Cytoplasmic TDP-43 De-mixing Independent of Stress Granules Drives Inhibition of Nuclear Import, Loss of Nuclear TDP-43, and Cell Death

Highlights

- Transient stress induces long-lasting phase separation of cytoplasmic TDP-43
- Formation/maintenance of phase separated TDP-43 is independent of stress granules
- Phase-separated TDP-43 inhibits nuclear transport by de-mixing importin-α and Nup62
- Cytoplasmic TDP-43 de-mixing depletes nuclear TDP-43 and induces cell death

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In Brief
TDP-43 aggregation is the major hallmark of multiple neurodegenerative diseases, including ALS and FTD. Gasset-Rosa et al. demonstrate that transient stress induces long-lasting cytoplasmic TDP-43 de-mixing independent of stress granules, driving nuclear import defects, nuclear TDP-43 clearance, and cell death.
Cytoplasmic TDP-43 De-mixing Independent of Stress Granules Drives Inhibition of Nuclear Import, Loss of Nuclear TDP-43, and Cell Death

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SUMMARY

While cytoplasmic aggregation of TDP-43 is a pathological hallmark of amyotrophic lateral sclerosis and frontotemporal dementia, how aggregates form and what drives its nuclear clearance have not been determined. Here we show that TDP-43 at its endogenous level undergoes liquid-liquid phase separation (LLPS) within nuclei in multiple cell types. Increased concentration of TDP-43 in the cytoplasm or transient exposure to sonicated amyloid-like fibrils is shown to provoke long-lived liquid droplets of cytosolic TDP-43 whose assembly and maintenance are independent of conventional stress granules. Cytosolic liquid droplets of TDP-43 accumulate phosphorylated TDP-43 and rapidly convert into gels/solids in response to transient, arsenite-mediated stress. Cytoplasmic TDP-43 droplets slowly recruit importin-α and Nup62 and induce mislocalization of RanGap1, Ran, and Nup107, thereby provoking inhibition of nucleocytoplasmic transport, clearance of nuclear TDP-43, and cell death. These findings identify a neuronal cell death mechanism that can be initiated by transient-stress-induced cytosolic demixing of TDP-43.

INTRODUCTION

Mislocalization, self-assembly, and aggregation of misfolded TAR DNA-binding protein 43 (TDP-43) in the cytoplasm of affected motor neurons is a common neuropathological hallmark of almost all cases of amyotrophic lateral sclerosis (ALS) (Neumann et al., 2006). Proteinaceous inclusions, containing misfolded aggregated proteins or fragments of them, are also found in each of the major neurodegenerative disorders, including Alzheimer’s (AD), Parkinson’s (PD), frontotemporal dementia (FTD), and Huntington’s (HD) diseases (Chiti and Dobson, 2006). Many of the aggregated proteins contain intrinsically disordered protein domains that are enriched in, or composed of, only a few amino acids and are referred to as low complexity (LC) domains. These domains display a sequence-intrinsic conformational heterogeneity (i.e., disorder) characteristic of intrinsically disordered proteins and/or regions (Boeynaems et al., 2018). LC domains are also present in yeast prion proteins, which have the ability to interconvert into amyloid fibers (King et al., 2012). Prion-like LC domains are particularly abundant in RNA- and DNA-binding proteins, and their amino acid composition has been conserved across evolution (King et al., 2012; Malinovska et al., 2013).

TDP-43 is an RNA-binding protein that localizes predominantly in the nucleus and is thought to shuttle between the cytoplasm and nucleus (Ayala et al., 2008). It forms abnormal cytoplasmic aggregates (Neumann et al., 2006) in neurons and glia in more than 90% of ALS and 45% of FTD cases. These two progressive neurodegenerative diseases, which share genetic and pathological features (Ling et al., 2013), are without effective treatments to slow fatal disease progression (Taylor et al., 2016). Discovery of missense mutations in TDP-43 in patients with ALS or FTD (Rutherford et al., 2008; Sreedharan et al., 2008) demonstrated a direct link between genetic variants and TDP-43 pathology. Many mechanisms have been proposed to explain the abnormal cytosolic accumulation of TDP-43 and the progressive spreading of TDP-43 pathology.

TDP-43 contains a prion-like, LC domain that is glycine, glutamine, and asparagine rich and is predominantly an intrinsically disordered region (IDR) (Conicella et al., 2016), which renders TDP-43 intrinsically aggregation prone (Johnson et al., 2009). Disordered domains of RNA-binding proteins can drive dynamic self-assembly into intracellular membraneless organelles, including P granules (paranuclear granules in germ cells of C. elegans), nucleoli, and stress granules, each of which has been reported to fuse, minimize surface tension, and dynamically exchange components with the solution, all properties indicative of liquid-like behavior (Brangwynne et al., 2009; Mitrea and Krilwacki, 2016). Correspondingly, phase separation might be the operational principle governing the formation of membraneless organelles. A key unresolved issue is whether (and, if so, how) assembly into membraneless organelles...
A  Endogenous TDP-43

- Mouse cortex
- U2OS
- SH-SY5Y
- Hippocampal neuron

TDP-43 C-terminal Ab
TDP-43 N-terminal Ab

B  GFP or mRuby2 tagged TDP-43

- U2OS
- SH-SY5Y

TDP-43mRuby2
Inducibly expressed C- or N-terminal tagged TDP-43WT
C-terminally tagged at both endogenous alleles

C  SH-SY5Y:
Induced EYFP TDP-43

D  Droplet intensity (%)

Relative intensity (%)

E  SH-SY5Y:
Induced EYFP TDP-43

F  Photo-bleaching

Relative intensity (%)

n=5

G  Half droplet bleaching

H  Nuclear TDP-43 droplet fusion

I  U2OS: TDP-43mRuby2

J  Control (n=4)

Unbleached (n=5)

Bleached (n=6)

(legend on next page)
regulates or affects biological function(s) of their constituent proteins. Beyond that, phase-separated droplets have increasingly been implicated as crucibles for nucleation of pathologic protein aggregation since protein concentration is expected to be sharply increased within the droplets (Li et al., 2013; Ramaswami et al., 2013).

Prion-like LC-domain-containing heterogeneous nuclear ribonucleoproteins (hnRNPs), such as FUS, hnRNPA1, and TIA1 (other proteins whose mutation is causative of ALS/FTD), exhibit liquid-liquid phase separation (LLPS) as purified proteins in vitro (Burke et al., 2015; Mackenzie et al., 2017; Mateju et al., 2017; Molièx et al., 2015; Patel et al., 2015). TDP-43 has been reported to display aspects (round-shaped morphology or fusion events) of liquid-phase separation in vitro (Conicella et al., 2016; Ryan et al., 2018; Wang et al., 2018). Prolonged LLPS of purified FUS or repeated cycles of temperature-dependent de-mixing of mutant hnRNPA1 can induce conversion to a solid phase (Molièx et al., 2015; Patel et al., 2015), while expression of FUS variants with decreased ability to bind RNA can form solid-like aggregates in a cancer cell line (Maharana et al., 2018). The relevance of this altered phase behavior is not established in disease; however, as in every reported instance, the presence of other proteins or post-translationally modified variants inhibits liquid-phase separation in vitro (Guo et al., 2018; Hofweber et al., 2018; Qamar et al., 2018; Yoshizawa et al., 2018). Only a handful of observations confirm LLPS properties of these RNPs in living cells, and in most examples, de-mixing requires extreme conditions, including transient overexpression or degradation of total cellular RNA or both (Gopal et al., 2017; Maharana et al., 2018; Wang et al., 2018).

No effort has succeeded in identifying whether TDP-43 undergoes liquid-liquid de-mixing in the cytoplasm—where pathological aggregates accumulate—and, if so, whether such de-mixing can trigger nuclear loss of TDP-43, conversion of TDP-43 to an aggregated state, or both. Here we identify intranuclear LLPS of native TDP-43 in multiple mammalian cells in cultured mouse cortical neurons and human iPSC-derived motor neurons. Arsenite-mediated stress is shown to induce TDP-43 de-mixing into cytoplasmic liquid droplets that are independent of stress granule components or formation and that rapidly convert to gels/solids that recruit phosphorylated TDP-43. Transient stress from exposure to fragmented amyloid-like fibrils induces endogenous TDP-43 de-mixing into cytosolic droplets that (1) form and persist for up to 1 month, (2) are independent of conventional stress granules, (3) accumulate phosphorylated TDP-43, (4) compromise nucleo-cytoplasmic transport, (5) slowly deplete nuclear TDP-43, and (6) ultimately elicit cell death.

RESULTS

Nuclear TDP-43 De-mixes under Physiological Conditions

While use of immunofluorescence has established that TDP-43 is primarily intranuclear, high-resolution inspection (using N- or C-terminal antibodies) of endogenous TDP-43 revealed that it is not distributed diffusely in nucleoplasm but rather forms rounded particles in both mouse and human cells, including cortical neurons from wild-type mouse brain, primary cultured mouse hippocampal neurons, human SH-SY5Y neuronal-like cells, U2OS, a human non-neuronal cell line, and motor neurons induced from human pluripotent stem cells (iPSCs) (Figure 1A). An apparently de-mixed distribution was confirmed by imaging directly fluorescently tagged TDP-43 expressed at levels equal to or below the normal endogenous level (Figure S1). Accumulation of (1) an amino-terminally EYFP-tagged TDP-43 (EYFP-TDP-43) to about one-third the level of endogenous TDP-43 (Figures S1A and S1B) or (2) a carboxy-terminally mRuby2-tagged TDP-43 (TDP-43mRuby2) replacing endogenous TDP-43 (Figure S1C) yielded between 5 and 50 rounded, droplet-like particles of 0.2–1 μm diameter in each nucleus (Figure 1B). Similar rounded particles were found (Figure 1B) in SH-SY5Y cells that had been genome edited at both TDP-43 alleles to encode fluorescently tagged TDP-43 (TDP-43mEGFP), a variant that confers the essential functions of TDP-43 (Figures S1D and S1E).

