TDP-43 oligomerization and RNA binding are codependent but their loss elicits distinct 1 pathologies 2

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23 **Short title**

- Oligomerization modulates TDP-43 aggregation patterns 24
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26 Abstract

Aggregation of the RNA-binding protein TDP-43 is the main common neuropathological feature 27 of TDP-43 proteinopathies. In physiological conditions, TDP-43 is predominantly nuclear and 28 29 contained in biomolecular condensates formed via liquid-liquid phase separation (LLPS). However, in disease, TDP-43 is depleted from these compartments and forms cytoplasmic or, sometimes, 30 intranuclear inclusions. How TDP-43 transitions from physiological to pathological states remains 31 poorly understood. Here, we show that self-oligomerization and RNA binding cooperatively govern 32 TDP-43 stability, functionality, LLPS and cellular localization. Importantly, our data reveal that 33 TDP-43 oligomerization is connected to, and conformationally modulated by, RNA binding. 34 35 Mimicking the impaired proteasomal activity observed in patients, we found that TDP-43 forms nuclear aggregates via LLPS and cytoplasmic aggregates via aggresome formation. The favored 36 aggregation pathway depended on the TDP-43 state -monomeric/oligomeric, RNA-bound/-37 38 unbound- and the subcellular environment -nucleus/cytoplasm. Our work unravels the origins of

heterogeneous pathological species occurring in TDP-43 proteinopathies. 39

40 MAIN TEXT

41 Introduction

Amyotrophic lateral sclerosis (ALS) and frontotemporal lobar degeneration (FTLD) are two 42 seemingly different, devastating adult-onset neurodegenerative diseases that exhibit a 43 significant genetic, clinical and pathological overlap (1). The vast majority of ALS patients 44 and up to half of FTLD cases are characterized by the accumulation of aggregated TAR 45 DNA-binding protein 43 (TDP-43) in affected neurons (1-3). Importantly, TDP-43 46 pathology is not an exclusive hallmark of ALS and FTLD, but is also the main pathological 47 feature in limbic-predominant age-related TDP-43 encephalopathy (LATE) (4) and a 48 concomitant pathology in a subset of patients with other neurodegenerative diseases 49 including Alzheimer's, Parkinson's and Huntington's disease (5). However, TDP-43 50 aggregates associated with different clinical subtypes present distinct subcellular 51 localization, namely cytoplasmic, intranuclear or axonal (6, 7), as well as morphological 52 and biochemical properties (8-10), indicating a distinct molecular origin of these diverse 53 pathological species. 54

TDP-43 is a ubiquitously expressed (11) and predominantly nuclear (12) nucleic acid-55 binding protein (11, 13) composed of an N-terminal domain (NTD, amino acids 1-80) 56 involved in self-oligomerization (14, 15), two tandem RNA-recognition motifs (RRMs, 57 amino acids 106-259) (13, 16) and an unstructured low complexity region (LCR, amino 58 acids 260-414). The latter contains a transient α -helix (amino acids 321-340) (17) that 59 associates with interaction partners (18) and was recently shown to coincide with the 60 aggregation core of pathological cytoplasmic TDP-43 in FTLD brains (19). Under 61 physiological conditions in the nucleus, TDP-43 mainly binds UG-rich intronic sites on pre-62 mRNA to regulate alternative splicing (20, 21) and undergoes liquid-liquid phase separation 63 (LLPS) (22) to form dynamic nuclear droplets (23-26), which were suggested to localize to 64 specific subnuclear membraneless compartments (26, 27). Albeit its predominantly nuclear 65 localization (12), TDP-43 shuttles between the nucleus and the cytoplasm (28), where it 66 plays roles in mRNA stability, transport and translation, miRNA processing, mitochondrial 67 and synaptic function and stress responses (1). In particular, TDP-43 was shown to 68 incorporate into and modulate the dynamics of stress granules (SGs) upon exposure to 69 different temperature, osmotic, oxidative and chemical stressors (29, 30). 70

RRM-mediated RNA binding is essential for TDP-43 to perform its physiological functions 71 in RNA metabolism (31). In addition, RNA binding precludes TDP-43 passive leakage out 72 of the nucleus (28, 32) and modulates its LLPS behavior (33, 34). In contrast, little is known 73 74 about the importance of TDP-43 NTD-driven self-oligomerization in physiology. Previous data have shown that nuclear TDP-43 oligomerization is required for alternative splicing of 75 at least a subset of its known RNA targets (15, 35-38). However, the role of TDP-43 76 oligomerization in its physiological properties, including its subcellular localization, 77 stability, LLPS behavior and cytoplasmic functions, remains poorly understood. Also, 78 whether -and if so, how- TDP-43 RNA binding and oligomerization impact each other in 79 cells is unknown. 80

The overexpression of TDP-43 in cellular and animal models results in its aggregation, a phenomenon that has been extensively explored in recent years to recapitulate the main neuropathological hallmark of ALS/FTLD (39). However, TDP-43 is a tightly autoregulated protein (20, 40) and overexpression can distort its subcellular (32) and subnuclear (24, 27) localization, and potentially its functions. We therefore aimed to study the physiological role of TDP-43 oligomerization and its interplay with RNA binding at

near-physiological protein levels, and to subsequently compare the pathways triggered by 87 88 their respective impairment. Using human neural cultures and single-copy expression systems in human cell lines, we show that NTD-driven TDP-43 oligomerization and RRMs-89 mediated RNA binding are intertwined and required to maintain the half-life, functionality 90 91 and localization of TDP-43. Upon failure of the ubiquitin-proteasome system (UPS), monomerization and impaired RNA binding triggered TDP-43 aggregation via distinct 92 pathways in the cytoplasm and nucleus. Our results underscore the relevance of loss of 93 94 oligomerization and RNA binding in the initiation of diverse TDP-43 pathologies and unravel the origins of heterogeneous pathological species occurring in human disease. 95

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97 **Results**

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Oligomerization and RNA-binding cooperatively stabilize the half-life of TDP-43

To systematically compare the properties of oligomeric, monomeric and RNA binding-99 deficient TDP-43, we introduced a single copy of an N-terminally green fluorescent protein 00 (GFP)-tagged TDP-43 coding sequence under the control of a doxycycline-inducible 01 promoter into HEK293 cells using the Flp-In T-REx technology (Figure S1A) (41). This 02 construct harbored previously described point mutations that disrupt TDP-43 03 oligomerization (termed 6M) (15), RNA binding through the RRMs (five F>A mutations in 04 the RRMs, referred to as RRMm) (13, 16) or both 6M&RRMm (Figure 1A). The resulting 05 four isogenic cell lines (WT, 6M, RRMm and 6M&RRMm) expressed equal levels of the 06 exogenous GFP-TDP-43 RNA (Figure S1B) and both RNA and protein were detectable 07 only upon addition of doxycycline (Figures 1B and S1C-D). Wild type (WT) GFP-TDP-08 43 protein levels displayed a mere 4-fold increase compared to endogenous TDP-43, as 09 determined by immunoblot analysis (Figure S1D). However, despite equal RNA levels 10 (Figure S1B), protein levels of the GFP-TDP-43 mutants were noticeably lower than their 11 WT counterpart (Figures 1B-C and S1E-G). A protein turnover analysis using the 12 translation inhibitor cycloheximide (CHX) showed that the half-life of the RNA-binding 13 TDP-43 mutant was reduced by >8 hours compared to WT GFP-TDP-43 (Figure 1D-E), 14 consistent with previous findings (42). Interestingly, oligomerization deficiency (6M) and 15 loss of RNA binding had a similar effect on the half-life of TDP-43. Furthermore, the 16 combined variant (6M&RRMm) presented a cumulative effect (Figure 1D-E). Since point 17 mutations can affect protein folding and thereby selectively target proteins for degradation 18 19 (43), we confirmed that the introduced mutations do not interfere with the folding of TDP-43 using circular dichroism (CD) (Figure 1F and S1H) and two-dimensional nuclear 20 magnetic resonance (2D-NMR) spectroscopy (Figure S1I-J) (15, 44), which revealed that 21 the mutated domains are properly folded. Therefore, our results indicate that loss of 22 oligomerization or RNA-binding ability similarly reduces the half-life of TDP-43. Since 23 incorporation of proteins into multimeric complexes has been reported to correlate with 24 longer half-lives in yeast (45) and mouse brain cells (46), these observations strengthen the 25 link between TDP-43 functionality and its half-life. 26

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TDP-43 oligomerization and RNA-binding preserve its nuclear localization

Due to an active nuclear import and its ability to passively diffuse out of the nucleus, TDP-43 is a nucleocytoplasmic shuttling protein (32, 47). RNA binding retains TDP-43 in the nucleus by forming bigger macromolecular complexes that slow down its diffusion (28, 32). We therefore wondered whether oligomerization is a greater driver of physiological TDP-43 nuclear localization. To address this question, we first measured the mean fluorescence

intensity of GFP-TDP-43 in the nucleus and the cytoplasm for all four GFP-TDP-43 34 35 variants. Using DNA and G3BP as nuclear and cytoplasmic markers, respectively (Figure S2A), we observed that monomeric TDP-43 (6M and 6M&RRMm) showed a significantly 36 increased cytoplasmic localization compared to WT GFP-TDP-43 (Figure 2A and S2A). 37 Mislocalization was exacerbated in combination with a loss of RNA binding, suggesting an 38 independent, additive contribution of both protein-protein and protein-RNA interactions to 39 the nuclear localization of TDP-43. Similar observations were obtained in human neurons 40 transduced with HA-tagged versions of the TDP-43 variants (Figures 2B-C). 41

We subsequently sought to confirm these results by nucleocytoplasmic fractionation. In line 42 43 with the immunocytochemistry results (Figure 1C and 2A-C), upon mild lysis and nuclei isolation by centrifugation, WT GFP-TDP-43 was mostly retained in the nuclear fraction 44 (Figure 2D-E), with a small cytoplasmic pool corresponding to the subset of TDP-43 45 performing functions in this compartment (1). In contrast, oligomerization-deficient 46 mutations (6M and 6M&RRMm) consistently shifted the majority of GFP-TDP-43 to the 47 cytoplasmic fraction (Figures 2D-E), to a larger extent than observed bv 48 immunocytochemistry (Figure 2A and 2C). Importantly, the endogenous protein in the 49 same samples remained predominantly nuclear (Figure S2B). This contrast was even more 50 pronounced in the case of RRMm GFP-TDP-43, which fully shifted its localization to the 51 cytoplasm upon fractionation (Figures 2D-E) as opposed to its nuclear localisation by 52 immunocytochemistry (Figure 2A and 2C). We therefore wondered whether monomeric 53 and RNA binding-deficient GFP-TDP-43 exhibit an increased passive diffusion rate and 54 diffuse out of the nucleus during the fractionation procedure, when active nuclear import is 55 absent. Indeed, stabilization of TDP-43 oligomers by protein-protein cross-linking with 56 disuccinimidyl glutarate (DSG) (15) before nucleocytoplasmic fractionation increased the 57 retention of endogenous TDP-43 in the nucleus (Figure 2F-G). Conversely, when we 58 pretreated the cells with actinomycin D (ActD) to block transcription, the localization of 59 endogenous TDP-43 shifted to the cytoplasm (Figure 2F-G and S2C-D), as previously 60 reported (28, 32). Notably, this efflux of TDP-43 upon a decrease in nuclear RNA levels 61 and subsequent sample fractionation (Figure 2F-G) was more pronounced than observed 62 by immunocytochemistry (Figure 2H-I). These observations suggest that active nuclear 63 import compensates for the abundant passive TDP-43 egress from the nucleus in the absence 64 of oligomerization or RNA binding. Similar results were observed in human neurons, where 65 the combined treatment of ActD and ivermeetin, a nuclear import inhibitor, increased the 66 cytoplasmic shift of endogenous TDP-43 as compared to treatment with ActD alone (Figure 67 **2J-K**). Collectively, these observations show that RNA binding and protein-protein 68 interactions, especially its self-oligomerization, involve TDP-43 in larger macromolecular 69 complexes that are retained in the nucleus. 70

