemblies had darkly refractile properties, angular edges, and/or spiculated appearances. We interpret these structures to represent further phase transitions to gel-like or solid assemblies. Spiculated assemblies were especially prevalent with the FUS(LC) constructs, where virtually all of the residual FUS(LC) material had this appearance (# in the graph). Error bars are SEM.

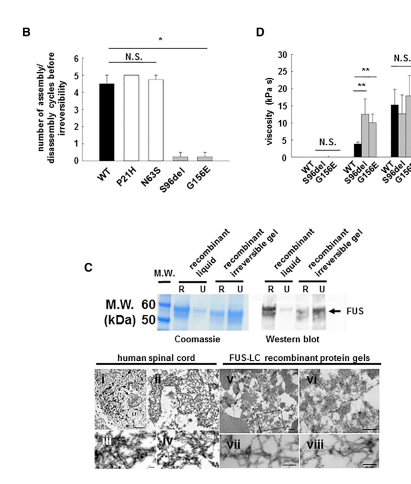
FUS assemblies modeled here accurately reflect those occurring in vivo in human tissue.

#### **Single Particle and GFP Tracking Gel Measurements**

To further investigate the biophysics of these phase transitions, we used single particle tracking methods (Crocker and Grier, 1996; Rees et al., 2013) to measure changes in the apparent viscosity between liquid/liquid droplet, reversible, and irreversible gel-like FUS states.

Briefly, 20-nm radius fluorescent beads were dispersed within a 15-µl volume of FUS solution, which was then cycled through different phases. The Brownian movement of the beads was then used to infer apparent viscosities of the FUS assemblies in each phase. Wild-type FUS(LC) assemblies in liquid state prior to, and immediately after a single round of assembly, displayed very low dynamic viscosity that is typical of a liquid (Figure 5D, left bars:  $0.4 \pm 0.03$  Pa s). When wild-type FUS(LC) formed a reversible gel, there was a modest increase in viscosity (Figure 5D, middle bars:  $3.8 \pm 0.4$  kPa s), which is similar to the viscosity reported for P-granules (~1.0 Pa s)(Brangwynne et al., 2009). In contrast, mutant FUS(LC) assemblies behaved





very differently. In initial liquid state, mutant FUS(LC) had very low dynamic viscosity similar to that of wild-type FUS(LC)  $((FUS(LC)(S96del) = 0.04 \pm 0.02 \text{ Pa s; } FUS(LC)(G156E) =$  $0.013 \pm 0.002$  Pa s; p = N.S.) (Figure 5D, left bars). However, mutant FUS(LC) that had been assembled once, but was still reversible, displayed a much greater viscosity compared to that of once-gelled wild-type FUS(LC) (FUS(LC)(S96del) =  $12 \pm 6 \text{ kPa s}$ ; FUS(LC)(G156E) =  $10 \pm 1.7 \text{ kPa s}$ ; p = 0.001) (Figure 5D, middle bars). Even more dramatically, irreversible assemblies, regardless of whether generated using wildtype or mutant FUS(LC), had extremely high viscosities  $(FUS(LC)(WT) = 15 \pm 3 \text{ kPa s}; FUS(LC)(S96del) = 12 \pm 8 \text{ kPa}$ s;  $FUS(LC)(G156E) = 17 \pm 2 \text{ kPa s}$ , p = N.S.) (Figure 5D, right bars). The very high viscosity values for irreversible FUS assemblies are fully consistent with the electron microscopic

#### Figure 5. Recombinant FUS Generates Gel-Like Assemblies, the Reversibility of which Is Reduced by ALS/FTD Mutants

(A) Purified recombinant FUS(LC) forms clear solution at 23°C (Top left panel) but forms reversible gel-like structures at 4°C (Top center panel). Rewarming to 23°C reverts FUS(LC) to a liquid state (top left panel). Repeated cycling of wild-type FUS(LC) eventually results in the formation of irreversible gels (top right panel).

(B)  $\sim$ 4 to 5 assembly/disassembly cycles can be achieved by wild-type FUS(LC) (black bar) and by FUS(LC) containing benign polymorphic variants (P21H and N63S, white bars), ALS/FTD mutations (gray bars) dramatically reduce the number of cycles to <1 before irreversibility (\*\*p < 0.01).

(C) Top panel: Coomassie gel (left) and western blot (right) of RIPA- and 8 M urea solubilisation of FUS(LC) reveal that liquid state FUS(LC) is fully soluble in RIPA buffer. Irreversible FUS(LC) is partly soluble in RIPA, but fully soluble in 8 M urea. Bottom panel: Transmission electron microscopy of human FUS inclusions ([Ci]-[Civ]) and irreversible recombinant protein gels ([Cv]-[Cviii]). Both are composed of loose networks of  $\sim$ 14 nm fibrils. Scale bars are 500 nm ([Cv] and [Cvi]) and 100 nm ([Cviil and [Cviii]).

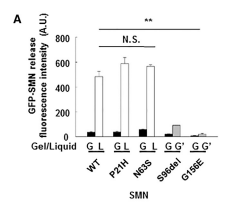
(D) Analysis of liquid, reversible gel, and irreversible gel using fluorescent particle tracking revealed that liquid (left cluster) and reversible gels (black bars, middle cluster) have viscosities similar to those of P-granules. Mutant reversible gels (gray bars, middle cluster) were more viscous than wildtype reversible gels. Irreversible gels, regardless of whether made from wild-type or ALS/FTD mutant FUS exhibit very high viscosities. N.S., not significant; \*\*p < 0.01. Error bars are SEM.

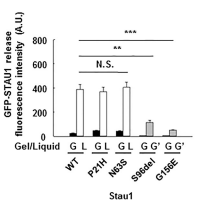
structure of these assemblies as loose meshworks of intertwined fibrils that fulfil the IUPAC definition of a hydrogel (Figure 5C). These results support the hypothesis that FUS assemblies can phase-transition between (i) soluble monomeric/liquid droplet like states, (ii)

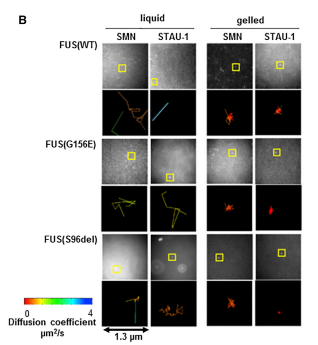
reversible hydrogel-like states, and (iii) irreversible, fibrillar hydrogel-like states.

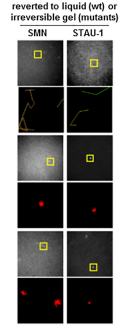
# **Mutant FUS Assemblies Trap Other RNA Binding**

We reasoned that condensation of mutant FUS into higher-order assemblies might alter binding and release of RNP granule cargo. To test this idea, we mixed mutant or wild-type FUS(LC) with GFP-tagged SMN (or in separate experiments, with GFP-STAU-1). The mixtures were cooled to induce phase transition and assembly, and buffer was layered on top. The diffusion of SMN (or STAU-1) into the overlying buffer layer was then measured by fluorescence. Very little SMN or STAU-1 was detected in the buffer above any of the gelled assemblies (black bars labeled G-gel in Figure 6A.), regardless of whether the









gels were composed of mutant or wild-type FUS(LC). However, SMN and STAU-1 were rapidly released into the buffer following melting of reversible gels compose of wild-type FUS(LC) or non-pathogenic FUS(LC) variants (P21H or N63S) (white bars labeled L-liquid in Figure 6A). However, gels composed of either of the pathogenic mutants (S96del or G156E) failed to revert to liquid states and, as a result, retained SMN and STAU-1 (gray bars labeled G', Figure 6A). These experiments reveal that cargo proteins can be retained by pathological irreversible FUS assemblies.