Live cell imaging was used to determine that these intranuclear TDP-43 particles have liquid-like properties. Fusion events...
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could be observed frequently within a 10 s period of monitoring in neuron-like SH-SY5Y cells (see, for example, the fusion of two EYFP-TDP-43 droplets labeled with green or red arrowheads in Figures 1G and 1D). Fluorescence recovery after photobleaching of whole EYFP-TDP-43-containing droplets in SH-SY5Y revealed rapid recovery (to ~70% of original intensity) in <10 s (Figures 1E and 1F), indicating fast molecular exchange of TDP-43 molecules between droplets and nucleoplasm. Internal molecular dynamics were five times faster, with fluorescence intensity re-equilibrated within 2 s following half-droplet bleaching (Figure 1G). Fusion events were also documented for TDP-43mRuby2 and TDP43EGFP-containing droplets in U2OS or SH-SY5Y cells (Figures 1H and S1F, respectively). Fusion events were also observed (Figure 1H, right panel), albeit at much lower frequency. As expected for LLPS, after photobleaching half of a nucleus, fluorescence intensity of TDP-43 recovered in the bleached half within 140 s, with a corresponding decrease in intensity in the unbleached half (Figures 1I and 1J). Thus, TDP-43 phase separates in normal physiological conditions into intranuclear liquid droplets in which TDP-43 molecules dynamically move into and out of the droplets.

Concentration-Dependent TDP-43 De-Mixing in the Cytoplasm

To mimic the increased accumulation of cytoplasmic TDP-43 during aging expected from the known age-dependent loss of nuclear pore components (D’Angelo et al., 2009), we expressed fluorescently tagged TDP-43 with a disrupted NLS (TDP-43NLs-Clover) using a doxycycline-inducible system in human U2OS cells that also stably express TDP-43mRuby2 (Figure 2A) at the endogenous TDP-43 level (Figure S1C). Induction of TDP-43NLs-Clover produced a range of accumulation of cytoplasmic TDP-43 in individual cells, with Clover (and mRuby2) fluorescence providing a direct readout of TDP-43 level (Figures 2B and 2C). Increased accumulation of cytoplasmic TDP-43 was sufficient to induce its de-mixing into 1–3 μm spherical, cytoplasmic particles (Figure 2B) that (1) over time recruited the initially nuclear TDP-43mRuby2 (Figure 2B) and (2) underwent fusions and fissions (Figure 2D). TDP-43 cytoplasmic droplets remained liquid over extended periods, with rapid and efficient (70%) recovery (within 2 or 4 min, respectively) after complete or partial photobleaching (Figures 2E and 2F). Thus, an increased level of cytoplasmic TDP-43 is sufficient to induce its de-mixing in the cytoplasm.

TDP-43 Cytoplasmic De-Mixing into Liquid Droplets

To determine whether nuclear TDP-43 could redistribute to the cytoplasm and de-mix, we genetically edited both TDP-43 alleles in SH-SY5Y cells to express physiological levels of wild-type TDP-43 fused to a green fluorescent protein (TDP-43EGFP) (Figures S1D and S1E). Non-dividing neuronal-like (SH-SY5Y) cells were then transiently exposed to fragmented amyloid-like-containing fibrils. mCherry-tagged recombinant TDP-43 (or FUS) was purified (Figure S2A) and incubated for 24 h at 20 °C. Both were self-assembled into fibrils whose appearance (Figures 2H and S2B) was highly similar to amyloid-like fibrils (Rambanan and Serpell, 2008). After sonication, the fragmented fibrils (~20 nm in size) were added to the culture media of SH-SY5Y cells expressing TDP-43EGFP from both endogenous TDP-43 alleles (Figures 2G and 2H and S2B). As expected, without fibril addition, TDP-43EGFP remained nuclear without cytoplasmic liquid-like structures in non-cycling cells arrested in the G1 cell-cycle phase (Figure S2C).

After transient addition of TDP-43mCherry sonicated fibrils (Figures 2G and 2H), time-lapse confocal imaging was used to (1) determine that fibril fragments were internalized within 12 h, (2) follow the fate of the fibrils, and (3) determine localization of TDP-43EGFP expressed at endogenous levels. At early times, fragmented TDP-43 fibrils accumulated in the cytoplasm but did not yield detectable TDP-43EGFP recruitment (Figure 2I).
Sonicated particles

His-TDP-43 or HA-FUS

1-3 days

Endogenous TDP-43 WT

1 day, 1 week, 1 month

Immunostaining (B, E, F, I)

HA-FUS/His-TDP-43 (E)

TDP-43

1 day, 1 week NaAsO₂ (60 min)

TIA 1 and G3BP1 (H, G)

TDP-43

A

Non-dividing Neuronal-like SH-SY5Y cells

B

Untreated

C

His-TDP-43

His-TDP-43

TDP-43

D

HA-FUS

HA-FUS

E

Exogenous His-TDP-43WT particles

Exogenous HA-FUSWT particles

F

% of cells with cytoplasmic TDP-43 LLPS

Number of TDP-43 LLPS per cell

1 week

1 month

1 week

1 month

1 day

1 week

1 month

1 day

1 week

1 month

Exogenous HA-FUSWT particles + Sodium arsenite treatment

G

Exogenous HA-FUSWT particles

TIA 1

TDP-43

TIA 1

TDP-43

TIA 1

TIA 1

1 day

1 week

1 day

1 week

1 day

1 week

1 day

1 week

1 day

1 week

(legend on next page)
However, within 10 days, TDP-43EGFP mislocalized to round, apparently de-mixed cytoplasmic droplets (Figure S2D), and by 1 month, most remaining cells accumulated TDP-43EGFP cytoplasmic particles accompanied by depletion of nuclear TDP-43 (Figure 2I). Cytoplasmic TDP-43 particles were confirmed by immunostaining with TDP-43 antibody (Figure S2E). A parallel experiment with addition of FUSmCherry particles (Figure S2B) yielded almost identical redistribution and apparent de-mixing of TDP-43EGFP expressed at endogenous levels (Figure 2J).

Recovery dynamics after photobleaching were used to determine that 1 month after their induction TDP-43 cytoplasmic particles represented LLPS, with rapid exchange with soluble TDP-43 in the cytoplasm driving fluorescence recovery within 25 s of full or focal photobleaching of single particles (Figures 2K and 2L). Moreover, these old cytoplasmic droplets containing TDP-43 underwent frequent fusion (two examples are shown in Figure 2M) and rarer fission events (Figure S2F), indicative of LLPS. Younger droplets (accumulated 10 days after fibril exposure) displayed ~2.5-fold faster recovery.

Cytoplasmic TDP-43 Liquid De-mixing Independent of Stress Granules

To determine whether endogenous untagged, wild-type TDP-43 can be induced to de-mix in the cytoplasm, we exposed non-dividing neuronal-like SH-SY5Y cells to fragmented TDP-43 or FUS particles (Figures 3A–3D and S3A). Within 24 h, a proportion of initially nuclear TDP-43 (Figure 3E) accumulated in spherical, cytoplasmic particles, many of which (see white arrows, top row, Figure 3E) had undetectable levels of the fragmented TDP-43 or FUS fibrils, respectively. Indeed, no fibril fragments of TDP-43 or FUS could be detected within 4 days after their removal from the media, although in the majority (60%) of cells, a proportion of endogenous TDP-43 remained in a series of round (~1 μm diameter) cytoplasmic particles (Figures 3E and 3F). This apparent de-mixing of TDP-43 was still present a month after fibril exposure, more than 3 weeks after fibrils could no longer be detected (Figure 3E), with the number of apparent droplets increasing 3-fold in a time-dependent manner (Figure 3F). In all cases and time points, TDP-43 remained nuclear and without cytoplasmic particles in untreated cells (Figure 3B) or cells treated with sonicated fragments of fibrils assembled from superoxide dismutase (SOD1) (Figures S3B and S3C).

The cytoplasmic, de-mixed TDP-43 droplets induced by transient exposure to TDP-43 or FUS particles were very different from conventional stress granules, which assemble in response to heat or oxidative stresses and have been proposed to be the crucibles for pathological inclusion formation (Li et al., 2013). With very rare exceptions, no classical stress granule marker (TIA1 and G3BP1) could be detected in fibril-induced TDP-43-containing droplets (Figures 3G and 3H). Further, when stress granules were induced by exposure of the cells to sodium arsenite (Figure S3D), little, if any, endogenously expressed TDP-43 was recruited to the resultant stress granules containing TIA1 or G3BP1 (Figure 3H). Similarly, no stress granule component (e.g., TIA1, Figures S3E and S3F) could be detected in de-mixed, fluorescently tagged TDP-43 (TDP-43EGFP) produced from the endogenous TDP-43 alleles even after cell exposure to arsenite (Figure S3F). Moreover, by 1 month, there was obvious nuclear clearing of TDP-43 in a proportion (~5%) of cells with fibril-induced cytoplasmic TDP-43 particles (Figure 3I), indicating that an external amyloid-like particle-induced re-localization of wild-type endogenous TDP-43 to spherical, cytoplasmic particles can drive loss of nuclear TDP-43 function.

We next followed the kinetics of TDP-43 de-mixing in response to a high (250 μM) dose of sodium arsenite. Conventional stress granules marked by UBAP2L (Figures 4A and 4B) or G3BP1 (Figures 4A and 4D) formed within 20 min and recruited a proportion of an initially diffusely localized cytoplasmic TDP-43SNLS-Clover (Figure 4B). By 40 min, most cells accumulated stress granules into which a small proportion of TDP-43 was recruited (Figures 4B and 4C). However, by 50 min, new rounded, cytoplasmic assemblies of TDP-43SNLS-Clover that did not contain UBAP2L began forming (see arrowheads, Figure 4B). By 90 min, most cytoplasmic TDP-43 (including the proportion initially recruited to stress granules) had redistributed into the new cytoplasmic droplets (Figure 4B) in ~90% of the cells (Figure 4C) despite continued presence of arsenite and UBAP2L- or G3BP1-containing stress granules (Figures 4B and 4D). These findings are consistent with observations of recruitment of a small proportion of full-length TDP-43 to stress granules and that this can be enhanced by poly(ADP-ribose) binding to the TDP-43 nuclear-localization sequence (McGurk et al., 2018).