Oligomerization is required for physiological phase separation of TDP-43 in the nucleus

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TDP-43 has been shown to undergo LLPS (22), a phenomenon visible in the nucleus as 74 small droplets that fuse and split at endogenous protein concentrations (23-26). Such 75 endogenous LLPS-driven droplets were also observed in our model (Figure 3A) as 76 confirmed by treatment with 1,6-hexanediol (1,6-HD), an LLPS-suppressing alcohol (48). 77 1,6-HD decreased both the number and size of endogenous TDP-43 droplets (Figure 3A-B 78 79 and S3A). Although high concentrations of the LCR of TDP-43 are sufficient for phase separation in vitro (17, 49), additional interactions must take place for full-length TDP-43 80 to undergo physiological LLPS at far lower concentrations (24, 38). Recent evidence points 81

towards self-interaction through the NTD as another driver of TDP-43 LLPS *in vitro* (37, 38) and in human cells (34). Indeed, at a reported physiological concentration of 10 μ M (25, 50) and using dextran as a crowding agent (25, 51) purified maltose binding protein (MBP)tagged full-length TDP-43 phase separated into droplets, which also dissolved in the presence of 1,6-HD (**Figure 3C-D**). In contrast, oligomerization-deficient TDP-43 (6M) did not form droplets under the same conditions, suggesting that NTD interactions are essential for TDP-43 LLPS (**Figure 3C-D**).

These findings were reproduced in our isogenic cell lines. At comparable protein levels, 89 oligomerization disruption virtually suppressed all GFP-TDP-43 droplet formation 90 91 (Figures 3E-F), pointing to an essential role for oligomerization in physiological TDP-43 LLPS in cells. Interestingly, RNA binding-deficient GFP-TDP-43 also formed nuclear 92 droplets (Figures 3E-F), albeit smaller in size (Figure S3B). Since local protein 93 concentration is a known driver of phase separation (22), the number of GFP-TDP-43 94 droplets for both WT and RRMm was proportional to their nuclear protein levels (Figure 95 **3F**). Disruption of both oligomerization and RNA binding in the combined GFP-TDP-43 96 97 variant drastically reduced the number of nuclear droplets (Figure 3E-F), indicating that TDP-43 droplet formation in the absence of RNA binding is mediated through NTD-driven 98 99 TDP-43 oligomerization. This was supported by biochemical analysis with DSG proteinprotein crosslinking which showed protein complexes at the expected size of GFP-TDP-43 200 dimers for both WT and RRMm, but not for the 6M-containing variants (Figures 3G-H 201 and S3C-D). Similar to its WT counterpart, stabilization of these protein complexes via 202 crosslinking retained GFP-TDP-43 RRMm predominantly in the nucleus in our isogenic 203 cell lines despite the lack of RNA binding (Figure S2C-D). Overall, our results indicate 204 that NTD-driven oligomerization - and not only LCR interactions- are essential for TDP-43 205 LLPS in cells, both in the presence and absence of RNA. 206

Loss of RNA binding leads to conformationally distinct TDP-43 oligomers

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The observed GFP-TDP-43 dimers in the RRMm variant were significantly reduced 209 compared to TDP-43 with retained RNA binding capability (Figures 3G-H), suggesting 210 that TDP-43 oligomerization is modulated by RNA binding. Indeed, treatment with ActD 211 decreased the level of endogenous TDP-43 dimers detected by protein-protein cross-linking :12 (Figure 4A-B). Concomitant with this reduction in oligomerization, ActD-treated cells also 213 displayed a reduced number of TDP-43 nuclear droplets (Figure 4C-D). To determine 214 whether TDP-43 oligomerization is exclusively confined to nuclear droplets, we first 215 assessed the exact subnuclear location of the oligomers. For this purpose, we employed 216 proximity ligation assay (PLA) to visualize TDP-43 dimers with a single monoclonal 217 antibody (mAb) conjugated to two different oligonucleotides. With this approach, only 218 TDP-43 molecules that come to close proximity (maximum 20 nm apart) allow 219 oligonucleotide hybridization and fluorescent signal amplification. Surprisingly, while 20 abundant PLA signal was detected in the nucleus, only a fraction overlapped with its nuclear 21 droplets, suggesting that TDP-43 dimerization is not restricted to nuclear droplets (Figure 22 4E), a result that was confirmed for WT GFP-TDP-43 using a mAb against GFP (Figure 23 4F and S4A). 224

In contrast, the oligomerization-deficient variants showed a markedly decreased PLA signal, even in cells with comparable protein levels, confirming that the observed PLA signal depends on oligomerization. In line with our DSG cross-linking results (**Figure 3G-H**), the RNA-binding GFP-TDP-43 mutant displayed a positive PLA signal (**Figure 4F**).

Interestingly, the mean intensity of PLA-positive foci was consistently higher than that of :29 :30 its WT counterpart (Figure 4F) at comparable protein levels (Figure 4G). These results were confirmed using a mAb targeting a different tag in the GFP-TDP-43 protein (Figure :31 S4B). We further validated this finding with an alternative approach using a GFP tripartite :32 fluorescence complementation (triFC) assay that measures physiological dimerization, as :33 we previously showed (31, 52) (Figure S4C). Co-transfection of T₁₀- and T₁₁-tagged TDP-:34 43 variants along with a nuclear-targeted GFP₁₋₉ in mouse motor neuron-like cells showed :35 positive GFP complementation signal for the WT protein, but none of the other variants :36 (Figure 4H-I), despite similar nuclear protein levels (Figure S4D-E). This suggests that :37 despite our biochemical (Figures 3G-H) and imaging (Figures 4F-G and S4B) :38 observations indicating that RNA binding-deficient TDP-43 dimerizes, these dimers do not :39 come in close enough contact and with the correct orientation to reconstitute GFP 240 fluorescence. This contrast to the WT protein supports the notion of a distinct conformation 241 of TDP-43 dimers in the absence of RNA. Of note, transient expression of the monomeric :42 GFP-TDP-43 variants in this cellular model also showed a noticeable cytoplasmic fraction 243 (Figures 4H and S4F), comparable to our observations in the isogenic cell lines and in 244 human neurons (Figure 2B and C). Overall, detection and quantification of dimeric TDP-245 43 species by a combination of different imaging and biochemical methodologies supports 246 the view that RNA binding is required for the proper orientation of TDP-43 dimers and 247 likely oligomers. 248

TDP-43 partitions in heterogenous nuclear bodies via oligomerization and RNA binding

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Since RNA promotes phase separation by providing a scaffold for many phase-separated :52 nuclear bodies (53, 54), we sought to determine whether all nuclear TDP-43 droplets were 253 of similar structure and composition. For this purpose, we determined the colocalization of 254 endogenous TDP-43 in HEK293 cells with a broad panel of nuclear membraneless 255 256 organelles (Figure 5A). This approach revealed that a fraction of TDP-43 droplets, in particular the largest in size, were labeled with markers of Cajal bodies (Figure 5A-B), the :57 maturation compartments of spliceosomal small nuclear ribonucleoprotein particles. :58 Interestingly, TDP-43 was present in all Caial bodies in HEK293 cells and neurons (Figure 259 5C-D), but Cajal bodies only comprised a small fraction of all TDP-43 droplets. Another 260 portion of TDP-43 was localized within paraspeckles, but, unlike Cajal bodies, not every 261 paraspeckle contained TDP-43. TDP-43 was absent from other RNA-nucleated 262 compartments such as nuclear speckles (54) and from protein-exclusive compartments like 263 promyelocytic leukemia (PML) bodies (Figure 5A-B) (55). In contrast to WT TDP-43, 264 RNA binding-deficient GFP-TDP-43 was incorporated into neither Cajal bodies nor 265 paraspeckles, in line with previous reports (27) (Figure 5E-F and S5A-B). Intriguingly, 266 also monomeric GFP-TDP-43 was largely absent from these nuclear compartments (Figure 267 5E-F and S5A-B), highlighting the requirement of TDP-43 oligomerization for its 268 incorporation within functional nuclear bodies. Collectively, these observations suggest that :69 at least a fraction of TDP-43 LLPS arises from an RNA scaffold, since TDP-43 has been 270 shown to bind small Cajal body-specific RNAs (56) and NEAT1, the architectural RNA of 271 paraspeckles (20, 21, 53). The remaining, unidentified nuclear TDP-43 droplets, present in :72 both the WT and RNA binding-deficient variant (Figure 5E-F and S5A-B), might represent :73 a precursor pool for Cajal bodies and paraspeckles, an inert droplet population or even be 274 linked to yet undefined bodies and functions. Whether any of these potentially "RNAless" 275 TDP-43 nuclear compartments formed by the RNA binding-deficient variant are identical :76 to the unidentified droplets occuring in physiological conditions and/or have functional :77

- roles remains unclear. Overall, the heterogeneous nature of nuclear TDP-43 droplets suggests that LLPS is required for a wide array of TDP-43 functions within the nucleus.
- 181TDP-43 oligomerization is required for the transcriptome-wide splicing regulation of182its RNA targets

NTD-mediated oligomerization is required for splicing of at least a subset of the RNA 283 targets of TDP-43 (15, 35-38), but given the broad role of TDP-43 in regulating splicing 284 events (20, 21, 57, 58), the question remained whether oligomerization is essential for all 285 its splicing targets. RNA-sequencing (RNA-seq) of our isogenic cell lines revealed that 286 expression of the WT variant resulted in alternative splicing of >70 genes when compared 287 to the expression of the GFP-TDP-43 RRMm, an established splicing-deficient protein 288 version (16, 18), including previously reported events of exon inclusion/exclusion, intron 289 retention and alternative polyadenylation site usage depending on TDP-43 binding (20, 21, 290 58) (Figures 6A and S6A). When the same comparison was performed between the 291 oligomerization-deficient and the RRMm GFP-TDP-43, no significant alternatively spliced :92 events were found (Figures 6A and S6A), suggesting a lack of splicing functionality of :93 monomeric GFP-TDP-43, which was also observable at the differentially expressed RNA 294 and protein levels (Figure S6B). A particularly interesting event modulated by TDP-43 295 binding is the splicing of an alternative intron (intron 7) in its own 3' UTR, which results in 296 autoregulation of the TDP-43 mRNA and protein levels (20, 40, 59) (Figure S6C). Analysis :97 of the 3' UTR of TDP-43 by RNA-seq and qPCR showed that, similar to other alternative :98 splicing events, 6M GFP-TDP-43 cannot promote the exclusion of intron 7 in the 299 endogenous TARDBP mRNA (Figure 6B and S6D-E), resulting in lack of autoregulation 00 at the protein level (Figure 6C-D). This lack of splicing activity by monomeric TDP-43 01 was not due to its reduced protein concentration, since WT GFP-TDP-43 autoregulated 602 endogenous TDP-43 at comparable expression levels (Figure S6D-E). Overall, our data 03 support the requirement of TDP-43 oligomerization for its broad role in splicing regulation. 604

