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Sidechain hydrogen bonding interactions within amyloid-like fibrils formed by the low-complexity domain of FUS: Evidence from solid state nuclear magnetic resonance spectroscopy

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Abstract

In aqueous solutions, the 214-residue low-complexity domain of the FUS protein (FUS-LC) is known to undergo liquid-liquid phase separation and also to self-assemble into amyloid-like fibrils. In previous work based on solid state nuclear magnetic resonance (ssNMR) methods, a structural model for the FUS-LC fibril core was developed, showing that residues 39–95 form the fibril core. Unlike fibrils formed by amyloid- β peptides, α -synuclein, and other amyloid-forming proteins, the FUS-LC core is largely devoid of purely hydrophobic amino acid sidechains. Instead, the core-forming segment contains numerous hydroxyl-bearing residues, including 18 serines, 6 threonines, and 8 tyrosines, suggesting that the FUS-LC fibril structure may be stabilized in part by inter-residue hydrogen bonds among sidechain hydroxyl groups. Here we describe ssNMR measurements, performed on ²H,¹⁵N,¹³C-labeled FUS-LC fibrils, that provide new information about the interactions of hydroxyl-bearing residues with one another and with water. The ssNMR data support the involvement of specific serine, threonine, and tyrosine residues in hydrogen-bonding interactions. The data also reveal differences in hydrogen exchange rates with water for different sidechain hydroxyl groups, providing information about solvent exposure and penetration of water into the FUS-LC fibril core.

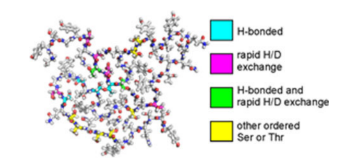
Graphical Abstract

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Supporting Information

Measurement conditions for ssNMR spectra (Table S1); signal tables for mcassign3a assignment calculations (Tables S2–S9); chemical shift assignments for ²H, ¹⁵N, ¹³C-FUS-LC fibrils (Table S10); Supporting Figures S1–S10.



Introduction

Numerous proteins that participate in DNA transcription and RNA processing possess lowcomplexity (LC) domains, in addition to nucleic acid-binding and other functional domains. The LC domains have simple amino acid compositions, can have quasi-repetitive sequences, and are unstructured as monomers. Examples of proteins with relatively long LC domains include FUS, TDP43, and members of the heterogeneous nuclear ribonucleoprotein (hnRNP) family.

The self-assembly behavior of proteins with LC domains is of current interest for several reasons. These proteins typically function as multimers, with their LC domains being essential for self-association and for association of different LC-domain proteins with one another^{2,3}. These proteins are also components of protein/nucleic acid condensates within cells, such as stress granules, P granules, and nuclear paraspeckles^{4,5}, with their LC domains contributing to localization to such condensates^{6–11}. In vitro, proteins containing long LC domains, or LC domains by themselves, exhibit liquid-liquid phase separation (LLPS), forming "droplets" of highly concentrated protein solution, surrounded by a dilute protein solution, at appropriate temperatures and total protein concentrations^{12–15}. The phaseseparated droplet state can be metastable, converting over time to a suspension of amyloidlike protein fibrils, which can behave as a hydrogel through entanglement of long fibrils^{9,16–18}. Finally, fibril formation by LC-domain proteins within cells can be a pathological protein aggregation event, possibly occurring by nucleation of fibril growth within otherwise functional phase-separated condensates^{10,19,20}. Intracellular aggregation of FUS, TDP43, and hnRNPs is associated with amyotrophic lateral sclerosis and other neurodegenerative diseases 21-26.

As a step towards understanding the self-assembly behavior of LC domains at the molecular level, we recently reported a molecular structural model for fibrils formed in vitro by the 214-residue LC domain of FUS (FUS-LC)¹, shown in Figs. 1a and 1b (see also Protein Data Bank file 5W3N). This model is based on a large set of solid state NMR (ssNMR) data, supplemented by information from transmission electron microscopy. The FUS-LC fibril structure has a number of intriguing properties: (i) A structurally ordered fibril core is formed by residues 39–95, while most segments outside this core remain dynamically disordered; (ii) FUS-LC fibrils are not polymorphic, with the same core structure being obtained in multiple independent sample preparations and with no evidence for detectable populations of alternative core structures or alternative core-forming segments; (iii)

Although residues 39–95 form the fibril core, the amino acid composition of this segment (21.0% Gly, 31.6% Ser, 17.5% Gln, 14.0% Tyr, 10.5% Thr, 1.8% Pro, 3.5% others) is nearly the same as the overall composition of FUS-LC (24.9% Gly, 23.9% Ser, 20.2% Gln, 12.6% Tyr, 5.1% Pro, 4.7% Thr, 9.0% others); (iv) Only one residue in the core, namely Pro72, has an entirely hydrophobic sidechain. These properties make FUS-LC fibrils qualitatively different from amyloid- β^{27-31} and α -synuclein fibrils^{32–34}, for example, which contain larger fractions of structurally ordered residues, are polymorphic, and have core-forming segments in which roughly 40% of the amino acids are hydrophobic (*i.e.*, Ala, Val, Leu, Ile, Phe, or Met).

For FUS-LC fibrils, it is currently unclear why segments other than residues 39–95, with nearly identical amino acid compositions, do not form alternative core structures with similar thermodynamic stabilities and kinetic accessibilities, leading to polymorphism. It is also unclear why residues 39–95 adopt a unique structure in FUS-LC fibril cores, with a unique backbone conformation and a unique set of sidechain-sidechain contacts. Finally, while hydrophobic interactions clearly stabilize the structures of amyloid- β and α -synuclein fibrils, interactions that stabilize the FUS-LC fibril core structure are unclear. More precisely, intermolecular hydrogen bonds between backbone carbonyl and amide groups provide stabilization along the fibril growth direction, as in any cross- β fibril structure³⁵. Intermolecular hydrogen bonds among sidechain amide groups of Gln and Asn residues, called polar zipper interactions^{36,37}, may also contribute to stability along the fibril growth direction in the in-register parallel cross- β structure that was established by experimental ssNMR data and mass-per-length measurements¹ (although it should be noted that residues 39-95 are not especially Gln- or Asn-rich). However, sidechain-sidechain interactions in planes perpendicular to the fibril growth direction, which ultimately determine the "fold" of the core, remain mysterious.

Insights into sidechain-sidechain interactions that may play a role in the FUS-LC core structure come from crystal structures of short peptides in cross- β motifs, including peptides from FUS-LC and other LC sequences^{38–41}. For example, a crystal structure of the hexapeptide SSTSAA (Fig. 1c) shows inter- β -sheet hydrogen bonding between sidechain hydroxyl groups of Ser1 and Ser2, as well as water-mediated hydrogen bonding between sidechain hydroxyl groups of Ser2 and Thr3 (*i.e.*, hydrogen bonds from Ser2 and Thr3 to the same single-file row of water molecules). A crystal structure of SYSGYS (Fig. 1d) shows water-mediated hydrogen bonding between sidechain hydroxyl groups of Ser39 and Tyr41. Additional examples of direct sidechain-sidechain, water-mediated sidechain-sidechain, and sidechain-backbone hydrogen bonding from cross- β crystal structures are shown in Fig. S2.

Do similar interactions exist in FUS-LC fibrils, thereby stabilizing the core structure in planes perpendicular to the fibril growth direction? In this paper, we present results from additional ssNMR experiments that provide information about the possible involvement of specific sidechain hydroxyl groups in stabilizing interactions within the FUS-LC fibril core, and that also indicate differential accessibilities of sidechain hydroxyl groups for hydrogen exchange with water. The ssNMR measurements were performed on a sample that was highly deuterated except at hydroxyl, amide, amine sites (*i.e.*, exchangeable sites), thereby allowing relatively sharp ¹H ssNMR signals to be observed and assigned.

