TDP-43 stabilizes G3BP1 mRNA: relevance to amyotrophic lateral sclerosis/frontotemporal dementia

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Abstract

TDP-43 nuclear depletion and concurrent cytoplasmic accumulation in vulnerable neurons is a hallmark feature of progressive neurodegenerative proteinopathies such as amyotrophic lateral sclerosis (ALS) and frontotemporal dementia (FTD). Cellular stress signalling and stress granule dynamics are now recognized to play a role in ALS/FTD pathogenesis. Defective stress granule assembly is associated with increased cellular vulnerability and death. Ras-GAP SH3-domain-binding protein 1 (G3BP1) is a critical stress granule assembly factor. Here, we define that TDP-43 stabilizes G3BP1 transcripts via direct binding of a highly conserved cis regulatory element within the 3’UTR. Moreover, we show in vitro and in vivo that nuclear TDP-43 depletion is sufficient to reduce G3BP1 protein levels. Finally, we establish that G3BP1 transcripts are reduced in ALS/FTD patient neurons bearing TDP-43 cytoplasmic inclusions/nuclear depletion. Thus, our data suggest that, in ALS/FTD, there is a compromised stress granule response in disease-affected neurons due to impaired G3BP1 mRNA stability caused by TDP-43 nuclear depletion. These data implicate TDP-43 and G3BP1 loss of function as contributors to disease.

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Abbreviations: ALS = amyotrophic lateral sclerosis; DIV = days in vitro; EMSA = electrophoretic mobility shift assay; FBS = fetal bovine serum; FTDP = frontotemporal dementia; FTLD = frontotemporal lobe degeneration; GTEx = Genotype-Tissue Expression project; hnRNP = heterogeneous nuclear ribonucleoprotein; iPSC = induced pluripotent stem cell; LLPS = liquid-liquid phase separation; NLS = nuclear localization sequence; ORF = open
reading frame; RBD = RNA binding domain; RRM = RNA recognition motif; smFISH = single molecule fluorescent in situ hybridization; SNP = single nucleotide polymorphism; UTR = untranslated region;

**Introduction**

TDP-43 is a ubiquitously expressed, mainly nuclear, RNA/DNA binding protein belonging to the heterogeneous nuclear ribonucleoprotein family (hnRNPs). It is a major regulator of RNA metabolism and is involved in several steps of RNA regulation (mRNA transport, nucleocytoplasmic export, stability, transcription, translation, splicing, etc.)¹. Like other hnRNPs, TDP-43 has an N-terminal domain containing a nuclear localization sequence (NLS), an RNA binding domain (RBD) composed of two RNA recognition motifs (RRM1 and RRM2) and a glycine rich C-terminal domain, containing intrinsically disordered regions¹. While the NLS dictates the localization of TDP-43 in the nucleus, the RRMs are essential for specific interaction with target RNAs, and the N and C-terminal domains serve as platforms to mediate TDP-43 self-assembly and interaction with other protein partners, respectively²⁻⁴.