To provide further support for the notion that irreversible FUS assemblies might trap cargo RNPs, we used single-molecule imaging to track movement of individual GFP-tagged SMN and GFP-tagged STAU-1 molecules within mutant and wild-type FUS assemblies. Both SMN and STAU-1 moved freely in never-gelled liquid preparations of wild-type and mutant FUS (diffusion coefficients: 1–3 μm<sup>2</sup>/s; Figure 6B, left two columns).

#### Figure 6. Mutant FUS Forms Poorly Reversible Gels, which Fail to Release Cargo such as SMN and STAU-1

(A) When recombinant GFP-tagged SMN (or STAU-1) protein is mixed with FUS(LC) protein and gel assembly is induced by cooling, little SMN (or STAU-1) is released into the buffer (as measured by fluorescence) regardless of whether the FUS gel is derived from wild-type or ALS/FTD mutant FUS(LC) (black bars). When the gel is reversed by warming to 23°C (white bars), significant SMN (or STAU-1) protein is released into the buffer from FUS(WT) hydrogels. Very little SMN (or STAU-1) is released by the irreversible ALS/FTD mutant FUS(LC) gels (gray bars;\*\*p < 0.001).

(B) Fluorescent molecule tracking in liquid, reversible gel, and irreversible gel assemblies reveals that in liquid state, SMN and STAU-1 diffuse rapidly at 1-4  $\mu\text{m/s}$  in both mutant and wild-type samples (left two columns). On formation of a gel (center two columns), movement of SMN and STAU-1 is constrained (<1 µm<sup>2</sup>/s). On rewarming, reversible FUS(WT) gels disassemble and release SMN and STAU-1. The irreversible mutant gels continue to constrain movement of SMN and STAU-1. Image stacks are captured at 29 Hz; pixel width corresponds to 160 nm on the specimen. Representative areas are boxed and visualized in a 1.3 µm<sup>2</sup> plot, color coded to the diffusion coefficient fitted to each track.

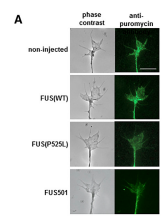
Error bars are SEM.

Following phase transition and assembly. the diffusion coefficients of both SMN and STAU-1 decreased substantially in reversible assemblies of both wild-type and mutant FUS(LC) (<<1 µm<sup>2</sup>/s) (Figure 6B, center two columns). Fast diffusion of SMN and STAU-1 returned when the reversible FUS(WT) assembly was melted at 23°C (Figure 6B, top right, columns). In contrast, despite re-warming, irreversible FUS assemblies comprised

of either twice-cycled mutant FUS or multiply cycled wild-type FUS, continued to significantly constrain movement of SMN and STAU-1 (Figure 6B, bottom right columns).

#### **Mutant FUS Reduce New Protein Synthesis in Neuronal Axons and Termini**

Taken together, our experiments demonstrate that mutant FUS has an increased propensity to form irreversible assemblies that could disrupt the function of FUS-containing RNP granules in several, non-mutually exclusive ways. For instance, irreversible RNP granules containing mutant FUS could disturb the intracellular transport of those RNP granules (Alami et al., 2014; Liu-Yesucevitz et al., 2014). Alternatively, mutant FUS assemblies could trap and titrate out key factors. In the nucleus, this could affect transcription and splicing. In the cytoplasm, and particularly in specific subcellular niches such as dendrites and axon terminals, the trapping could impair local RNA metabolism and



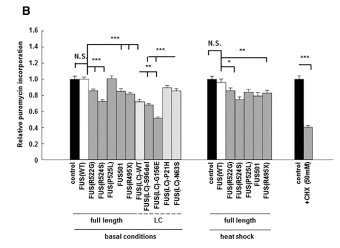


Figure 7. Mutant FUS and FUS(LC) Domain, but Not Full-Length Wild-Type FUS, Cause Reduced New Protein Synthesis When Expressed in Cultured Neurons

(A) New protein synthesis was detected by puromycylation in axon terminals of cultured Xenopus retinal ganglion neurons (left panel: phase contrast; right panel: anti-puromycin fluorescent images) expressing equivalent quantities of full-length FUS(WT) or various mutant and truncated human FUS proteins. Scale bar, 10 um.

(B) Under basal conditions, de novo protein synthesis is equivalent in non-injected neurons, and in neurons expressing FUS(WT). However, de novo protein synthesis was significantly reduced in neurons expressing ALS/FTD mutants including: FUS(R522G), FUS(R524S), FUS501, FUS(R495X), and in gellingprone FUS(LC)(WT), FUS(LC)(S96del), FUS(LC)(G156E), FUS(LC)(P21H), and FUS(LC)(N63S). In good agreement with the in vivo studies in C. elegans, heat shock causes significant reduction of new protein synthesis in neurons expressing mutant FUS, but minimal impact on non-transfected and FUS(WT) expressing neurons (right panel). N.S., not significant; \*p < 0.05, \*\*p < 0.01, and \*\*\*p < 0.001.

Error bars are SEM.

translation, thereby converting irreversible FUS RNP granules into local translation repressors. This possibility is distinctly likely because both TIA-1 and mutant FUS are present in stress granules, which act as translational repressors (Anderson and Kedersha, 2002; Bosco et al., 2010; Decker and Parker, 2012).

To assess the latter hypothesis, we investigated de novo protein synthesis in axon terminals of primary Xenopus retinal neuron cultures expressing equivalent quantities of human full-length FUS(WT), full-length mutant FUS, wild-type FUS(LC), pathogenic mutant FUS(LC), or non-pathogenic FUS(LC) variants (Figure S15). Newly synthesized proteins were detected by pulse labeling with puromycin. The released truncated puromycinylated proteins were imaged and quantified by immunofluorescence using an anti-puromycin antibody (Lin et al., 2009; Schmidt et al., 2009; tom Dieck et al., 2015) (Figure 7A). Under basal conditions, new protein synthesis was significantly reduced in axons expressing full-length FUS(R522G) (0.86 ± 0.02), FUS(R524S) (0.72  $\pm$  0.02), FUS501 (0.85  $\pm$  0.03), and FUS(R495X)  $(0.82 \pm 0.02)$  (Figure 7B, left panel, p < 0.001). Following heat shock using the identical paradigm employed in C. elegans, new protein synthesis was further reduced in all neurons expressing mutant FUS, but not in non-injected neurons or neurons expressing FUS(WT) (Figure 7B, center panel, p < 0.01). This result precisely mirrors the effect of the same heat shock paradigm on in vivo toxicity of mutant FUS in C. elegans. Additionally, and again in good agreement with the in vivo neurotoxicity studies in C. elegans, new protein synthesis was significantly decreased in neurons expressing FUS(LC), and this was further worsened by the S96del or G156E ALS/FTD mutants (FUS(LC)(WT)  $(0.72 \pm 0.02)$ , FUS(LC)(S96del)  $(0.68 \pm 0.02)$ , and FUS(LC)(G165E)  $(0.52 \pm 0.02)$ , FUS(LC)(P21H)  $(0.89 \pm 0.03)$ , FUS(LC)(N63S)  $(0.86 \pm 0.03)$ , Figure 7B, left panel, p < 0.001). Treatment of neurons expressing wild-type or mutant FUS with cyclohexamide was used as a positive control (Figure 7B, right panel).