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Cytoplasmic TDP-43 oligomerization is required for its incorporation into stress granules

To understand whether oligomerization is also required for TDP-43 functions outside of the 608 nucleus, we studied TDP-43 incorporation into SGs in the cytoplasm (29). Since relocation :09 of TDP-43 into SGs depends on the cell type and the stressor applied (29), we resorted to 510 the GFP triFC assay (Figure S2L) to investigate TDP-43 complementation in the well-511 established oxidative stress model in HeLa cells (29, 30) T₁₀- and T₁₁-tagged TDP-43 12 variants and incubation with recombinant GFP₁₋₉ revealed that, unlike the WT protein -but 13 similarly to the RNA binding-deficient mutant-, monomeric TDP-43 did not incorporate 514 into SGs (Figure 6G-H), suggesting that TDP-43 oligomerization also takes place in the :15 cytoplasm. To confirm the presence of cytoplasmic TDP-43 oligomers in physiological 16 conditions, we developed isogenic HEK293 Flp-In T-REx lines harboring one copy of each 517 of the GFP-TDP-43 variants in combination with previously published mutations in the ;18 nuclear localization signal that abolish nuclear import (GFP-TDP-43 mutNLS) (60). Protein 19 levels of the GFP-TDP-43 mutNLS variants were similar to that of their nuclear 20 counterparts (Figure S6F and 1D). Interestingly, the localization of the GFP-TDP-43 21 mutNLS variants differed between the four cell lines. WT and RRMm GFP-TDP-43 22 mutNLS were predominantly present in the cytoplasm, whereas their monomeric 23 counterparts significantly shifted their localization to the nucleus (Figure S6G-H). This 24

suggests that WT and RRMm GFP-TDP-43 mutNLS oligomerize in the cytoplasm, which :25 hinders their passive diffusion back into the nucleus. To confirm this, we performed DSG :26 cross-linking of protein-protein interactions in the GFP-TDP-43 mutNLS lines and found :27 that both WT and RRMm GFP-TDP-43 mutNLS formed oligomers, albeit their presence 28 was reduced in the RRMm variant (Figure 61-J). Moreover, DSG cross-linking followed 129 by nucleocytoplasmic fractionation enabled the detection of WT and RRM GFP-TDP-43 30 oligomers in the cytoplasmic fraction (Figure 6K). Albeit less abundant than nuclear 31 oligomerization, cytoplasmic oligomerization was also observed at endogenous TDP-43 :32 levels by PLA (Figure 4E). Altogether, our observations suggest that oligomerization is :33 essential in both the nucleus and the cytoplasm for TDP-43 to perform its functions in RNA :34 metabolism. 135

Loss of RNA binding or oligomerization differentially modulate the subcellular localization of TDP-43 inclusions

Decline of the cellular proteostasis capacity with age contributes to protein misfolding in :39 neurodegenerative diseases (43), often resulting in the accumulation of ubiquitinated 640 inclusions in affected tissues, including TDP-43 proteinopathies (2, 3). Since monomeric 41 and RNA binding-deficient TDP-43 showed shorter half-lives at physiological levels :42 (Figure 1D-E), we sought to determine how failure of the UPS machinery affects the 43 accumulation of these species. By blocking the proteasome with the inhibitor MG132, we 44 observed that both monomeric and RNA binding-deficient GFP-TDP-43 formed protein 45 inclusions in the isogenic cell lines, in contrast to the WT counterpart, which remained 646 largely diffuse (Figure 7A-B and S7A). The vast majority of aggregates formed by 47 monomeric GFP-TDP-43 (6M and 6M&RRMm) localized to the cytoplasm, in line with 48 previous results showing that high overexpression of oligomerization-deficient GFP-TDP-:49 43 by transient transfection triggers cytoplasmic TDP-43 aggregation (15). Interestingly, in 50 addition to cytoplasmic inclusions, MG132 treatment resulted in nuclear aggregation of 51 :52 RNA binding-deficient GFP-TDP-43 in >70% of the cells. The combined loss of oligomerization and RNA binding shifted the aggregation to the cytoplasm, suggesting that :53 nuclear TDP-43 aggregation depended on NTD interactions. The observed TDP-43 54 aggregation patterns were specific to the inhibition of the UPS degradation pathway, as :55 several classes of proteasome inhibitors, but not an autophagy one, yielded similar outcomes 56 in the isogenic cell lines (Figure S7B-C). 57

In human neurons, monomeric TDP-43-HA variants also predominantly aggregated in the :58 cytoplasm, whereas RRMm TDP-43-HA additionally presented nuclear inclusions in >50% :59 of transduced neurons (Figure 7C-D), thus reproducing the distinct TDP-43 aggregation 60 patterns observed in isogenic cell lines. Upon MG132 treatment, WT TDP-43-HA also 61 formed inclusions in neurons, both in the nucleus and cytoplasm, likely due to higher 62 transgene protein levels in transduced neurons compared to the isogenic GFP-TDP-43 lines. 63 Interestingly, and in line with higher protein levels, RRMm TDP-43-HA already formed 64 nuclear inclusions in the absence of proteasome inhibition in a subset of transduced human 65 neurons (Figure S7D). Taken together, our data show that loss of oligomerization shifts 666 TDP-43 aggregate formation from the nucleus to the cytoplasm. 67

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TDP-43 aggregates in an LLPS- or an aggresome-dependent manner in the nucleus and cytoplasm, respectively

To understand the origin of cytoplasmic and nuclear TDP-43 inclusions, we performed live 71 ;72 cell imaging of the GFP-TDP-43 isogenic lines during the treatment with the proteasome inhibitor. While WT GFP-TDP-43 droplets merely changed position, fused and split within ;73 the nucleus upon MG132 addition, monomeric GFP-TDP-43 formed a single cytoplasmic ;74 inclusion whose size increased over time, accompanied by gradual decrease in the diffused :75 nuclear TDP-43 signal (Figure 8A) resembling the nuclear clearance that has been widely ;76 reported in neurons with TDP-43 pathology in ALS and FTLD patients (1-3). Fluorescence :77 ;78 recovery after photobleaching (FRAP) experiments revealed that whereas WT GFP-TDP-43 remained diffuse throughout the treatment, cytoplasmic inclusions comprising ;79 monomeric TDP-43 were immobile structures, (Figure 8B and S8A-B). A similar 80 aggregation pathway –a single focus expanding in size yielding one solid cytoplasmic 81 inclusion- was also observed for the RRMm GFP-TDP-43 in the cytoplasm (Figure S8C-82 **D**), suggesting that the formation of cytoplasmic aggregates upon proteasomal failure does 183 require neither oligomerization nor RNA binding. In contrast, in the nucleus, the elevated 84 protein levels of RRMm GFP-TDP-43 caused by proteasome inhibition induced the 185 formation of many initially dynamic droplets, which eventually fused to form a single solid 86 inclusion (Figure 8B-C), reminiscent of the single intranuclear inclusions found in patients 87 with specific FTLD subtypes (6, 61). FRAP analysis showed that GFP-TDP-43 RRMm in 188 the final inclusion had lost its fluid behavior (Figure 8B). In a subset of cells, RRMm GFP-189 TDP-43 deposited in the nucleoli (Figure S8C), in line with the protein quality control 90 properties of this phase-separated compartment (62). Importantly, also in human neurons 91 the formation of nuclear aggregates in a subset of transduced cells expressing RRMm TDP-192 43-HA (Figure S7B) was accompanied by the pronounced presence of nuclear droplets in 193 transduced cells without inclusion (Figure S8F). Together, these data suggest that TDP-43 94 aggregates via LLPS in the nucleus. 195

Cytoplasmic TDP-43 inclusions were consistently found adjacent to the nucleus, in a 96 location occupied by the aggresome, a juxtanuclear accumulation of misfolded proteins 97 resulting from saturation of the chaperone refolding system and/or the UPS degradation 198 pathway (Figure S8G) which has been linked to the origin of protein aggregates in 99 neurodegenerative diseases (63). Indeed, cytoplasmic inclusions, formed by 6M, RRMm -00 or 6M&RRMm GFP-TDP-43, were surrounded by a vimentin cage, a characteristic 01 feature of aggresomes (Figure 8D and S8H) (63). Additionally, cytoplasmic, but not 02 nuclear, TDP-43 aggregates were positive for p62 (Figure 8E), a critical component of 03 aggresomes (64) and a pathological aggregate marker in certain FTLD subtypes (6, 61). 04 These observations suggest that distinct pathways towards TDP-43 aggregation are at play 105 in the nucleus and cytoplasm and that monomerization increases TDP-43 incorporation in 06 cytoplasmic aggresomes upon proteasomal failure, thereby potentially triggering +07cytoplasmic TDP-43 aggregation in disease. 108

10 **Discussion**

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In this study, we describe the interconnection between NTD-driven TDP-43 oligomerization 11 and RNA binding, and show that they cooperatively retain TDP-43 in the nucleus, instruct 12 its functionality and therefore slow down its turnover. We demonstrate that oligomerization 13 is essential for the broad function of TDP-43 in splicing regulation and allows its LLPS-14 mediated incorporation into nuclear membraneless compartments, including Caial bodies 15 and paraspeckles. Our work describes for the first time that, under physiological conditions, 16 TDP-43 oligomers exist in the cytoplasm, where they are required for LLPS-dependent 17 incorporation into SGs. Moreover, we show that, in the absence of RNA binding, TDP-43 18

oligomerization is reduced, but the dimers that do form adopt a spatial conformation that is
 different from the RNA-bound oligomeric state. Importantly, our results shed light on the
 molecular mechanism of two distinct and independent pathways triggering TDP-43
 aggregation, which highlight the importance of TDP-43 monomerization and/or loss of
 RNA binding as key early events in the development of TDP-43 proteinopathies (Figure
 9).

