Natural Selection on the Phase-Separation Properties of FUS during 160 My of Mammalian Evolution

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Abstract

Protein phase separation can help explain the formation of many nonmembranous organelles. However, we know little about its ability to change in evolution. Here we studied the evolution of the mammalian RNA-binding protein Fused in Sarcoma (FUS), a protein whose prion-like domain (PLD) contributes to the formation of stress granules through liquid–liquid phase separation. Although the PLD evolves three times as rapidly as the remainder of FUS, it harbors absolutely conserved tyrosine residues that are crucial for phase separation. Ancestral reconstruction shows that the phosphorylation sites within the PLD are subject to stabilizing selection. They toggle among a small number of amino acid states. One exception to this pattern is primates, where the number of such phosphosites has increased through positive selection. In addition, we find frequent glutamine to proline changes that help maintain the unstructured state of FUS that is necessary for phase separation. Our work provides evidence that natural selection has stabilized the liquid forming potential of FUS and minimized the propensity of cytotoxic liquid-to-solid phase transitions during 160 My of mammalian evolution.

Key words: FUS, prion-like domain, liquid–liquid phase separation.

Introduction

Protein phase separation is a thermodynamic process through which a well-mixed solution of proteins and DNA/RNA molecules demixes into two phases of high and low densities (Shin and Brangwynne 2017; Boeynaems et al. 2018). It has emerged as a biophysical mechanism to explain the formation of several membrane-less organelles and biomolecular condensates in living cells. Examples of these condensates include the nucleolus, which is involved in ribosome assembly (Mitrea et al. 2018), cytoplasmic granules that form when cells are stressed (Bergeron-Sandoval et al. 2016; Banani et al. 2017; Riback et al. 2017; Franzmann et al. 2018), and RNA processing bodies involved in mRNA metabolism and turnover (Kroschwald et al. 2015).

One mammalian protein whose phase separation is well-studied is the RNA-binding protein Fused in Sarcoma (FUS) (Vance et al. 2009; Mackenzie et al. 2010; Kapeli et al. 2016). FUS is predominantly a nuclear protein and regulates the mRNA life cycle at different stages, from pre-mRNA splicing to mRNA transport and translation. In addition, FUS directly interacts with Poly-ADP-ribose polymerase and mediates DNA damage response in the cell (Naumann et al. 2018). FUS self-assembles through the interaction of its N-terminal prion-like domain (PLD) and the arginine–glycine–glycine domains (Kato and McKnight 2017; Murray et al. 2017).

FUS requires this self-assembly for its nuclear functions, such as binding to chromatin and recruitment to the sites of DNA damage (Mastrocola et al. 2013; Altmeyer et al. 2015). FUS most likely behaves as a multivalent heteropolymer composed of simple sequence motifs, also called stickers (Wang et al. 2018). These motifs consist of arginine residues in the arginine–glycine–glycine domain, and of tyrosine residues in the low complexity PLD. Their interaction is the major driving force for the phase separation of FUS (Wang et al. 2018).

Sticker amino acids are interspersed with sequence segments called spacers. Spacers can modify the properties of polypeptide chains and regulate the formation kinetics or the material states of phase-separated condensates (Harmon et al. 2017; Choi et al. 2019). In addition to this polymerization behavior, the formation of amyloid-like cross-beta sheet structures might play a role in the self-assembly of FUS droplets. Within the PLD, a region of 100 amino acid residues is especially prone to form these labile beta-sheet structures (Murray et al. 2017). Phosphorylation of several serine and threonine residues within this region disrupts these aggregates and regulates FUS assembly in the cell (Monahan et al. 2017; Murray et al. 2017).

The liquid-like state of FUS is also stabilized by a diverse set of molecular interactions within the PLD (Murthy et al. 2019). This dynamic state is exquisitely sensitive to single-point...
mutations. In fact, missense mutations in FUS occur in patients with the neurodegenerative diseases amyotrophic lateral sclerosis (ALS) and frontotemporal lobar degeneration (Mackenzie et al. 2010). For instance, a single ALS-associated mutation, G156E, facilitates a liquid-to-solid phase transition of FUS into irreversible aggregates (Patel et al. 2015).

The importance of FUS in the life of cells, together with the sensitivity of FUS assemblies to point mutations, raises the possibility that natural selection must actively maintain the ability of FUS to form the liquid-droplet state. We thus hypothesized that evolution has preserved the phase-separation propensity of FUS, and avoids the pathological liquid-to-solid phase separation in FUS, just like it maintains folding stability and reduces misfolding in proteins with structured domains (Zhang and Yang 2015).

Results
To identify the forces and mechanisms that govern the evolution of liquid–liquid phase separation, we studied how FUS evolved in mammals. We first identified 105 mammalian FUS orthologs, aligned them, and computed each residue’s sequence entropy, a widely used measure of sequence divergence. The PLD, which is central for FUS phase separation, has the highest sequence entropy of all FUS domains, with a median sequence entropy 3-fold higher than that of the other FUS domains. The first row corresponds to the PLD sequence of human FUS color-coded in the CLUSTAL format. The following rows show the sequences of mammalian PLDs compared with the sequence of the human PLD in the first row. If, at any position, the amino acid is different from that of the human PLD, the new amino acid is shown by colored boxes. Identical amino acids are shown as blue dots. Amino acid substitutions involve changes to serine (forest green), threonine (green), glycine (orange), alanine (blue), proline (yellow), and glutamine (Emerald green). Ancestral state mapping of substitutions in the evolution of PLD. Each box represents a substitution with its color saturation proportional to the number of such replacements. The first and second amino acids in a substitution are shown on the X and Y axes, respectively. Arrows highlight the especially frequent A to T, G to S, S to G, and T to A substitutions. Ancestral state mapping from F represented as a directed graph. Each circle or node represents one amino acid, and substitutions are shown as edges that connect these nodes. The thickness of each edge corresponds to the number of substitutions between the two incident nodes. Substitutions with more than Ancestral state mapping ten occurrences are shown in red.
domain of FUS is essential for phase separation (Wang et al. 2018), this strong conservation might indicate that evolution has maintained the phase-separation propensity of FUS in mammals