Figure 3. Transient Stress Induces Long-Lasting Untagged, Cytosolic TDP-43 Particles Independent of Conventional Stress Granules

(A) Schematic of experimental design to assess endogenous wild-type TDP-43 de-mixing in the cytoplasm of neuronal-like SH-SY5Y cells (non-cycling).
(B) Representative images of endogenous nuclear TDP-43 in non-fibril-treated SH-SY5Y cells at day 1 (left) and after 1 month of culture (right).
(C and D) Electron micrographs of amyloid-like fibrils of full-length wild-type His-TDP-43 (C) or HA-FUS (D) purified from bacteria. Respective right panels illustrate the His-TDP-43 or HA-FUS fibrils after sonication before inoculating them into cell media.
(E) Representative images of neuronal-like SH-SY5Y cells after 1 day, 1 week, or 1 month after inoculated sonicated His-TDP-43 (left) or HA-FUS (right) fibrils at a final concentration of 200 nM and immunostained with His- or HA-tag (red) and TDP-43 (green) antibodies. Media were changed after 3 days. White arrows indicate cytoplasmic particles containing endogenous wild-type TDP-43 (green).
(F) Left: quantification of the number of cells with cytoplasmic de-mixed TDP-43 over time. Right: quantification of cytoplasmic particles per cell with time (represented as mean ± SEM from 200 cells in a total of three independent experiments).
(G and H) Representative images of SH-SY5Y cells incubated with sonicated HA-FUS particles (G) and immunostained after 1 day (top panel) or 1 week (bottom panel) with stress granule marker TIA1 (red) and TDP-43 (green) antibodies or (H) after treatment with NaAsO2 (0.5 mM) for 1 h and immunostained with TIA1 or G3BP1 (red) and TDP-43 (green) antibodies.
(I) Representative images of SH-SY5Y cells with apparent cytoplasmic TDP-43 de-mixing accompanied by depletion of nuclear TDP-43 1 month post treatment with sonicated HA-FUS particles. Higher magnification of the white boxed area in the left panel (right panel). The white dashed line outlines the nucleus.
Add doxycycline to induce cytoplasmic TDP-43 (TDP-43ΔNLS-Clover)

Live cell imaging (B, F)
or
Immunostaining for G3BP (D, E)

Determine molecular dynamics with fluorescence recovery after photo-bleaching (FRAP) (G-J, 5A-E)

B

<table>
<thead>
<tr>
<th>0 min</th>
<th>20 min</th>
<th>40 min</th>
<th>50 min</th>
<th>60 min</th>
<th>90 min</th>
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<tr>
<td>TDP-43ΔNLS-Clover + NaAsO2 (250 μM)</td>
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D

NaAsO2 (250 μM) 80 min

TDP-43ΔNLS-Clover G3BP1 MERGE

E

Pre-treated with cycloheximide

TDP-43ΔNLS-Clover G3BP1 MERGE

C

With TDP-43 assemblies independent of stress granules
With TDP-43 localized in stress granules

F

+ low dose NaAsO2 (50 μM)

0 hr 1 hr 2 hr 3 hr 3.6 hr

G

Merge

TDP-43 in stress granules

H

Relative intensity (%)

TDP-43ΔNLS-Clover

UBAP2LmRuby2

I

Initial formation of stress-granule independent TDP-43 assemblies

J

Relative intensity (%)

Full bleach

(legend on next page)
Untagged cytoplasmic TDP-43^SNLS showed the same apparent arsenite-induced de-mixing behavior without recruitment of stress granule components (Figures S4A and S4B). Furthermore, application of cycloheximide before arsenite treatment abolished stress granule formation (as expected) but had no effect on the timing or extent of TDP-43 assembly into rounded cytoplasmic droplets (Figure 4E). Addition of arsenite to a level too low to induce stress granules still provoked droplets of TDP-43^SNLS-NLS-Clover (Figure 4F), consistent with extended low-dose arsenite treatment producing cytoplasmic TDP-43-containing particles that are independent of stress granules and their components (McGurk et al., 2018).

Photobleaching was used to test molecular exchange within the TDP-43 spherical particles that do or do not contain stress granule markers and that formed in the cytoplasm in response to arsenite stress (Figures 4G and 4H). Like UBAP2L itself, about half of the TDP-43 initially recruited into rounded, UBAP2L-positive stress granules recovered rapidly after photobleaching (Figures 4G and 4H), with kinetics similar to the previously reported dynamics of components of conventional stress granules (Mollie et al., 2015). Consistent with droplets formed by LLPS, similarly rapid recovery after photobleaching was seen for about half of the TDP-43 recruited into the rounded TDP-43 assemblies that formed 50 min after exposure to arsenite and that did not contain stress granule components (Figures 4I and 4J).

Arsenite-Induced Gelling of De-mixed Cytoplasmic TDP-43 Droplets

While arsenite-induced stress granules (marked by UBAP2L) maintained liquid properties (Figures 5A and 5B) throughout prolonged arsenite exposure, the initially liquid-like cytoplasmic TDP-43^SNLS-NLS-Clover droplets that formed without stress granule components (Figures 4I and 4J) converted within an additional 30 min of arsenite exposure into gels/solids, with almost no intraparticle molecular exchange following partial or complete photobleaching (Figures 5A–5D). The particles did not contain detectable mRNA (Figure S5C), EDC4-containing P body components (Figure S5D), or p62 (Figure S5E) but did acquire TDP-43 phosphorylated at serines 409 and 410 (Figure S5B), as is found in human ALS/FTD pathology. Consistently, the proportion of insoluble TDP-43 increased with time of exposure to arsenite (Figure S5A). Once converted, the particles remained gel/solid-like even after removal of arsenite (Figures S5E and S5F) and recruited p62 with time (Figure S5F).

TDP-43^EGFP droplets induced by transient exposure to sonicated fibrils of TDP-43 or FUS remained liquid for at least 3 weeks after initial droplet formation (Figures 2K–2M) without acquisition of amyloid oligomers detectable with the widely used A11 antibody (Figure S5G, left panel) that recognizes a peptide backbone epitope common to amyloid oligomers (Kayed et al., 2007) and also recognizes cytoplasmic TDP-43-containing “myogranules” in regenerating muscle cells (Vogler et al., 2018). However, with arsenite, even after 30 min of exposure to sodium arsenite, the TDP-43^EGFP droplets quickly converted into a gelled state (Figures 5G and 5H), remaining rounded but with almost no recruitment of new molecules from the cytoplasm after full-particle photobleaching (Figure 5H). Furthermore, 80% of cytoplasmic gel/solid-like TDP-43 particles acquired some amyloid-like character measured with the A11 antibody that recognizes a pre-amyloid oligomeric conformation (Figure S5G, right lower panel). In contrast, almost all of the cytoplasmic TDP-43 de-mixed particles that persisted over 1 month without exposure to arsenite did not acquire detectable amyloid-oligomers (Figure S5G, right upper panel). Hence, arsenite mediates a structural conversion within TDP-43 droplets from weak self-interactions to gels/solids containing β sheet structure, whose incorporation is likely to slow dynamics of exchange with soluble TDP-43 and facilitate TDP-43 aggregation.

**TDP-43 De-mixing in iPSC-Derived Motor Neurons Independent of Stress Granules**

We next tested whether an increased level of cytoplasmic TDP-43 was sufficient to drive its de-mixing within human motor neurons (Figure 4F). Representation of experimental design to assess the relationship of cytoplasmic TDP-43 with stress granules (TDP-43^SNLS-NLS-Clover) in the presence of sodium arsenite.

(A) Schematic of experimental design to assess the relationship of cytoplasmic TDP-43 with stress granules (TDP-43^SNLS-NLS-Clover) in the presence of sodium arsenite.

(B) Representative images of U2OS cells co-expressing stress granule protein UBAP2L^mRuby2 and cytoplasmic TDP-43 (TDP-43^SNLS-NLS-Clover) after addition of 250 μM NaAsO2. White arrowheads indicate round TDP-43 particles independent of stress granules after 50 min of NaAsO2 treatment.

(C) Quantification of TDP-43 recruitment to stress granules and cells that show formation of stress-granule-independent TDP-43 assemblies (from five independent frames in a live-imaging experiment, total cell number from each frame = 25, 12, 30, 16, 10, respectively). Data are shown as box whisker plot (showing mean, min, and max values). ***p < 0.001, pared t test.

(D) Endogenous stress granules (using G3BP1 antibody; red) in cells expressing cytoplasmic TDP-43 (TDP-43^SNLS-NLS-Clover; green) after treatment with NaAsO2 for 80 min.

(E) Endogenous G3BP1 (red) in cells expressing TDP-43^SNLS-NLS-Clover (green) after pre-treatment with cycloheximide (10 μg/mL) followed by NaAsO2 treatment for 80 min.

(F) Representative images of U2OS cells co-expressing stress granule protein UBAP2L^mRuby2 and cytoplasmic TDP-43 (TDP-43^SNLS-NLS-Clover) after addition of 50 μM NaAsO2.

(G) FRAP of TDP-43^SNLS-NLS-Clover (green) and UBAP2L^mRuby2 (red) in a stress granule.

(H) Mean fluorescence intensity of TDP-43^SNLS-NLS-Clover and UBAP2L^mRuby2 plotted over time (normalized to average intensity of a droplet before photobleaching and represented as mean ± SEM from the recovery curves of four granules from two independent FRAP experiments).

(I) FRAP example of an initial stage formation of a cytoplasmic TDP-43^SNLS-NLS-Clover (green) droplet independent of the stress granules UBAP2L^mRuby2 (red).

(J) Mean fluorescence intensity of TDP-43^SNLS-NLS-Clover droplet (indicated by white arrowhead in I) plotted over time (lower graph). Data are normalized to the average intensity of a droplet before photobleaching and are represented as mean ± SEM from the recovery curves of four droplets in a total of three independent experiments.