TDP-43 oligomerization is a dynamic event that controls the relative amounts of TDP-43 25 monomers, dimers and oligomers in the cell. However, the specialized roles of the 26 individual TDP-43 species in health and disease remained unknown. Expression of different 27 28 TDP-43 variants at near-physiological levels in our human cell line and neural system revealed that, in comparison to its oligometric counterpart, monometric TDP-43 lacks 29 functionality, becomes more prone to escape the nucleus -likely by passive diffusion (32, 30 47)- and is rapidly degraded. In the event of proteasomal failure, these otherwise short-lived 31 TDP-43 monomers are deposited into aggresomes, whose expansion is accompanied by a 32 progressive decrease in nuclear TDP-43. This cytoplasmic aggregation observed in our 33 34 human cellular models recapitulates the key pathological TDP-43 features observed in affected neurons of patients with TDP-43 proteinopathies, namely its nuclear clearance, loss 35 of function and cytoplasmic aggregation (2, 3). Interestingly, the aggresome pathway was 36 previously linked to sporadic ALS/FTLD (65) through the disturbance of p62 (65, 66), a 37 key player in aggresome formation (64). Furthermore, aggresome markers HDAC6 and p62 38 have been reported to colocalize with a subset of cytoplasmic TDP-43 aggregates in 39 ALS/FTLD patients (6, 61, 67). When the conditions that favor the monomeric state of 40 TDP-43 and hamper its proteasomal degradation persist, TDP-43 monomers contained in 41 the aggresomes may resist clearance by aggrephagy and rather further mature into compact 42 aggregates due to the high concentration of unfolded LCRs in the absence of NTD-driven 43 organization that keeps them apart (15). Since monomeric TDP-43 is unable to autoregulate 44 its own levels, the continued production of more TDP-43 to compensate for its loss of 45 function will only exacerbate this pathological transition. Our data signifies that loss of 46 TDP-43 oligomerization ignites a pathological cascade that culminates in the formation of 47 cytoplasmic TDP-43 inclusions via the aggresome pathway. 48

In addition to loss of oligomerization, our data indicate that also the disturbance of TDP-43 49 oligomer conformation triggered by loss of RNA binding plays a role in pathology through 50 a distinct molecular pathway in the nucleus. While previous studies have addressed the role 51 of RNA in TDP-43 pathogenesis, our work provides insights into the molecular mechanism 52 underlying the aberrant phase transition of NTD-driven RNAless TDP-43 oligomers into 53 nuclear immobile inclusions. Using a combination of imaging and biochemical assays, we 54 observed that TDP-43 oligomers present a different conformation in an RNA-bound or -55 unbound state. RNA-bound TDP-43 oligomers enact its physiological functions, maintain 56 its localization and antagonize the formation of pathological aggregates (15 and this study), 57 58 while RNAless TDP-43 oligomers, that may or may not have functional roles, undergo aberrant phase separation leading to nuclear aggregation (37 and this study). These 59 observations clarify the apparently controversial role of the NTD in aggregate formation, 60 which has been found both synergistic (35, 68-70) and antagonistic (15). Based on our 61 observations, we propose that NTD-mediated TDP-43 self-interaction is a double-edged 62 sword: in the presence of RNA, it is essential for TDP-43 to perform its functions and 63 undergo physiological LLPS, while in the absence of RNA binding it promotes aberrant 64 LLPS that leads to aggregation. 65

In addition to the cytoplasmic aggregation of TDP-43, neuronal intranuclear inclusions have 66 67 been reported in two of the five recognized FTLD-TDP subtypes (2, 3, 6, 71-73) and are particularly abundant in FTLD cases linked to mutations in valosin-containing protein 68 (VCP) (71, 72), which is involved in nuclear protein quality control degradation (74). In our 69 human cellular models, we observed LLPS-driven intranuclear aggregation of RNA 170 binding-deficient TDP-43 oligomers upon inhibition of proteasomal degradation. Similarly, 171 nuclear inclusions have also been reported in previous studies employing strong +72 173 overexpression of TDP-43 RNA binding mutants in cells and neurons (23, 33, 36, 42, 60). In addition to its LLPS-mediated nuclear aggregation, RNA binding-deficient TDP-43 74 formed cytoplasmic inclusions via the aggresome pathway. This route is likely favored by 175 the increase of the monomeric state in the absence of RNA binding, accompanied by nuclear 176 efflux, as previously shown upon proteasome inhibition in cultured neurons (75). 177 Altogether, our data show that when a cell encounters proteostatic stress, TDP-43 takes 78 179 different routes towards inclusion formation, and the selection of the pathway depends on both the protein state (monomeric versus RNAless) and the subcellular environment 80 (cytoplasm versus nucleus). While our study provides evidence for the importance of two 81 such routes (nuclear LLPS-dependent and cytoplasmic aggresome-mediated), additional 82 aggregation pathways - for example, cytoplasmic LLPS-mediated (24, 33, 67)- likely exist 83 and may be triggered under different circumstances and involve other TDP-43 states. 84 Collectively these distinct pathways may account for the spectrum of cytoplasmic 85 aggregates observed in patients with TDP-43 proteinopathies. 186

Why would the majority of TDP-43 aggregates reside in the cytoplasm in ALS/FTLD 87 patients? Our data indicates that TDP-43 monomerization and its subsequent nuclear efflux 88 is a more frequent or potent event than decreased RNA binding affinity in these cases. 89 Alternatively, the high nuclear RNA concentrations that have been shown to prevent LLPS 90 of RNA-binding proteins (50) may counteract aberrant LLPS and aggregation, even in the 91 absence of specific RRM-RNA interactions for TDP-43. This balancing mechanism may be -92 reinforced by the upregulation of some architectural TDP-43 RNA targets in the nucleus, 193 including NEAT1 (26, 50). This binding could instruct the proper, RNA-loaded orientation 94 of oligomers with a concomitant increase in physiological LLPS in the form of nuclear 95 bodies. In fact, elevated *NEAT1* levels and paraspeckle formation have been amply reported 96 in TDP-43 proteinopathies (21, 26, 76, 77). 197

The cellular machinery that regulates TDP-43 oligomerization remains unknown. While -98 99 protein concentration is a determinant of TDP-43 oligomerization (14, 15), the conditions that keep the balance between monomeric and oligomeric TDP-43 species in healthy cells 500 and, most importantly, increase TDP-43 monomerization in disease require further 501 investigation. Post-translational modifications (PTMs) are excellent candidates for such ;02 physiological regulation of TDP-43 oligomerization and a recent study identified a single ;03 serine phosphorylation event within the NTD interface that decreases oligomerization in 504vitro (38), albeit its effects within cells or occurrence in disease have not yet been tested. ;05 Similarly, specific acetylation events within the RRMs that lower the affinity of TDP-43 for ;06 RNA have been detected in ALS patients (78) and were subsequently reported to trigger ;07 nuclear TDP-43 aggregation via aberrant LLPS (34). Supporting the link between reduced ;08 RNA affinity and disease, ALS/FTD-associated mutations within the RRMs have been ;09 shown to disrupt RNA binding and enhance TDP-43 proteinopathy (79). Moreover, since ;10 ATP was recently shown to directly bind the NTD of TDP-43, thereby enhancing its ;11 oligomerization (44), the decrease in cellular ATP levels with age (80) could also act as a ;12 monomerization-inducing trigger. It is also conceivable that additional, yet unknown, ;13 protein interactors of TDP-43 may act as modifiers of oligomerization. Future work should ;14

focus on determining the molecular switches within TDP-43 and/or protein partners that regulate its oligomerization and RNA binding and on validating the occurrence of diminished TDP-43 oligomerization and RNA affinity in patient tissue. Such insights will be valuable to inform drug design targeting TDP-43 oligomerization and RNA binding, including dimer stabilization or recovery of RRM-RNA interactions, among others.

In conclusion, oligomerization and RNA binding allow TDP-43 to maintain its localization and function in physiology and their disruption drives distinct aggregation pathways in the nucleus and cytoplasm, indicating that distinct molecular origins may account for the plethora of TDP-43 aggregation types observed in ALS and FTLD subtypes.

524

Materials and Methods

Plasmids

The pcDNA5 plasmid containing the GFP-tagged human TDP-43 cDNA sequence was a ;27 kind gift of Dr. Shuo-Chien Ling (41). The mutations to introduce a siRNA resistance in the ;28 TDP-43 coding region without altering the amino acid sequence and the oligomerization-;29 disrupting mutations in the N-terminal domain were previously described (15). RNA-;30 binding disruption mutations were introduced by site-directed mutagenesis PCR as ;31 described before (15), using high fidelity Phusion DNA Polymerase (New England Biolabs, ;32 M0530) with primers detailed in Table S1 followed by DpnI (New England Biolabs, ;33 R0176) digestion before bacterial transformation and colony selection. ;34

Plasmids encoding T_{10} - and T_{11} -tagged TDP-43 and TDP-43 6M have already been reported ;35 (15, 52). The sequence encoding TDP-43 RRMm was amplified by PCR from the pcDNA5 ;36 plasmids described above by adding BamHI/XhoI restriction sites using the primers detailed ;37 in Table S1. Amplified products were cloned into the T₁₀ and T₁₁pcDNA3 parental plasmids ;38 ;39 (52) between BamHI and XhoI in order to obtain plasmids encoding T₁₀-HA-TDP43 RRMm and T_{11} - β 1-TDP-43 RRMm. The same amplified sequence was cloned into the plasmids ;40 encoding T₁₀- and T₁₁-tagged TDP-43 6M (52) between EcoRI and XhoI in order to obtain ;41 plasmids encoding for T₁₀-HA-TDP43 6M&RRMm and T₁₁-β1-TDP-43 6M&RRMm. ;42

- TDP-43 with a C-terminal HA tag was directly amplified from the GFP-TDP-43 construct described above and inserted into an autoregulatory all-in-one TetON cassette previously inserted into pLVX backbone (*81*) by Gibson cloning using the NEBuilder HiFi DNA Assembly Cloning Kit (New England Biolabs, E5520S) according to the manufacturer's instructions.
- The plasmid encoding the His6-tagged RRMs of TDP-43 for bacterial protein expression ;48 was previously published (16) and mutations disrupting RNA binding were introduced as ;49 described above. The plasmid encoding TDP-43-MBP-His6 for bacterial protein expression 50 was a gift from Nicolas Fawzi (Addgene plasmid #104480; http://n2t.net/addgene:104480; ;51 RRID:Addgene 104480) (38). Point mutations for 6M, S2C, C39S and C50S were ;52 introduced with site-directed mutagenesis PCR with high fidelity Phusion DNA Polymerase ;53 (New England Biolabs, M0530) using primers described in Table S1. The plasmid encoding ;54 the MBP-His6 was generated using deletion cloning by PCR with high fidelity Phusion ;55 DNA Polymerase (New England Biolabs, M0530) using designed primers described in ;56 Table S1. ;57
- **Recombinant protein expression and purification**

Production of TDP-43 RRMs for NMR studies was performed as previously described (16). ;59 Production of full-length TDP-43 was performed as previously reported (7). For more 60 details, see Supplementary materials & methods. The MBP-His6 was purified as the full-61 length TDP-43 but as purity was reached after the Ni Sepharose Excel material the protein 62 was subsequently dialysed against a storage buffer (20 mM Tris pH 8.0, 300 mM NaCl, 1 63 mM TCEP) overnight at 4°C, concentrated the next day using Amicon Ultra-15 64 concentrators MWCO 10 kDa (Merck Millipore, UFC901024), flash frozen and kept at -80 65 °C. ;66

67 **Circular dichroism experiments**

TDP-43-MBP constructs were thawed on ice and centrifuged for 15 min at 17 100 g and ;68 4°C. The buffer was exchanged to 10 mM sodium phosphate pH 7.4 (0.036 % (w/v) sodium ;69 phosphate monobasic (Sigma Aldrich, S3139), 0.2 % (w/v) sodium phosphate dibasic ;70 (Sigma Aldrich, S5136)) using 500 µl concentrators with a MWCO of 10 kDa (Merck ;71 Millipore, UFC501024) 5 x (11 000 g, 10 min, 4°C). To ensure the solubility of the isolated ;72 protein, the concentrate was again centrifuged as described above. Protein concentration ;73 was determined by the extinction coefficients and molecular weights of the constructs ;74 (ExPASy ProtParam software) and absorbance at 280 nm (NanoDrop, Thermo Scientific). ;75 CD spectra of 200 µg/ml recombinant protein were recorded at 20 °C from 180-250 nm ;76 with a bandwidth of 1 nm and a sampling period of 25 s on a Chirascan V100 (Applied ;77 Photophysics). Ellipticity (in mdeg) was transformed to molar ellipticity (in deg cm² dmol⁻ ;78 ¹) with the following formula, where c (in M) is the concentration and L (in cm) represents ;79 ;80 the pathlength of the cuvette:

;81

;95

$[\theta] = \frac{\theta}{10 \text{ x c x L}}$

¹⁸² Phase separation experiments

Pure concentrated fractions of recombinant protein MBP-tagged full-length WT and 6M ;83 TDP-43 were desalted in a buffer of 20 mM sodium phosphate pH 7.2 (0.072 % (w/v) ;84 sodium phosphate monobasic, 0.4 %(w/v) sodium phosphate dibasic), 300 mM NaCl, ;85 0.001% (v/v) TWEEN 20, 50 mM L-arginine (Merck Millipore, A5006), 50 mM L-glutamic ;86 acid (Merck Millipore, G1251) on a HiTrap desalting column (Cytavia, 17140801). An ;87 ;88 equimolar fluorophore labeling reaction was set up with CF660R maleimide (Biotum Inc., Fremont, CA) previously dissolved in dimethylformamide (DMF) in an N2-hood and ;89 incubated for 16 h at 4°C under constant rotation. Reaction was stopped with 10 mM DTT ;90 and subsequently passed through a HiLoad 10/300 Superdex 200 pg SEC (Cytiva, ;91 28990944) in SEC buffer on an Äkta pure system (Cytavia). Labelling position was ;92 confirmed by mass spectrometry and labelling ratio was determined by first determining the ;93 ;94 correct protein concentration:

Protein concentration (M) =
$$\frac{A_{280} - (A_{max CF660R} \times CF)}{\epsilon}$$

where A_{280} is the absorbance of the protein at 280 nm, $A_{max CF660R}$ is the CF660R dye absorbance at 663 nm (absorbance maximum of CF660R), CF is the correction factor for the amount of absorbance at A_{280} caused by the dye and ε is the protein molar extinction coefficient.