Our interpretation of these results is that mutant FUS assemblies perturb RNP granule function by trapping key components of the RNA metabolism and translation machinery. However, an alternate possibility is that the inhibition of new protein synthesis arises from cellular stress due to accumulation of misfolded FUS proteins. While there are several non-specific forms of stress that might differentially occur in animals and cells expressing mutant FUS, the type of stress most clearly associated with the accumulation of misfolded proteins and with inhibition of new protein synthesis is the unfolded protein response (UPR), also known as the endoplasmic reticulum (ER) stress response. Traditionally, the UPR has only been associated with accumulation of misfolded proteins in the ER. However, in some models of neurodegenerative diseases, intracellular misfolded non-ER resident proteins have also been noted to activate the UPR and attenuate protein synthesis (Moreno et al., 2013; Nishitoh et al., 2002).

To address this possibility, we undertook three experiments: the first two, in vivo in our C. elegans model; the third, in cultured Xenopus neurons.

We have previously noted that C. elegans expressing either full-length wild-type FUS or full-length mutant FUS had very few intra-cytoplasmic stress granules (zero to two per neuron; p = N.S.) (Murakami et al., 2012). However, animals expressing FUS (WT) and animals expressing mutant FUS are all clearly capable of mounting a vigorous stress response after heat shock, displaying abundant stress granules (five to six granules/cell) (Figure 8, left panel; p = N.S.). These in vivo observations suggest that (1) animals expressing mutant or wild-type FUS are capable of mounting a stress response and (2) mutant-expressing animals are not more stressed by accumulation of misfolded mutant FUS than are animals expressing wildtype FUS.

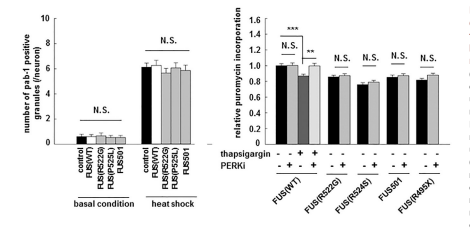


Figure 8. Neurotoxicity of Mutant FUS in C. elegans and Reduced New Protein Synthesis in Xenopus Neurons Are Unlikely to be Due to Activation of the UPR due to Accumulation of Misfolded FUS

(A) In vivo stress granule density measured by counting pab-l-positive granules in C. elegans under basal conditions and after heat shock were not different between control (empty vector); FUS(WT) and mutant FUS animals, indicating that mutant FUS animals were not under greater unfolded protein stress.

(B) Thapsigargin treatment induced UPR and reduced new protein synthesis in Xenopus neurons expressing FUS(WT). PERK1 inhibitors rescued reduced protein synthesis in thapsigargin-treated neurons expressing FUS(WT), but not mutant FUSexpressing neurons.

Error bars are SEM.

Second, we examined the activity of the UPR in our C elegans model. Translation repression in the UPR is mediated by the eIF2a kinase PERK, and it can be substantially reversed by PERK kinase inhibitors or by PERK deletion. We used RNAi to completely ablate detectable mRNA expression for the endogenous C. elegans PERK homolog (pek-1) and the other eIF2a kinase of worms (gcn-2) (Mao and Crowder, 2010; Scheuner et al., 2001). Despite the complete loss of pek-1 and gcn-2, there were no changes in the motor function and lifespan in pek-1 or gcn-2 RNAi-treated mutant FUS-expressing animals (Figures 2A and 2B). These results confirm that neurotoxicity induced by mutant FUS is not directly attributable to activation of PERK (or of eIF2a phosphorylation by other means), and hence, neurotoxicity is unlikely to reflect activation of the UPR.

Third, we undertook PERK inhibitor studies in the Xenopus neuron culture model. We compared the effect of the PERK kinase inhibitor GSK2606414 (Axten et al., 2012) on new protein synthesis in neurons expressing (i) wild-type FUS, (ii) wildtype FUS plus thapsigargin (a well-established activator of the UPR), and (iii) ALS/FTD mutant FUS: FUS(R522G), FUS(R524S), FUS501, or FUS(R495X). The UPR was induced, and new protein synthesis was stalled by thapsigargin treatment in neurons expressing full-length FUS(WT) (Figure 8, right panel, lane 3). The UPR was blocked, and new protein synthesis was restored in FUS(WT) neurons treated with GSK2606414 (Figure 8, right panel, lane 4). However, equivalent doses of the inhibitor failed to reverse the impaired new protein synthesis in neurons expressing mutant FUS (Figure 8, right panel, lanes 6, 8, 10, and 12).

#### **DISCUSSION**

A detailed explanation of how ALS/FTD-associated mutations in RBPs might impact neuron biology would increase our understanding both of the molecular origins of these neurodegenerative disorders and of the biology of nuclear and cytoplasmic RNP granules. Our work provides several important insights into these issues.

First, the biochemical and biophysical experiments reported here support a physiological model in which reversible phase transition of LC-containing proteins such as FUS can be used to control the assembly and the function of membrane-free structures such as nuclear and cytoplasmic RNP granules. Thus, as we have shown, various cargo elements such as other RNP proteins can be concentrated into a dimensionally constrained space during liquid droplet and reversible hydrogel formation and then released upon their dissolution. The biophysics of conversion from liquid droplet to reversible hydrogel is not yet clear, but they differ only slightly in viscosity, and reversible hydrogels might reflect the partial polymerization into a loose but reversible fibrous network that provides more stability for longer-lived granules.

However, the biophysical propensity of LC-containing proteins to undergo phase transitions to liquid droplets and hydrogels also renders them vulnerable to further condensation into hyper-stable structures. Thus, we have shown that ALS/FTD mutations in FUS dramatically increase the propensity to undergo phase transition into stable "irreversible" fibrillar hydrogel-like states. Indeed, even wild-type FUS can eventually be induced to phase shift into fibrillar hydrogel form—an observation that provides a mechanism for the occurrence of irreversible FUS assemblies in cases of sporadic ALS/FTD with FUS pathology, but without mutations in FUS.

We show that FUS assembly is largely driven by the LC domain. However, our biophysical modeling suggests that other parts of the protein (especially the C terminus) likely buffer this tendency and render the overall protein less prone to form higher order assemblies (Figures S1-S4). Our biophysical modeling also suggests that mutations in the LC domain might directly increase the propensity of the LC domain to form irreversible gels, while mutations in the C terminus are likely to indirectly increase the risk of phase transition by reducing the anti-assembly property of the C terminus (Figures S1-S4). Still other mutations may affect this process by altering binding of RNA or other proteins, which are known to nucleate FUS assembly (Schwartz et al., 2013).