Materials and Methods

Preparation of ²H,¹⁵N,¹³C-labeled FUS-LC fibrils

Unless indicated otherwise, chemicals were obtained from Sigma-Aldrich and solutions were 0.22 μ m sterile filtered. N-terminally His₆-tagged residues 2–214 of FUS-LC was expressed in BL21(DE3) pLysS *E. coli* cells (ThermoFisher Scientific) using a pHis-parallel vector.⁴² The sequence of the N-terminal tag was

MSYYHHHHHHDYDIPTTENLYFQGAMDP, which was followed by the sequence in Fig. 1. (The tag is dynamically disordered in FUS-LC fibrils and does not contribute to ssNMR spectra.) Cells were incubated quiescently in 5 ml unfiltered Luria-Bertani medium at 37° C for 4 h. Then 200 μ l of the culture was transferred to 2 ml of M9 medium and shaken at 37° C for 3.5 h. The 2.2 ml culture was then added to 50 ml of M9 medium and shaken overnight at 37° C. The following day, the 50 ml overnight culture was added to 950 ml of M9 medium and shaken at 37° C for 5 h. The cells were transferred to 20° C and shaken for 1 h (optical density at 600 nm equal to 1.0 ± 0.1) before adding isopropyl β -D-1thiogalactopyranoside (IPTG) to a final concentration of 0.5 mM. The culture was then shaken for 21 h at 20° C before harvesting the cells, flash freezing them in liquid nitrogen, and storing them at -80° C. All culture media contained 100 µl/ml ampicillin. For isotopic labeling of FUS-LC, M9 salts were prepared in 99.9% D₂O (Cambridge Isotope Labs) supplemented with 1x MEM vitamin mix (Corning), 1.0 g/l¹⁵N ammonium chloride (Cambridge Isotope Labs), 2.0 g/l ¹³C₆,²H₇-D-glucose, and 0.3 g/l ²H,¹³C, ¹⁵N-Isogro. The MEM vitamin mix was lyophilized and resuspended in 99.9 % D₂O before use. Ampicillin and IPTG stock solutions were prepared at 1000-fold higher concentration than the working solution in 99.9% D₂O. Unfiltered 100x MEM stock solutions were frozen in liquid nitrogen, lyophilized overnight, and resuspended in 99.9% D₂O prior to use.

The frozen cell pellet was resuspended in 40 ml of pH 7.9 lysis buffer containing 50 mM tris(hydroxymethyl)aminomethane (KD Medical), 500 mM sodium chloride (KD Medical), 6 M guanidine HCl, 1% w/w Triton X-100 (American Bio), and 4 cOmplete Protease Inhibitor Cocktail tablets (Roche). Then 25 mg of hen egg white lysozyme powder and 4 μ l of Pierce Universal Nuclease were added and the suspension was sonicated on ice with Branson 250 Sonifier using a 1/4" diameter tip, an output power of 0.3 and 10% duty cycle. The mixture was rotated at 4° C for 10 min and then centrifuged at 207,000 × g for 30 min at 4° C. The supernatant was mixed with 5 ml of NiNTA resin (Qiagen) in pH 7.4 equilibration buffer containing 20 mM sodium phosphate, 8 M urea, 500 mM sodium chloride, and 5 mM 2-mercaptoethanol before rotating at 4 °C for 30 min. The resin was loaded onto a gravity flow column (Kimble-Chase) and washed with equilibration buffer containing 0.1 mM phenylmethanesulfonyl fluoride and 20 mM imidazole until the 280 nm absorbance of the elute reached baseline. The protein was eluted using equilibration buffer with 200 mM imidazole and 7.6 M urea.

The elution fraction containing $34 \ \mu\text{M}^2\text{H}$, ^{15}N , ^{13}C -FUS-LC was concentrated to $300 \ \mu\text{M}$ using a 0.5 ml Amicon Ultra centrifugal filter with 10 kDa cutoff (Millipore). About 175 μ l of the protein solution was loaded into a 0.5 ml Slide-A-Lyzer dialysis cassette with 3.5 kDa cutoff (ThermoFisher Scientific) and dialyzed against 2.0 l of 20 mM sodium phosphate, pH

7.4. After dialysis for 19 hr about 300 μ l of the protein solution was collected and allowed to sit quiescently at room temperature (approximately 22° C). After a seven-day incubation period, a mixture of fibrillar and amorphous assemblies was observed by transmission electron microscopy (TEM). The solution was then flash frozen in liquid nitrogen and thawed seven times to break up the protein aggregates, creating fragments that serve as fibril seeds. An additional 2.3 ml of ²H,¹⁵N,¹³C-FUS-LC protein was concentrated from 31 μ M to 42 μ M, then dialyzed in a 3 ml Slide-A-Lyzer dialysis cassette with 3.5 kDa cutoff against 2.0 l of 20 mM sodium phosphate, pH 7.4, for 24 h. A 1.7 ml volume of protein solution was recovered from the dialysis cassette and mixed with the seeds. The mixture was incubated quiescently at room temperature for eight days. The sample then contained almost exclusively fibrillar FUS-LC assemblies, with no significant fraction of amorphous aggregates in TEM images. Finally, the fibrils were harvested by centrifugation at 279,000 × g for 20.5 h and packed into a magic-angle spinning (MAS) NMR rotor with 1.8 mm outer diameter and 10 μ l sample volume by centrifugation at 15,000 × g for 1.0 h at 4° C in a home-made centrifugal funnel device.

Electron microscopy

TEM images were obtained with a FEI Morgagni microscope, equipped with a side-mounted Advantage HR camera (Advanced Microscopy Techniques) and operating at 80 keV. Grids were home-made carbon films supported on lacey carbon on 300 mesh copper (Electron Microscopy Sciences) and were glow-discharged immediately before sample application. A 5 μ l aliquot of fibril solution, diluted by a factor of 10, was adsorbed to the carbon film for two minutes, blotted, washed with 5 μ l of water for 10 s, blotted, washed again, blotted, stained with 5 μ l of 3% uranyl acetate for 10 s, blotted, and dried in air.

Solid state NMR

ssNMR measurements were performed at 17.5 T (746.2 MHz ¹H NMR frequency) and 14.1 T (599.2 MHz ¹H NMR frequency), using Varian Infinity and InfinityPlus spectrometers, respectively, and triple-channel MAS ssNMR probes with 1.8 mm rotor diameters produced by the laboratory of Prof. Ago Samoson (NMR Institute and Tallinn Institute of Technology, Estonia). Except as noted below, ssNMR data were acquired at a MAS frequency of 40.00 kHz. Sample temperatures were maintained at approximately 28° C, as indicated by the temperature-dependent ¹H chemical shift of water in the sample, using -50° C nitrogen gas at the cooling gas input of the probe (17 standard liters per minute flow rate) to counteract frictional heating from high-frequency MAS. Radio-frequency (RF) pulse sequences for three-dimensional (3D) spectra are shown in Fig. S1. ¹H, ¹³C, or ¹⁵N decoupling used the XiX technique⁴³, with RF fields of 6-10 kHz. ¹³C-¹³C and ¹H-¹H polarization transfers used the finite-pulse radio-frequency-driven recoupling (fpRFDR) technique^{44,45}. ¹H-¹⁵N. ¹⁵N-¹H, and ¹H-¹³C polarization transfers used cross-polarization (CP)⁴⁶, with ramped RF amplitudes as depicted in Fig. S1. A 3D ¹⁵N-¹³C-¹³C spectrum used non-selective ¹⁵N-¹³C CP transfers, so that NCACX and NCOCX crosspeaks were contained in the same spectrum. 3D CANH, CAN(H)H, CONH, CON(H)H, and HN(CO)CX spectra⁴⁷⁻⁴⁹ used frequencyselective ¹³C-¹⁵N or ¹⁵N-¹³C CP transfers, with the ¹³C carrier frequency shifted to the 13 CO or 13 C_g regions of the spectrum.⁵⁰ In ¹H-detected measurements, strong signals from water were attenuated by a train of $\pi/2$ pulses before the final CP transfer to ¹H nuclei.