See also Figure S4.
TDP-43 completely separated from stress granules (80 min after NaAsO₂ treatment)

A

UBAP2LmRuby2

TDP-43NLS-Clover

80 min NaAsO₂

B

UBAP2LmRuby2

TDP-43NLS-Clover

Full bleach

C

Full bleach

D

E

Full bleach

F

TDP-43NLS-Clover

G

Non-dividing Neuronal-like SH-SY5Y cells

Endogenous TDP-43EGFP 2 alleles

NaAsO₂

Live Imaging & FRAP

TDP-43EGFP

15 days post-particles

Before NaAsO₂

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neurons (Figure 6A). Forced expression of cytoplasmic TDP-43 in iPSC-derived motor neurons produced rounded cytoplasmic structures in the absence (Figure 6B) or presence of a low dose of sodium arsenite (Figure 6C), in both cases without stress granule assembly or recruitment of stress granule proteins (e.g., G3BP1, Figures 6B and 6C). In the absence of arsenite, the rounded TDP-43 particles were liquid-like (as indicated by fast fluorescence recovery after photobleaching, Figures 6D and 6E). Low-dose arsenite addition rapidly converted the TDP-43-containing droplets into gel/solid-like particles that did not recover after photobleaching (Figures 6F and 6G). Thus, elevation of cytoplasmic TDP-43 level in human motor neurons is sufficient to drive cytoplasmic LLPS of TDP-43 independently of the principal components of stress granules or stress granule assembly.

**Cytoplasmic De-mixing Progressively Depletes Nuclear TDP-43 and Induces Cell Death**

We next evaluated the kinetics and longevity of cytoplasmic TDP-43 LLPS induced in the cytoplasm by transient exposure to sonicated amyloid-like fibrils and whether continued de-mixing affected cell viability. Fragmentated fibrils were transiently added to the culture media of non-cycling, neuronal SH-SY5Y cells genome edited to express TDP-43EGFP from both endogenous TDP-43 alleles (Figure 7A). Within 4 days, TDP-43EGFP partially re-localized into rounded cytoplasmic particles, many of which had undetectable fibrils (e.g., see white arrow, column 2, Figure 7B). The number and size of the cytoplasmic TDP-43-containing particles increased with time (Figures 3F and 7B). Seven days after transient fibril exposure, 50% of the cells had rounded cytoplasmic TDP-43 droplets, but no remaining fibrils could be detected in any cell (Figure 7B). Beginning ~10 days after fibril exposure, cytoplasmic LLPS of TDP-43 was accompanied by nuclear clearing of TDP-43 in some cells. By 1 month, most remaining cells accumulated TDP-43EGFP cytoplasmic particles with nuclear TDP-43 depletion (Figure 7B). The TDP-43 droplets did not contain detectable RNA (Figures S6A, S6B, and S6D) and were independent of arsenite-induced poly-A RNA-containing stress granules (Figure S6C). The 1-month-old cytoplasmic TDP-43 LLPS droplets induced by amyloid-like particles recruited phospho-TDP-43 (Figure 7E), but not ubiquitin or the autophagy adaptor protein p62 (Brady et al., 2011) (Figure S6F). While no cell death was seen within the first week, within 2 weeks after fibril exposure, 60% of cells had died, with most of the remaining cells displaying both cytoplasmic LLPS and nuclear clearance of TDP-43. By 6 weeks, there were almost no cell survivors (Figures 7C and 7D).

**Disrupted Nucleocytoplasmic Transport Enhances Nuclear Depletion of TDP-43**

Recognizing that defects in nuclear membrane structure and/or nucleocytoplasmic transport have been reported in the nervous systems of patients with ALS caused by hexanucleotide repeat expansion in C9orf72 (Freibaum et al., 2015; Jovičić et al., 2015; Zhang et al., 2015) or mutation in SOD1 (Kinoshita et al., 2009) and in mouse models of TDP-43 or SOD1 mutant-mediated disease (Chou et al., 2018; Ditsworth et al., 2017; Zhang et al., 2006), we examined the integrity of the nuclear membrane and nuclear import in non-cycling SH-SY5Y neuronal cells with cytoplasmic LLPS of TDP-43. Transient exposure to amyloid-like particles provoked gradual retention in the cytoplasm of components critical for nucleocytoplasmic transport (Figures 8A–8D). Cytoplasmic mislocalization of RanGAP1, the RAN GTPase-activating protein 1 required for Ran-dependent nuclear import and export, was found in some cells within 1 week (Figure 8B). RanGAP1 mislocalization increased in a time-dependent manner, with 70% of cells accumulating RanGAP1 in large, cytoplasmic inclusions within a month after fibril exposure (Figure 8C), along with progressive accumulation of de-mixed TDP-43 in the cytoplasm.

Imported cargo is released into the nucleus when importin transporters interact with intranuclear Ran-GTP. Correspondingly, high levels of intranuclear Ran-GTP are essential for active transport through nuclear pore complexes (NPCs) and for defining nucleocytoplasmic transport directionality. Consistent with disruption in nuclear import, continuing LLPS of fragmented fibril-induced TDP-43 was accompanied by Ran accumulation into small cytoplasmic inclusions (Figure 8D). Furthermore, after
Figure 6. De-mixing of Cytoplasmic TDP-43 in Human iPSC-Derived Motor Neurons Forms Liquid-like Droplets Independent of Stress Granules, which Convert into a Gel/Solid-like State after Arsenite Induced Stress

(A) Schematic of experimental design to assess the properties of de-mixed cytoplasmic TDP-43 particles in human iPSC-derived motor neurons with or without sodium arsenite treatment. Human iPSC-derived motor neuron precursor cells were differentiated for 7 days, and then the differentiated motor neurons were infected with a lentivirus, driving expression of Ubi::TDP-43ΔNLS-GFP. After 1–2 weeks of expression, the cells were analyzed.

(B and C) Immunostaining of G3BP1 of mature motor neurons expressing cytoplasmic TDP-43 in the absence of sodium arsenite treatment (B) or with 50 μM sodium arsenite (C). MAP2 was stained for neuron marker.

(D and E) Representative images of cytoplasmic TDP-43ΔNLS-GFP particles formed in the absence of stress. FRAP example of TDP-43ΔNLS-GFP particles in a neuron after a complete bleaching (D). Mean fluorescence intensity of the fully bleached TDP-43ΔNLS-GFP particles over time (E). Data are normalized to the average intensity of a particle before photobleaching and are represented as mean ± SEM from the recovery curves of eight particles in a total of four independent experiments.

(F and G) Representative images of cytoplasmic TDP-43ΔNLS-GFP particles after 3 h of 50 μM sodium arsenite treatment. FRAP example of TDP-43ΔNLS-GFP particles in a motor neuron after a complete bleaching (F). Mean fluorescence intensity of the fully bleached TDP-43ΔNLS-GFP particles over time (G). Data are normalized to the average intensity of a particle before photobleaching and are represented as mean ± SEM from the recovery curves of nine particles in a total of four independent experiments.
Figure 7. Endogenous EGFP-Tagged TDP-43 De-mixes in the Cytoplasm, Forming Liquid-like droplets that Slowly Deplete Nuclear TDP-43 and Compromise Cell Survival

(A) Schematic of experimental design to assess LLPS properties of endogenously EGFP-tagged TDP-43 in non-cycling SH-SY5Y cells (knockin in both alleles), which were incubated with fluorescently labeled wild-type FUS-mCherry fibrils and visualized over time by live imaging or immunofluorescence.

(B) Representative images using confocal microscopy of neuronal-like SH-SY5Y cells inoculated with sonicated His-FUS mCherry particles at a final concentration of 200 nM and further imaged for FUS mCherry fibrils (red) and TDP-43 EGFP (green) up to 1 month. Media were changed after 3 days. White arrows indicate cytoplasmic particles containing mislocalized endogenous TDP-43 EGFP (green). Dashed white line outlines nuclei.

(C) Cell survival quantification of non-treated (black) and fibril-treated (red) neuronal-like cells over time. Data are represented as mean ± SEM from a total of three independent experiments.

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1 month of TDP-43 LLPS, two components of nuclear pores (Nup107 and Nup62) were found accumulated in cytoplasmic particles, with one (Nup62) co-recruited into rounded TDP-43-containing droplets (see white arrows, Figure 6D), indicative of phase separation of Nup62 together with TDP-43. Similarly, FUS and hnRNPA1 also redistributed to the cytoplasm (Figures S7A and S7B), with hnRNPA1 also recruited into TDP-43 de-mixed particles (Figure S7B, lower panel). The evidence that hnRNPA1 is co-recruited with TDP-43 LLPS is in accordance with a report that the LCD (low complexity domain) of TDP-43 phase separates with the LCD of hnRNPA2 in vitro (Ryan et al., 2018). Finally, the nuclear import transporter importin-β1 can dissolve TDP-43 amyloid-like fibrils in vitro (Guo et al., 2018), was also phase separated into large, cytoplasmic TDP-43-containing droplets (see white arrows, Figure 6D, lower panels). Overall, induction of cytoplasmic TDP-43 LLPS is sufficient over a timescale of weeks to drive a feedforward mechanism compromising nucleo-cytoplasm transport and exacerbating loss of nuclear TDP-43 (Figure 8E).

DISCUSSION

Here we demonstrate that LLPS reflects a normal TDP-43 behavior in vivo, as we show that it is apparently de-mixed intranuclearly in cortical neurons in the mouse brain and multiple examples in cell culture. TDP-43 molecules exhibit fast (within seconds) dynamic exchange within the liquid droplets as well as between the droplets and an aqueous pool. Intranuclear TDP-43 thus recapitulates the characteristics of what has been defined as a liquid produced by LLPS (Boynaaemis et al., 2018; Brangwynne et al., 2009). We also show that beyond intranuclear de-mixing, TDP-43 can be induced to de-mix in the cytoplasm (1) when cytoplasmic TDP-43 levels are elevated, (2) in response to transient exposure to amyloid-like fibrillar TDP-43 or FUS, or (3) following exposure to sodium arsenite. In all cases, TDP-43 de-mixes initially into liquid droplets (with rapid molecular exchange with soluble TDP-43 molecules) but converts to rounded gel/solid-like structures with arsenite exposure.