Our work suggests that stable fibrillar hydrogels induce toxicity by mechanisms different from conventional amyloids, namely by disrupting RNP granule function. This conclusion is supported by the fact that downregulating the expression of other RNP granule components destabilizes the pathological RNP granules and rescues toxicity.

Perturbation of RNP granule function could induce neurotoxicity through several mechanisms. Our results strongly suggest that one, but not necessarily the only, neurotoxic effect of phase transition into irreversible higher order assemblies is to disrupt the function of cytoplasmic RNP granules, thereby reducing new protein synthesis in intracellular compartments, such as axon terminals, that are heavily dependent on local translation control by RNP granules (Akbalik and Schuman, 2014; Holt and Schuman, 2013; Jung et al., 2014). This effect would be predicted to be most severe in dendrites and axon terminals of long projection neurons (e.g., motor neurons affected in ALS). The reduction in local new protein synthesis could arise through several molecular mechanisms that are not mutually exclusive, including: altering intracellular transport of FUS granules (Alami et al., 2014; Liu-Yesucevitz et al., 2014), and, as we have shown here, trapping cargo proteins (e.g., TIAR-1) involved in regulating local RNA metabolism and translation.

It is quite probable that similar phase transitions in nuclear FUS might also induce changes in RNA transcription and splicing, which could contribute to toxicity.

Looking forward, it will be important to catalog the differential RNA and protein components of normal and pathological FUS assemblies in ALS and in FTD and to discover the signaling events that govern their assembly and disassembly. Understanding the basis for these processes may provide an explanation for the different clinical phenotypes induced by FUS assemblies. More importantly, as we have shown, manipulation of molecular components of pathological RNP granules (like STAU-1 and SMN) to destabilize pathological assemblies and manipulation of the signaling pathways that govern assembly and disassembly of physiological and pathological RNP granules might be tractable therapeutic targets for these diseases.

The observations reported here are likely to pertain to neurodegenerative diseases associated with TDP-43, TAF15, EWS, hnRNPA1, and B2. Impaired degradation and removal of these assembly-prone proteins could also explain the occurrence of pathological assemblies of these proteins (particularly TDP-43) in ALS/FTD caused by mutations in autophagy pathway genes such as p62, optineurin, TBK-1, and valosin-containing protein-1.

#### **EXPERIMENTAL PROCEDURES**

#### Transgenic FUS animals

C. elegans stably expressing FUS(WT), FUS(R522G), FUS(R524S), FUS(P525L), FUS(R495X), and FUS501 were generated as described (Murakami et al., 2012). FUS501 mimics human pathogenic truncation mutants such as FUS G497AfsX527(Yan et al., 2010), K510WfsX517X (Yan et al., 2010), G472X (Hara et al., 2012), G466VfsX15 (DeJesus-Hernandez et al., 2010), and R495X(Bosco et al., 2010; Yan et al., 2010) G504Wfs (Kent et al., 2014). Strains lacking the LC motif or expressing LC motif alone (wild-type, S96del and G156E) were made similarly. Strains had similar levels of transgene expression. C. elegans hsp-70, pab-1, stau-1, smn-1, tiar-1 and hrp-1 cDNAs were injected into lin-15 animals together with FUS constructs to generate double transgenics.

#### **Animals and Human Tissues**

Experiments on C. elegans and Xenopus were conducted using approved protocols at the University of Toronto and University of Cambridge. Post-mortem human brain and spinal cord tissues were obtained from four unrelated subjects with FUS(P525L) using institutionally approved protocols at Mayo Clinic (D. Dickson) and Columbia University (N. Shneider).

#### Thrashing Assay, Heat Shock Treatment, and Small Molecule **Compound Treatment**

Motor activity was assessed by counting body bending for 1 min in 4-day-old adult C. elegans (Gao and Zhen, 2011). Heat shock involved incubation at 30°C for 6 hr, and then at 22°C, repeated daily for 3 days (Murakami et al., 2012). Scyllo-inositol (100  $\mu$ M, IO631, TCI America) was added to the media.

#### **Electron Microscopy**

Anterior horns of formalin-fixed spinal cord were removed, infiltrated, and embedded in LR White resin (Polysciences). Sections were stained with 2% uranyl acetate and lead citrate and examined with a Philips 208S electron microscope. Recombinant FUS(LC) was diluted to 0.2 mM, blotted onto Formvar grids, stained with 1% phosphotungstic acid (PTA, pH 7.0), and examined with a Hitachi 7000 electron microscope at 75 kV (Morita et al., 2009).

#### **Live Cell Imaging**

Human Neuroblastoma cell line, SH-SY5Y, was transfected using Neon Transfection System (Life Technologies). After 12 hr incubation, cells were stained and visualized with 710 Zeiss confocal microscope.

#### RNAi Assay of C. elegans Strains

RNAi was performed as previously (Dixon et al., 2006; Kamath et al., 2003).

#### **Sequential Fractionation and Western Blot Analysis**

Animals were lysed in ten volumes (v/w) of 50 mM Tris-HCl, 750 mM NaCl, 10 mM NaF, and 5 mM EDTA (pH 7.4) with sonication. The lysate was centrifuged and supernatant was collected (high-salt-soluble fraction). The procedure was repeated sequentially in RIPA buffer, 2% SDS buffer, and 8 M urea buffer as described (Murakami et al., 2012).

#### RNA Granule Colocalization and GFP Intensity Measurement

Cytoplasmic RFP-positive, GFP-positive, and double-positive granules were counted in tail neurons of double transgenic animals co-expressing GFPtagged-FUS and RFP-tagged stau-1, smn-1, tiar-1, or hrp-1.

#### Single Particle Tracking Analysis in FUS Droplets and Gels

Samples were imaged on a Nikon TE-300 inverted wide-field fluorescence microscope and laser illumination (Cobalt, 488 nm wavelength laser attenuated to 2 mW/cm<sup>2</sup> intensity on the specimen for imaging GFP, and Toptica iBeam smart 640 nm laser at 20 mW/cm<sup>2</sup> for imaging fluorescent spheres). Images were captured from a focal plane 15 um above the coverslip to ensure the observations measured the particles in the bulk specimen and not any adhering to the coverslip. Particle track analysis was performed as described (Crocker and Grier, 1996; Rees et al., 2013). Each repeat measurement of viscosity was obtained from a separate specimen, and at least four image stacks were measured for each.

#### Expression, Purification, and Gelling of Recombinant FUS LC **Domain Constructs**

cDNAs for FUS LC residues 2-214 and its variants were cloned (pOPINS vector) and expressed in E. coli BL21(DE3), purified on Ni-NTA resin, buffer exchanged into either droplet buffer (50 mM Tris and 150 mM NaCl [pH 7.5]) or gelation buffer (50 mM Tris and 500 mM NaCl [pH 7.5]). For gelling experiments. FUS samples were concentrated to 1 mM, transferred to thin-walled PCR tubes and incubated at 4°C, and transferred to 23°C to induce melting. For liquid droplet assays, samples were adjusted to 1–5  $\mu$ M FUS, and 20  $\mu$ l aliquots were spotted onto siliconised coverslips, cooled to 4°C, and imaged at 40× on an EVOS FL microscope.