TDP-43 proteinopathies refer to a group of neurological disorders characterized by the pathological accumulation of TDP-43 in the cytoplasm of affected neurons and, less frequently, oligodendroglia⁵. The fatal neurodegenerative diseases ALS and frontotemporal dementia (FTD) are included in this group. FTD, the second most common cause of dementia in those <65 years, is caused by the degeneration of the frontal and temporal lobes (referred to as frontotemporal lobe degeneration or FTLD). ALS shares clinical, neuropathological and genetic overlaps with FTD. In addition to the neuromuscular component of ALS, ~50% of ALS patients exhibit cognitive impairment caused by FTLD, with 15% fulfilling the clinical diagnostic criteria of FTD⁶. TDP-43 nuclear depletion and concomitant accumulation in the cytoplasm, often phosphorylated and as a component of skein-like or round inclusions, in motor and cortical neurons is observed in 97% of all ALS cases and 50% of FTD patients¹. Given these shared clinical and pathological features, ALS and FTD with TDP-43 pathology are now considered to be part of the same disease spectrum¹. However, it remains debated as to whether nuclear depletion of TDP-43 with simultaneous cytoplasmic accumulation, which may be of another conformation, induces a loss or gain of function effect, or some combination thereof.
The interplay between genetics and the environment is suspected to play a major role in the development and/or progression of ALS and FTD\textsuperscript{7}. Recent work places stress granules, a core aspect of the integrated stress response that facilitates cellular recovery from adverse environmental exposures, as a critical component of disease pathogenesis\textsuperscript{8-11}. These micron-sized cytoplasmic foci form concomitant with translational arrest following stress exposure and are proposed to protect and/or sort non-translating mRNAs\textsuperscript{12, 13}. The key protein for stress granule assembly is G3BP1 (Ras-GAP SH3-domain-binding protein 1)\textsuperscript{14, 15}. We have previously demonstrated that G3BP1 protein and transcript levels are decreased as TDP-43 levels are lowered, which compromises cellular viability post-stress\textsuperscript{16-19}. This TDP-43-induced G3BP1 depletion disrupts stress granule dynamics, which can be rescued by the reintroduction of G3BP1\textsuperscript{18}. Recent work reinforces that G3BP1 is central to stress granule formation as it is a tunable switch regulating the liquid-liquid phase separation (LLPS) required for stress granule formation\textsuperscript{16, 18, 20-22}. Thus, we reason that defining the molecular mechanism underlying TDP-43 modulation of G3BP1 is essential to understanding the contribution of stress granule biology to ALS and FTD pathogenesis.

Herein, we demonstrate that nuclear TDP-43 depletion, as observed in the neurons of ALS/FTD patients, as well as expression of pathological TDP-43 variants associated with ALS/FTD, impact G3BP1 mRNA metabolism. Additionally, we have uncovered that while there are two G3BP1 transcripts encoding the same protein, only one of them is regulated by TDP-43. TDP-43 binds and stabilizes the short G3BP1 transcript via an evolutionary conserved sequence. Finally, we demonstrate that this transcript is the predominant transcript expressed in adult human motor neurons and G3BP1 mRNA is decreased in ALS/FTD patient neurons featuring TDP-43 nuclear depletion/cytoplasmic aggregation. Taken together, our work suggests that compromised stress granule assembly due to the loss of TDP-43 nuclear function in G3BP1 stability may be relevant to disease pathogenesis.

Materials and methods

Constructs

Flag-TDP-43\textsuperscript{\textsubscript{ANLSiRes}} was generated using the QuickChange\textsuperscript{\textregistered} II Site-Directed mutagenesis kit (Agilent Technologies) on pCS2-Flag-TDP-43-WT\textsuperscript{23} with the forward primer 5'
CAACTATCCAAAGATAACGCAGCAGCAATGGATGAGACAGATGC-3’ and its complementary reverse to mutate the NLS, as previously published\textsuperscript{24}. Modifications to render TDP-43 plasmids siRNA-resistant were performed with the forward primer 5’-CTTCCTAATTCTAAGCAGAGCGAGCAGAGCCTTGTAGAAGC-3’ and its complementary reverse. The His-G3BP1cDNA-3’UTR construct was subcloned into pTRE-tight which was provided by Dr. Yves Berthiaume (Institut de recherche cliniques de Montreal). His-G3BP1 was generated with the forward primer coding for His-tag: forward 5’-CTAGGATCCGCAATGCACCACCACCACCACGTGATGGAGAAGCCTAGTCC CTG-3’; reverse 5’-ATCGCAGCTAGCGACCTGCGCTGGCGCAAGC-3’) and inserted using BamHI and NheI restriction sites. G3BP1 3’UTR was generated using PCR amplification with the following primers: forward 5’-ATCGCAGCTAGCGACCTGCAATGCACCACCACCACGTGATGGAGAAGCCTAGTCCC CTG-3’; reverse 5’-CTTAGATCCGCAATGCACCACCACCACCGATGGAGAAGCCTAGTCCC CTG-3’; and inserted using BamHI and NheI restriction sites. Deletion of nt319-372 (GRCh38/hg38 chr5: 151,804,204-151,804,257 was made using the following phosphorylated primers: forward 5’-5/phos/CTTAAGCAGTTTATAACAGACTGGGGTCATA-3’ and reverse 5’-5/phos/GCACTAATCGATGCAAACAAAACCTTCACCCCATCTCAC-3’ and inserted using NheI and ClaI restriction sites. Deletion of nt319-372 (GRCh38/hg38 chr5: 151,804,204-151,804,257 was made using the following phosphorylated primers: forward 5’-5/phos/CTTAAGCAGTTTATAACAGACTGGGGTCATA-3’ and reverse 5’-5/phos/GCACTAATCGATGCAAACAAAACCTTCACCCCATCTCAC-3’ and inserted using NheI and ClaI restriction sites.