Typically, $n_{sat} = 20$ pulses separated by 15–25 ms delays were used, as depicted in Fig. S1. Other experimental parameters for individual 3D spectra are given in Table S1 and in figure captions. Two-dimensional (2D) and 3D data were processed with NMRpipe software⁵¹ and analyzed, displayed, and plotted with Sparky software (available at https://www.cgl.ucsf.edu/home/sparky/).

Water-filtered 2D ¹³C-¹³C spectra,^{52,53} used to characterize hydrogen exchange between water and sidechain hydroxyl groups of Ser and Thr residues, were acquired at 14. 1 T with MAS at 13.60 kHz. The sample temperature was maintained at 26° C. The RF pulse sequence for these measurements is shown in Fig. S9a. A ¹H spin echo period $\tau_{echo} = 2.0$ ms was used to suppress ¹H polarization at immobilized sites on the protein, which have short T₂ relaxation times at the relatively low MAS frequency, and to select ¹H polarization of water, which has a much longer T₂ value. The spin echo period was followed by a variable ¹H-¹H exchange period t_{exch} during which polarization could be transferred to sidechain hydroxyl sites (and other exchangeable sites). Hydroxyl ¹H polarization was then measured through the intensities of ${}^{13}C_{\beta}/{}^{13}C_{\alpha}$ (and other) crosspeaks, which were quantified with the seriesTab program of NMRpipe. Two-pulse phase-modulated (TPPM) proton decoupling⁴⁴ with an RF amplitude of 85 kHz was applied during t₁ and t₂ periods. A 50 ms ${}^{13}C_{-13}C$ mixing period with dipolar-assisted rotational resonance⁵⁴ (DARR) was used.

The reference for ¹³C chemical shifts in this paper is nominally DSS (4,4-dimethyl-4silapentane-1-sulfonic acid). However, the ¹³C chemical shift reference was adjusted to make ¹³C_a shifts in ²H,¹⁵N,¹³C-FUS-LC fibrils agree optimally with our published ¹³C_a shifts for ¹⁵N,¹³C-FUS-LC fibrils¹. Due to deuterium isotope effects⁵⁵, ¹³C chemical shifts for carbonyl, carboxyl, methylene, and methyl sites in ²H,¹⁵N,¹³C-FUS-LC fibrils are then expected to differ from those in ¹⁵N,¹³C-FUS-LC fibrils by approximately +0.3 ppm, +0.3 ppm, -0.3 ppm, and -0.6 ppm, respectively. Our ¹H chemical shift reference is such that the chemical shift of water is 4.50 ppm. The nominal ¹⁵N chemical shift reference is liquid ammonia.

Chemical shift assignment calculations

After manual peak-picking of 3D solid ssNMR spectra in Sparky, assignments were generated with the Monte Carlo/simulated annealing approach described previously⁵⁶. A modified version of the assignment program, called mcassign3a, was used. The modified program allows inter-residue crosspeaks with ambiguous connection directions to be used. This capability was necessary for inclusion of 3D CAN(H)H, CON(H)H, and NHH spectra in the assignment calculations, since 1 H- 1 H spin polarization transfers can in principle occur with equal likelihood from the amide proton of residue *k* to the amide protons of either residue *k*+1 or residue *k*-1. The modified program also allows residue type assignments to be specified for two successive residues that are involved in an inter-residue crosspeak signal, so that the signal can only be assigned to a given residue *j* if both residue *j* and residue *j*+1 match the specified residue type assignments. This second modification in mcassign3a was not used in the current work but is helpful in other contexts.

Assignment calculations proceeded in an iterative fashion, as described previously¹, with a gradual increase in the number of signals with definite assignments. Signal assignments

were restricted to residues 31-110 of the full-length FUS-LC sequence, based on our previous identification of residues 39-95 as the structurally ordered core. The initial calculations used signal tables from 3D NCACX, NCOCX, HN(CO)CX, CANH, and CONH spectra, with 60, 52, 62, 81, and 104 signals in each table, respectively. Since 3D CANH and CONH spectra were recorded at both 14.1 T and 17.5 T, the signal tables for these spectra were generated from crosspeaks in spectra at both field strengths. The relatively large numbers of signals in these two tables is attributable to non-negligible interresidue CA-N, non-sequential CO-N, and inter-residue N-H polarization transfers, which produce extra crosspeaks. All other signal tables were generated from spectra at 14.1 T. Signal tables were updated after each set of calculations to include any new definite assignments (defined as the same assignment for a given signal in 10 out of 10 independent mcassign3a runs). After three sets of assignment calculations, signal tables from 3D CAN(H)H, CON(H)H, and NHH spectra were added, with 28, 17, and 39 inter-residue crosspeak signals, respectively. Each of these spectra was represented by two signal tables, which for example allowed two NHH crosspeaks (one from residue k to residue k-1, the other from residue k to residue k+1) to be assigned to the same residue. After one set of calculations with the full set of signal tables, signals that were never assigned (in 10 out of 10 independent runs) were removed from all tables, leaving 60, 52, 59, 67, 64, 28, 17, and 39 signals from 3D NCACX, NCOCX, HN(CO)CX, CANH, CONH, CAN(H)H, CON(H)H, and NHH spectra. The final set of calculations was then performed, with 50 simulating annealing steps in each of 10 runs, with 10⁸ Monte Carlo moves (*i.e.*, attempted assignment changes) in each step. The final signal tables are given in Tables S2–S9.

A significant number of signals remained unassigned, specifically 9, 10, 13, 17, 10, 5, 2, and 11 signals from the final 3D NCACX, NCOCX, HN(CO)CX, CANH, CONH, CAN(H)H, CON(H)H, and NHH signal tables, respectively. We can attribute these signals to short segments or individual residues outside the main FUS-LC fibril core segment that are immobilized or partially immobilized through contacts with the core, as discussed previously¹. The repetitive, low-complexity nature of the FUS-LC sequence makes signals from isolated short segments difficult to assign unambiguously.

Results

Chemical shift assignments for ²H,¹⁵N,¹³C-FUS-LC fibrils

Chemical shift assignments were determined from ¹³C-detected and ¹H-detected 3D ssNMR spectra, recorded with MAS at 40.00 kHz, using a Monte Carlo/simulated annealing algorithm to assign multidimensional signals from the 3D spectra to specific amino acids (see Materials and Methods). Although ¹⁵N and ¹³C chemical shift assignments for ¹⁵N,¹³C-FUS-LC fibrils were reported previously, the assignment process was repeated independently for ²H,¹⁵N,¹³C-FUS-LC fibrils to obtain ¹H chemical shift assignments and to determine the changes in ¹³C chemical shifts that are induced by deuteration.