The interaction of TDP-43 (Afroz et al., 2017; Becker et al., 2017; Khalifallah et al., 2018; Li et al., 2013; Ramaswami et al., 2013) or FUS (Guo et al., 2018; Hofweber et al., 2018; Li et al., 2013; Marrone et al., 2018; Zhang et al., 2018) with stress granules has repeatedly been proposed to be a means to enhance their subsequent aggregation. In contrast, we have found that cytoplasmic LLPS of TDP-43 can be independent of components of stress granules and stress granule assembly despite initial recruitment of a small proportion of TDP-43 into stress granules induced with a high dose of sodium arsenite. Perhaps most provocatively, simply increasing cytoplasmic TDP-43 (as would be driven by the known age-dependent reduction in nuclear import; D’Angelo et al., 2009; Mertens et al., 2015) or transient exposure to fibrillar fragments of aggregated TDP-43 (or FUS) recruits TDP-43 into de-mixed cytoplasmic liquid droplets that are independent of stress granules and that slowly deplete nuclear TDP-43 and induce cell death over a 6-week period. Added to the complexity of TDP-43 de-mixing, it is not established whether the cytoplasmic particles we identify here are similar to axonally transported TDP-43-containing droplets that form in cortical neurons following TDP-43 overexpression (Gopal et al., 2017). Identifying the constituents of each of these TDP-43-containing compartments is an unsolved question of high interest for elucidating possible functions carried out within such compartments.

A prominent hallmark of ALS and FTD is nuclear clearance and cytoplasmic TDP-43 aggregation in neurons and glia within the central nervous system (Neumann et al., 2006). Here, we have established that increased TDP-43 concentration in the cytoplasm or application of either of two transient stresses can induce long-lasting LLPS of endogenous TDP-43. While much attention has focused on the relationship of stress granules with disease-related RNA-binding protein aggregation (Becker et al., 2017; Jain et al., 2018; Mackenzie et al., 2017; Markmiller et al., 2018), our findings demonstrate that cytoplasmic de-mixing of TDP-43 can be independent of stress granule formation, with at most a small proportion of TDP-43 transiently recruited to stress granules. Recognizing that other RNA-binding proteins have previously been reported to indirectly associate with TDP-43 through simultaneous binding to a common RNA (Coyne et al., 2015; Elden et al., 2010), it is plausible that transient association of a minority of TDP-43 molecules with stress granules is driven indirectly through TDP-43 binding to RNAs that are also bound to proteins directly recruited to stress granules. In addition, TDP-43 can bind to poly(ADP-ribose) via its nuclear-localization sequence. Binding promotes TDP-43 recruitment to stress granules and delays the initial accumulation of disease-associated TDP-43 phosphorylation that accompanies sustained arsenite-mediated stress (McGurk et al., 2018). TDP-43 phase separation into droplets without conventional stress granule components is consistent with the near absence of evidence using immunohistochemistry and immunofluorescence of stress granule formation or co-localization of stress granule proteins with cytoplasmic TDP-43 aggregates in tissues from patients (Colombrita et al., 2009; Dormann et al., 2010; Liu-Yesucevitz et al., 2010; Mackenzie et al., 2017; McGurk et al., 2014; Neumann et al., 2007) or, more recently, from proteomic analysis of insoluble TDP-43-containing homogenates from ALS or FTD patients (Laferrière et al., 2019).

We have established that transient exposure to amyloid-like TDP-43 (or FUS) particles is sufficient to induce TDP-43 de-mixing into spherical liquid-liquid cytoplasmic droplets (1) that can be stable for weeks and exhibit dynamic internal molecules, (2) that exchange with TDP-43 in the cytoplasm and fusion and fission.
Figure 8. TDP-43 LLPS Is Accompanied by Disruption of Nucleocytoplasmic Transport
(A) Schematic of experimental design to assess LLPS properties of endogenous TDP-43EGFP in cells and their impact on nucleocytoplasmic transport.
(B) Representative images using confocal microscopy of neuronal-like SH-SY5Y cells 7 days and 15 days post incubation of sonicated FUSmCherry fibrils and further immunostained with RanGap1 (red) and cytoplasmic TDP-43 EGFP (green).
(C) Immunostained RanGap1 (red) and cytoplasmic TDP-43EGFP (green) 1 month after incubation with FUSmCherry sonicated fibrils (left panel) or in absence of fibril treatment (right panel). Quantification of the percentage of cells with cytoplasmic RanGap1 inclusions 1 month post treatment (from 200 cells in a total of three independent experiments).

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events, and (3) whose presence slowly depletes nuclear TDP-43 accompanied by reduced cell survival. Nuclear TDP-43 depletion has also been reported when TDP-43 is co-recruited into polyglutamine repeat inclusions (Fuentealba et al., 2010). Induced LLPS of TDP-43 is potentially mediated through modifications of it or its binding partners that affect the multivalent interactions that are generally thought to be driving the forces of protein phase separation (Dao et al., 2018; Li et al., 2012; Mollieix et al., 2015). Evidence from multiple investigators has supported post-translational modifications and/or binding partners that can act to suppress or enhance de-mixing of FUS (Guo et al., 2018; Hofweber et al., 2018; Qamar et al., 2018; Yoshizawa et al., 2018). Here we show that a portion of TDP-43 within liquid cytoplasmic droplets is phosphorylated, suggesting that LLPS may represent the initial event that then matures to a gel-like solid state that ultimately nucleates TDP-43 aggregation.

A likely, disease-related stress is the reduction in nuclear import during normal aging (D’Angelo et al., 2009; Mertens et al., 2015; Scaffidi and Misteli, 2006), coupled with further import inhibition in different neurodegenerative diseases (Freibaum et al., 2015; Gasset-Rosa et al., 2017), including ALS (Boeynaems et al., 2016; Ditsworth et al., 2017; Freibaum et al., 2015; Jovičić et al., 2015; Zhang et al., 2015, 2018). Here we demonstrate that major regulators of nucleocytoplasmic trafficking (RanGap1, Ran, Nup107, and Nup62) are abnormally localized as a consequence of induced cytoplasmic LLPS of TDP-43. While nuclear-import receptors, including importin-α and karyopherin-β1, have been reported to prevent or reverse TDP-43 aggregation in vitro (Guo et al., 2018) (in addition to directly mediating nuclear import of TDP-43), we have found that induced LLPS of TDP-43 in the cytoplasm recruits importin-α, thereby producing an importin-α loss of function that will exacerbate nuclear import deficits. Co-recruitment of Nup62 and importin-α into TDP-43 droplets points to TDP-43 phase separation to be at least one driving force inducing inhibited nuclear import.

We note that reduced nuclear import will obligatorily disrupt the TDP-43 autoregulation pathway (Ayala et al., 2011; Polymenidou et al., 2011), which sets the level of new TDP-43 synthesis through the nuclear action of TDP-43 in the processing of its own pre-mRNA (Polymenidou et al., 2011). Diminished nuclear TDP-43 function, coupled with increased cytoplasmic accumulation from inhibition of nuclear import, produces a feedforward loop for driving increasing cytoplasmic TDP-43 accumulation, further exacerbating the disruption of its autoregulation pathway and producing even higher levels of cytoplasmic TDP-43. During aging, this would be expected to be sufficient to drive concentration-dependent cytoplasmic phase separation of TDP-43, which, together with the loss of TDP-43 nuclear function, ultimately drives cell death.

Lastly, our evidence has shown that beyond age-dependent loss of nuclear import, other stresses—including transient, extracellular contact with aggregated, amyloid-like fibrils of assembled TDP-43 or FUS—can trigger or contribute to the de-mixing of TDP-43 in the cytoplasm. One of these stresses is likely to be a proteome imbalance generated by the altered protein abundances that arise from changes in the levels and/or splicing of >1,500 mRNAs in the nervous system when TDP-43 is diminished (Polymenidou et al., 2011; Tollervy et al., 2011). Efforts to identify additional biological sources of such stress are now of high importance for establishing the cascade of events underlying age-dependent neurodegenerative diseases that have TDP-43 mis-accumulation and aggregation as prominent features.

**STAR METHODS**

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**SUPPLEMENTAL INFORMATION**

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AUTHOR CONTRIBUTIONS


DECLARATION OF INTERESTS

The authors declare no competing interests.

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# STAR★METHODS

## KEY RESOURCES TABLE

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CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Don Cleveland: dcleveland@ucsd.edu.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Mice

C57BL/6 mice were obtained from Charles River Laboratories (Cat# C57BL/6NCrl). Males were perfused in a 4% paraformaldehyde solution and tissues collected for imaging. Maintenance and experimental procedures were approved by the Institutional Animal Care and Use Committee of the University of California, San Diego.

Cells Lines

iPSC-derived motor neurons

Human iPSC cells, derived from peripheral blood mononuclear cells donated by a 58-year-old healthy Caucasian male, were generated by iXCells Biotechnologies and selected for normal karyotype with normal self-renewal and differentiation ability (Melamed et al., 2019). iPSC cells were first differentiated into motor neuron precursors in 21 days and then further differentiated into motor neurons for another 7 days using a differentiation protocol patented by iXCells Biotechnologies (Provisional Application no. 14359-001-888) (Melamed et al., 2019). Then the cells were infected by lentivirus to express cytoplasmic TDP-43<sub>DNLS-GFP</sub> under ubiquitin promoter as summarized in Figure 6. Human iPSCs were obtained from iXCells and their use further approved by the Embryonic Stem Cell Research Oversight (ESCR0) Committee and Institutional Review Boards (IRB) of the University of California San Diego in accordance with the requirements of the Code of Regulations on the Protection of Human Subjects.

Stable cell line construction: lentivirus infection and selection

HEK293T cells were used for packaging lentiviruses. Briefly, 0.5 x 10^6 per well of 293T cells were seeded in a 6-well plate and cultured with DMEM medium (Thermo Fisher) supplemented with 10% FBS (Omega Scientific), 100 units/mL of penicillin, 100 µg/mL of streptomycin, and 250 ng/mL of Gibco Amphotericin B (Thermo Fisher). For lentiviral transfection, 1.5 µg of the lentiviral plasmid (constructs are labeled as lentivirus in Key Resources Table), 1 µg of pMD2.G and 0.5 µg of pSAX2 were inoculated to each well using Mirus transIT-X2 transfection reagent. Culture medium was changed to fresh medium at 4-24 h post transfection. Two days after transfection, the culture medium was filtered through a 0.45 µm syringe filter to generate the viral soup. The viral soup containing 10-50 µg/mL protamine sulfate was added to U2OS or SH-SY5Y cells cultured in DMEM medium or in DMEM/F12 medium (Thermo Fisher) for infection. The viral soup was removed 24 h after infection, and cells were passaged at least once before selection. Infected cells are selected based on the selection markers encoded by the lentivirus. For U2OS, the concentrations of the antibiotics used for selection were 200 µg/mL for neomycin (Thermo Fisher), 20 µg/mL for blasticidin (Thermo Fisher), 1 µg/mL for puromycin (Thermo Fisher). For SH-SY5Y cells, the concentrations were 400 µg/mL for neomycin, 10 µg/mL for blasticidin and 3 µg/mL for puromycin.