Cell culture and transfection

HeLa, HEK293, U2OS and Mo3.13 cells were cultured in Dulbecco’s high glucose modified Eagle medium (DMEM, GE Healthcare) supplemented with 10% fetal bovine serum (FBS, Wisent) and 2 mM L-glutamine (Sigma). HeLa Tet-off cells were cultured in DMEM supplemented with 10% tetracycline-free FBS (Wisent) and 2 mM L-glutamine. SH-SY5Y and SK-N-SH cells were cultured in Dulbecco’s high glucose modified Eagle medium/Nutrient Mixture F-12 Ham (DMEM-F12, ThermoFisher Scientific) supplemented with 10% FBS (Gibco), 2 mM L-glutamine and 1% MEM non-essential amino acids (ThermoFisher Scientific). Cells were collected 72 h after transfection with 125 pmol Stealth siRNA using Lipofectamine® 2000 (Invitrogen) and 24h after plasmid transfection using Lipofectamine® LTX and Plus reagent (Invitrogen). siRNA sequences used were: Control (siCTL): 5′ AAGCAAAAGCCCAAGAGAGCCUUUGA-3′, as previously published\textsuperscript{16-18, 25}. 

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**Human neuron differentiation**

Human induced pluripotent stem cells (iPSCs) from a healthy 54 year old female were reprogrammed from fibroblasts via transfection with episomal vectors encoding seven reprogramming factors, as previously described. A polycistronic construct containing a doxycycline-inducible cassette driving expression of the transcription factors neurogenins 1 and 2, Dendra2-tagged TDP-43 (WT or M337V) and eIF3-iRFP was integrated into the *CLYBL* safe harbour locus on chromosome 13. For creation of TDP-43(M337V), the homologous recombination vector used to insert Dendra2 was modified to include the corresponding mutation in the *TARDBP* ORF (c. 1009A>G). All insertions and base pair mutations were verified by PCR and Sanger sequencing. As described in prior work, integration of this cassette enables rapid, robust and consistent differentiations, and produces homogenous cultures of forebrain-like, excitatory, glutamatergic neurons optimal for subsequent biochemical or genetic studies. Cells were harvested in Trizol at DIV10. Lines are verified mycoplasma-free on a monthly basis.

**Immunoblotting**

Proteins were extracted with RIPA buffer (150 mM NaCl, 50 mM Tris pH 7.4, 1% Triton X-100, 0.1% SDS, 1% sodium deoxycholate) with protease inhibitors (10 µg/ml leupeptin, 10 µg/ml pepstatin A, 10 µg/ml chymostatin), separated by SDS-PAGE, transferred to nitrocellulose and blocked with 5% powdered milk in PBS-T (137mM NaCl, 2.7mM KCl, 8mM Na$_2$HPO$_4$, 1.5mM KH$_2$PO$_4$, 1% Tween-20). Membranes were incubated with rabbit anti-TDP-43 (10789-2-AP; Proteintech), mouse anti-Actin (69100; MP Biomedicals), and mouse anti-G3BP1 (sc-81940; Santa Cruz) followed by HRP-conjugated secondary antibodies (Jackson Immunoresearch) and labeled with ECL chemoluminescence (ThermoFisher Scientific). Films were quantified by densitometry using Photoshop (Adobe, CS4).