Fig. 2 shows examples of 2D planes from ¹³C-detected 3D spectra to illustrate the resolution and signal-to-noise ratios of these spectra. ¹H, ¹³C, and ¹⁵N linewidths (full widths at half-maximum, FWHM) were 0.20–0.25 ppm, 0.6–1.0 ppm, and 0.9–1.3 ppm, respectively. 3D NCACX and NCOCX spectra in Figs. 2a and 2b were obtained from a single 3D

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¹⁵N-¹³C-¹³C measurement, using the RF pulse sequence in Fig. S1a with non-selective ¹⁵N-¹³C cross-polarization between t₁ and t₂ periods. These spectra primarily correlate the backbone amide ¹⁵N chemical shift of residue *k* with the ¹³C_a or ¹³CO chemical shift of residue *k* or *k* – 1, and then with sidechain ¹³C chemical shifts of residue *k* or *k* – 1. The 3D HN(CO)CX spectrum in Fig. 2c, obtained with the RF pulse sequence in Fig. S1b, primarily correlates the backbone amide ¹H chemical shift of residue *k* with the backbone amide ¹⁵N chemical shift of residue *k* and then with backbone ¹³CO, ¹³C_a, and sidechain ¹³C chemical shifts of residue *k* – 1. Additional 2D planes from these 3D spectra are shown in Fig. S3.

Fig. 3 shows examples of 2D planes from ¹H-detected 3D spectra. 3D CANH and CONH spectra in Figs. 3a and 3b (left side) primarily correlate the ¹³C_a chemical shift of residue k or the ¹³CO chemical shift of residue k - 1 with the backbone amide ¹⁵N chemical shift of residue k, and then with the backbone amide ¹H chemical shift of residue k. 3D CAN(H)H and CON(H)H spectra in Figs. 3a and 3b (right side) allow correlations to backbone amide ¹H chemical shifts of residues k - 1 and k + 1 in the third dimension. These spectra were obtained with the RF pulse sequence in Fig. S1c. The 3D NHH spectrum in Fig. 3c, obtained with the RF pulse sequence in Fig. S1d, primarily correlates the backbone amide ¹⁵N chemical shift of residue k, and then with the backbone amide ¹H chemical shift of residue k, and then with the backbone amide ¹H chemical shift of residue k and then spectrum in Fig. 3c, obtained with the RF pulse sequence in Fig. S1d, primarily correlates the backbone amide ¹⁵N chemical shift of residue k with the backbone amide ¹H chemical shift of residue k, and then with the backbone amide ¹H chemical shift of residue k, and then with the backbone amide ¹H chemical shift of residue k, and then with the backbone amide ¹H chemical shift of residue k, and then with the backbone amide ¹H chemical shift of residue k, and then with the backbone amide ¹H chemical shift of residue k, and then with the backbone amide ¹H chemical shift of residue k. 3D planes from these 3D spectra are shown in Fig. S4.

A simple 2D ¹H-¹⁵N spectrum of ²H,¹⁵N,¹³C-FUS-LC fibrils is poorly resolved, showing only a small number of interpretable crosspeaks (see Fig. S5). With our fibril growth conditions, we expect all exchangeable hydrogen sites (*i.e.*, amide, amine, and hydroxyl sites) to be protonated. This means that the overall deuteration level in residues 39–95 of FUS-LC is roughly 68%. Our MAS frequency of 40.00 kHz is then not sufficient to reduce ¹H ssNMR linewidths to the point where a well-resolved 2D ¹H-¹⁵N spectrum is obtained.

From the 3D spectra, definite assignments were found for residues 41–55, 63, 65–70, 72–87, 89–94, and 107–110, as given in Table S10 (Biological Magnetic Resonance Bank code 50026). For the most part, chemical shift assignments for ²H, ¹⁵N, ¹³C-FUS-LC fibrils are in good agreement with previously published assignments from spectra of ¹⁵N, ¹³C-FUS-LC fibrils¹, which included residues 39–40, 44–54, and 63–95. Differences in the identities of residues for which assignments were obtained arise from differences in the types of 3D spectra that were used and the signal-to-noise ratios of the spectra. On the average, ¹⁵N chemical shifts in Table S10 are about 0.4 ppm lower than in previously published assignments. ¹³C chemical shifts also differ somewhat due to isotope effects and differences in referencing.

Chemical shifts assigned to ¹⁵N of G82 and to ¹⁵N, ¹³C_{β}, and ¹³CO of S89 in Table S10 differ significantly from previously published assignments¹. The previously published ¹⁵N assignment for G82 is incorrect. However, S89 signals with chemical shifts in Table S10 were not present in the previously published 3D spectra of ¹⁵N, ¹³C-FUS-LC fibrils, and S89 signals with the previously published chemical shifts are not present in 3D spectra of ²H, ¹⁵N, ¹³C-FUS-LC fibrils. Chemical shift differences for S89 apparently reflect minor structural differences.

Overall, the ssNMR spectra of ²H,¹⁵N,¹³C-FUS-LC fibrils support our earlier determination that the structurally ordered core of FUS-LC fibrils is formed by the segment depicted in Fig. 1b, with most of the rest of the 214-residue LC domain remaining dynamically disordered. These data indicate that residues 107–110 are rigid in the FUS-LC fibrils. Since long-range distance restraints involving residues 107–110 were not obtained, the structural role of this segment is not yet known. Secondary ¹³C chemical shifts do not support β -strand or α -helical secondary structure in this segment. As in our earlier work¹, the ssNMR spectra include signals from outside the core-forming segment, which we can not assign definitively to specific residues. We attribute these signals to short segments (perhaps single residues or pairs of residues) that are partially ordered through interactions with the fibril core, or possibly through interactions between fibrils that are packed together densely in the MAS rotor.

Site-specific correlations with water

Correlations between the ¹H chemical shift of water and certain ¹³C chemical shifts of ²H, ¹⁵N, ¹³C-FUS-LC are apparent in ¹³C-detected 2D ¹H-¹³C and 3D ¹H-¹³C-¹³C spectra. For detecting these correlations, ¹³C detection is preferred over ¹H detection because ¹H-¹³C crosspeaks at or near the ¹H chemical shift of water are unavoidably obscured by large residual water signals in ¹H-detected spectra. In the 2D spectrum (Fig. S6), correlations between the ¹H chemical shift of water and ¹³C chemical shifts of T68 and/or T78, T45 and/or T47, S42 and/or S48, D46, G49, Y75 and/or Y81, P72, and Gln (unassigned) are observed. In 3D spectra, additional correlations with water are detected and ambiguities between T68 and T78 and between T45 and T47 are resolved. The 2D plane from a 3D spectrum with $\tau_{HH} = 0$ (*i.e.*, 3D HCC spectrum, pulse sequence in Fig. S1e), taken at the ¹H chemical shift of water, is shown in Figs. 4a and 4c. Crosspeaks assigned to S42, S44, T45, D46, S48, T68, S83, and S110 are apparent in this 2D plane. Crosspeaks to aromatic carbons of at least one Tyr residue are also observed (Fig. S7). Crosspeak intensities are obviously not symmetric about the ¹³C-¹³C diagonal, indicating stronger nuclear spin polarization transfers from ¹H of water to sidechain carbon sites, including the hydroxyl-bearing ¹³C_B sites of Ser and Thr residues and the carboxyl ${}^{13}C_{\gamma}$ site of D46, than to backbone ${}^{13}C_{\alpha}$ and ¹³CO sites. In the case of Tyr, the strongest polarization transfer is to the hydroxyl-bearing ¹³C_c site (Fig. S7). Similar effects have been seen in previous studies of water/protein interactions by ssNMR^{53,57-60}. We did not observe correlations to distinct water populations as reported by Wang *et al.* in studies of amyloid- β fibrils⁵², possibly because the resolution in the ¹H dimension of our 3D ¹H-¹³C-¹³C spectrum (about 0.2 ppm, limited by the maximum t_1 value) was insufficient to distinguish signals from distinct populations.