Within the PLD, we observed two evolutionary hotspots, which are the regions S30 to S86, and A105 to Q147 (all site numbers and amino acids refer to human FUS). These regions are subject to multiple substitutions that involve the amino acids glycine, serine, alanine, threonine, asparagine, proline, and glutamine (fig. 1D). By reconstructing ancestral FUS proteins (supplementary tables S1 and S2, Supplementary Material online, see Materials and Methods for details), we found that changes where the PLD sites toggle forth and back between G and S (113 changes), as well as between A and T (34 changes), are especially prevalent (fig. 1E and F). Together, these changes account for ~60% of all changes in the evolution of the PLD. In addition, we found 32 switches between serine and asparagine, and 20 switches between glutamine and proline in these evolutionary hotspots (supplementary tables S1 and S2, Supplementary Material online). To exclude the possibility that the high rates of substitutions between glycine and serine as well as alanine and threonine are caused by inaccurate ancestral reconstruction of the intrinsically disordered PLD, we repeated our analysis, but only counted substitutions in residues with a reconstruction accuracy of >90%. We found that these substitutions still account for ~55% of all substitutions (supplementary fig. S4 and table S9, Supplementary Material online). We observed the same percentage when we used a substitution matrix developed to model the evolution of intrinsically disordered mammalian proteins (Szalkowski and Anisimova 2011) (supplementary fig. S5 and table S9, Supplementary Material online). In sum, the high rate of G to S and A to T substitution is an inherent feature of PLD evolution rather than an artifact of ancestral sequence reconstruction.

To understand whether these amino acid switches are caused by neutral evolution or positive selection, we estimated how strongly evolutionary rates vary across the amino acid sites within the PLD, and along the branches of its phylogenetic tree (see Materials and Methods for details) (Yang and Dos Reis 2011). We detected positive selection in 10 branches (P value < 0.05) and 12 sites (probability > 0.90) (fig. 2A and supplementary table S3, Supplementary Material online) and in three types of substitutions: G to S, S to G, and Q to P. We observed the highest likelihood of positive selection for serine at the sites 42, 119, 129, and 131 and threonine at the sites 40 and 71 which occurred in the branches leading to primates and greater apes (supplementary information, Supplementary Material online).

The positively selected residues in primates (i.e., sites 42, 71, 78, 129, and 131) are among the sites in human FUS that are phosphorylated at serine and threonine (Monahan et al. 2017; Rhoads et al. 2018). Their phosphorylation not only increases the recruitment of FUS to the sites of DNA damage (Wang et al. 2013) but also inhibits the time-dependent changes in the material properties from liquid-like to solid-like states (Monahan et al. 2017). From a total of 32 phosphosites, only nine sites (i.e., sites 3, 7, 11, 26, 57, 77, 87, 96, and 148) were fully conserved, but the rest (24 sites) switched forth and back between only two pairs of amino acids (G-S and A-T) (fig. 2B). These sites occurred in both evolutionary hotspots, and their evolutionary rates were significantly higher than for the rest of the PLD residues (fig. 2E; Wilcoxon rank-sum test, $P = 7.9 \times 10^{-4}$, supplementary fig. S3, Supplementary Material online). The PLD sequences of primates and great apes harbor an exceptionally large number of phosphosites. That is, they harbor 29 and 31 phosphosites, respectively, which is 3 and 6 sites more than the average number of mammalian PLD phosphosites (fig. 2E). We found that the ratio of phosphosite-creating substitutions to phosphosite-destroying substitutions (i.e., $[G \text{ to } S + A \text{ to } T]/[S \text{ to } G + T \text{ to } A]$) is 0.67 in the whole phylogenetic tree, but significantly increases to 2.75 in the primate clade (Fisher’s exact test, $P = 0.011$; fig. 2C, 2D and supplementary information, Supplementary Material online). In other words, positively selected G to S and A to T substitutions have significantly increased the total number of phosphosites in the PLD sequences of primates.