TDP-43 alleles were genetically modified in SH-SY5Y cells. A single guided RNA targeting TDP-43 was designed (Benchlin webtool) and cloned into pSpCas9 plasmid (px330-Addgene) (Cong et al., 2013) using the BbsWE restriction site. The sequence for guided RNA targeting TDP-43 is: GCTGGGGAATGTAGACAGTG. To promote homologous recombination, pSpCas9-gRNA plasmid was electroporated using Amaxa Nucleofactor (assay A-023) along with a template plasmid, containing two TDP-43 homology arms, each 800bp long, separated by AID-GFP segment that is adjacent upstream to the TDP-43 terminal codon. 48 h following electroporation, cells were collected and dissociated using Accutase (Innovative Cell Technologies). GFP-positive cells were sorted and single-cell seeded into 96 wells plates using the SH800S Sony cell sorter. Individual clones were expanded and DNA was extracted for PCR amplification of TDP-43 genomic locus. Insertion of AID-GFP was finally confirmed by immunoblotting. The entire TDP-43 coding region was then sequenced to verify the absence of any additional DNA alteration.
METHOD DETAILS

Mice surgery and tissue preservation
Anesthetized animals (C57BL/6) were subjected to transcardial perfusion with room temperature Sorenson's phosphate buffer (SPB), and fixed with ice-cold 4% paraformaldehyde (PFA) in phosphate buffer (PBS). Brains were removed, trimmed with coronal cuts immediately rostral to the forebrain, post-fixed in 30% sucrose, embedded in HistoPrep (TM) (SH75-125D, Fisher Chemical) followed by a rapid incubation in 4°C 2-Methylbutane (Fisher Scientific), and kept at −80°C. Fixed brains were cut using the Leica 2800E Frigocut cryostat at −20°C. 35 μm thick free-floating sections were preserved in PBS containing Sodium Azide (0.02%) at 4°C.

Immunofluorescence (free-floating OCT-embedded sections)
The sections were washed in PBS (3 times, 5 min each), permeabilized and blocked in a PBS/0.5% Tween 20/1.5% BSA for 1 h at room temperature, followed by an overnight incubation with the primary antibody anti-TDP-43 (12892-1-AP, Proteintech; 1:500) diluted in a PBS/0.3% Triton X-100 solution at room temperature. Next, sections were washed in PBS (3 times, 10 min each) and incubated with the secondary antibody (diluted in PBS) for 1 h, washed twice with PBS (10 min each), and then incubated for 10 min with PBS/DAPI (Thermo Fisher Scientific, 100ng/mL) solution. Sections were mounted on Fisherbrand Superfrost Plus Microscope Slides (Thermo Fisher Scientific) with ProLong Gold antifade reagent with DAPI (Thermo Fisher Scientific).

Cell culture
All cell lines were cultured at 37°C in a humidified atmosphere at 5% CO2. U2OS cells were cultured in DMEM (GIBCO) supplemented with 10% fetal bovine serum (FBS) and Antibiotic-Antimycotic (GIBCO). SH-SY5Y cells were cultured in DMEM+F12 (GIBCO) supplemented with 10% FBS and Antibiotic-Antimycotic. When the cell cycle was arrested in G1, Palbociclib (Apexbio) was added into the media. Cell culture media was changed every 3 days.

Primary hippocampal neuronal cultures were prepared as described previously (Seibenhener and Wooten, 2012) with some modifications. In brief, timed pregnant C57BL/6 mice (Charles River) were sacrificed, and embryos were collected at embryonic day 18 (E18). Primary hippocampal neurons were isolated from both male and female embryos and pooled. Cells were dissociated with 0.125% trypsin, and plated on 96-well glass-bottom plates coated with poly-D-lysine (Sigma-Aldrich) at a cell density of 60,000/mL in neurobasal medium (GIBCO) supplemented with 2% B27 (Sigma-Aldrich) and 0.25% glutamine (Sigma-Aldrich). Thereafter, half of the medium was replaced twice a week. Neurons were used 6 to 8 days after plating.

Biochemical fractionation of soluble and insoluble proteins
Cells from six wells were treated as indicated and then lysed in 300 μL of 1x RIPA buffer (50 mM Tris pH 8.0, 150 mM NaCl, 1% NP-40, 5 mM EDTA, 0.5% sodium deoxycholate, 0.1% SDS), supplemented with Halt Protease and Phosphatase Inhibitor Cocktail (Thermo-Fisher Scientific). After sonication (13% Amp, 5s, Branson digital Sonifier), cell lysates were centrifuged at 21,100 g for 30 min at 4°C and the supernatant was collected as the soluble fraction. The pellet fraction was washed with 1x RIPA buffer once before solubilizing in 75 μL of urea buffer (7M urea, 2 M Thiourea, 4% CHAPS, 30 mM Tris, pH 8.5) and cleared by centrifugation at 21,100 x g for 30 min at 4°C. Then the soluble and insoluble fractions were subsequently analyzed by western blot.

Immunoblotting
Whole-cell extracts (from 5x10⁶ cells) were directly lysed in SDS sample buffer and boiled for 10 min. Samples were resolved by SDS-PAGE, transferred to PVDF, and blocked with 5% milk in TBST (TBS, 0.1% Tween-20). The following primary antibodies were used at 1:1,000 dilution (unless noted) in TBST: anti-TDP-43 (10782-2-AP, Proteintech), anti-tubulin (ab6160, Abcam), anti-GFP (632381, Clontech) and 1:2,000 anti-GAPDH (Ab8245, Abcam), anti-phospho-TDP-43 (829901, Biolegend; 1:500). Blots were probed with 1:5,000 dilutions of HRP-conjugated secondary antibodies (GE Healthcare) and exposed to film.

Live cell imaging
U2OS cells were plated on a density of 8,000 cells per well onto a glass-bottom 96-well plate. TDP-43ΔNLS-clover proteins were induced for 24 h by adding 1 μg/mL of doxycycline. TDP-43ΔNLS-clover was induced for 48 h due to the low-level expression of this variant. Live cell imaging was performed on CQ1 Confocal Quantitative Image Cytometer with a 40x objective and environmental control system of 37°C, 95% humidity and 5% of CO2. Nine z stack images (1 μm per stack) were collected per frame, and the maximum intensity projection was used to construct the final image series. To detect the fast fusion behavior of TDP-43 particles, images were collected at 150 s or 30 s intervals. For sodium arsenite treatment, 0.25 mM or 50 μM of sodium arsenite was added to the cells and images were collected at 2-min, 5-min or 10-min intervals as indicated. A maximal intensity projection image was generated per frame.

SH-SY5Y TDP-43ΔEGFP cells were plated on 8-well chamber (Ibidi) at 25,000 cells per well. After 24 h cells were arrested in G1 with Palbociclib (Apexbio) and maintained arrested during the whole length of the experiment. Sonicated fibrils were added at final...
Fluorescence recovery after photo bleach (FRAP) analysis
FRAP experiments on U2OS cells were performed on Zeiss LSM880 Aryscop microscope with 40x/1.2 W objective. The intensity of the fluorescent signal is controlled in the detection range through changing the laser power, digital gain and off-set. For the green channel, bleaching was conducted by a 488-nm line from an argon laser at ~80%-100% intensity with ~10-20 iterations. FRAP experiments on SH-SY5Y TDP-43EGFP cells were performed on Olympus FV1000 Spectral Confocal using SIM Scanner with 100x oil immersion objective, bleaching was conducted scanning a region of 1x1 μm for 300ms at 8% of laser intensity at 405-nm. Fluorescence recovery was monitored at 0.5 s, 2 s, 5 s or 10 s intervals for 3 min. In the focal-bleach experiment roughly half of a particle is bleached or fully photobleached, and then the distribution of the fluorescence within the photo-manipulated particle is determined over time. During the experiment cells were maintained in Leibovitz’s L-15 medium (CO2 independent).

Immunofluorescence from cells
Primary hippocampal neurons, U2OS and SH-SY5Y cells were cultured on 8-well glass-bottom chamber slides (Ibidi). SH-SY5Y cells were plated on 8-well chamber (Ibidi) at 25,000 cells per well. After 24 h cells were arrested in G1 with Palbociclib (Apexbio) and maintained arrested during the whole length of the experiment. Sonicated fibrils were added at final concentration of 200 nM and media was removed after 3 days. Media was changed every 3 days.

At the indicated time points, cells were fixed with 4% PFA for 10 min at room temperature After two washes with PBS, cells were permeabilized and blocked with blocking solution (0.1% Triton, 2% BSA in PBS) for 1 h at room temperature. Cells were then incubated overnight with the primary antibody in PBS/0.3% Tween 20. The primary antibodies used for staining were: anti-G3BP1 (ab56574, Abcam; 1:1000), anti-TDP-43 (10782-2-AP, Proteintech; 1:500), anti-TDP-43 (12892-1-AP, Proteintech; 1:500), anti-TIA1 (sc-1751, Santa Cruz; 1:300), anti-HA (mms-101P, Covance; 1:500), anti-His (H1029, Sigma; 1:2000), anti-RanGap1 (sc-1882, Santa Cruz; 1:500), anti-Ran (bd-610340, BD Bioscience; 1:500) anti-Nup107 (Ab73290, Abcam; 1:300), anti-Nup62 (MABE1043, Millipore; 1:500), anti-FUS (A303-839A, Bethlehem Laboratories; 1:500), anti-phospho-TDP-43 (829901, Biolegend; 1:500), anti-hnRNPA1 (sc-374053, Santa Cruz; 1:500), anti-amyloid A1 (AHB0052, ThermoFisher; 1:500), anti-p62 (GP62-C-WBC, Progen; 1:500), anti-EDC4 (sc-376382, Santa Cruz; 1:100) and anti-MAP2 (NB300-213, Novus Biologicals; 1:500). After three washes with PBS, the cells were subsequently incubated with fluorescently labeled secondary antibodies diluted at 1:500 in PBS/0.3% Tween 20 for 1 h at room temperature. After one wash with PBS, one wash with 300 nM DAPWE (PBS) and then one wash with PBS, the cells are preserved in PBS for imaging. SH-SY5Y cells imaging was performed on FV1000 Spectral Confocal (Olympus) at 60-100x magnification.