#### **GFP-SMN/STAU-1** Release Fluorescence Assay

Human SMN or STAU-1 with N-terminal Hisg-GFP tag were expressed and purified as for FUS constructs, including a gel filtration step. 5  $\mu$ l of 10  $\mu$ M GFP-SMN or GFP-Stau1 protein was mixed with 100 µl of 1 mM wild-type or mutant FUS LC and then incubated at 4°C until gelling occurred. Gels were overlaid with 100  $\mu$ l of 50 mM Tris with 500 mM NaCl (pH 7.5) buffer; transferred to 23°C to melt; and monitored by fluorescence for release of GFP-SMN or GFP-Stau1.

#### **Retinal Cultures and Puromycin Labeling**

Xenopus laevis embryos were fertilized in vitro and raised in 0.1× Modified Barth's Saline at 14°C–18°C. Capped mRNAs of wild-type or mutant FUS were made (SP6 kit (Ambion)), polyadenylated using Poly(A)-tailing kit, and injected into the two dorsal blastomeres at four-cell stage. Eye primodia from stage 24–26 embryos were dissected and cultured in 60% L–15 on laminin-coated coverslips at 20°C for 48 hr (basal condition), or at 20°C for 42 hr followed by 6 hr at 30°C (heat shock treatment). For puromycin labeling, cultures were treated with 1.8 μM puromycin for 15 min, labeled with Alexa Fluor 488 conjugated anti-puromycin antibody (1:250, Millipore), and imaged as described (Lin et al., 2009; Schmidt et al., 2009; tom Dieck et al., 2015). The fluorescent intensities of 80–250 growth cones per sample were collected. Experiments were performed in three to four replicates.

#### **Statistical Analysis**

Unpaired Student's t tests, Fisher's exact test, and ANOVA were performed with SigmaPlot version 11.0.

#### SUPPLEMENTAL INFORMATION

Supplemental Information includes fifteen figures and can be found with this article online at http://dx.doi.org/10.1016/j.neuron.2015.10.030.

#### **AUTHOR CONTRIBUTIONS**

T.M., A.M., S.-P.Y., R.R.F, Y.W., and M.Z. performed the *C. elegans* experiments. S.Q., A.R.C., R.B.D., D.K.-V., Y.L., W.M., and L.D. performed the recombinant protein experiments. E.R., G.S.K.S., F.T.S.C., and C.H.M. did the single particle experiments. G.G.T., G.F., and M.V. did bioinformatics and computational analyses. W.-L.L., D.W.D., and P.E.F. did to the electron microscopy. J.Q.L. and C.H. did the new protein synthesis experiments. P.S.G.-H., M.V., C.H., P.E.F., N.A.S., M.Z., C.F.K., E.R., G.S.K.S., D.R., and D.W.D. designed the experiments. All authors contributed to the writing of the manuscript.

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# **Supplemental Information**

## **ALS/FTD Mutation-Induced Phase Transition of FUS**

# **Liquid Droplets and Reversible Hydrogels into**

# **Irreversible Hydrogels Impairs RNP Granule Function**

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# **Supplemental Data inventory**

Supplemental   Topic   Related to   Comments			
Item	-	Main Text item	
Supplemental Figure 1-10	Biophysical studies of FUS. Correlation of 8M urea soluble FUS with toxicity	Introduction and main Figure 1	<ul> <li>1-3: absence of amyloid characteristics</li> <li>4: aggregation propensity</li> <li>5: correlation of aggregation propensity with disease severity in humans and C. elegans.</li> <li>6-10: basal characteristics of transgenic FUS strains (6-8) and detailed data items (9, 10) correlating abundance of 8M urea soluble FUS species with toxicity (lifespan and motor function). Showing that reducing 8M urea soluble species rescues toxicity, while increasing 8M urea soluble species increases toxicity.</li> </ul>
Supplemental Figure 11, 12	Demonstration that the low complexity (LC) domain is necessary and sufficient for toxicity	Main Figure 1	<ul> <li>11: experimental details from experiments using deletion of LC domain to demonstrate decreases in 8M urea and toxicity</li> <li>12: experimental details from experiments using overexpression of wild-type or mutant LC domain to demonstrate increases in 8M urea and toxicity.</li> </ul>
Supplemental Figure 13, 14	Demonstration that other RNA binding proteins colocalise with FUS in pathological aggregates, and some help stabilise these aggregates	Main Figure 2, 3	13, 14: experimental details from experiments using RNAi treatment against SMN and STAU-1.
Supplemental Figure 15	Demonstration that expression levels of human FUS protein in Xenopus retinal neurons are equivalent	Main Figure 7	15: western blotting of Xenopus neurons expressing human FUS

## Supplemental Figure 1-3: FUS aggregates are not amyloids in vivo or in vitro

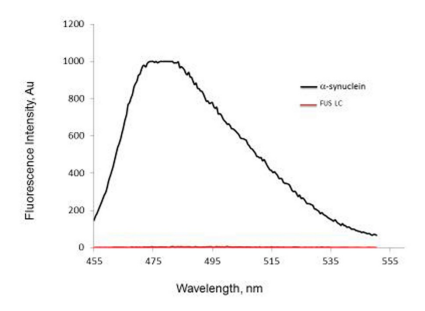


Figure 1: FUS LC hydrogel show no features characteristic of amyloids: No Thioflavin T (ThT) binding: 0.4 mg/ml irreversible hydrogel of FUS LC (red) does not bind Thioflavin T. 0.4 mg/ml of fibrillar  $\alpha$ -synuclein (black) shows fluorescence enhancement upon Tht binding indicating the presence of  $\beta$ -sheet structures characteristic of amyloids. Fluorescence spectra were collected using an LS 55 fluorometer following incubation of gelled proteins samples and  $\alpha$ -synuclein with 50  $\mu$ M ThT. Samples were excited at 442 nm and a fluorescence spectrum collected from 465-520 nm.

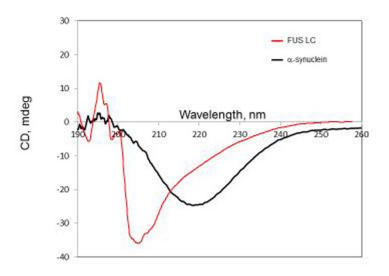


Figure 2: FUS LC hydrogel show no features characteristic of amyloids: Non  $\beta$ -sheet Circular dichroism spectra: FUS LC hydrogel spectra (red) shows no discernable features of the  $\beta$ -sheet secondary structure typical of amyloids. The  $\beta$ -sheet spectra of fibrillar  $\alpha$ -synculein (black) is shown for comparison. Circular dichroism spectra were acquired using a Jasco J-810 spectropolarimeter at a scan speed of 50 nm/min with a response time of 2 s and averaged over 20 scans at 25 °C.