We interpret the observed correlations to water in Figs. S6, 4, and S7 as being primarily the result of hydrogen exchange between water and sidechain hydroxyl (or carboxyl) sites during the 4.0 ms τ_{HC} period of the 3D HCC pulse sequence. Near neutral pH and 30° C, intrinsic hydrogen exchange rates for sidechain hydroxyl groups of Ser and Thr are reported to be approximately 400 s⁻¹, while the intrinsic hydrogen exchange rate for the sidechain hydroxyl group of Tyr is much greater⁶¹. Thus, we expect solvent-exposed, relatively "free" sidechain hydroxyl groups of the FUS-LC fibril core structure to exhibit correlations to water. In contrast, sidechain hydroxyl groups that are involved in hydrogen-bonding

interactions within the core are expected to have slower hydrogen exchange rates, leading to weaker correlations to the water ¹H chemical shift for the relevant residues^{57,58,62}. Importantly, we do not observe correlations to water for certain Ser and Thr residues that have ¹³C chemical shift assignments, namely T47, S53, S54, S70, S77, T78, S84, S86, S87, S89, S90, S107, S108, and T109. Of these, S87, S89, and T109 have relatively weak crosspeaks in other types of 3D spectra, suggesting time scales or amplitudes of motion that attenuate their signals. The absence of detectable correlations to water for S87, S89, and T109 is therefore attributable to the overall weakness of signals from these three residues. On the other hand, the absence of detectable correlations to water for T47, S53, S54, S70, S77, T78, S84, S86, S90, S107, and S108 (which have relatively strong signals in other 3D spectra) is attributable to participation of these 11 residues in hydrogen-bonding interactions that reduce the rates of hydrogen exchange rates with water of their sidechain hydroxyl groups. Additional measurements of hydrogen exchange rates are described below.

Figs. 4b and 4d show the 2D plane at the ¹H chemical shift of water from a 3D spectrum with $\tau_{HH} = 2.0$ ms (*i.e.*, 3D H(H)CC spectrum, pulse sequence in Fig. S1e). Compared to results with $\tau_{HH} = 0$, the pattern of ¹³C-¹³C crosspeaks is changed by ¹H-¹H polarization transfers during the τ_{HH} period, especially polarization transfers from sidechain hydroxyl protons to backbone amide protons that lead to stronger ¹³C_a and ¹³CO signals in the t₂ dimension of the 3D spectrum. Crosspeaks from additional residues, namely Q43, G49, Y50, Q52, S53, S54, G82, and Q94, appear. Crosspeaks from the water ¹H chemical shift to ¹³C chemical shifts of Q43, G49, and G82 can be attributed to polarization transfers from the rapidly-exchanging sidechain hydroxyl protons of S42, S48, and S83 discussed above to backbone amide protons, followed by transfers from amide protons to ¹³C_a and ¹³CO sites during the τ_{HC} period. Crosspeaks in the 3D H(H)CC spectrum from the water ¹H chemical shift to ¹³C chemical shifts of Y50, Q52, S53, S54, and Q94 can not be attributed similarly to polarization transfer pathways that involve sidechain hydroxyl sites of preceding residues. Polarization transfer pathways that produce these crosspeaks are unclear.

Correlations to sidechain hydroxyl ¹H chemical shifts

Sidechain hydroxyl ¹H chemical shifts of certain Ser and Thr residues in the FUS-LC fibril core can also be identified from intra-residue crosspeaks in the ¹³C-detected 3D ¹H-¹³C-¹³C spectra. Fig. 5 shows 2D planes from these spectra at sidechain hydroxyl ¹H chemical shifts of S84 (Figs 5a and 5b), S77 (Figs. 5c and 5d), S53 and T45 (Figs. 5e and 5f), and S48 (Figs. 5g and 5h). Additional 2D planes at sidechain hydroxyl ¹H chemical shifts of S83, S86, T47, T78, and T68 are shown in Fig. S8. As discussed above, T45, S48, S83, and T68 also have detectable correlations with the ¹H chemical shift of water, arising from hydrogen exchange between water and their sidechain hydroxyl groups, but the exchange rate is not large enough to prevent observation of the sidechain hydroxyl ¹H chemical shift as a separate signal.

Inter-residue crosspeaks involving sidechain hydroxyl ¹H chemical shifts can provide information about long-range contacts in the FUS-LC fibril core. With $\tau_{HH} = 0$, correlations from the S84 hydroxyl proton to G79 and from the S77 hydroxyl proton to T47 are observed (Figs. 5a, and 5c). With $\tau_{HH} = 2.0$ ms, additional correlations from the S84 hydroxyl to T78,

from the T45 hydroxyl to a Tyr ${}^{13}C_{e}$ site (unassigned), and from the S48 hydroxyl to a Tyr ${}^{13}C_{\zeta}$ (unassigned) are observed (Figs. 5b, 5f, and 5h). These inter-residue crosspeaks are discussed further below.

Hydrogen exchange rates from water-filtered 2D ¹³C-¹³C spectra

Fig. 6a shows the region of water-filtered 2D ${}^{13}C{}^{-13}C$ spectra of ${}^{2}H{},{}^{15}N{},{}^{13}C{}$ -FUS-LC fibrils that contains C_{β}/C_{α} crosspeaks of Thr and Ser residues, with ${}^{1}H{}^{-1}H$ exchange periods t_{exch} equal to 0.1 ms and 100 ms. The RF pulse sequence in Fig. S9a was used to obtain these spectra, in which ${}^{13}C{}$ ssNMR signals arise from ${}^{1}H{}$ polarization that was associated with water molecules before the t_{exch} period and transferred to exchangeable sites on the protein during t_{exch} (or during the 1.5 ms ${}^{1}H{}^{-13}C{}$ cross-polarization period τ_{HC}). 52,53 The primary mechanism for rapid build-up of C_{β}/C_{α} crosspeaks of Thr and Ser residues during t_{exch} is expected to be hydrogen exchange from water to sidechain hydroxyl sites. Patterns of crosspeak intensities are clearly different for the two values of t_{exch} in Fig. 6a, indicating that different residues have different hydrogen exchange rates for their sidechain hydroxyl sites.

Build-up curves for resolved and partially-resolved C_{β}/C_{α} crosspeaks are shown in Fig. 6b and are fit with single-exponential functions $I(t_{exch}) = I_0 + (I_{\infty} - I_0)[1 - \exp(t_{exch}/\tau)]$. Hydrogen exchange rates are then indicated by the fitted values of $1/\tau$. For most Thr and Ser residues, exchange rates determined in this way are 4–10 times slower than the intrinsic rates near neutral pH and 30° C,⁶¹ indicating significant protection from hydrogen exchange. Measured hydrogen exchange rates for S48 and S83 are close to intrinsic rates, consistent with an absence of stable inter-residue hydrogen bonding for sidechain hydroxyl sites of these residues.

Build-up curves for other resolved crosspeaks are shown in Fig. S9b. As expected,⁶¹ buildup of Tyr sidechain crosspeaks (unassigned) is very rapid due to the large intrinsic exchange rates for Tyr sidechain hydroxyl groups. Best-fit values of τ for most CO/C_a and CO/C_β crosspeaks are in the 15–20 ms range, similar to the τ values for sidechain hydroxyl sites of protected Ser and Thr residues (*e.g.*, S84 and S70/S90 in Fig. 6b). Under our experimental conditions, build-up times for ¹³C-¹³C crosspeak intensities in the 15–20 ms range may indicate an overall growth of ¹H polarization within the FUS-LC fibril core, which is likely to result from hydrogen exchange at unprotected sidechain hydroxyl and backbone amide sites combined with ¹H-¹H spin diffusion within the fibril core. This overall growth of ¹H polarization sets an upper limit on τ . Consequently, time scales for hydrogen exchange that may exceed 15–20 ms at specific, highly protected sites can not be determined accurately from our water-filtered 2D ¹³C-¹³C data.