**Immunofluorescence**

Coverslips were fixed in 1% formaldehyde/PBS, washed with PBS, permeabilized in 0.1% Triton X-100/PBS, and blocked in 0.1% bovine serum albumin/PBS. Coverslips were incubated with primary antibodies: mouse anti-Flag (F1804; Sigma) and rabbit anti-TDP-43 (10789-2-AP; Proteintech) diluted in 0.1% BSA/PBS; washed once with 0.1% Triton X-100/PBS and then twice with 0.1% BSA/PBS. Coverslips were then incubated with secondary
antibodies: donkey anti-mouse Alexa 488 (Jackson Immunoresearch) and donkey anti-rabbit Alexa 594 (Jackson Immunoresearch) diluted in 0.1% BSA/PBS, washed, labelled with TO-PRO-3 iodide (ThermoFisher Scientific), and mounted using ProLong Antifade (ThermoFisher Scientific). Images were collected on a Leica TCS SP5 confocal microscope equipped with 40x (1.25 N.A.) oil objective and the Leica Application Suite imaging software.

**smFISH**

Custom DNA probe sets (20mers) were designed using Stellaris® Probe Designer for total *G3BP1* (targeting *G3BP1* ORF and a portion of 3’UTR) and the long *G3BP1* (targeting a unique region in the long *G3BP1* 3’UTR) (Supplementary Table I). Probes were purchased from Biosearch Technologies, with the total *G3BP1* probes labelled with Quasar 570 and long *G3BP1* probes labelled with Quasar 670. For smFISH, the protocol was adapted from S. Rahman *et al.* Briefly, HeLa cells were plated on glass coverslips and treated with control or TDP-43 siRNA (see above). Coverslips were then washed with 1× PBS, fixed with 4% PFA for 10 min at room temperature, washed twice is wash buffer (10% formamide, 2xSSC) and stored overnight in 70% ethanol at −20°C. Cells were washed in wash buffer and then hybridized with 125 nM of each probe set (total and long) with 40 ug ssDNA/tRNA resuspended in hybridization buffer (10% dextran sulfate, 10% formamide, 2× SSC, 2 mM Vanadyl Ribonucleoside Complexes, 0.1 mg/ml BSA) for 3 h in the dark at 37°C in a humid hybridization chamber. Cells were then washed twice with wash buffer at 37°C for 30 min each. The second wash was supplemented with 1:10000 Hoechst. Cells were washed twice with 1x PBS and mounted with ProLong Gold antifade reagent. Images were acquired using a confocal Leica TCS SP5 MP microscope at 63× oil (1.5x zoom) objective and using a single z-plane focused on the channel for nuclear Hoechst. At least 3 - 5 frames were captured per condition and repeated for three biological replicates. 15 cells were counted per experiment for each condition using ImageJ/FIJI. The Region of Interest tool was used to draw the boundary of each cell and foci per cell were counted for each channel using the Find maxima tool at >80 prominence to generate a mask of foci. A nuclear mask was generated in the Hoechst channel by adjusting the threshold. Colocalization between magenta and green foci, and the nucleus, were calculated using the masks generated and the Object based methods in the JACoP plugin in ImageJ. Foci were plotted using Prism and statistical analysis was performed using an unpaired t-test with Welch correction.
Cell lines qRT-PCR