RNA Fluorescence in situ hybridization (FISH)
All hybridization steps were performed under RNase-free conditions. Cells were fixed in 3.7% formaldehyde for 10 min and permeabilized with ethanol 70% for 1 h at 4°C. Then, cells were washed with Wash Buffer (1:5 Wash Buffer A, Biosearch technologies Cat SMF-WA1-60 plus 1:10 deionized Formamide) and blocked in the dark at 37°C for 4 h in a humidified chamber with Hybridization Buffer (9:10 RNA FISH Hybridization Buffer Cat SMF-HB1-10 plus 1:10 deionized Formamide) containing 5'-labeled Cy5-(d)T20 oligonucleotides (gift from Dr. J. Paul Taylor, St. Jude Children Hospital) (1ng/μl). After cells were washed with wash buffer in the dark at 37°C for 30 min were stained with DAPI and proceed to imaging.

SYTO RNA Select Green Fluorescent Cell Stain
All the staining steps were performed under RNase-free conditions. SH-SY5Y cells were fixed with cold methanol at −20°C for 10 min and immunostained with anti-TDP-43 (12892-1-AP, Proteintech; 1:500) following the protocol described. Then, cells were washed with PBS two times and stained with freshly prepared RNASelect (Molecular Probes) (500 nM) at room temperature for 20 min. Finally, cells were washed for 5 min in PBS and stained with DAPI.

Recombinant protein purification
All proteins were expressed and purified from E. colwe BL21 under native conditions. Protein expression was induced with 1 mM IPTG for 16 h at 16°C. E.colwe cells were lysed by sonication on ice in PBS with protease inhibitors (cOmplete, EDTA-free, Roche Applied Science). HA-FUS expression constructs were generated in pGST-Duet to contain a TEV-cleavable site, resulting in GST-TEV-HA-FUS construct. The protein was purified over pre-packed Glutathione Sepharose High Performance resin (GSTrap HP columns, GE) with one-step purification of glutathione S-Transferase (GST) tagged FUS protein using Akta Pure fast protein liquid chromatography (FPLC) system (GE) at 4°C. GST-HA-FUS protein was eluted in 50 mM Tris-HCI, pH 8, 200 mM trehalose, and 20 mM glutathione. His-TDP-43 WT, His-TDP-43 WT-mCherry, His-TDP-43 G298S-mCherry, His-SOD1 WT-mCherry and His-FUS WT-mCherry proteins were purified over pre-packed Ni Sepharose High Performance HisTrap HP (GE) using an AKTA pure chromatography system at 4°C and eluted with 50 mM Tris pH 7.4, 100 mM NaCl and 400 mM Imidazole. In the case of His-TDP-43 WT-mCherry Ni-IMAC eluted fractions were followed by gel Filtration Chromatography (Superdex 75 10/300, GE). The following Molecular Weight Markers were concentration of 200 nM and media was removed after 3 days. Live cell imaging was performed on Olympus FV1000 Spectral Confocal (Olympus) at 60x magnification at environmental control system of 37°C, and 5% of CO2 at the time points indicated.
used: Carbonic Anhydrase from bovine erythrocytes (29 KDa, Sigma), Albumin, bovine serum (66 KDa, Sigma) and β-Amylase from sweet potato, (200 KDa, Sigma). Eluted proteins (GST-HA-FUS, His-TDP-43G93A-mCherry, His-SOD1WT-mCherry and His-FUSWT-mCherry) with the expected size were collected and concentrated to final concentration of 12 μM using Amicon Ultra centrifugal filter units (10 KDa molecular weight cut-off; Millipore). All proteins after purification were centrifuged for 15 min at 14,000 rpm at 4°C to remove any aggregated material. Protein concentration was calculated by Coomassie Blue with BSA protein as standard, and by colorimetric Bradford assay (Bio-Rad). For protein storage at −80°C glycerol was added (30%)

**Amyloid-fibrils reconstruction**

HA-FUSWT protein was buffer exchanged at 4°C into FUS assembly buffer (50 mM Tris–HCl, pH 8, 200 mM trehalose, 1 mM DTT, 20 mM glutathione). Seed formation was initiated by addition of TEV protease to GST-TEV-HA-FUS (4 μM) in FUS assembly buffer for 3 h and then high salt storage buffer (40 mM HEPES pH 7.4, 500 mM KCl, 20 mM MgCl2, 10% glycerol, 1 mM DTT) was added for 3 h (Sun et al., 2011). FUS fibrilization was initiated by adding 1:20 of FUS seeds to GST-TEV-FUS (4 μM) and TEV protease in FUS assembly buffer for 24 h at 22°C. The fibrils generated were sonicated at 45% for 45 s. Sonicated fibrils were used as seeds on the following FUS fibrilization assays. His-FUSmCherry was initially dialyzed with FUSWT-mCherry assembly buffer (Tris 50 mM pH 7.4, 100 mM NaCl, 1 mM DTT). Fibrils were generated by adding sonicated HA-FUSWT fibrils to His-FUSWT-mCherry (4 μM) for 24 h at 22°C.

His-TDP-43 protein (4 μM) was initially dialyzed at 4°C with Tris 50 mM pH 7.4, 100 mM NaCl, 1 mM DTT buffer. Fibrils were induced by adding polyethylene glycol (PEG)-4000 (7%) and 2-methyl-2,4-pentanediol (MPD) (3%) for 16 h at 22°C. Sonicated His-TDP-43 fibrils were used as seeds on the following TDP-43 fibrilization assays. His-TDP-43WT-mCherry fibrils were generated by adding sonicated His-TDP-43 fibrils as seeds to soluble His-TDP-43WT-mCherry in TDP-43 assembly buffer supplemented with 7% PEG and 3% MPD.

His-SOD1G93A-mCherry fibrilization was induced by adding SOD1A4V aggregates produced in bacteria. Briefly, SOD1A4V protein aggregates were induced in BL21 cells, bacteria cell pellet was collected, lysed, and centrifuged/resuspended for 5 cycles (Molina-García et al., 2018). Then a sucrose discontinuous gradient was prepared: in a 2 mL Eppendorf tube, successively was displayed 200 μL of 60% (bottom)>50 > 40 > 30 > 20% (top) sucrose, freezing each layer at −80°C before adding the next one. Resuspended aggregates were laid on the sucrose gradient and centrifuged at 10,000 rpm 1 h at 4°C. Fractions were carefully collected and a Coomassie Blue gel was performed to identify the fraction(s) that contained SOD1A4V protein. SOD1A4V containing fractions were further dialyzed with PBS and sonicated. The final product was used as seeds for His-SOD1G93A-mCherry fibrilization. Seeds were added to His-SOD1G93A-mCherry (4 μM) in SOD1 assembly buffer (100 mM Na2SO4, 30 mM Tris–HCl pH 8, 4 mM MgSO4, 7% polyethylene glycol (PEG)-4000 and 3% 2-methyl-2,4-pentanediol (MPD)) at 22°C during 16 h. Finally, fibrils were dialyzed using slide-A-Lyzer MINWE Dialysis Units (10 kDa molecular weight cut-off; Thermo) in PBS for 3 h and sonicated at 45% 45 s just before adding them to the cell media.

**Transmission Electron Microscopy**

Fibril protein reactions (10 μl) were adsorbed onto glow-discharged 300-mesh Formvar/carbon coated copper grids (Ted Pella) and stained with 2% (w/v) aqueous uranyl acetate (Ladd Research Industries, Williston, VT). Excess liquid was removed, and grids were allowed to air dry. Grids were examined using a Tecnaw G2 Spirit BioTWIN transmission electron microscope equipped with an Eagle 4k HS digital camera (FEI, Hillsboro, OR).

**Survival curve**

For survival curve analysis, SH-SY5Y TDP-43EGFP cells were plated on a 24 well-plate and cytoplasmic LLPS were induced with His-FUSmCherry fibrils as described. At indicated time points, cells were collected and counted using a haemocytometer (Fisher Scientific).

**QUANTITATIVE AND STATISTICAL ANALYSIS**

All quantitative analyses relied on systematic uniform random sampling.

1. **FRAP**

   The FRAP data were quantified using ImageJ. The time series of the granule fluorescence intensity was calculated and the intensity of the background (area with no cells) was subtracted from the granule intensity. The intensity of the granule during the whole experiment was normalized to one before bleaching and the intensity of the granule just after bleaching was normalized to zero. An average of at least 4-20 particles per condition was used to calculate the mean and standard deviation. The averaged relative intensity and standard error were plotted to calculate the dynamics of the particles.

2. **Cytoplasmic TDP-43 LLPS**

   Number of cells with cytoplasmic TDP-43 particles and number of cytoplasmic particles were counted on 60x images taken on the Olympus FV1000 Spectral Confocal. DAPI staining was used to address cytoplasmic localization. ~200 cells were counted per condition and time point. Quantification was performed in at least three independent replicates. Arsenite induced cytoplasmic TDP-43...
3. Fluorescence quantification of Clover in living cells
U2OS were plated on 8-well glass-bottom chamber slides (Ibidi). Imaging of TDP-43<sup>DNLs-clover</sup> were done using the 20x objective (0.8. air. 0.55 mm) on Zeiss LSM880 confocal microscope with settings: scan resolution = 1024 × 1024; scan speed level = 8; pinhole = 20; Gain was set at 500 and offset was set to 600 so that all the signal was in the linear range. 488 nm were used for excitation and the signals from 500 nm to 600 nm were collected. Mean intensity of TDP-43<sup>DNLs-clover</sup> in the cytoplasm was measured using FIJI (ImageJ) by circling the area of the cells with fluorescence. Dot plotting of the intensity from 23 cells with TDP-43 granules, and 27 cells without granules from nine images.