Outside of the primates, the average number of phosphosites in the mammalian PLD sequences is $\pm 26 \pm 2$ sites. This small variation might indicate that the total number of S/T amino acids in these sites is stabilized in the evolution of the FUS PLD in mammals. To find out whether stabilizing selection has acted on our FUS sequences, we compared the likelihood that genetic drift alone or drift together with selection acted on the total number of phosphosites using an Ornstein–Uhlenbeck (OU) process (Butler and King 2004). This process has been used to compare the likelihood of drift alone with that of drift and selection in the evolution of different traits and characters (Hansen et al. 2000) (fig. 2F; see Materials and Methods for details). We found that stabilizing selection better explains the evolution of phosphosites than pure drift (likelihood ratio test, $P = 0.015$, fig. 2G; supplementary table S4, Supplementary Material online). We also asked whether two regimes (primate/nonprimate) of phosphosite evolution can better explain evolutionary rates than a single regime. We did so by comparing the likelihood of a two-optimum OU process given the data with that of a one-optimum OU process. The likelihood of the two-optimum process was significantly greater than the one-optimum process (supplementary table S4, Supplementary Material online; likelihood ratio test, $P = 0.045$). To ensure that the small size of one data set (22 species in the primate clade) did not bias our calculations, as has been observed in other applications of OU processes (Cooper et al. 2016), we further calculated the likelihood of stabilizing selection in the evolution of mammals without primates. The signature of stabilizing selection increases dramatically when primates are removed from this analysis (fig. 2H; likelihood ratio test, $P = 2.28 \times 10^{-7}$; supplementary table S4, Supplementary Material online). Together with our phylogenetic analysis of positive selection (fig. 2A), these observations suggest two regimes in the evolution of FUS phosphosites. In mammals except for primates, the number of phosphosites is under stabilizing selection. In primates and, in particular in great apes, positive selection has further increased the number of phosphorylation sites.
The disordered domains of proteins, in particular proteins that undergo phase separation, preserve key amino acid features such as charge and sequence composition throughout their evolution (Zarin et al. 2017, 2019). We thus examined the physicochemical properties that are either conserved or positively selected in the evolution of the PLD in mammalian FUS (see Materials and Methods for details). We found that amino acid substitutions in the PLD have significantly conserved polarity, flexibility, and solvation free energy (fig. 3 and supplementary table S5, Supplementary Material online; chi-square goodness-of-fit, \( P < 10^{-7} \)). We also found several properties whose changes were more frequent than expected from strict neutrality, and that had diversified in the evolution of the PLD (fig. 3). The most significantly diversified property is the average occurrence of amino acids in a tetrapeptide unit in protein structures (Rackovsky and Scheraga, 1982). This property quantifies the nucleation propensity of amino acids in segments of four residues and divides the amino acids into two groups. Tetrapeptides with amino acids in the first group (Pro, Gly, His, Tyr, Cys, Asn, and Trp) are more likely to...
Proline substitutions disrupt the formation of secondary structure in an evolutionary PLD hotspot. (A) In total, 108 different amino acid properties showing radical changes in the evolution of the PLD, ranked by selection’s tendency to either help diversify them (positive values, red) or conserve them (negative values, blue) during PLD evolution. Properties falling above and below the dashed line are selected for or against at a high significance level of 0.001. Phosphosites are excluded from this analysis. Because several amino properties correlate with each other, we averaged the selection’s tendency between highly correlated properties (Spearman’s correlation $R > 0.8$). Error bars show one standard deviation of selection strength between highly correlated amino acid properties. Selection tends to diversify properties that change the structure-formation propensity, whereas it tends to conserve amino acid polarity and flexibility. (B) Selection strength of the property “fractional occurrence in tetrapeptides in protein structures (RACS820101; Rackovsky and Scheraga 1982)” over the length of the PLD (horizontal axis). The area highlighted in blue shows the region S117 to S147, where selection to diversify this property is maximal. (C) Multiple sequence alignment of the segment S117 to S147 for selected mammalian PLD sequences. Color saturation represents the frequency of the amino acid in each column, and ranges from dark blue ($>80\%$) to white ($<40\%$). (D) PEP-FOLD predicted structure of the glutamine-rich, and (E) the proline-rich variant of the region S117 to S147. The sites 126, 132, 140, 141, 146, and 147 were computationally substituted to glutamine or proline to create the Q-rich or the P-rich variants, respectively. Both structures were generated by the PEP-FOLD webserver (Maupetit et al. 2009). (F) The radius of gyration per simulation time for the P-rich (shown in blue) and the Q-rich (shown in red) variants. (G) The solvent accessible surface area (Eisenhaber et al. 1995) of tyrosine sticker residues per simulation time for the P-rich and the Q-rich variants. (H) Boxplots comparing the radius of gyration of the P-rich and the Q-rich variants. (I) Boxplots comparing the solvent accessible area of tyrosine sticker residues. All P values were calculated using the Wilcoxon test.
adopt extended structures. Tetrapeptides with amino acids in the second group (all other amino acids) are more likely to form helical and bend conformations.

For this property, amino acid substitutions in the region S117 to S147 had the maximum strength of positive selection (fig. 3B and supplementary table S6, Supplementary Material online). This region is enriched in substitutions involving proline (i.e., in residues 105, 117, 126, 132, 134, 140, 141, 146, and 147; fig. 3C). Importantly, our analysis of positive selection had shown that Q to P substitutions are positively selected in different branches of the phylogenetic tree (e.g., Q141P in Gorilla (Gorilla gorilla) and S134P in the branch leading to primates; supplementary information, Supplementary Material online).

To study the effect of proline substitutions in this evolutionary hotspot, we created in silico a glutamine-rich and a proline-rich variant from the region S117 to S147 by selecting Q and P in all residues that had experienced Q to P substitutions in different mammalian sequences (residues 126, 132, 140, 141, 146, and 147), respectively. We then predicted the secondary structure content of the two variants using the PEP-FOLD algorithm (Maupetit et al. 2009) and validated the stability of these predictions using molecular dynamics simulations (see Materials and Methods for details). The glutamine-rich variant forms three short helices that extend from residue Q118 to S121, from S129 to S135, and from Q139 to G144 (fig. 3D). In contrast, the proline-rich variant is mostly unstructured and retains only partially the middle of the three helices (Q126 to Y129; fig. 3E). We further performed molecular dynamics simulations for 120 ns on these initial structures. Importantly, the Q-rich variant showed higher helical content (supplementary fig. S6A, Supplementary Material online) and, on average, five more side-chain hydrogen bonds compared with the P-rich variant (supplementary fig. S6B, Supplementary Material online). The P-rich variant had an increased radius of gyration compared with the Q-rich variant (fig. 3F and H; Wilcoxon rank-sum test, $P < 2.2 \times 10^{-16}$), and its tyrosine residues were on average more exposed to solvent (fig. 3F and I; Wilcoxon rank-sum test, $P < 2.2 \times 10^{-16}$). We thus conclude that Q to P substitutions help maintain an unstructured state and increase the likelihood of the interactions of the sticker tyrosine residues with arginines in the RNA-binding domain of FUS.