Discussion

Fig. 7 shows two representations of our structural model for the FUS-LC fibril core, with residues color-coded to indicate the presence or absence of correlations to the ¹H chemical shift of water or their involvement in inter-residue crosspeaks. In Fig. 7a, sidechains that lack correlations to water in a 3D HCC spectrum of ²H, ¹⁵N, ¹³C-FUS-LC fibrils are colored cyan. A subset of these residues, including sidechains of T47, S53, S70, S77, T78, S84, S85, and S90, are shielded within the fibril core and may participate in inter-residue hydrogen

bond interactions that would reduce their rates of hydrogen exchange with water. Data in Fig. 6b also indicate protection from hydrogen exchange for T47, S70, S77, S84, and S90. (S53, T78, and S85 do not have sufficiently well-resolved C_{β}/C_{α} crosspeaks to be included in Fig. 6b.) However, sidechains of other residues that lack correlations to water, including sidechains of S54, S87, and S89, are apparently exposed on the FUS-LC fibril surface. As mentioned above, crosspeak signals of S87 and S89 are relatively weak in all 3D spectra, so the absence of detectable correlations to water for these residues may be due to the overall weakness of their ¹³C ssNMR signals, which is presumably a consequence of molecular motions in residues 87–89.

For the most part, sidechains of residues that do exhibit correlations to water (yellow in Fig. 7a) are exposed on the FUS-LC fibril surface or would be exposed after minor adjustments of the structural model depicted in Fig. 7. In this context, it should be noted that sidechains of S44, T45, and D46 are fully accessible to solvent in some models within the bundle of FUS-LC fibril core structures that were found to be consistent with all ssNMR restraints (see Protein Data Bank file 5W3N). Moreover, in simple molecular dynamic simulations with explicit solvent⁶³ that start with the structure in Fig. 7, sidechains of S44 and D46 become solvated within 20 ns (see Fig. S10). Water molecules also enter a cavity that develops between S48 and T71, accounting for the correlations to water exhibited by S48. In addition, data in Fig. 6b indicate a lack of protection from hydrogen exchange for S48. Thus, the observed correlations to water (and absence thereof) for Ser and Thr residues that have assigned ¹³C ssNMR signals are consistent with the structural model in Fig. 7 after including adjustments that occur upon immersion of this model in water.

In Fig. 7b, carbon atoms of S84, T78, and G79 are colored cyan to illustrate the observed correlations from the S84 hydroxyl ¹H signal to ¹³C signals of T78 and G79 in Figs. 5a, 5b, and S6a. The greater intensity of S84-G79 crosspeaks and the fact that these crosspeaks are observed with $\tau_{HH} = 0$ (Fig. 5a) suggests the presence of a hydrogen bond between the S84 sidechain and the backbone carbonyl group of G79. Data in Fig. 6b also indicate protection from hydrogen exchange for S84. The S84-T78 crosspeak with $\tau_{HH} = 2.0$ ms (Fig. 5b) may result from ¹H-¹H polarization transfers from G79 to T78 during the τ_{HH} period. Interestingly, two hydroxyl ¹H chemical shifts (5.89 ppm and 6.24 ppm) can be assigned to T78, since both ¹H chemical shifts correlate with the ¹³C_a and ¹³C_β chemical shifts of T78, as shown in Fig. S8. Neither T78 hydroxyl ¹H signal has detectable correlations with ¹³C signals of S84. We interpret these observations as evidence that the T78 hydroxyl group does not form hydrogen bonds with S84, but may interact with other sidechains, such as that of Y75, with two distinct configurations that are in slow exchange. As discussed above, T78 does not show correlations to water in the 3D HCC spectrum, and therefore is apparently protected from rapid hydrogen exchange with water.

Carbon atoms of S77 and T47 are colored yellow in Fig. 7b to illustrate the observed correlations from the S77 hydroxyl ¹H signal to ¹³C signals of T47 (Figs. 5c and 5d). We attribute these crosspeaks to hydrogen bonding between S77 and T47 sidechains. In water-filtered 2D ¹³C-¹³C spectra, crosspeaks from S77 and T47 are not resolved from those of S87 and T45. Nonetheless, values of the apparent hydrogen exchange time τ for these crosspeaks are approximately 10 ms, greater than the intrinsic hydrogen exchange time of an

unprotected Ser or Thr hydroxyl group.⁶¹ Although the T47 hydroxyl ¹H signal is relatively strong, crosspeaks from the T47 hydroxyl ¹H signal to ¹³C signals of S77 can not be observed conclusively in our 3D HCC and H(H)CC spectra, as such inter-residue crosspeaks are only partially resolved from intra-residue crosspeaks of S87 (see Table S10).

Carbon atoms of T45 and Y81 are colored orange and carbon atoms of S48 and Y66 are colored green in Fig. 7b to illustrate the observed correlations from the T45 and S48 hydroxyl ¹H signals to ¹³C signals of Tyr sidechains (Figs. 5f and 5h). Assignments to Y81 and Y66 are based tentatively on the structural model itself, because the Tyr ¹³C chemical shifts do not have site-specific assignments. These correlations may result from T45-Y81 and S48-Y66 hydrogen bonds.

As described in the Introduction above, evidence for the importance of sidechain hydrogen bonding in self-assembly of low complexity sequences was provided previously by structural studies of peptides in amyloid-like crystalline states.^{38–41} In particular, Luo *et al.* identified segments of FUS-LC that form amyloid fibrils in a thermally reversible manner, which they call RAC1 (residues 37–42 of FUS-LC, sequence SYSGYS) and RAC2 (residues 54–59 of FUS-LC, sequence SYSSYG). High-resolution structures of crystalline RAC1 and RAC2 were solved by micro-electron diffraction (PDB 5XSG) and x-ray diffraction (PDB 5XRR, see Fig. S2d), respectively.³⁸ A similar crystal structure of RAC1 (PDB 6BWZ) was also reported by Hughes *et al.*³⁹ For RAC1, stabilizing interactions include hydrogen bonding of S39 and Y41 sidechains with an ordered row of water molecules (Fig. 1d), as well as intermolecular hydrogen bonding between sidechains of Y38 and S42 within each cross- β layer. For RAC2, stabilizing interactions include water-mediated hydrogen bonding among Y58 sidechains in different cross- β layers, as well as hydrogen bonding of Y55 and S57 sidechains with ordered rows of water molecules.

The specific sidechain interactions observed in structures of RAC1 and RAC2 are not directly transferable to the FUS-LC fibril structure for several reasons: (i) FUS-LC fibrils are noncrystalline; (ii) FUS-LC fibrils contain the full low-complexity sequence, so that a given segment of the sequence interacts both with the same segment in neighboring molecules and with different segments in the same and neighboring molecules; (iii) residues 37–42 and 54–59 are not fully structurally ordered in FUS-LC fibrils, as indicated by the ssNMR data of Murray *et al.*¹ as well as the ssNMR data for ²H,¹⁵N,¹³C-FUS-LC fibrils presented above (see Table S10). Nonetheless, the data for ²H,¹⁵N,¹³C-FUS-LC fibrils indicate that similar interactions are present in FUS-LC fibrils.