### In vivo fluorescent life times

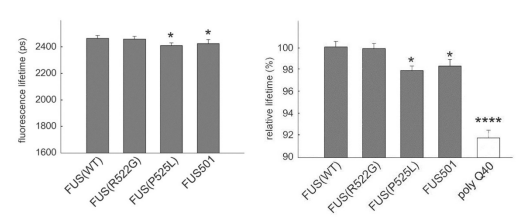
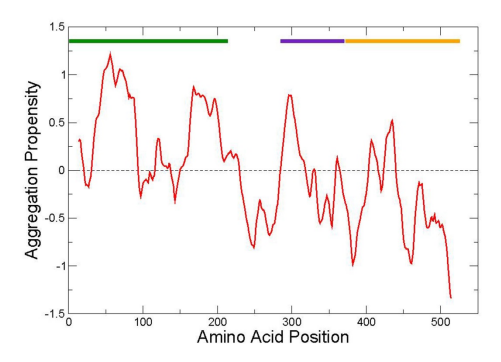


Figure 3: in vivo Mean fluorescence lifetimes are not strongly decreased in FUS gels, but are in conventional amyloid aggregates. Fluorescence lifetimes were measured *in vivo* in wild-type and mutant FUS animals plotted as percentage of GFP-only containing control (blue bars). FUS(P525L) and FUS501 mutant animals are significantly different to

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FUS(WT) animals (mean lifetime  $\pm$  SEM, ANOVA with Scheffe's post hoc test).\*P < 0.05. In comparison, YFP-labelled classical amyloids such as  $\alpha$ -synuclein (not shown) and polyQ40 display >20% fluorescence lifetime drops between soluble and aggregated states (2 way ANOVA, Tukey's multiple comparisons \*\*\*\*P< 0.0001).

# Supplemental Figure 4: predicted aggregation propensity of FUS holoprotein



**Figure 4.** Aggregation propensity profile of FUS (UniProt code: P35637) calculated with the Zyggregator method<sup>1</sup>. The horizontal bars indicate structurally disordered regions located at positions 1-214 (low complexity domain, Gln/Gly/Ser/Tyr-rich; green) and 371-526 (Arg/Gly-rich; orange), as well as the RNA-binding domain at position 285-371 (RRM; purple). The N-terminus shows higher aggregation propensity (residues 1-100) than the C-terminus (residues 450-526). The intrinsic profile is smoothed using a 25-residue window.

## **Aggregation propensity predictions**

The Zyggregator score computes the intrinsic aggregation propensity of proteins, i.e. the propensity of aggregation in the unfolded state<sup>1</sup>. The aggregation propensity score is calculated using the position-dependent score  $p_i^{agg}$ . For a given residue i, the  $p_i^{agg}$  value is calculated as

$$p_i^{agg} = \alpha_h p_h + \alpha_s p_s + \alpha_{hyd} p_{hyd}$$

where  $p_h$  and  $p_i$  are the propensities for  $\alpha$ -helix and  $\beta$ -sheet formation, respectively, and  $p_{hyd}$  is the hydrophobicity<sup>1</sup>. The  $p_i^{agg}$  values are combined to provide a profile,  $A_i^p$ , which

describes the intrinsic propensity for aggregation as a function of the amino acid sequence<sup>2,3</sup>. At each position i along the sequence we define the profile  $A_i^p$  as an average over a window of seven residues

$$A_{i}^{p} = \frac{1}{7} \sum_{j=-3}^{3} p_{i+j}^{agg} + \alpha_{pat} I_{i}^{pat} + \alpha_{gk} I_{i}^{gk}$$

where  $I_i^{pat}$  is the term that takes into account the presence of specific patterns of alternating hydrophobic and hydrophilic residues<sup>4</sup> and  $I_i^{gk}$  is a term that takes into account the gatekeeping effect of individual charges  $c_i^1$ 

$$I_i^{gk} = \sum_{j=-10}^{10} c_{i+j}$$

The parameters  $\alpha$  are fitted according to the procedure described by DuBay *et al.* <sup>1,2</sup>. In order to compare the intrinsic propensity profiles we normalize  $A_i^p$  by considering the average  $(\sigma_A)$  and the standard deviation  $(\sigma_A)$  of  $A_i^p$  at each position *i* for random sequences. We thus obtain the normalized intrinsic aggregation propensity profile

$$Z_i^{agg} = \frac{A_i^p - \mu}{\sigma}$$

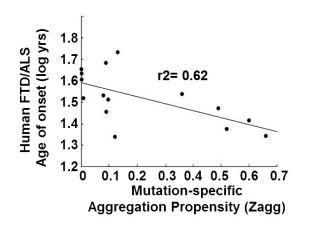
where we calculated the average  $\mu$  and the standard deviation  $\sigma$  over random sequences

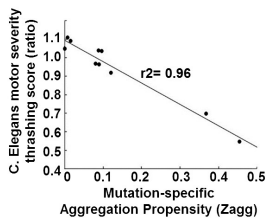
$$\mu = \frac{1}{(N-8)N_s} \sum_{k=1}^{N_s} \sum_{i=4}^{N-4} A_i^p(S_k) \qquad \sigma^2 = \frac{1}{(N-8)N_s} \sum_{k=1}^{N_s} \sum_{i=4}^{N-4} \left( A_i^p(S_k) - \mu \right)^2$$

In these formulas we used  $N_s$  random sequences of length N, and we verified that  $\mu$  and  $\sigma$  are essentially constant for values of N ranging from 50 to 1000.

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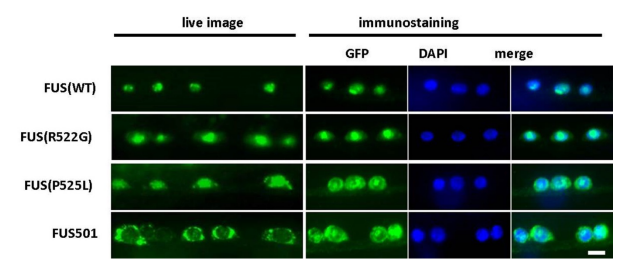
# Supplemental Figure 5: Correlations of aggregation propensity and disease severity



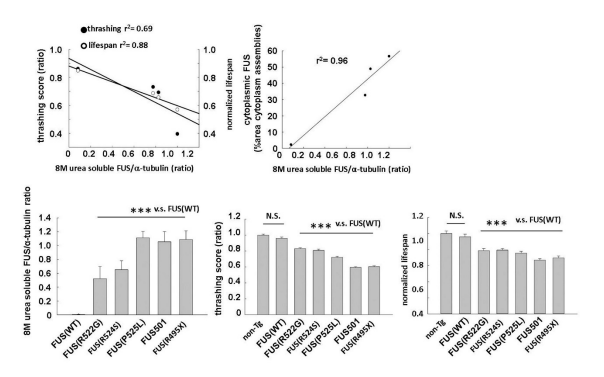


**Left panel:** The severity of FTLD/MND induced by each FUS mutation (as measured by the age-of-onset) shows a weak correlation with the aggregation propensity of that mutation (rR=0.62). The weak correlation likely reflects both the effects of the outbred genetic background of human populations, and differences in presentation (bulbar versus limb weakness etc), which profoundly affect the clinical outcome in humans with ALS. **Right panel:** The severity of neuronal dysfunction induced by each FUS mutation (as measured by the thrashing score), is strongly correlated with the aggregation propensity of that mutation (r=0.98).