Finally, we examined the changes in the propensity of fibril formation in the evolution of the PLD in mammalian FUS. The first hotspot in the PLD, from S30 to S86, corresponds to a region that forms a fibrillar beta-sheet structure at high concentrations (fig. 4A) (Murray et al. 2017). The formation of these cross-beta sheet structures has been proposed (Murray et al. 2017) (but also disputed; Burke et al. 2015; Franzmann and Alberti 2019) to drive phase separation of FUS. Strikingly, we observed hydrogen-bond breaking substitutions in these regions that abolish side-chain hydrogen bonding and likely destabilize the fibril core (fig. 4B). For example, T78 and S84, which form inter-residue hydrogen bonds in the structure of the fibril core, are repeatedly substituted to alanine and glycine in different mammals. Other examples include alanine or proline substitutions in the residues S48, Q69, and T71, which hydrogen-bond and join the segment S44 to Y50 with T64 to G80. We further predicted the stability of fibril cores in different mammals and found a substantial variation in free energy of folding (fig. 4C and supplementary table S7, Supplementary Material online) which is more likely caused by pure drift than stabilizing selection (likelihood ratio test, $P = 0.019$; supplementary table S8, Supplementary Material online). In line with this observation, evolutionary rates of the PLD sequences were not significantly higher along the branches leading to fibrils with higher stabilities (fig. 4D and F; Spearman’s rank correlation, $R = 0.047, P = 0.71$).

We also pursued a complementary analysis to study the coevolution of fibril core residues. If the fibril core has remained intact during mammalian evolution, then we would expect that residues in the fibril core coevolve with each other, and possibly also with the rest of the PLD. To find out, we used Bayesian Graphical Models (Poon et al. 2008) to estimate the probability of coevolution between pairs of residues in the fibril core (see Materials and Methods). Despite the presence of pairs of coevolving residues (i.e., S39-T45, T68-S86, S68-S108, and G79-S116), we found no correlation between the probability of coevolution and the distance of coevolving residues on the 3D structure (fig. 4F; Spearman’s rank correlation, $R = -0.061, P = 0.13$). The coevolving residues T68 and S86 are phosphorylated in human FUS, showing that coevolution has modulated the phosphorylation potential of the PLD (fig. 4G and H). Altogether, our analyses reveal that the stability and structural integrity of the fibril core are not likely to be maintained during PLD evolution in mammalian FUS.

**Discussion**

Our study reveals that the PLD is the most rapidly evolving domain of mammalian FUS. Within the PLD, the total number of phosphorylation sites has generally evolved under stabilizing selection and has been maintained at ~26 sites per protein. We found that positive selection has significantly increased the number of phosphorylation sites in primates, particularly in great apes. Stabilizing selection has previously been proposed as a major process governing the evolution of phosphosites (Landry et al. 2009, 2014). According to this proposal, the cluster of phosphosites acts as a functional unit and is best described as a complex trait consisting of individual residues. Evolution then ensures that enough phosphorylation sites are retained for proper protein function (Landry et al. 2014). The evolution of mammalian FUS sequences provides a compelling case for this proposal and suggests that the total number of phosphosites can adapt to new regimes.

We also found that positive selection has favored glutamine to proline substitutions in an evolutionary hotspot of the PLD, likely to avoid secondary structure formation and maintain an unstructured state in the liquid state of FUS. Previously, proline substitutions had been shown to reduce the formation of irreversible aggregates of the huntingtin protein in Huntington disease (Popiel et al. 2004). Our
analysis extends this observation to the much greater evolutionary time scale of mammalian evolution, which unfolded over 160 My.

The evolutionary pattern of the PLD in mammals supports the polymer-based "molecular grammar" of phase separation in FUS (Wang et al. 2018). Specifically, the complete conservation of the number as well as position of tyrosines is in line with the role of these residues as stickers in associative polymers. This observation further extends a recent observation that conserved patterning of aromatic residues is important for the phase-separation behavior of PLDs (Martin et al. 2020). Also, the strong conservation of polarity, flexibility, and solvation free energy in the regions that separate the tyrosine residues ensures that these regions can act as flexible spacers to keep the PLD chain flexible in the liquid state of FUS. Whether this conservation pattern in sticker number and patterning extends to other phase-separating proteins is an important question for future research.

We also examined the relevance of cross-beta sheet fibril formation in the evolution of the PLD. We did not observe selection to either maintain the stability of the fibril core or destabilize its structure, suggesting that fibril core stability has evolved under genetic drift alone. Consistent with this notion is our observation that the amino acids that contact each other in the fibril core of human FUS do not coevolve. Because naturally occurring substitutions disrupt the hydrogen bonding that is necessary for the stability of the fibril core, it is unlikely that the formation of this core is a universal driver of FUS self-assembly in mammals. However, our results do not reject the possibility that the fibril core forms, because we found fibril core structures in mammals that are more stable than the structure of the fibril core in human FUS. Also, we...
performed our computations on the 3D structure of human FUS, such that the absence of biochemical and structural data on nonhuman FUS orthologs might affect our results. To assess the importance of hydrogen-bond breaking substitutions on the structure of FUS more generally, it will be necessary to resolve the structure of nonhuman FUS orthologs with fibril cores that are less stable than the human fibril core, using methods such as solid-state nuclear magnetic resonance. Likewise, we cannot rule out that the PLD in different FUS orthologs forms amyloid polymorphs. Cryogenic transmission electron microscopy (cryo-EM) can help determine whether amyloid morphology varies among mammalian FUS orthologs. If fibril formation is not observed experimentally in FUS orthologs, we predict that phosphorylation or frequent core-destabilizing substitutions (mainly S to G and T to A) will help avoid the formation of fibrils, which can mature into solid-like and irreversible aggregates.