RNA was extracted using RNAeasy Minikit (Qiagen) and treated with DNase I (Qiagen) according to the manufacturer’s instructions. Equal amounts of RNA were reverse transcribed using the QuantiTect Reverse Transcription kit (Qiagen). The QuantStudio 7 Flex Real-Time PCR System (Life Technologies) was used for qPCR. PrimeTime Standard qPCR assays (IDT) were: GAPDH (targeting exon 1-2) Hs.PT.39a.22214836; G3BP1 total (targeting exon 6 and 7) Hs.PT.58.20396264; G3BP1 long forward: 5’TCTTACTGGACACTCAACCTTG-3’; G3BP1 long reverse: 5’TGCCATAACCTTTTGACTTCATG-3’; G3BP1 long detection probe 5’-56-FAM/AGCTTCCCC/Zen/AGTGCTTTTCTGTCA/3IABkFQ-3’ 18S forward: 5’-CCAGTAAGTGGTCPATAAG-3’; 18S reverse: 5’-GGCCCTCAATAACACCTCCCA-3’; 18S detection probe 5’-56-FAM/TGCGTTGAT/Zen/TAAGTCCCTGCCCT/3IABkFQ-3’. Data was analyzed using the 2-ΔΔCt method. The genes of interest were standardized to the geometric mean of 2 housekeeping genes (GAPDH and 18S).

RNA immunoprecipitation

HeLa cells were lysed in 50 mM Tris, pH 7.5, 150 mM NaCl, 0.5% NP-40, 1mM MgCl₂, with protease inhibitors and RNAse inhibitors, triturated subsequently through 18G and 25G needle syringes, incubated for 30 min at room temperature, centrifuged at 13 000g and the supernatant collected. Aliquots of 10 mg of pre-cleared lysate were immunoprecipitated at 4°C for 6 h with rabbit anti-TDP-43 (10789-2-AP; Proteintech) or rabbit anti-Flag as control IgG (F1804; Sigma) pre-bound to Protein G Dynabeads (ThermoFisher Scientific). Immuno-precipitates were treated with DNase (Qiagen) and RNA was recovered with TRIzol (Invitrogen) and chloroform (Fisher Scientific). Equal amounts of RNA were reverse transcribed using the QuantiTect Reverse Transcription kit (Qiagen). cDNA was amplified by standard PCR with the following primers: total G3BP1 forward 5’-AAGCCCCCTCCACTCCAA -3’; total G3BP1 reverse 5’-TAATCGCCTTCGGGACCTG-3’ (targeting exon 12); long G3BP1 forward 5’-CAGTGAGGTGCCTCCGATA-3’; long G3BP1 reverse 5’-CCAGAACCATGGGAGACACCT-3’; CAMKII forward 5’-CCACAGGGGCTTTAGGAGA-3’; CAMKII reverse 5’-GCTGCTGCGCTTTTGTGA-3’; HSPA1A forward 5’TGAAGGAGACAGGCGAAAGGAA-3’; HSPA1A reverse 5’TGCACTCAGTATAGAAACAGGAGA-3’.
Luciferase assays

HeLa cells were transfected with siRNA for 48 h, and then subsequently transfected using FuGENE (Promega) for 24 h with reporter plasmids including the promoter and 3’ untranslated region (UTR) of human G3BP1 (SwitchGear Genomics) and relevant controls, as recommended by the manufacturer (for promoter assay: GAPDH and a vector containing random sequence, R01; for 3’ UTR: GAPDH, HDAC6 and vector containing random sequence, R03). The LightSwitch Assay (Promega) reagents were added according to the manufacturer’s instructions and luciferase activity was assessed with a Synergy H4 Hybrid Multi-Mode Microplate reader (Biotek).

RNA stability assay

HeLa Tet-off cells were subjected to siRNA using Lipofectamine 2000 (Invitrogen) for 48 h then transfected for 24 h with 1 µg His-G3BP1 3’UTR constructs using Lipofectamine LTX and Plus reagent (Invitrogen). To determine mRNA stability, HeLa cells were treated with 1 µg/ml doxycycline (Sigma) for 0.5, 1, 1.5, 2, 4, or 6 h prior to RNA extraction with the RNeasy Minikit (Qiagen). Equal amounts were reverse transcribed via the QuantiTect Reverse Transcription kit (Qiagen), according to the manufacturer’s instructions. The QuantStudio 7 Flex Real-Time PCR System (Life Technologies) was used for qPCR. PrimeTime Standard qPCR assays (IDT) for His-G3BP1 forward 5’-ACGTGATGGAGAAGCCTAGT-3’; His-G3BP1 reverse 5’-TGACATCACCCTCTGTGGATTTT-3’; His-G3BP1 detection probe 5’-56-FAM/CGGGCGGGA/Zen/ATTTGTGAGACAGTA/3IABkFQ-3’; 18S forward 5’-CCAGTAAGTGGGTCATAAAG-3’; 18S reverse 5’-GGCCTCACTAAACCATCCAA-3’; 18S detection probe 5’-56-FAM/TGCGTTGAT/3IABkFQ-3’.