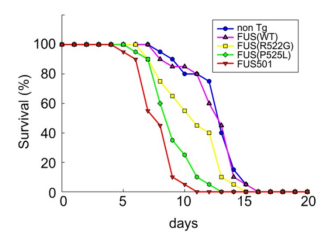
# Supplemental Figure 6: mutant FUS protein mislocalises in the cytoplasm in *C. elegans*.



# Supplemental Figure 7 and 8: Basal data of wild-type and mutant FUS strains.



Supplemental Figure 7: Basal data of Wild-type and mutant FUS strains.

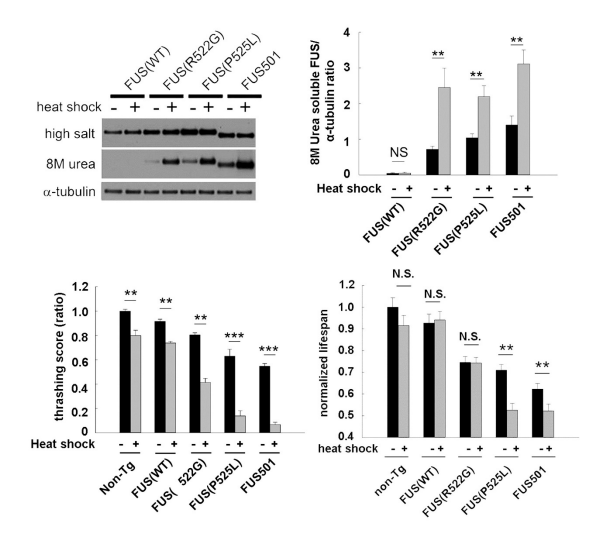


**Supplemental Figure 8: Baseline survival curve.** All mutant FUS strains show shorter life spans compared to FUS(WT) strain, whereas lifespans of FUS(WT) are not different from N2 strain.

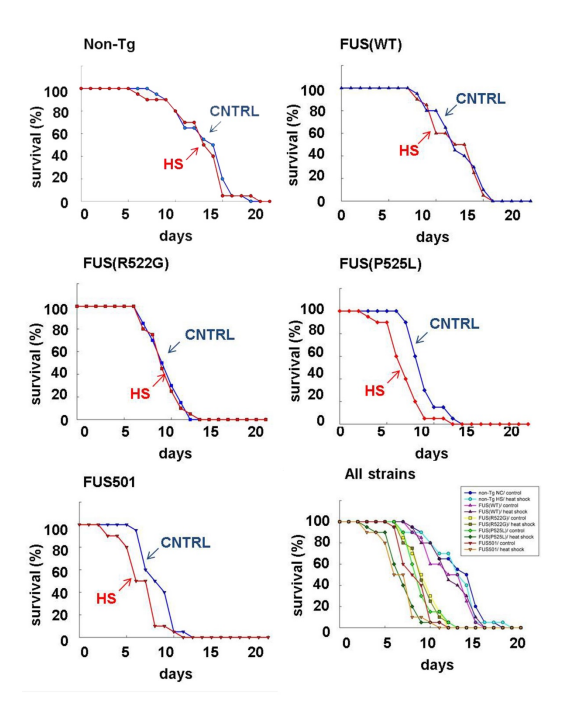
# Supplemental Figure 9 and 10: The severity of neurotoxic phenotypes can be increased or decreased by manipulating the abundance of 8M Urea soluble assemblies in *C. elegans*

Repeated mild heat stress induces increased abundance of FUS assemblies in neuronal cytoplasm (not shown) and of 8M urea soluble FUS assemblies in animals expressing mutant FUS but not wild-type FUS. The presence of more FUS assemblies is accompanied by a marked increase in neurotoxicity as manifest by significantly decreased thrashing scores compared to genotypic controls and to wild-type controls and reduced longevity. Note that heat shock induces a mild, non-specific decrease in motor performance in both non-transgenic and FUS(WT) animals. However, this reduction is proportionately much less than the decrements observed in animals expressing mutant FUS. Similarly, although the increase in 8M urea soluble FUS assemblies in the FUS(R522G) animals is sufficient to cause an increase in motor dysfunction, the phenotype in these animals is milder and it does not increase mortality. N.S. = not significant, \*\*P < 0.01, \*\*\*P < 0.001. Non-transgenic animals, and animals expressing wild-type or mutant FUS were treated by the addition of a small molecule anti-aggregant (scylloinositol) to the culture plates (grey bars) or buffer (black bars). Scylloinositol had no effect on FUS localisation, 8M urea soluble FUS assembly or motor function and longevity in non-transgenic animals or expressing wildtype FUS. In animals expressing mutant FUS, scylloinositol reduces cytoplasmic FUS (not shown), and reduces the abundance of 8M urea soluble FUS assemblies. This reduction is accompanied by significantly improved motor function, which in the case of FUS (R522G) and FUS(P525L) expressing animals almost approximates the motor performance of FUS(WT) and N2 animals. The reduction in 8M urea soluble FUS assemblies is accompanied by improved lifespan in the FUS(P525L) and FUS501 mutant expressing animals. Scylloinositol had no effect on the already essentially normal lifespan of the mildly affected FUS (R522G) animals. FUS501 animals were crossbred with animals expressing hsp-70. Overexpression of hsp-70 in the FUS501 animals resulted in significant reduction in 8M urea soluble FUS assemblies (3C, left panel) accompanied by significant improvement in motor function (3C, right panel) and longevity (3D). Overexpression of hsp-70 alone had no effect on these parameters. Because of the site of integration of the hsp-70 transgene on the same chromosome as the FUS(WT) and FUS(P525L) transgenes, we were unable to generate double transgenic animals for these lines. N.S. = not significant, \*\*P < 0.01, \*\*\*P < 0.001.

Figure 9: Heat Shock increases 8M urea soluble assemblies and neurotoxicity



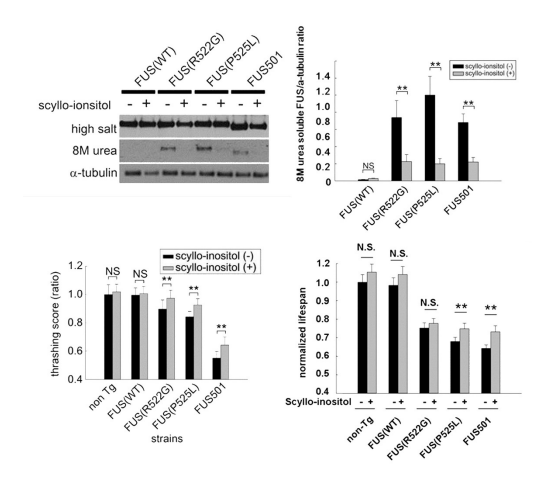
## Kaplan-Meier curves for heat-shock



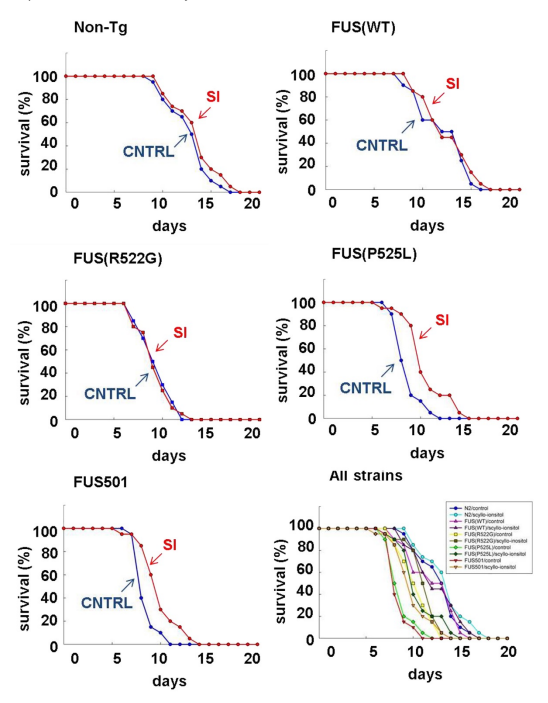
\*CNTRL; control, HS; heat shock

Figure 10: Scyllo-inositol and over-expression of HSP70 decrease 8M urea assemblies and reduce neurotoxicity