Altogether, our work suggests that properties affecting phase separation may be evolvable in FUS, an essential protein that is also involved in neurological diseases. To prove this evolvability will require experimental studies. We propose the following experiments to validate our computational predictions. First and foremost, phase-separation assays with purified proteins can assess whether positively selected serine residues are functional and influence the time-dependent changes in the material states of FUS assemblies. For example, one can quantify the aggregation propensity of phosphomimetic FUS mutants in these residues (Monahan et al. 2017). One can also quantify and compare aggregation propensity of proline substitutions in positively selected proline residues among different FUS orthologs. Second, fluorescence recovery after photobleaching can help find out whether the hardening and softening of FUS droplets are substantially affected by substitutions between glycine and serine, alanine and threonine, and proline and glutamine (Wang et al. 2018). Third, small-angle X-ray scattering of the truncated evolutionary hotspot sequences, as well as hydrogen-deuterium exchange measurements of tyrosine residues in these segments can help determine whether evolution has altered the radii of gyration, as well as the accessibility of tyrosine residues for interaction with arginines in the RNA-binding domain of FUS.

The lack of protein structure in the PLD of FUS leads to substantial sequence variation, a feature that is common in intrinsically disordered proteins (Brown et al. 2002; Lin et al. 2007; Chen et al. 2011; Xue et al. 2013). Given this high overall divergence, stabilizing the condensed liquid-droplet state requires evolutionary mechanisms to maintain the disordered nature of the protein and avoid liquid-to-solid phase transitions. Our observations show that stabilizing selection of phosphorylation sites and positive selection of proline substitutions are two primary mechanisms to maintain the phase-separation propensity of FUS in mammals. We expect these mechanisms to hold for the evolution of the large set of ~2,600 proteins in the human proteome with similar sequence architecture to FUS. These proteins, including the members of the FUS-like family of proteins such as EWS and TAF15, have prion-like and RNA-binding domains that are similar in length and composition to these domains in FUS, and might function as scaffolds for biomolecular condensates in the cell (Wang et al. 2018).

Materials and Methods

Data Compilation

We retrieved 105 coding sequences of mammalian FUS genes from the NCBI (Pruitt et al. 2007) and ENSEMBL (Hubbard 2002) databases. We subdivided these sequences into three subsets. We used the first of these subsets, which comprised 105 mammalian sequences, to build a multiple sequence alignment for the calculation of sequence entropy. We used the second subset of 85 sequences, which had confident phylogenetic support from the TimeTree database (Hedges et al. 2006) to 1) estimate the likelihood of drift and selection with the aid of an OU process, 2) correlate the evolutionary rates of the PLD with the stability of the fibril core’s structure, and 3) to analyze coevolution of the PLD residues. This sequence alignment was highly conserved and indels were infrequent. From 160 residues in 85 mammals, only 14 positions were deleted in less than 3 species. The average conservation per residue, calculated from the BLOSUM 62 substitution matrix, was 0.92 ± 0.13. Finally, we used a third subset of 50 sequences with diverse taxonomic sampling in our analysis of positive selection. (The accession numbers of all sequences used in this work, as well as the sequences of reconstructed ancestors are available in the supplementary information, Supplementary Material online.)

Estimating Evolution Rate and Detecting Positive Evolution

We prepared protein sequence alignments with the codon-based CLUSTAL algorithm (Sievers et al. 2011) implemented in MEGA (Kumar et al. 2016) and Aliview (Larsson 2014) using the default parameters. We used the codeml program within the PAML suite (Yang 2007) to obtain maximum-likelihood estimates of the ratio dN/dS, that is, the ratio of the number of nonsynonymous substitutions per nonsynonymous site to the number of synonymous substitutions per synonymous site. This ratio is a widely used measure of selection strength on an evolving sequence (Yang 2006). For the estimation of dN/dS, we used the equilibrium codon frequencies from the products of the average observed frequencies in the three codon positions using the F3X4 model (Yang 2007). We tested the likelihood of positive selection in our sequences using the branch-site test for positive selection (Yang and Dos Reis 2011). In this model, a phylogenetic tree is partitioned into the foreground and background branches. The likelihoods of dN/dS > 1 and dN/dS = 1 along the foreground branches are compared using likelihood ratio tests. We determined the posterior probabilities that specific sites (amino acids) are subject to positive selection using the Bayes Empirical Bayes (Yang et al. 2005) method implemented in PAML (Yang 2007).

To mitigate the high rates of false positives in the detection of positively selected residues when indels are present, we performed two complementary sets of calculations. In the first set, we treated gaps as ambiguous data using the

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cleandata = 0 command in PAML. In the second set, we removed all positions that contained gap using the cleandata = 1 command. Overall, we only observed 6 out of 97 branches whose likelihood values were significantly affected by the removal of indels. Importantly, the likelihood of positive selection along the branches leading to primates was unaffected (supplementary table S10, Supplementary Material online).

Ancestral Sequence Reconstruction
To reconstruct ancestral sequences, we fitted different substitution models to our data (PLD sequences and the mammalian phylogenetic tree), allowing that evolutionary rates may vary among protein sites. The substitution model JTT (Jones et al. 1992) with the gamma distribution of evolutionary rates had the highest Bayesian Information Criterion score (Burke et al. 2015). We thus used this model and inferred ancestral PLD sequences using the maximum-likelihood method implemented in MEGA7 (Kumar et al. 2016). The probabilities of the most likely inferred amino acids were on average 0.93 ± 0.16. We also reconstructed ancestors using a substitution matrix developed to study the evolution of intrinsically disordered mammalian proteins (Szalkowski and Anisimova 2011). For this matrix too, the probabilities of reconstructed ancestral sequences were 0.92 ± 0.19.