Conclusions

Data presented above provide evidence that the FUS-LC fibril core structure is stabilized in part by hydrogen bonds involving sidechains of hydroxyl-bearing residues, or more precisely that such hydrogen bonds exist within the core structure. In particular, the sidechain hydroxyl group of S84 appears to form a hydrogen bond to the backbone carbonyl group of G79, bridging a β -turn structure in residues 79–84. The S84 sidechain may also interact with the sidechain of T78, although evidence for a S84-T78 hydrogen bond is not clear. In addition, the sidechain hydroxyl group of S77 appears to form a hydrogen bond to

the sidechain of T47, contributing to energetically favorable stacking of a cross- β layer formed by residues 44–47 on a layer formed by residues 76–79. Data presented above also provide information about interactions of Thr and Ser hydroxyl groups with water. In particular, sidechain hydroxyl groups of specific residues (*i.e.*, T47, S53, S70, S77, T78, S84, S86, and S90) are relatively protected from hydrogen exchange with water, consistent with their involvement in hydrogen bonds that may contribute to structural stabilization. Protection from hydrogen exchange with water is indicated by the absence of correlations with water in 3D ¹H-¹³C-¹³C spectra, as well as by crosspeak build-up rates in water-filtered 2D ¹³C-¹³C spectra that are less than intrinsic hydrogen exchange rates.

Additionally, data presented above demonstrate how multidimensional ssNMR measurements on deuterated, back-exchanged samples can provide site-specific insights into hydrogen bonding, hydrogen exchange, and internal water locations within protein assemblies.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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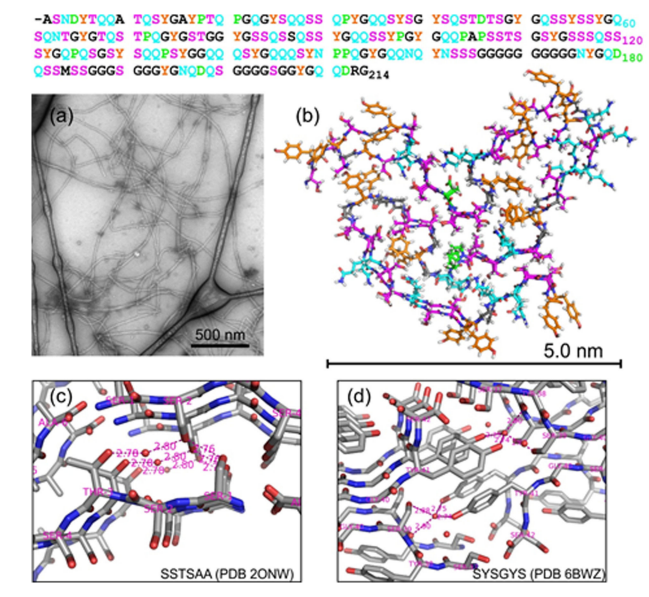


Figure 1:

(a) TEM image of negatively stained ²H,¹⁵N,¹³C-FUS-LC fibrils. (b) Molecular structural model for the FUS-LC fibril core, as determined by Murray *et al.*¹, viewed with the fibril growth axis perpendicular to the page and including residues 37–97. To illustrate the precision of the model, one molecule from model 1 of Protein Data Bank file 5W3N is superimposed on one molecule from model 16, with Ser and Thr residues in magenta, Gln and Asn residues in cyan, Tyr residues in orange, Pro and Asp residues in green, and all other residues in black. The full sequence of FUS-LC is shown above with the same color scheme. (c) Portion of the crystal structure of the hexapeptide SSTSAA, from Protein Data Bank file 2ONW. Short distances involving oxygen atoms of sidechain hydroxyl groups and water molecules are indicated in Å units. (d) Portion of the crystal structure of the hexapeptide SYSGYS, from Protein Data Bank file 6BWZ.

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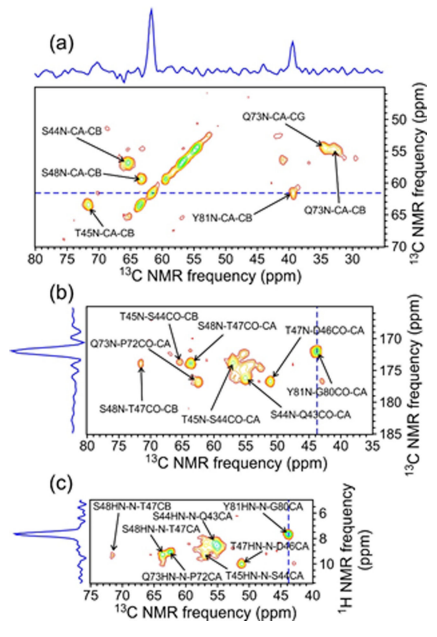


Figure 2:

2D planes from 3D NCACX (a), NCOCX (b), and HN(CO)CX (c) spectra of 2 H, 15 N, 13 C-FUS-LC fibrils, taken at a 15 N chemical shift of 121.4 ppm. Assignments are shown for crosspeaks with maxima within ±0.8 ppm of 121.4 ppm in the 15 N dimension. Contour levels increase by successive factors of 1.4. 1D slices shown above or on the left side of the 2D planes correspond to the dashed blue lines and illustrate the signal-to-noise ratios and ssNMR linewidths in the 3D spectra. Spectra were acquired at 14.1 T with 40.00 kHz MAS. (Note that the NCACX and NCOCX planes are different sections of the same 3D 15 N- 13 C- 13 C spectrum.)

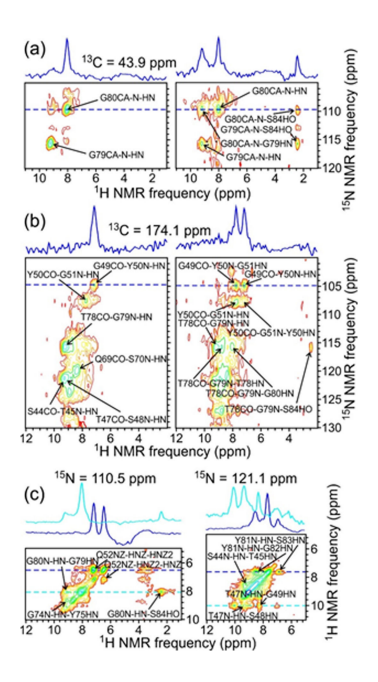


Figure 3:

2D planes from 3D CANH and CAN(H)H spectra (a, left and right, respectively), 3D CONH and CON(H)H spectra (b, left and right, respectively), and a 3D NHH spectrum (c) of ²H, ¹⁵N,¹³C-FUS-LC fibrils. Assignments are shown for crosspeaks with maxima within ± 0.3 ppm of 43.9 ppm in the ¹³C dimension in panel a, within ± 0.4 ppm of 174.1 ppm in the ¹³C dimension in panel b, and within ± 0.6 ppm of 110.5 ppm (left) or 121.1 ppm (right) in the ¹⁵N dimension in panel c. Contour levels increase by successive factors of 1.4. 1D slices shown above the 2D sections correspond to the dashed blue and cyan lines. In panels a and b, these slices illustrate the development of additional inter-residue crosspeaks from ¹H-¹H spin polarization transfers in the CAN(H)H and CON(H)H measurements. Spectra were acquired at 14.1 T with 40.00 kHz MAS.