The labelling ratio was subsequently calculated with following formula:

Dye/protein ratio =
$$\frac{A_{max CF660R}}{\epsilon' x \text{ protein concentration (M)}}$$

where, ε ' is the molar extinction coefficient of the CF660R fluorescent dye.

Correctly labelled protein with a labeling efficiency of >75% was snap frozen and stored at 503 -20°C. TDP-43-MBP constructs were thawn and centrifuged (17100 g, 15 min, 4°C) before 504 exchanging the buffer to a to phase separation buffer consisting of 20 mM HEPES pH 7.4 605 (Biosolve, 0008042359BS), 150 mM NaCl, 1 mM TCEP. The samples were centrifuged 5 606 x (11 000 g, 10 min, 4°C) and protein concentration determined as described above. Phase 507 separation of 10 µM TDP-43-MBP containing CF660R-labelled TDP-43-MBP at a ratio of 508 1/200 was induced using dextran (Sigma, D8906) at a final concentration of 10% (w/v) in 509 phase separation buffer and incubation for 2 h at 22°C in u-Slide Angiogenesis glass-bottom 510 coverslips (ibidi, 81507). Images were acquired on a fluorescence microscope (Nanoimager 511 S, ONI) with an Olympus 100x objective (1.4 NA) using a wavelength of 640 nm and 13% 512 513 laser power. Reversibility was tested by adding 1,6-hexanediol (Sigma, 240117) to a final concentration of 8% (w/v) in phase separation buffer for 10 min at 22°C and subsequent 514 imaging. Image analysis and droplet counts were performed with ImageJ (1.52k). 515

616Cell culture

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All cells were cultured at 37°C with saturated humidity and an atmosphere of 5% CO2. HeLa cells were cultured in DMEM (Gibco, 61965-059) supplemented with 1% nonessential amino acids (Gibco, 11140035) and 10% fetal bovine serum (FBS; Gibco, 10270-106).

Mouse motor neuron-like hybrid cells NSC-34 (Bioconcept, CLU140) were proliferated on Matrigel (Corning, 354234)-coated dishes in Dulbecco's modified Eagle medium (DMEM; Sigma, D5671) supplemented with 10% FBS, 1X GlutaMAX (Gibco, 35050-061), 100 U/ml penicillin and 100 μ g/ml streptomycin (Gibco, 15140-122). For experiments, differentiation was induced by switching DMEM/F12 medium (Gibco, 21331-020) supplemented with 1X GlutaMAX, 1X B27+ supplement (Gibco, 17504-044), 1X N2 supplement (Gibco, 17502-048), 100 U/ml penicillin and 100 μ g/ml streptomycin.

- HEK293 Flp-In T-REx stable lines were cultured in DMEM (Sigma, D5671) supplemented 528 with 10% FBS, 1X GlutaMAX, 15 µg/ml blasticidin S (Gibco, R21001; Invivogen, ant-bl-529 10p), 100 µg/ml hygromycin B (Gibco, 10687010), 100 U/ml penicillin and 100 µg/ml 630 streptomycin. For details regarding live development, see Supplementary materials and 531 **methods**. When required, cells were treated with the appropriate chemicals at the following 632 concentrations: 1 µg/ml doxycycline (Clontech, 631311), 2.5 µM MG132 (APExBIO, *i*33 A2585), 1.25 µM MLN9708 (Selleck Chemicals, S2181), 500 nM bortezomib (BTZ; 534 APExBIO, A2614), 100 nM bafilomycin A1 (Sigma-Aldrich, SML1661), 100 µg/ml 635 cycloheximide (Sigma-Aldrich, C4859), 5 µg/ml actinomycin D (Sigma-Aldrich, A1410), *i*36 10% 1,6-hexanediol, 1mM sodium arsenite (Sigma-Aldrich, 35000-1L-R), dimethyl 537 sulfoxide (DMSO; Sigma-Aldrich, D2650). **i**38
- Human neural networks were differentiated from an in-house-developed iPSC-derived selfrenewing human neural stem cell line (iCoMoNSCs) obtained from control human skin fibroblast, as described previously (81). For details regarding the culture and differentiation, see **Supplementary materials and methods**. When required, neural cultures were treated with the appropriate chemicals at the following concentrations: 1 μ g/mL doxycycline, 10 μ M MG132, 10 μ g/ml actinomycin D, 25 μ M ivermectin (Sigma-Aldrich, I8898).

i45Lentiviral vector production

TDP-43-HA variants were packaged into lentivirus, harvested and concentrated as described previously (*81*). Lentiviral pellets were resuspended in neural maturation medium (containing all supplements but forskolin and cAMP, see **Supplementary materials and methods**), achieving 10x concentrated preparations of which the lentiviral titer was determined using Lenti-X GoStix Plus (Takara, 631280). Lentiviral preparations were aliquoted and stored at -80°C until use.

Quantitative PCR (qPCR)

HEK stable lines were plated at a density of 3 x 105 cells/well in a 6-well plate (TPP, 53 92406). Expression of GFP-TDP-43 was induced with 1 µg/ml doxycycline after 24 hours 554 and cells were harvested 48 h later. Total RNA from cells in a single well of a 6-well plate 655 was isolated using the RNeasy Plus Mini Kit (Qiagen, 74134) according to the 556 manufacturer's instructions. Complementary DNA (cDNA) corresponding to 1 µg was 57 generated using oligo(dT)20 primers and the SuperScript[™] III First-Strand Synthesis 558 SuperMix (Invitrogen, 18080400) according to the manufacturer's instructions. A qPCR of 559 50 cycles was performed with 10 ng of cDNA and 6.6 pmol of each of the primers (Table 660 S2) per reaction using Fast SYBRTM Green Master Mix (Applied Biosystems, 4385612) in 61 a AriaMx Real-Time PCR System (Agilent, G8830A). Relative fold gene expression was 62 calculated with the $2-\Delta\Delta Ct$ method. 663

GFP complementation assay

Tripartite GFP complementation experiments were performed as described before (15, 52). For more details, see **Supplementary materials and methods**.

67 **Protein-protein cross-linking in cells**

Native protein-protein interactions were stabilized by crosslinking with disuccinimidyl 68 glutarate (DSG; Thermo Scientific, 20593 or A35392) as previously reported (15). In brief, 669 570 cells grown to 80% confluency in 6-well or 10 cm plates were washed once in cell culturegrade PBS (Gibco, 10010-015), scrapped in ice-cold cell culture-grade PBS and collected 571 at 300 g and 4°C for 5 min in a 1.5 ml microfuge tube. Cells were resuspended in 100/600 572 µl ice-cold cell culture-grade PBS containing protease inhibitors and a freshly prepared 20 573 mM or 100 mM DSG solution in DMSO was added to the suspension to a final 574 concentration of 1 mM. After incubation at 25 °C and 1500 rpm for 30 min in a 575 Thermomixer (Eppendorf, 2230000048), the reaction was quenched by addition of Tris base 576 to a final concentration of 20 mM and further incubation for 15 min. Cells were collected 577 by centrifugation at 300 g for 5 min. 578

Nucleocytoplasmic fractionation of cells

Nucleocytoplasmic fractionation of the GFP-TDP-43 (mutNLS) isogenic HEK293 lines 680 was performed following a previously published protocol (82) with the following changes. 581 Cells in 10-cm dishes at 80-90% confluency were scrapped in 1 ml cell culture-grade PBS, 582 collected by centrifugation at 300 g for 5 min and resuspended in 880 µl. The final pellet 683 fraction containing the isolated nuclei was resuspended in 380 µl of 1X Laemmli buffer and 584 1X reducing agent (Invitrogen, B0009) in cell culture-grade PBS supplemented with 685 protease inhibitors, and 10 μ l of each three of the final fractions were loaded onto the **i**86 polyacrylamide gel for western blot analysis. 587

588 Immunocytochemistry

For immunocytochemistry experiments, cell lines were plated onto poly-D-lysine (Sigma**i**89 Aldrich, P6407)-coated 96- or 24-well plates (Greiner Bio-One, 655090; ibidi, 82426) or 8*i*90 *5*91 well glass chambers (ibidi, 80827). Unless stated otherwise, cell line cultures were fixed in 4% methanol-free formaldehyde (Thermo Scientific, 28908) in warm medium for 15 min, 592 washed with cell culture-grade PBS, permeabilized and blocked in 10% donkey serum **i**93 (Sigma-Aldrich, S30-M) and 0.1% Triton X-100 in cell culture-grade PBS for 10 min at *i*94 RT. Neural cultures were plated onto Matrigel-coated 96-well plates (135 000 cells/well) or *i*95 8-well glass-bottom chambers (between 240 0000 and 425 000 cells/well). Neural cultures *i*96 **i**97 were fixed in 4% methanol-free formaldehyde in cell culture-grade PBS for 25 min, washed with cell culture-grade PBS, permeabilized in 0.5% Triton X-100 in cell culture-grade PBS **i**98 for 5 min and blocked in 10% donkey serum and 0.1% Triton X-100 in cell culture-grade ;99 PBS for 30 min at RT. Primary antibodies (Table S3) were diluted in the same buffer and '00' incubated with the samples overnight at 4°C. After three washes with cell culture-grade '01 PBS, Alexa Fluor-conjugated donkey secondary antibodies (Table S3) were incubated for '02 1h at RT and subsequently further washed with cell culture-grade PBS. Nuclei were '03 counterstained with 1 µg/ml DAPI (Thermo Scientific, 62248) and samples imaged in cell '04 culture-grade PBS. '05