Detection of Amino Acid Properties under Selection
We used the TreeSAAP method (Woolley et al. 2003) to infer how natural selection may change amino acid properties in the evolution of the PLD. Briefly, this method compares the distribution of changes in amino acid properties along the branches of a phylogenetic tree with an expected distribution, using the codon composition of a set of extant sequences. Changes in amino acid properties are divided into eight categories, from the most conserved (category 1) to the most radical changes (category 8). The method then calculates the goodness-of-fit (χ²-distribution) between the expected and the observed frequencies and tests the hypothesis that these distributions are equal for each amino acid property. For a specific property, the deviation between observed and expected frequencies in each category is calculated using a Z-score. We refer to this Z-score as the deviation from neutrality or the selection strength throughout this paper. A highly significant Z-score (Z > 3.09, P < 0.01) shows that more nonsynonymous substitutions change the property of interest than expected from neutral evolution. Although TreeSAAP was originally devised to study the evolution of structured proteins, the conserved sequence alignment of the PLD in mammals justifies the use of this approach in this work. The limited usage of amino acids in low complexity regions renders the expected distribution of properties dramatically different between structured and unstructured proteins. Nonetheless, if the expected distribution is built from the extant sequences, the inference of selection should be robust.

Detection of Coevolution
To infer the coevolutionary history of protein sites within the PLD, we used Bayesian Graphical Models (Poon et al. 2008) implemented in the HyPhy package (Pond and Muse 2005). We first used our ancestral reconstruction of FUS to construct a binary matrix representing the presence and absence of substitutions on each branch (rows) of the phylogenetic tree and in each site of the protein (columns). The joint distribution of all substitutions was then inferred using Bayesian networks and Markov Chain Monte Carlo sampling with default parameters in the SpiderMonkey method (Poon et al. 2008). We avoided the use of mutual information to infer coevolution because it leads to a high rate of false positives in the detection of coevolving sites when sequences are substantially similar (Harmon et al. 2017) (~>62%), as in our case.

Prediction of Folding Free Energy
We predicted the stability of the structure of the fibril core of the PLD in different mammals. We generated the 3D structures of the fibril core using its structure in human (PDB ID: 5W3N; Murray et al. 2017) as a template. We then calculated the free energy of folding of the fibril core made from the mammalian PLD sequences using the FoldX algorithm, which uses an empirical force field for the prediction of the free energy change of protein structures upon mutations (Schymkowitz et al. 2005; Buß et al. 2018). We first minimized the free energy of this structure using the Repair command in FoldX (Schymkowitz et al. 2005). We then created in silico mutants of this structure to create different mammalian PLD orthologs using the BuildModel command of FoldX (Schymkowitz et al. 2005).

OU Processes
To estimate the significance of stabilizing selection versus pure drift, we used OU processes that are corrected for phylogenetic dependence of species (Hansen et al. 2000). These models have been used to test various evolutionary hypotheses in the evolution of different characters and traits (Butler and King 2004), gene expression level (Rohlfs et al. 2014), and protein structure (Challis and Schmidler 2012). In brief, these models assume that the character of interest, $X(t)$, evolves in time unit $(t)$, according to an OU process:

$$dX_i(t) = \alpha(\theta - X_i(t))dt + \sigma dB_i(t)$$  \hspace{1cm} (1)

The parameter $\theta$ is the optimum value of $X(t)$ in the $i$th lineage and the parameters $\alpha$, $\sigma$, and $T$ represent the strength of selection and drift, respectively. The term $dB_i(t)$ is a white noise term, with mean 0 and variance $dt$. Equation (1) defines a Gaussian process whose moments depend on the parameters $\theta$, $\alpha$, $\sigma$ and the total time spent for a character in the lineage of interest, $t = T$. $T$ is either defined as the evolutionary time or equal to the branch length, as in our case. The values of the optimum parameter of the evolving trait ($\theta$) can vary according to the assumed evolutionary process. For example, if the character of interest evolves under a single optimum, $\theta$ is the same for all lineages. These parameters are
estimated by minimizing the logarithm of a likelihood function that assumes multivariate normality of all characters at the terminal taxa, \( X(t = 7) \).

We used the total number of phosphosites and the folding free energy as the traits of interest in evolution. We then fitted the models for pure drift and stabilizing selection using the BROWN and HANSEN commands in the OUCH package (Butler and King 2004), respectively. The input to these commands was the mammalian phylogenetic tree in Newick format, together with a data vector of the trait of interest, either the total number of phosphosites or the stability of the fibril core. We used the initial values of \( \alpha = 1 \) and \( \sigma = 1 \) to initialize the optimization process using the Nelder and Mead simplex algorithm (Nelder and Mead 1965). For modeling stabilizing selection, we assumed that all nodes belong to a single selective regime. We used likelihood ratio tests to compare the likelihoods of drift and selection.

**Supplementary Material**

**Supplementary data** are available at *Molecular Biology and Evolution* online.

**Data Availability**

All data generated or analyzed during this study are available in the repository: https://github.com/dasmeh/FUSEVOL.

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**References**

Altmeyer M, Neelsen KJ, Teloni F, Pozdnayakova L, Pellegrino S, Grafe M, Rask M-BD, Streicher W, Jungmichel S, Nielsen ML, et al. 2015. Liquid demixing of intrinsically disordered proteins is seeded by poly (ADP-ribose). *Nat Commun.* 6(1):8088.

Banani SF, Lee HO, Hyman AA, Rosen MK. 2017. Biomolecular condensates: organizers of cellular biochemistry. *Nat Rev Mol Cell Biol.* 18(5):285–298.

Bergeron-Sandoval L-P, Safaei N, Michnick SW. 2016. Mechanisms and consequences of macromolecular phase separation. *Cell* 165(5):1067–1079.

Boezaars S, Alberti S, Fawzi NL, Mittag T, Polymenidou M, Rousseau F, Schymkowitz J, Shorter J, Wolozin B, Van Den Bosch L, et al. 2018. Protein phase separation: a new phase in cell biology. *Trends Cell Biol.* 28(6):420–435.

Bonferroni CE. 1936. Teoria statistica delle classi e calcolo delle probabilita. *Pubblicazioni del R Istituto Superiore di Scienze Economiche e Commerziali di Firenze* 8–3–62.

Brown CJ, Takayama S, Campbell AM, Vise P, Marshall TW, Oldfield CJ, Williams CJ, Keith Dunker A. 2002. Evolutionary rate heterogeneity in proteins with long disordered regions. *J Mol Evol.* 55(1):104–110.