C. elegans RT–qPCR

Some strains were provided by the CGC, which is funded by NIH Office of Research Infrastructure Programs (P40 OD010440). RNA from N2 and tdp-1(ok803) strains was extracted using TRIzol (Invitrogen) and chloroform (Fisher Scientific). Equal amounts of RNA were reverse transcribed using the QuantiTect Reverse Transcription kit (Qiagen). cDNA was amplified with the following primer sets: ama-1: Ce02462726_m1 (targeting exon boundary 12-13), gtpb-1: Ce02458711_g1 (targeting exon boundary 5-6).
RNA–pull down

Lysates were prepared in pulldown buffer (10 mM Tris pH 7.4, 50 mM NaCl, 0.5% Triton X-100) with protease inhibitors (10 µg/ml leupeptin, 10 µg/ml pepstatin A, 10 µg/ml chymostatin) and subsequently triturated with a 25G needle. Protein G Dynabeads (ThermoFisher) were prepared according to the manufacturer’s instructions. 100 µg of lysates or 100 ng of TDP-43 recombinant protein (NM_007375, OriGene) was incubated with 30 pmol of biotin-labelled RNA for 1 h at 4°C. Proteins were eluted with 2.5× Laemmli buffer and immediately analysed by immunoblotting with rabbit anti-TDP-43 (10789-2-AP; Proteintech) and rabbit anti-hnRNPL (156682; Abcam). Probes: positive control GRN (nt262-288): biotin-UGUGUGUGUGUGCGCGUGUGUGUG; negative control (AC)_{12}: biotin-ACACACACACACACACACACAC; G3BP1 (nt334-358): biotin-UUUUUUGUGUGUAAUGUGUGUG.

Recombinant protein expression and purification

The RRM1 and RRM2 domains of TDP-43 (aa 102-269) in the WT and F147L/F149L mutant forms were subcloned from the pCS2 vector\textsuperscript{23} to the pET21b vector (Novagen) modified to include a TEV cleavage site, and the resulting plasmids were transformed into \textit{E. coli} host strain BL21 (DE3). The bacteria were grown at 37°C in Luria-Bertani (LB) media, and protein expression was induced with 1 mM isopropyl-\textbeta-D-1-galactopyranoside (IPTG) for 4 h at 30°C. The cells were harvested by centrifugation and resuspended in lysis buffer (25 mM HEPES pH 8.0, 500 mM NaCl, 1 mM DTT) supplemented with 0.15% (w/v) Protease Inhibitor Cocktail (Sigma-Aldrich). The cells were lysed by French press, sonicated for 10 sec, and centrifuged at 11,000 g for 30 min at 4°C. The supernatant was incubated at 4°C for 1 h with Ni-charged IMAC Sepharose 6 Fast Flow (GE Healthcare). The resin was then washed three times with His-A buffer (lysis buffer + 20 mM imidazole). The bound His_{6}-tagged TDP-43\textsubscript{102-269} fusion protein was eluted from the washed resin by two 10-min incubations at room temperature with His-B buffer (lysis buffer + 30 mM imidazole) followed by centrifugation. The supernatant containing the protein of interest was dialysed overnight at 4°C in FPLC-A buffer (25 mM HEPES pH 8.0, 100 mM NaCl, 1 mM DTT) supplemented with TEV protease (kindly provided by J.G. Omichinski, Université de Montréal). The retentate was then applied to a Q Sepharose High-Performance column (GE Healthcare; 60-mL bed volume) equilibrated with FPLC-A buffer. The protein was eluted from the column using a gradient (from 0% to 50% over 450
mL) of FPLC-B buffer (FPLC-A + 1 M NaCl). The pooled fractions containing the protein of interest were dialyzed overnight at 4°C in storage buffer (25 mM HEPES pH 8.0, 100 mM NaCl, 2 mM DTT, and 20% glycerol) and the protein was stored at -80°C. The purity of the protein (>98%) was assessed by Coomassie-stained SDS-PAGE.