'06 **Proximity ligation assay (PLA)**

Cell lines plated on 96-well plates were fixed in pure ice-cold methanol for 7 minutes at -'07 20°C, followed by three cell culture-grade PBS washes. PLA assay was performed using '08 custom labelled antibodies and the Duolink In Situ Detection Reagents Red (Sigma-Aldrich, '09 DUO92008) as described by the manufacturer but with slight modifications. In brief, '10 antibodies of choice (anti-GFP and anti-HA in Table S3) in a carrier-free buffer were '11 labelled with MINUS and PLUS probes using the Duolink In Situ Probemaker MINUS '12 (Sigma-Aldrich, DUO92010) and Duolink In Situ Probemaker PLUS (Sigma-Aldrich, '13 DUO92009) according to the manufacturer's instructions. Cells grown in 96-well plates '14 were fixed in pure ice-cold methanol for 7.5 min at -20° C, followed by two washes with '15 cell culture-grade PBS. After blocking for at least 15 min with blocking solution at room '16 temperature, probe-conjugate antibodies were incubated overnight at 4°C. After two washes '17 of 5 minutes each with buffer A (10 M Tris, 150 mM NaCl 0.05% Tween 20), ligation '18 reaction was performed as indicated by the manufacturer. After an additional two washes '19 '20 of 5 min each with buffer A, amplification reaction was performed as indicated by the manufacturer. The reaction was quenched with buffer B (100 mM Tris, 100 mM NaCl). If '21 required, secondary antibodies were incubated in cell culture-grade PBS overnight at 4°C, '22 followed by DAPI counterstaining and imaging in cell culture-grade PBS. '23

²²⁴ Fluorescence in situ hybridization (FISH)

FISH assay was performed using target probes for NEAT1 (Invitrogen, VX-01) and the 125 '26 View RNA Cell Plus Assay (Invitrogen, 88-19000) according to the manufacturer's instructions with slight modifications. HEK stable lines were plated at a density of 10×10^4 '27 cells/well onto poly-D-lysine-coated 96-well plates. After 24 h, expression of GFP-TDP-43 '28 was induced and, after further 48 h, cells were fixed in ViewRNA Cell Plus '29 '30 Fixation/Permeabilization Solution for 30 min at room temperature. After five washes with 1x PBS with RNase Inhibitor, ViewRNA Cell Plus Probe Solution containing the target '31 probes was incubated for 2 h at 40°C. Following, the cells were washed with ViewRNA '32 Cell Plus Wash Buffer Solution and Signal amplification was continued as indicated by the '33

manufacturer. Nuclei were counterstained with DAPI and samples were imaged in cellculture-grade PBS.

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075 Acknowledgments

We thank Prof. Benjamin Schuler (Department of Biochemistry, University of Zurich) for)76 critical feedback on this work and Dr. Matthias Gstaiger (Institute of Molecular Systems)77 Biology, ETH Zurich, Switzerland) for sharing the Flp-In T-REx HEK293 cell line and the)78 pOG44 plasmid. We gratefully acknowledge the support from the Proteomics team at the)79 Functional Genomics Center Zurich (FGCZ) from the University of Zurich (UZH),)80 especially Dr. Paolo Nanni, Dr. Serge Chesnov and Dr. Witold Wolski. We also like to)81 thank Dr. Jana Doehner, Dr. Joana Delgado Martins, Dr. José María Mateos Melero and 182 Johannes Riemann from the Center for Microscopy and Image Analysis (ZMB) at the 183 University of Zurich UZH for their kind help with image acquisition and analysis. We are)84 grateful to Dr. Erik Slabber for his technical advice on cloning and PLA and help during 185 manuscript preparation, and to our fellow Polymenidou lab members for the continuous and 186 fruitful discussions on the project. 187

- Funding: This work was supported by the Swiss National Science Foundation (grants PP00P3_144862 and PP00P3_176966 to MP), the National Centre for Competence in Research (NCCR) RNA & Disease and a Sinergia grant (CRSII5_170976 11). M.P.-B. received a Candoc Grant (Forschungskredit) from the University of Zurich. V.I.W. is supported by the FEBS Long-Term Fellowship. P.D.R. received a Career Development Award from the Stiftung Synapsis – Alzheimer Forschung Schweiz. S.S. was supported by a Swiss Government Excellence Scholarship for Foreign Scholars.
- Author contributions: M.P.-B performed plasmid cloning unless stated otherwise below, 195 developed all stable isogenic HEK cell lines and carried out the corresponding experiments 196 on them, including sample preparation for transcriptomics and proteomics. V.I.W. cultured 197 the human neural networks, cloned LV transfer plasmids, produced lentiviral vectors and 198 performed experiments on neurons and mutNLS isogenic HEK lines. A.Z. performed the 199 full-length TDP-43-MBP protein purification, its corresponding plasmid cloning and in 00 vitro experiments. L.D.V. provided cell culture support and performed 01 immunocytochemistry experiments on isogenic HEK lines. C.F. performed triFC 02 experiments on HeLa cells and its corresponding plasmid cloning. U.W. performed the 03 RNA-seq analysis. I.M. and K.M.B. supported the RNA-seq analysis. A.C. performed the 04 purification of TDP-43 RRMs and the NMR experiments. J.W. and Z.G. provided cell 05 culture support. P.D.R. helped with the colocalization studies of nuclear markers and E.T. 06 with the PLA. R.M. supported plasmid cloning and R.R. protein purification and in vitro 07 08 experiments. S.S. and O.S. provided technical feedback. M.H.-P. provided samples of human neural networks, overall cell culture and experimental support and critical input on 09 the study. F.H.-T.A., P.P. and M.P. provided supervision. M.P.-B, V.I.M., A.Z. and M.P. 10

- wrote and edited the manuscript and prepared the figures. M.P. directed the study. All authors read, edited and approved the final manuscript.
- 13 **Conflict of interests:** The authors declare that they have no conflict of interests.

Data and materials availability: All data needed to evaluate the conclusions in the paper are present in the paper and/or the Supplementary Materials. Proteomics data will be deposited to the ProteomeXchange Consortium via the PRIDE partner repository and the transcriptomics data will be deposited to the Gene Expression Omnibus (GEO) genomics data repository.

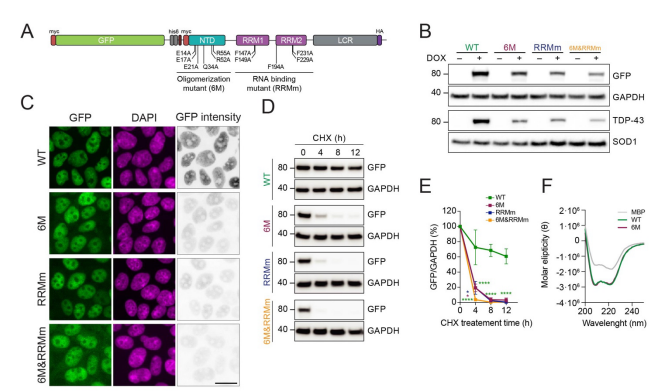


Figure 1. Oligomerization and RNA-binding cooperatively stabilize the half-life of **TDP-43.** (A) Schematic representation depicting the specific mutations used to disrupt oligomerization and/or RNA binding on the GFP-TDP-43 variants, used to develop the inducible, isogenic cell lines. (B) Western blot analysis of the generated isogenic cell lines described in A after inducing GFP-TDP-43 expression for 48 h showing the tightness of the doxycycline (DOX)-modulated expression system. Note also the different protein levels of the expressed variants. (C) Representative images of widefield fluorescence microscopy of the isogenic HEK293 cell lines depicted in B. GFP brightness is adjusted in each condition for optimal visualization of GFP-TDP-43 localization. Original intensity values are represented in the right column using grayscale. Scale bar: 20 µm. (D) GFP-TDP-43 expression was induced with DOX for 24 h before cycloheximide (CHX) treatment for the indicated times and western blot analysis. (E) Quantification of the GFP signal from D. N=3 independent experiments. Two-way ANOVA with Tukey's multiple comparisons post hoc test. (F) Average far-UV CD spectra of purified TDP-43-MBP variants from N=3 independent experiments. **** p<0.0001. Graph bars represent mean \pm SD.

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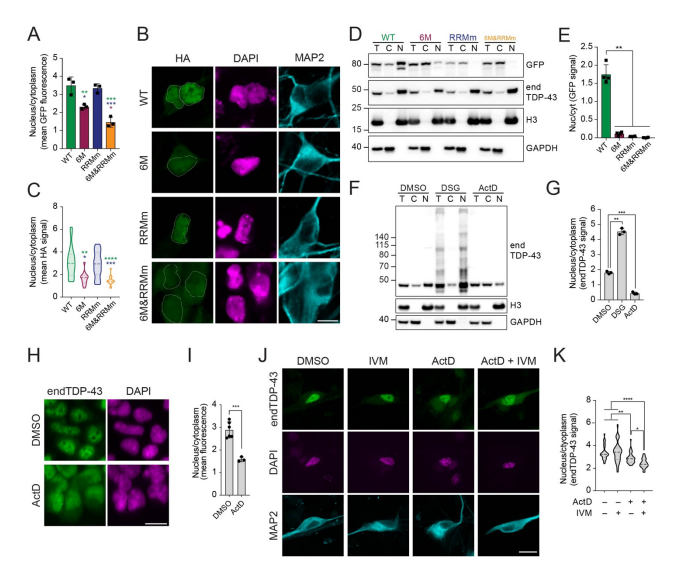


Figure 2. TDP-43 oligomerization and RNA-binding preserve its nuclear localization.

(A) Quantification of nucleocytoplasmic levels of GFP-TDP-43 in the immunocytochemistry images shown in Figure 1C. N=3 independent experiments. One-way ANOVA with Tukey's multiple comparisons post hoc test. (B) Representative image of confocal fluorescence imaging of human neurons transduced with TDP-43-HA variants. Scale bar: 10 µm. (C) Quantification of nucleocytoplasmic levels of TDP-43-HA in the immunocytochemistry images shown in B. N=14-20 cells. Kruskal-Wallis test with Dunn's multiple comparisons post hoc test. (D) GFP-TDP-43 expression was induced with doxycycline (DOX) for 4 h before nucleocytoplasmic fractionation and subsequent analysis of GFP-TDP-43 and endogenous TDP-43 (endDP-43) by western blot. T: total lysate, C: cytoplasmic fraction, N: nuclear fraction. (E) Quantification of the GFP signal from Figure 2D. Repeated measures one-way ANOVA with Greenhouse-Geisser correction and Tukey's multiple comparisons post hoc test. cvt: cvtoplasm, nuc: nucleus. (F) Representative images of widefield fluorescence microscopy of HEK293 cells treated with ActD for 4 h. Scale bar: 20 µm. (G) Quantification of F. N=3 independent experiments. Unpaired two-tailed t-test. (H) HEK293 cells were treated with ActD to inhibit transcription or subjected to protein-protein cross-link with DSG followed by nucleocytoplasmic fractionation and western blot analysis. T: total lysate, C: cytoplasmic fraction, N: nuclear fraction. (I) Quantification of

endTDP-43 signal shown in H. N=3 independent experiments. Repeated measures 41 one-way ANOVA with Greenhouse-Geisser correction and Dunnett's multiple 42 comparisons post hoc test. (J) Representative images of confocal fluorescence 43 microscopy of human neural cultures treated with actinomycin D (ActD) and 44 ivermectin (IVM). Scale bar: 20 µm. (K) Quantification of nucleocytoplasmic levels 45 of endTDP-43 in the immunocytochemistry images shown in J. Kruskal-Wallis test 46 with Dunn's multiple comparisons post hoc test. N=23-48 fields corresponding to a 47 total of 351-569 neurons per condition. * p<0.05, ** p<0.01, *** p<0.001, 48 **** p<0.0001. Graph bars represent mean \pm SD. Violin plots show mean and 49 quartiles. 50

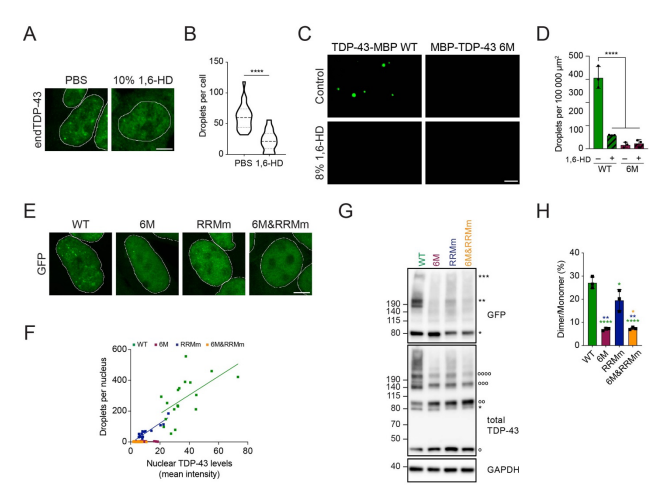


Figure 3. Oligomerization is required for physiological LLPS of TDP-43 in the nucleus.