Burke KA, Janke AM, Rhine CL, Fawzi NL. 2015. Residue-by-residue view of in vitro FUS granules that bind the C-terminal domain of RNA polymerase II. *Mol Cell.* 60(2):231–241.

Buò B, Rudat J, Ochsnerreither K. 2018. FoldX as protein engineering tool: better than random based approaches? *Comput Struct Biotechnol J.* 16:25–33.

Butler MA, King AA. 2004. Phylogenetic comparative analysis: a modeling approach for adaptive evolution. *Am Nat.* 164(6):683–695.

Challis CJ, Schmidler SC. 2012. A stochastic evolutionary model for protein structure alignment and phylogeny. *Mol Biol Evol.* 29(11):3575–3587.

Chen SC-C, Chuang T-J, Li W-H. 2011. The relationships among microRNA regulation, intrinsically disordered regions, and other indicators of protein evolutionary rate. *Mol Biol Evol.* 28(9):2513–2520.

Choi J-M, Dar F, Pappu RV. 2019. LASSI: a lattice model for simulating phase transitions of multivalent proteins. *PLoS Comput Biol.* 15(10):e1007028.

Cooper N, Thomas GH, Venditti C, Meade A, Freckleton RP. 2016. A cautionary note on the use of Ornstein Uhlenbeck models in macroevolutionary studies. *Biol J Linn Soc.* 118(1):64–77.

Damas JM, Filipe LCS, Campos SRR, Lousa D, Victor BL, Baptista AM, Franca-Oliveira R, Oroszlan S, Pozo A. 2012. Helix-forming cyclic peptides: organizers of cellular biochemistry. *Mol Cell Biol.* 32(11):2461–2474.

Eisenhaber F, Lijnzaad P, Argos P, Sander C, Scharf M. 1995. The double indicator of protein evolutionary rate. *J Mol Evol.* 45(1):104–110.

Franzmann TM, Alberti S. 2019. Priors-like low-complexity sequences: key regulators of protein solubility and phase behavior. *J Biol Chem.* 294(18):7128–7136.

Franzmann TM, Jahnel M, Pozniakovsky A, Mahamid J, Holehouse AS, Nüske E, Richter D, Baumeister W, Grill SW, Pappu RV, et al. 2018.
Phase separation of a yeast prion protein promotes cellular fitness. Science 359(6371):eaao5654.

Hansen TF, Armbruster WS, Antonsen L. 2000. Comparative analysis of character displacement and spatial adaptations as illustrated by the evolution of Dalechampia blossoms. Am Nat 156(5):S57–S54.

Harmon, TS, Holehouse AS, Rosen MK, Pappu RV. 2017. Intrinsically disordered linkers determine the interplay between phase separation and gelation in multivalent proteins. Elife 6:e30294.

Hedges SB, Dudley J, Kumar S. 2006. TimeTree: a public knowledge-base of divergence times among organisms. Bioinformatics 22(23):2971–2972.

Hoober WG. 1986. Constant-pressure equations of motion. Phys Rev A 34(3):2499–2500.

Hubbard T. 2002. The Ensembl genome database project. Nucleic Acids Res 30(1):38–41.

Jones DT, Taylor WR, Thornton JM. 1992. The rapid generation of mutation data matrices from protein sequences. Bioinformatics 8(3):275–282.

Kapeli K, Pratt CA, Vu AQ, Hutt KR, Martinez FJ, Sundaramaram B, Batra R, Freese P, Lambert NJ, Huelga SC, et al. 2016. Distinct and shared functions of ALS-associated proteins TDP-43, FUS and TARF revealed by multisystem analyses. Nat Commun 7(1):12143.

Kato M, McKnight SL. 2017. Cross-β polymerization of low complexity sequence domains. Cold Spring Harb Perspect Biol 9(3):a023598.

Kroschwald S, Maharana S, Mateju D, Malinovska L, Núñez E, Poser I, Richter D, Alberti S. 2015. Promiscuous interactions and protein dis-aggregates determine the material state of stress-inducible RNP granules. Elife 4:e06807.

Kumar S, Stecher G, Tamura K. 2016. MEGA7: molecular evolutionary genetics analysis version 7.0 for bigger datasets. Mol Biol Evol 33(7):1870–1874.

Landry CR, Freschi L, Zarin T, Moses AM. 2014. Turnover of protein phosphorylation during stabilizing selection. Front Genet 5:2542.

Landry CR, Levy ED, Michnick SW. 2009. Weak functional constraints on phosphoproteomes. Trends Genet 25(5):193–197.

Larsson A. 2014. AllView: a fast and lightweight alignment viewer and editor for large datasets. Bioinformatics 30(22):3276–3278.

Lin Y-S, Hsu W-L, Hwang J-K, Li W-H. 2007. Proportion of solvent-exposed amino acids in a protein and rate of protein evolution. Mol Biol Evol 24(4):1005–1011.

Mackenzi ER, Rademakers R, Neumann M. 2010. TDP-43 and FUS in amyotrophic lateral sclerosis and frontotemporal dementia. Cell 142(2):277–290.

Mastrocola AS, Kim SH, Trinh AT, Rodenkirch LA, Tibbetts RS. 2013. The RNA-binding protein fused in sarcoma (FUS) functions downstream of poly (ADP-ribose) polymerase (PARP) in response to DNA damage. J Biol Chem 288(34):24731–24741.

Maugeretti J, Derreumaux P, Tuffery P. 2009. PEP-FOLD: an online resource for de novo peptide structure prediction. Nucleic Acids Res 37(Web Server):W498–W503.

Mitrea DM, Cika JA, Stanley CB, Nourse A, Onuchic JL, Banerjee PR, Phillips AH, Park C-G, Deniz AA, Kriwacki RW, et al. 2018. Self-interaction of NPM1 modulates multiple mechanisms of liquid–liquid phase separation. Nat Commun 9(1):1–13.