# <Scyllo-inositol>

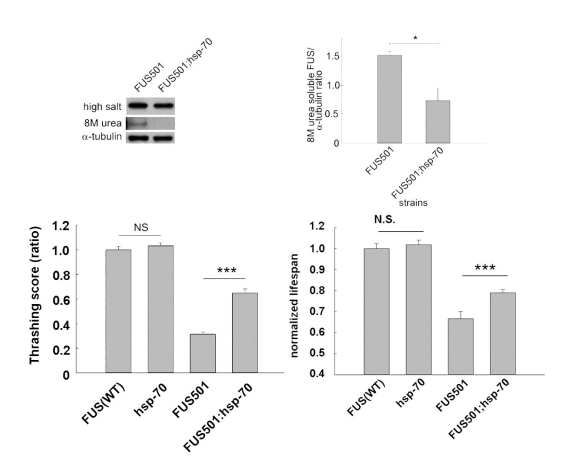


# Kaplan-Meier curves for scyllo-inositol

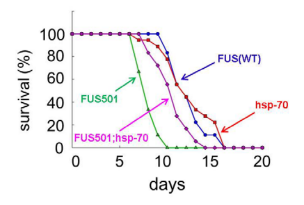


\*CNTRL; control, SI; scyllo-inositol

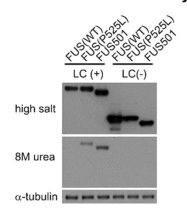
## <hsp-70>

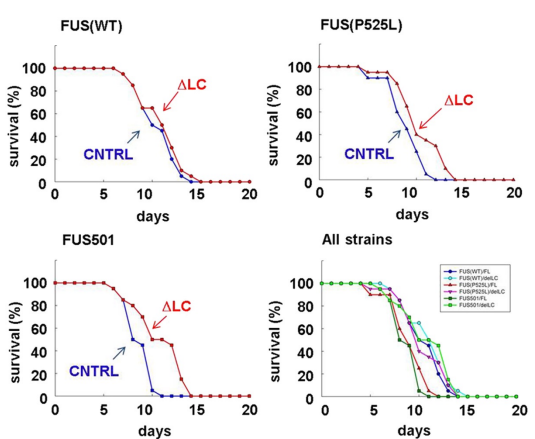


# Kaplan-Meier curves for hsp-70



# Supplemental Figure 11: Deletion of LC domain decrease 8M urea assemblies and reduce neurotoxicity

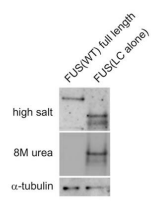




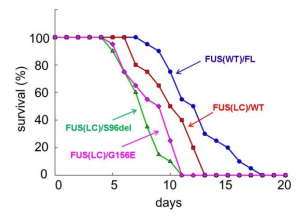
Kaplan-Meier curves for delLC strains

\*CNTRL; control,  $\Delta$ LC; strains overexpressing delLC motif.

# Supplemental Figure 12: Overexpresison of LC domain increases 8M urea soluble assemblies and neurotoxicity

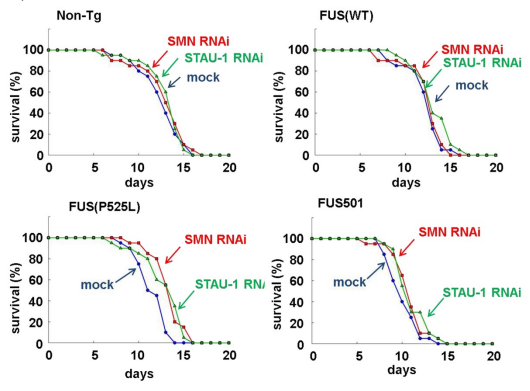


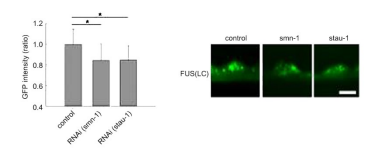
# Kaplan-Meier curves for LC mutants



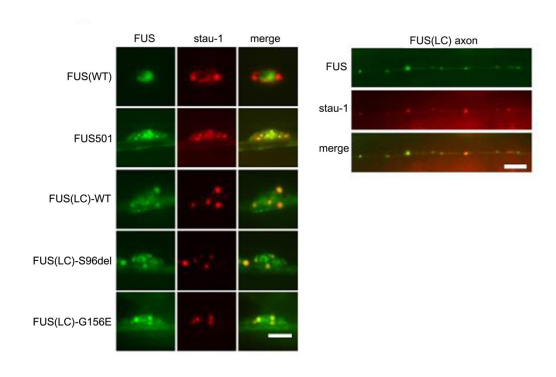
# Supplemental Figure 13: smn1 and stau-1 are structural and stabilizing components of mutant pathological FUS granules

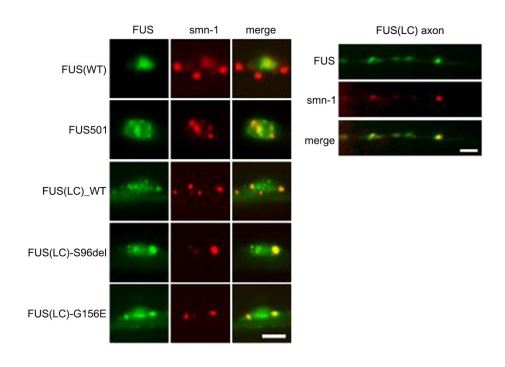
Kaplan-Meier curves for smn1 and sta1 RNAi



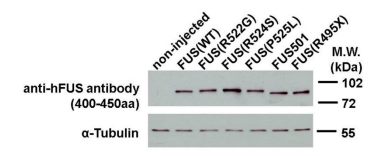


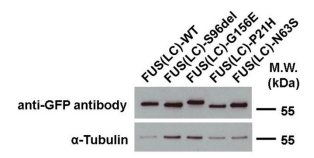
# Supplmental Figures 14: SMN, STAU-1 co-localise with pathological mutant FUS aggregates in the cytoplasm of C elegans neurons.





# Supplemental Figure 15: Mutant FUs reduces new protein synthesis in neuronal axons and termini.





Western blotting of *Xenopus* retinal neurons overexpressing wild-type and mutant FUS proteins.