4. Cell survival and TDP-43 LLPS with nuclear depletion
Cell survival was quantified manually with using a haemocytometer. The number of survival cells was normalized to day 1 post-fibril treatment. Non-fibrils treated cells were added as control. Survival was measured in at least three independent replicates.

Quantifications of the percentage of cells with cytoplasmic LLPS (with or without nuclear TDP-43 clearance), or with cytoplasmic LLPS accompanied by nuclear clearance were quantified manually on 60x images taken on the Olympus FV1000 Spectral Confocal. DAPI staining was used to address cytoplasmic localization. Data are represented as mean ± SEM from a total of three independent experiments.

5. RanGap1 aggregates
Number of cells with cytoplasmic RanGap1 aggregates were counted manually on 60x images taken on the Olympus FV1000 Spectral Confocal. 200 cells were counted. Quantification was performed in at least three independent replicates.

Statistical Analysis
Statistical analyses were performed using GraphPad Prism. All data are shown as the box whisker plot (mean, min, and max value) or mean ± standard error of the mean (SEM). The statistical significance of the differences between two groups was investigated by paired t test. Statistical tests are indicated in each figure legend along with the corresponding significance level (p value). The number of cells analyzed per experiment is provided in the corresponding figure legends.
Supplemental Information

Cytoplasmic TDP-43 De-mixing Independent of Stress Granules Drives Inhibition of Nuclear Import,
Loss of Nuclear TDP-43, and Cell Death

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Figure S1: Expression of fluorescently-tagged TDP-43 at physiological levels in SH-SY5Y cells. (Related to Figure 1) (A) Scheme outlining doxycycline inducible expression of TDP-43 with N-terminally tagged EYFP (EYFP-TDP-43) in SH-SY5Y cells. (B) Immunoblot showing EYFP-TDP-43 levels in SH-SY5Y cells after 72 hours of induction using a TDP-43 antibody. GAPDH antibody was used as loading control. (C) Immunoblotting of TDP-43 levels in U2OS cells expressing TDP-43mRuby2 upon doxycycline induction compared to endogenous TDP-43. GAPDH was used as loading control. (D) Scheme illustrating the genome editing of TDP-43 locus to introduce AID-EGFP in both TDP-43 alleles before the stop codons in SH-SY5Y cells. (E) Immunoblot showing that TDP-43*EGFP replaces endogenous TDP-43 with TDP-43 total level being maintained (compared to the parental untagged cells) using TDP-43 antibody. Tubulin antibody was used as loading control. (F) Representative fusion event of endogenous TDP-43*EGFP droplets in SH-SY5Y cells. Red arrowheads indicate the fusing droplets.
Figure S2: After 10 days of TDP-43\textsuperscript{mCherry} particles treatment, TDP-43 mislocalizes in the cytoplasm and TDP-43 LLPS presents fission events. (Related to Figure 2) (A) Coomassie blue staining of recombinant TDP-43\textsuperscript{mCherry} protein purified from bacteria at the expected size of 70 KDa (left panel), and FUS\textsuperscript{mCherry} protein at the expected size of 85 KDa (right panel). (B) Electron micrograph of amyloid-like fibrils of FUS\textsuperscript{mCherry} recombinant protein purified from bacteria. Right panel illustrates the FUS\textsuperscript{mCherry} fibrils after sonication before inoculating them into cell media. (C) Representative images using confocal microscopy of endogenous nuclear TDP-43\textsuperscript{EGFP} in non-fibril-treated SH-SY5Y cells at day 1 (left) and after 1 month of culture (right). (D) Representative images using confocal microscopy of neuronal-like SH-SY5Y cells inoculated with sonicated His-TDP-43\textsuperscript{mCherry} particles at a final concentration of 200 nM and further imaged for TDP-43\textsuperscript{mCherry} fibrils (red) and TDP-43\textsuperscript{EGFP} (green) at 10 days. Media was changed after 3 days. White arrows indicate cytoplasmic particles containing mislocalized endogenous TDP-43\textsuperscript{EGFP} (green). Dashed white line outlines nuclei. (E) Representative images using confocal microscopy of neuronal-like SH-SY5Y cells inoculated with sonicated FUS\textsuperscript{mCherry} fibrils and immunostained with TDP-43 antibody (red) with direct GFP fluorescence signal from TDP-43\textsuperscript{EGFP} (green). (F) A representative fission event of cytoplasmic TDP-43\textsuperscript{EGFP} LLPS is shown with thick yellow arrow pointing to the particle that is divided into two new particles (two thinner arrows) at minute 4.
Supplemental information

Figure S3: TDP-43 droplets are induced by FUS particles, but not SOD1 particles and are independent of stress granules. (Related to Figure 3) (A) Coomassie blue staining of recombinant HA-FUS-(TEV cleavage site)-GST protein purified from bacteria at the expected size of 82 KDa (left panel), and His-TDP-43 protein at the expected size of 43 KDa (right panel). (B) Electron micrograph of amyloid-like fibrils of SOD1\textsuperscript{mCherry} recombinant protein purified from bacteria. Right panel illustrates the SOD1\textsuperscript{mCherry} fibrils after sonication before inoculating them into cell media. (C) Representative images using confocal microscopy of neuronal-like SH-SY5Y cells inoculated with sonicated SOD1\textsuperscript{mCherry} particles (red) and immunostained after 1 day TDP-43 antibody. D) Confocal representative images of neuronal-like cells treated with NaAsO\textsubscript{2} (500 mM) for 1 hour and immunostained with TIA1 (red) and TDP-43 (green) antibodies to induce TIA1 positive stress granules. (E) Scheme of the experimental design to assess stress granule dependency of endogenously EGFP tagged TDP-43 in non-cycling SH-SY5Y cells (knock-in in both alleles), which were incubated with fluorescently labelled FUS\textsuperscript{mCherry} particles and visualized over time by live-imaging or immunofluorescence. (F) Representative images using confocal microscopy of neuronal-like SH-SY5Y cells inoculated with sonicated FUS\textsuperscript{mCherry} particles in absence (left panel) or presence of NaAsO\textsubscript{2}, for 1 hour (right panel) and further immunostained with stress granules marker TIA1 (red) and cytoplasmic TDP-43\textsuperscript{EGFP} (green) after 1 month of fibril treatment.
Figure S4: Untagged TDP-43\textsuperscript{ΔNLS} forms stress granule-independent particles. (Related to Figure 4) (A) Experimental design and (B) immunostaining of untagged cytoplasmic TDP-43 forming stress granule-independent particles after 250 µM sodium arsenite treatment of U2OS cells. G3BP1 was used as a stress granule marker.
Figure S5: Stress induces the formation of detergent-insoluble TDP-43 inclusions with the gel-like particles exhibiting amyloid-like features. (Related to Figure 5) (A) Western blot analysis of the proportion of TDP-43 and phospho-TDP-43 in the soluble (RIPA buffer) and insoluble (Urea buffer) fractions without or with 30 min or 60 min of 250 μM sodium arsenite treatment. GAPDH was used as loading control. (B) Representative images of phospho-TDP-43 immunostaining (red) in U2OS cells that accumulate cytoplasmic TDP-43ΔNLS-Clover particles (green) after 60 min of 250 μM sodium arsenite treatment. (C) Representative images of mRNA (red) using FISH and cytoplasmic TDP-43ΔNLS-Clover particles (green) after 90 min of 250 μM sodium arsenite treatment (red). (D) Representative images of EDC4 (P body) immunostaining (blue) in U2OS cells that form stress granule-independent cytoplasmic TDP-43ΔNLS-Clover particles (green) and stress granules indicated by UBAP2LmRuby2 (red) after 90 min of 250 μM sodium arsenite treatment. (E-F) Representative images of p62 immunostaining (red) with cytoplasmic TDP-43ΔNLS-Clover particles after 60 min of 250 μM sodium arsenite treatment (E) or in the cells with remaining cytoplasmic TDP-43ΔNLS-Clover particles after a four hour of wash off of 250 μM sodium arsenite (F). (G) Representative images using confocal microscopy of neuronal-like SH-SY5Y cells inoculated with sonicated FUSmCherry particles in absence or presence of NaAsO₂ for 30 minutes (lower panel) and further immunostained with amyloid-oligomers antibody A11 (red) and cytoplasmic TDP-43EGFP (green) after 1 month of fibril treatment or in absence of fibril treatment (left panel).
**Figure S6:** Cytoplasmic TDP-43 de-mixed droplets remain liquid for long periods and do not co-localize with polyA-containing RNAs. (Related to Figure 7)  
(A) Scheme of the experimental design to assess LLPS properties of endogenous TDP-43EGFP in cells and determine recruitment of RNA within the droplets. (B-C) Representative images using confocal microscopy of neuronal-like SH-SY5Y cells (endogenously EGFP tagged TDP-43) 1 month after inoculation of sonicated FUSmCherry particles revealing Poly-A-RNA using fluorescence in situ hybridization (FISH) (grey) and cytoplasmic TDP-43EGFP (green) or (C) subsequently treated for one hour of NaAsO2 (0.5 mM). (D) Neuronal-like SH-SY5Y cells 1 month after inoculation of sonicated FUSmCherry fibrils further stained with SYTO RNA (green) and TDP-43 (red). (E) Scheme of the experimental design to assess LLPS properties of endogenous TDP-43EGFP in cells and determine LLPS features. (F) Representative images using confocal microscopy of neuronal-like SH-SY5Y cells 1 month after inoculation of sonicated FUSmCherry fibrils and further immunostained with p62 (upper panel) or ubiquitin (lower panel) (red) and cytoplasmic TDP-43EGFP (green). Left panel illustrates cells in absence of fibril treatment.
Figure S7: TDP-43 LLPS induces mislocalization of hnRNPA1 and FUS into the cytoplasm due to nuclear import defects. (Related to Figure 8) (A-B) Representative images using confocal microscopy of neuronal-like SH-SY5Y cells in absence of fibril treatment (A) or 1 month after inoculation of sonicated FUS-mCherry fibrils (B) and further immunostained with FUS (upper panel), hnRNPA1 (lower panel) (red) and cytoplasmic TDP-43 EGFP (green).