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(A) Representative maximum intensity Z-projections from confocal fluorescence imaging (thickness of $\sim 10 \,\mu\text{m}$, in steps of 0.21 μm) of WT HEK293 cells after mock or 1,6-HD treatment for 15 min stained for endogenous TDP-43 (endTDP-43). Scale bar: 5 µm. (B) Quantification of three-dimensional (3D) reconstructions from the Zstack confocal microscopy images shown in A depicting the number of nuclear droplets per cell in the conditions described in A. N=14-23 cells. Unpaired twotailed t-test. (C) Fluorescence microscopy images of 10 µM purified MBP-tagged full-length TDP-43 and its oligomerization-deficient counterpart showing different abilities to undergo LLPS and its disruption by 1,6-HD treatment for 10 min. Scale bar: 10 µm. (D) Quantification of the number of droplets in the conditions shown in C per 100 000 µm² field. N=3 independent experiments. One-way ANOVA with Tukey's multiple comparisons post hoc test. (E) Representative maximum intensity Z-projections (thickness of $\sim 10 \,\mu$ m, in steps of 0.21 μ m) from confocal fluorescence microscopy of the isogenic cell lines expressing GFP-TDP-43 for 48 h with doxycycline (DOX). Scale bar: 5 µm. (F) 3D quantification of the number of nuclear droplets per cell shown in E. N=16-22 cells. (G) GFP-TDP-43 expression was induced with DOX for 4 h before crosslinking protein-protein interactions with DSG and subsequent analysis by western blot. *, ** and *** indicate GFP-TDP-43 monomers, dimers and trimers, respectively. °, °°, °°° and °°° indicate endTDP-43 monomers, dimers, trimers and tetramers. (H) Quantification of GFP-TDP-43 dimer/monomer ratio based on the GFP signal from G. N=3 independent experiments. Repeated measures one-way ANOVA with Greenhouse-Geisser correction and Tukey's multiple comparisons post hoc test. * p<0.05, ** p<0.01,

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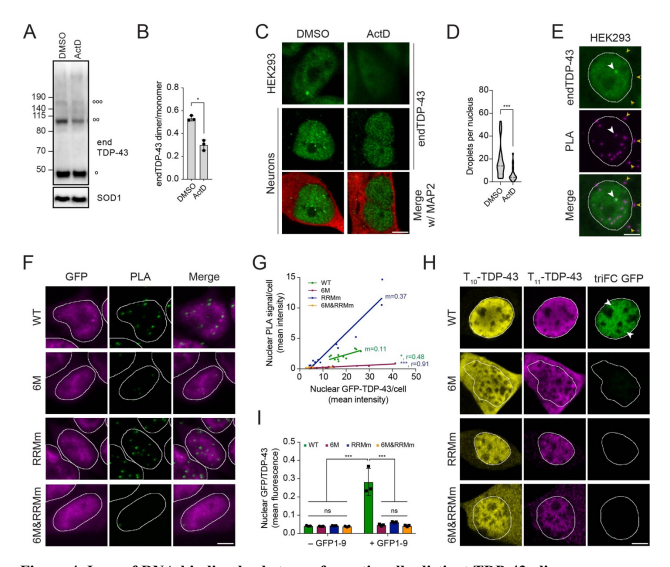


Figure 4. Loss of RNA binding leads to conformationally distinct TDP-43 oligomers. (A) HEK293 cells were treated with ActD for 4 h to inhibit transcription before treatment with the protein-protein cross-linker DSG and western blot analysis. °, °° and ⁰⁰⁰⁰ indicate endogenous TDP-43 (endTDP-43) monomers, dimers and trimers, respectively. (B) Quantification of the endTDP-43 signal from A. N=3 independent experiments. Paired two-tailed t-test. (C) Representative image of confocal fluorescence microscopy of HEK293 cells and neurons treated with ActD. Scale bar: $5 \,\mu m.$ (D) Single-plane quantification of the number of nuclear droplets per neuron in the conditions described in C. N=25-26 cells. Mann-Whitney U test. (E) Proximity ligation assay (PLA) using a monoclonal anti-TDP-43 antibody reveals nuclear and cytoplasmic localization of endTDP-43 dimers in physiological conditions. Big white arrowheads indicate overlapping GFP-TDP-43 droplets and PLA signal. Small yellow arrowheads indicate cytoplasmic PLA signal. Scale bar: 5 µm. (F) PLA using a monoclonal anti-GFP antibody reveals the localization of GFP-TDP-43 dimers in the isogenic cell lines upon protein expression with doxycycline (DOX) for 48 h. Note the absence of dimers in the oligomerizationdeficient variant (6M). Scale bar: 5 µm. (G) Quantification of the nuclear PLA signal shown in G correlated to the protein expression levels of the different TDP-43 variants, measured as the mean GFP fluorescence. N=11-13 cells. (H) Tripartite GFP complementation assay using a pair of N-terminally T₁₀- and T₁₁-tagged TDP-43 constructs co-transfected with GFP_{1-9} in motoneuron-like NSC-34 cells. triFC:

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.00	trimolecular fluorescence complementation. Scale bar: 5 µm (I) Quantification of
.01	the GFP fluorescence levels relative to the T_{10}/T_{11} -TDP-43 expression levels as
.02	shown in H. N=3 replicates, with N=6-35 cells per replicate. Two-way ANOVA
.03	with Tukey's multiple comparisons post hoc test. * p<0.05, *** p<0.001. Graph bars
.04	represent mean \pm SD. Violin plots show mean and quartiles.

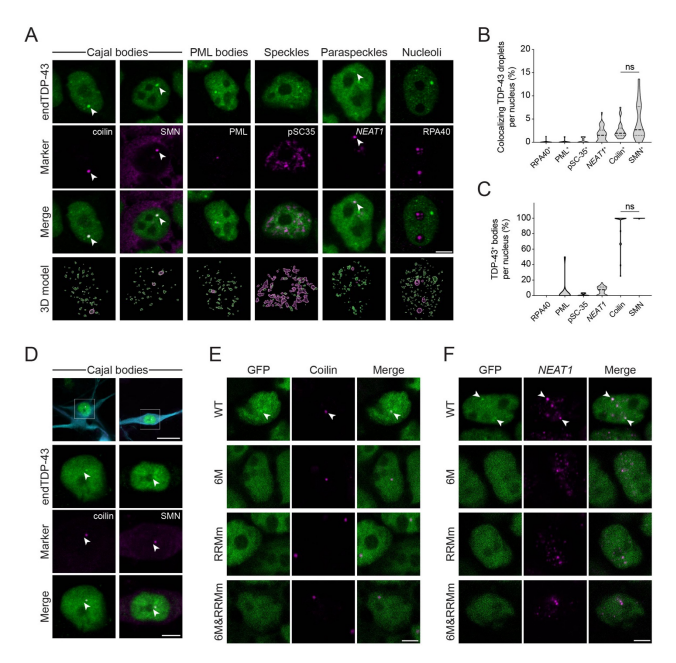


Figure 5. Cajal bodies and paraspeckles are the main TDP-43-containing nuclear bodies. (A) Representative confocal microscopy images of HEK293 cells probed for endogenous TDP-43 (endTDP-43) and different subnuclear compartment markers. Lower panel shows the three-dimensional (3D) reconstruction of the endogenous TDP-43 (endTDP-43) droplets and the indicated nuclear body obtained from the confocal Z-stacks. Scale bar: 5 µm. (B) Quantification of the 3D reconstructions shown in A depicting the percentage of nuclear TDP-43 droplets colocalizing with markers of subnuclear compartments. N=14-38 cells. (C) Quantification of 3D reconstructions shown in A depicting the percentage of each of the analyzed subnuclear compartments that colocalize with endogenous TDP-43. N=14-38 cells. (D) Representative confocal microscopy images of human neurons showing the presence of TDP-43 in Cajal bodies. Scale bar: 20 µm (inset: 5 µm). (E) Representative confocal microscopy images of the isogenic HEK293 lines expressing the different GFP-TDP-43 variants for 24 h and stained for the Cajal body marker coilin. Scale bar: 5 µm. (F) Representative confocal microscopy images of the isogenic HEK293 lines expressing the different GFP-TDP-43 variants

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for 24 h and hybridized with a fluorescent *NEAT1* probe to mark the paraspeckles. Scale bar: $5 \mu m$.

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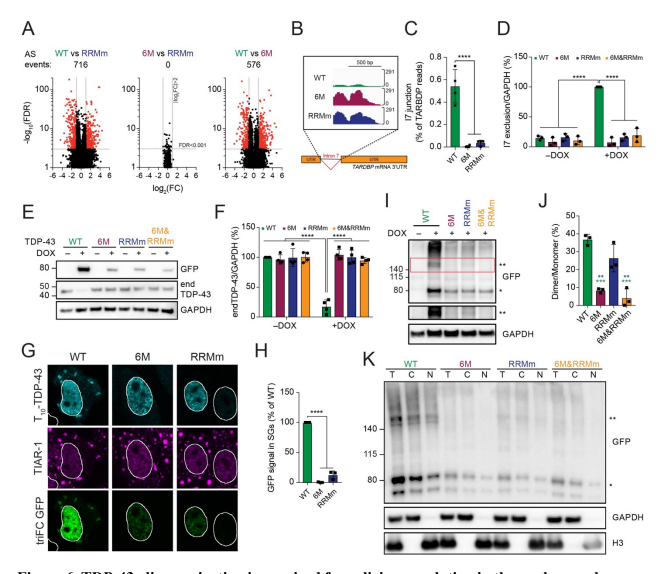
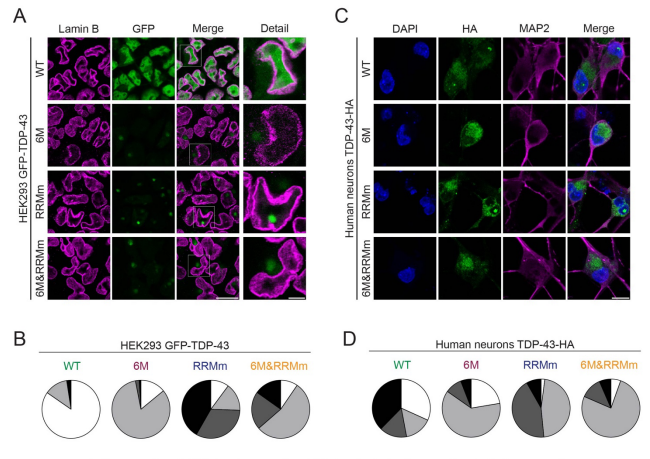