Monahan Z, Ryan VH, Janke AM, Burke KA, Rhoads SN, Zerze GH, Mahamid J, Saha S, Franzmann TM, et al. 2015. A liquid-to-solid phase transition of the ALS protein FUS accelerated by disease mutation. Cell 162(5):1066–1077.

Pond SLK, Muse SV. 2005. HyPhy: hypothesis testing using phylogenies. In: Nielsen R, editor. Statistical methods in molecular evolution. New York (NY): Springer. p. 125–181.

Poon AF, Lewis FL, Frost SD, Kosakovksy Pond SL. 2008. Spidermonkey: rapid detection of co-evolving sites using Bayesian graphical models. Bioinformatics 24(17):1949–1950.

Popiel HA, Nagai Y, Onodera O, Inui T, Fujikake N, Urade Y, Strittmatter WJ, Burke JR, Ichikawa A, Toda T, et al. 2004. Disruption of the toxic conformation of the expanded polyglutamine stretch leads to suppression of aggregate formation and cytotoxicity. Biochem Biophys Res Commun 317(4):1200–1206.

Pruitt KD, Tatusova T, Maglott DR. 2007. NCBI reference sequences (RefSeq): a curated non-redundant sequence database of genomes, transcripts and proteins. Nucleic Acids Res 35(Database):D61–D65.

Rackovsky S, Schera H. 1982. Differential geometry and polymer conformation. 4. Conformational and nucleation properties of individual amino acids. Macromolecules 15(5):1340–1346.

Rhoads SN, Monahan ZT, Yee DS, Shewmaker FP. 2018. The role of post-translational modifications on prion-like aggregation and liquid-phase separation of FUS. Int J Mol Sci. 19(3):8868.

Riback JA, Katsanski CD, Kearn-Scott JT, Pilipenko EV, Rojek AE, Sosnir TK, Drummond DA. 2017. Stress-triggered phase separation is an adaptive, evolutionarily tuned response. Cell 168(6):1026–1040.e19.

Rohlf RV, Harigan P, Nielsen R. 2014. Modeling gene expression evolution with an extended Ornstein–Uhlenbeck process accounting for within-species variation. Mol Biol Evol 31(1):201–211.

Schmid N, Eichenberger AP, Chotuok A, Rinker S, Wingier M, Mark AE, van Gunsteren WF. 2011. Definition and testing of the GROMOS force-field versions 54A7 and 54B7. Eur Biophys J. 40(7):843–856.

Schymkowitz J, Borg J, Stricher F, Nys R, Rousseau F, Serrano L. 2005. The propensity of a protein to form different structures: an empirical force field approach. J Mol Biol 347(4):305–318.

Shin Y, Brangwynne CP. 2017. Liquid phase condensation in cell physiology and disease. Science 357(6357):eaaf4382.

Sievers F, Wilm A, Dineen D, Gibson TJ, Karplus K, Li W, Lopez R, McWilliam H, Remmert M, Soding J, et al. 2011. Fast, scalable generation of high-quality protein multiple sequence alignments using Clustal Omega. Mol Syst Biol 7(1):539.

Szalkowski AM, Anisimova M. 2011. Markov models of amino acid substitution to study proteins with intrinsically disordered regions. PLoS One 6(5):e20488.

Van Der Spoel D, Lindahl E, Hess B, Groenhof G, Mark AE, Berendsen HJC. 2005. GROMACS: fast, flexible, and free. J Comput Chem. 26(16):1701–1718.

Vance C, Rogelj B, Hortobagyi T, De Vos KJ, Nishimura AL, Sreedharan J, Hu X, Eichenberger AP, Chotuok A, Rinker S, Wingier M, Mark AE, van Gunsteren WF. 2018. A molecular grammar governing the driving forces for phase separation of prion-like RNA binding proteins. Cell 174(3):688–699.e16.

Wang W-Y, Pan L, Su SC, Quinn EJ, Sasaki M, Jimenez JC, Mackenzie IRA, Huang EJ, Tsai L-H. 2013. Interaction of FUS and HDAC1 regulates...
DNA damage response and repair in neurons. *Nat Neurosci.* 16(10):1383–1391.

Woolley S, Johnson J, Smith MJ, Crandall KA, McClellan DA. 2003. TreeSAAP: selection on amino acid properties using phylogenetic trees. *Bioinformatics* 19(5):671–672.

Xue B, Brown CJ, Dunker AK, Uversky VN. 2013. Intrinsically disordered regions of p53 family are highly diversified in evolution. *Biochim Biophys Acta.* 1834(4):725–738.

Yang Z. 2006. Computational molecular evolution. New York: Oxford University Press View Article.

Yang Z. 2007. PAML 4: phylogenetic analysis by maximum likelihood. *Mol Biol Evol.* 24(8):1586–1591.

Yang Z, Dos Reis M. 2011. Statistical properties of the branch-site test of positive selection. *Mol Biol Evol.* 28(3):1217–1228.

Yang Z, Wong WS, Nielsen R. 2005. Bayes empirical Bayes inference of amino acid sites under positive selection. *Mol Biol Evol.* 22(4):1107–1118.

Zarin T, Strome B, Nguyen Ba AN, Alberti S, Forman-Kay JD, Moses AM. 2019. Proteome-wide signatures of function in highly diverged intrinsically disordered regions. *Elife.* 8:e46883.

Zarin T, Tsai CN, Ba ANN, Moses AM. 2017. Selection maintains signaling function of a highly diverged intrinsically disordered region. *Proc Natl Acad Sci U S A.* 114(8):E1450–E1459.

Zhang J, Yang J-R. 2015. Determinants of the rate of protein sequence evolution. *Nat Rev Genet.* 16(7):409–420.