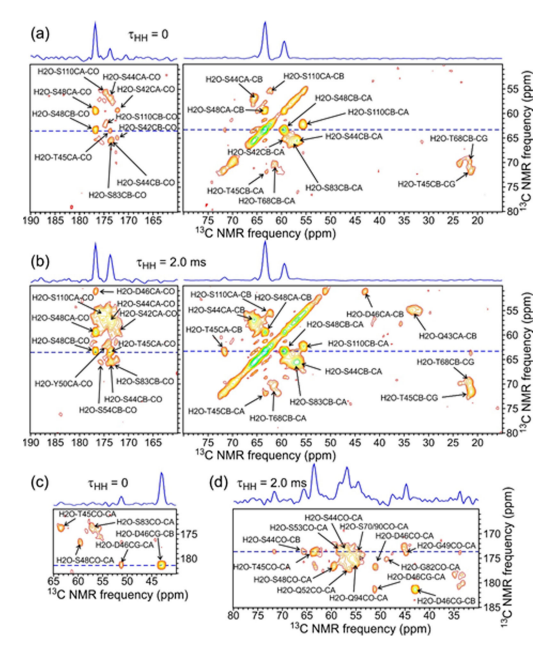


Figure 4:

2D planes from 3D HCC (a and c; $\tau_{HH} = 0$) and H(H)CC (b and d; $\tau_{HH} = 2.0$ ms) spectra of ²H,¹⁵N,¹³C-FUS-LC fibrils at the ¹H chemical shift of water (4.50 ppm). Assignments for crosspeaks within ±0.1 ppm of the water shift are shown. Contour levels increase by successive factors of 1.4. 1D slices above the 2D section correspond to dashed blue lines. Spectra were acquired at 14.1 T with 40.00 kHz MAS.

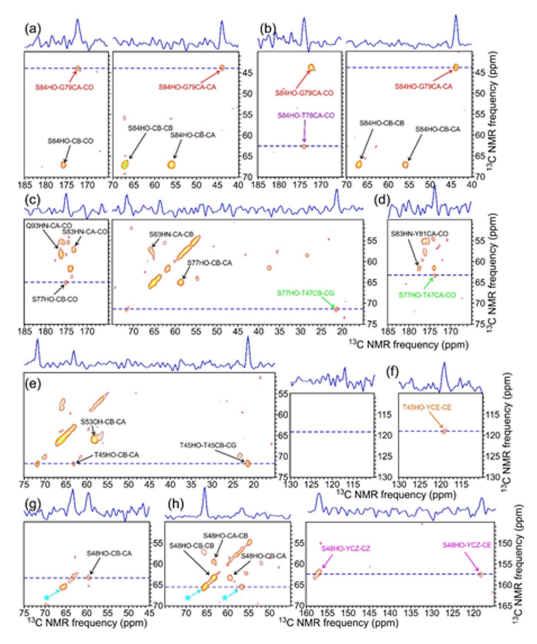


Figure 5:

2D planes from 3D HCC (a,c,e,g) and H(H)CC (b,d,f,h) spectra of 2 H, 15 N, 13 C-FUS-LC fibrils at 1 H chemical shifts of 2.39 ppm (a,b), 6.91 ppm (c,d), 4.93 ppm (e,f), and 4.00 ppm (g,h), corresponding to sidechain hydroxyl protons of S84, S77, S53 and T45, and S48, respectively. Contour levels increase by successive factors of 1.4, and assignments are shown for crosspeaks within ±0.1 ppm of the nominal 1 H chemical shift in each panel. (Asterisks in panels g and h indicate apparent crosspeaks arising from truncation artifacts from strong crosspeaks between the water 1 H chemical shift and 13 C_a and 13 C_b chemical shifts of S44.)

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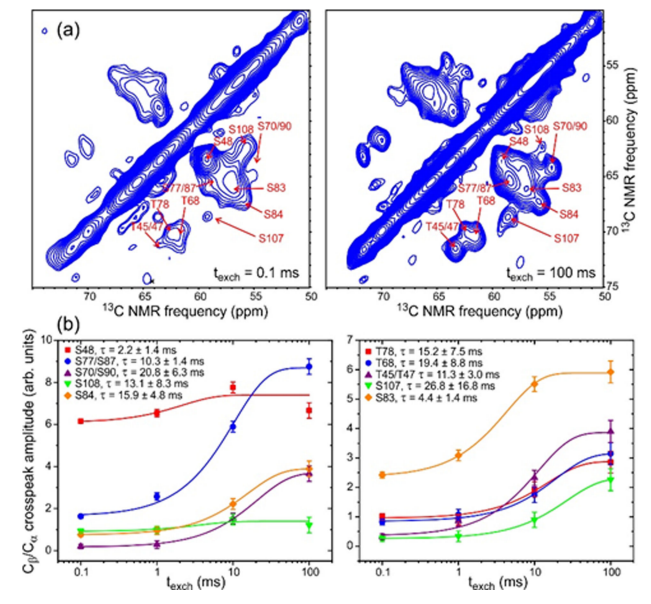


Figure 6:

(a) Water-filtered 2D ¹³C-¹³C spectra of ²H,¹⁵N,¹³C-FUS-LC fibrils with ¹H-¹H exchange periods $t_{exch} = 0.1$ ms (left) and $t_{exch} = 100$ ms (right). Contour levels increase by successive factors of 1.2. 2D spectra were obtained with 1600 (left) and 192 (right) scans per t_1 point. (b) Build-up curves for C_{β}/C_{α} crosspeak intensities of Ser and Thr residues. Error bars represent the root-mean-squared noise in the 2D spectra. Color-coded lines are fits with single-exponential functions, yielding the indicated values of τ .



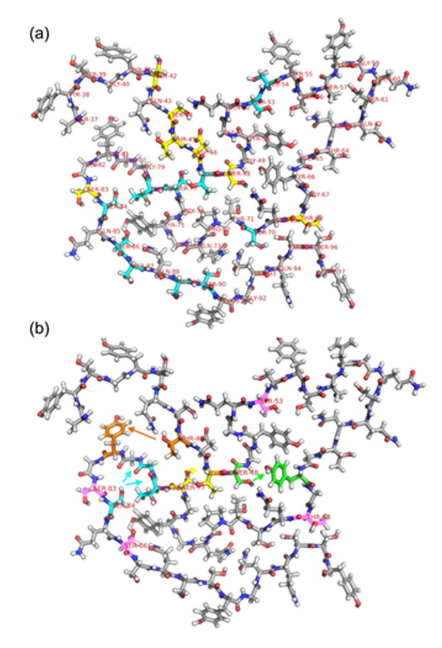


Figure 7:

(a) One molecule from Protein Data Bank file 5W3N, with carbons of Ser, Thr, Asp residues that have correlations to the ¹H chemical shift of water in the 3D HCC spectrum of ²H,¹⁵N, ¹³C-FUS-LC fibrils (and also have ¹³C chemical shift assignments) shown in yellow. Carbons of Ser and Thr residues that have ¹³C chemical shift assignments, but do not have correlations to water, are shown in cyan. (b) One molecule from Protein Data Bank file 5W3N, shown with residue number labels on all Ser and Thr residues that are involved in inter-residue crosspeaks in 3D HCC or 3D H(H)CC spectra are colored orange, cyan, yellow, and green, with arrows indicating the direction of these crosspeaks from hydroxyl ¹H chemical shifts to ¹³C chemical shifts. Based on the structural model, crosspeaks from

hydroxyl ¹H chemical shifts of T48 and S48 to aromatic ¹³C chemical shifts of Tyr residues are depicted as T48-Y81 and S48- Y66 crosspeaks. Carbons of Ser and Thr residues that have hydroxyl ¹H chemical shift assignments but do not exhibit detectable inter-residue crosspeaks are colored pink.