Figure 6. TDP-43 oligomerization is required for splicing regulation in the nucleus and stress granule incorporation in the cytoplasm. (A) Volcano plots showing alternative splicing events upon expression of GFP-TDP-43 variants for 48 h. (B) RNA sequencing (RNA-seq) coverage across the intron 7 of the TARDBP gene, showing a strong decrease in the WT, but not the mutant, GFP-TDP-43-expressing cells. AS events: alternative splicing events. (C) Quantification of the RNA-seq reads spanning the intron 7 junction. One-way ANOVA with Tukey's multiple comparisons post hoc test. (D) Endogenous TDP-43 (endTDP-43) intron 7 exclusion levels after expression of the GFP-TDP-43 variants for 48 h in the isogenic cell lines measured by qPCR with primers specifically targeted to the transcripts excluding this region. Two-way ANOVA with Tukey's multiple comparisons post hoc test. (E) Western blot analysis of the isogenic HEK293 after GFP-TDP-43 expression for 48 h showing that only the WT variant regulates endTDP-43 levels. (F) Quantification of the endTDP-43 signal from E. N=4 independent experiments. Two-way ANOVA with Tukey's multiple comparisons post hoc test. (G) Tripartite GFP complementation assay using a pair of N-terminally T₁₀- and T₁₁-tagged TDP-43 constructs co-transfected with $GFP_{1.9}$ in HeLa cells and subjected to arsenite stress for 30 min. TriFC: trimolecular fluorescence complementation. Scale bar: 10 μm. (H) Quantification of the signal trimolecular fluorescence complementation (triFC) of GFP in the TIA-1-marked SGs from the immunocytochemistry images shown in G. N=3 independent experiments. Repeated measures one-way ANOVA

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.46	with Greenhouse-Geisser correction and Tukey's multiple comparisons post hoc
.47	test. (I) Expression of GDP-TDP-43 mutNLS variants was induced with
.48	doxycycline (DOX) for 4 h before crosslinking protein-protein interactions with
.49	DSG and subsequent analysis by western blot. * and ** indicate GFP-TDP-43
.50	monomers and dimers, respectively. (J) Quantification of the GFP signal from I.
.51	N=3 independent experiments. Repeated measures one-way ANOVA with
.52	Greenhouse-Geisser correction and Tukey's multiple comparisons post hoc test. (K)
.53	After expression of GFP-TDP-43 mutNLS variants for 48 h, HEK293 cells were
.54	treated with DSG to cross-link protein-protein interactions before performing
.55	nucleocytoplasmic fractionation and western blot analysis. * and ** indicate GFP-
.56	TDP-43 monomers and dimers, respectively. ** p<0.01, *** p<0.001,
.57	**** p<0.001. Graph bars represent mean \pm SD.



□ No aggregation □ Cytoplasmic aggregation □ Nuclear & cytoplasmic aggregation □ Nuclear aggregation

- Figure 7. Loss of RNA binding or oligomerization differentially modulate the subcellular localization of TDP-43 inclusions. (A) Representative confocal microscopy images of the isogenic GFP-TDP-43 lines after 48 h of expression, treated with the proteasome inhibitor MG132 for the last 24 h and stained for lamin B to mark the nuclear envelope. Note the different localization of TDP-43 inclusions in the oligomerization- (6M and 6M&RRMm, cytoplasmic) versus RNA bindingdeficient (RRMm, nuclear) variants. Scale bar: 20 µm (5 µm for inset). (B) Ouantification of the differentially localized GFP-TDP-43 inclusions after MG132 treatment for the different variants in the isogenic HEK lines as shown in A. Represented values are averages from N=3 replicates, with N=189-497 cells quantified per condition and replicate. (C) Representative maximum intensity Zprojections from confocal fluorescence imaging (thickness of 4 µm, in steps of 1 µm) of human neurons transduced with TDP-43-HA variants and treated overnight with the proteasome inhibitor MG132. Scale bar: $10 \,\mu\text{m}$. (D) Quantification of the differentially localized TDP-43-HA inclusions in human neurons as described in C. Represented values correspond to the quantification of N=85-97 neurons from two independent experiments.
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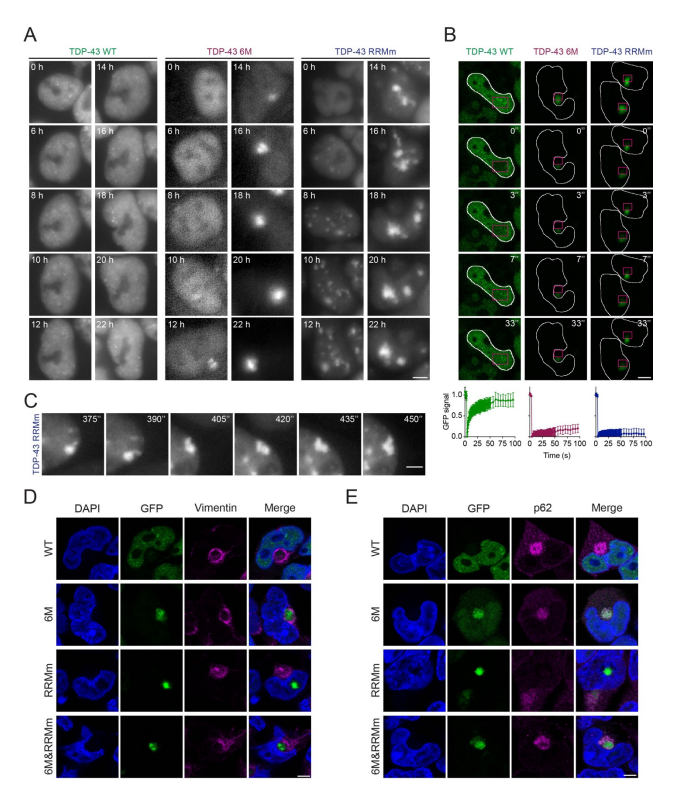


Figure 8. TDP-43 aggregates in an LLPS- or an aggresome-dependent manner in the nucleus and cytoplasm, respectively. (A) Representative images of live widefield fluorescence microscopy over the course of the MG132 treatment of the isogenic GFP-TDP-43 lines in the conditions described in Figure 7A. Numbers in images indicate the experimental time point in hours (h) of MG132 treatment. Scale bar: 5 μm. (B) Representative fluorescent confocal microscopy images of FRAP experiments in the GFP-TDP-43 aggregates originated upon MG132 treatment as described in Figure 7A. FRAP was performed in the areas highlighted in magenta.

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Bottom panel: Measured GFP values are expressed as a fraction of the average prebleach fluorescence levels. Scale bar: 5 μ m. (C) Live widefield fluorescence microscopy depicting a fusion event and aberrant phase transition of RRMm GFP-TDP-43 droplets upon MG132 treatment in the conditions described in Figure 7A. Scale bar: 3 μ m. (D) Representative confocal microscopy images of the isogenic GFP-TDP-43 lines in the conditions described in figure 7A and stained for vimentin. Scale bar: 5 μ m. (E) Representative confocal microscopy images of the isogenic GFP-TDP-43 lines in the conditions described in Figure 7A and stained for p62. Scale bar: 5 μ m.

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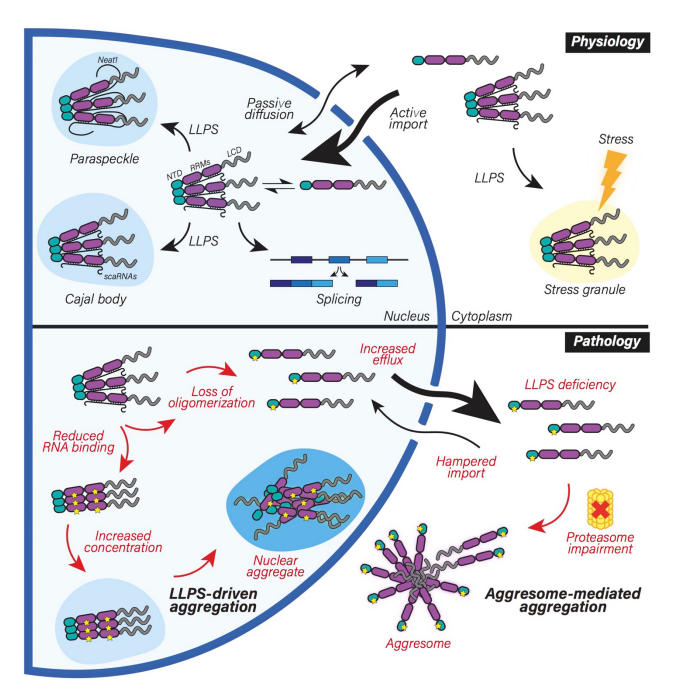


Figure 9. Oligomerization and RNA-binding enable TDP-43 physiological functions and their disruption drives nuclear and cytoplasmic aggregate formation via distinct pathways. Schematic representation of the role of NTD-driven oligomerization and RNA binding in TDP-43 physiology and pathology. Upper panel: In healthy cells, TDP-43 monomers and oligomers are in a dynamic equilibrium. TDP-43 is actively imported into the nucleus (12), where oligomerization and RNA binding retain it in large macromolecular complexes, limiting passive outflow. In the nucleus, oligomerization and RNA binding are key for the essential roles of TDP-43 in RNA processing, including alternative splicing. Furthermore, oligomerization enables the LLPS of TDP-43 and in conjunction with binding to specific RNAs –such as small Cajal body-specific RNAs (scaRNAs) (56) and *NEAT1* (20, 21)– allows its localization to distinct subnuclear compartments, primarily Cajal bodies and paraspeckles. TDP-43 oligomers are also detected in the cytoplasm, where its LLPS-mediated incorporation into SGs under stress conditions

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depends on both oligomerization and RNA binding. Lower panel: In disease, loss of TDP-43 oligomerization or RNA binding increases the nuclear efflux of LLPSdeficient monomers, disrupts its inclusion into nuclear bodies, leads to transcriptome-wide splicing alterations (including TDP-43 auto-regulation) and drives TDP-43 aggregation via two independent pathways. Upon failure of the ubiquitin-proteasome degradation machinery observed with aging and in ALS/FTLD patients (2, 3, 43), monomeric TDP-43 aggregates in an aggresomedependent manner in the cytoplasm. Notably, the known decline in active nuclear import in disease (83) would further exacerbate TDP-43 cytoplasmic accumulation. In the nucleus, loss of TDP-43 RNA-binding results in enhanced formation of both monomers that escape to the cytoplasm and conformationally distinct TDP-43 oligomers. When the concentration rises (e.g. due to the aforementioned proteasomal failure), RNA-binding deficiency modulates TDP-43 LLPS, culminating in the formation of nuclear aggregates via an LLPS-mediated pathway. Taken together, RNA binding and oligomerization allow TDP-43 to maintain its localization and function in physiology and their disruption drives LLPS-dependent and aggresome-dependent aggregation pathways in the nucleus and cytoplasm, respectively. LCD: low complexity domain, LLPS: liquid-liquid phase separation, NTD: N-terminal domain, RRMs: RNA recognition motifs.

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