RNA preparation

The TDP-43 binding sequence of human G3BP1 3'-UTR (G3BP1-RNA32: 5'-TTGTGTGTTAATGGTGTGTGCTCCCTCTCCA-3') was first cloned into the pARiBo4 plasmid. After plasmid linearization with EcoRI, the ARiBo-fusion RNA was in vitro transcribed for 3 h at 37°C using standard conditions. The RNA was then purified using the ARiBo affinity purification method under non-denaturing condition, as previously described. The purified RNA was concentrated with Amicon Ultra-4 centrifugal filter devices (Millipore), and exchanged in H2O. For radioactive labeling of the RNA, a phosphatase alkaline treatment was first performed for 1 h at 37°C, followed by inactivation of the enzyme at 65°C for 15 min. The RNA was then 5'-end-labeled with γ-(32P) ATP (PerkinElmer) using T4 polynucleotide kinase (New England Biolabs) according to the manufacturer’s instructions and then further purified by 15% denaturing gel electrophoresis. After gel extraction, the 32P-labeled RNA was resuspended in TE buffer (10 mM Tris pH 7.6, 1 mM EDTA) and stored at -20°C.

Electrophoretic mobility shift assay (EMSA)

The 32P-labeled RNA was first refolded by heating 2 min at 95°C and snap-cooling on ice for 5 min. The protein samples were diluted in EMSA buffer (50 mM Tris pH 7.5, 50 mM NaCl, 0.05% NP40, 2 mM DTT, and 20% glycerol), and the concentrations were adjusted to span from 0.01X to 5X of the estimated \( K_d \). The binding reactions (20 μL) were initiated by mixing 10 pM of 32P-labeled RNA with the diluted protein and incubating at 4°C for 30 min. For each \( K_d \) determination, binding reactions were loaded on a 10% native polyacrylamide gel (37.5:1 polyacrylamide/bisacrylamide) and run in Tris-Glycine buffer (25 mM Tris-base, 200 mM glycine) at 200 V for 2 h, with active water cooling in a cold room. The gels were then dried and exposed overnight to a storage phosphor screen. The 32P-labeled RNA was visualized with a Bio-Rad Personal Molecular Imager and band intensities were quantified using the ImageLab software (version 5.2.1, Bio-Rad).
concentration, and the data fitted to the Hill equation by non-linear regression analysis within the OriginPro 2015 software (OriginLab). Four independent $K_d$ determination experiments were performed for the WT TDP-43$_{102-269}$. The reported $K_d$ reflects the average values and the standard deviations from these multiple experiments.

**Sciatic axotomy and immunofluorescence**

The use of animals and all procedures were performed according to the guidelines of the Canadian Council on Animal Care and were approved by the CRCHUM Institutional Committee for the Protection of Animals. Female C57BL/6 mice (6-8 weeks, ~20 g) were purchased from Charles River Laboratories (Kingston, NY, USA). Our experiments were based on a previously published protocol for axotomy$^{37}$ with modifications. Mice were weighed and anesthetized by exposure to 1.5 l/min oxygen and 4% isoflurane. After loss of limb reflexes, animals were transferred to a mask system and maintained with 1 l/min oxygen and 2% isoflurane. For a medial axotomy, mice were shaved and the site of surgery sterilized. The right sciatic nerve was exposed with an incision 1 cm below the exit from the pelvic bone. The nerve was then cut 1 cm distal from the exit point of spinal nerve roots. A surgical sterile sponge soaked in 5% fluorogold (Fluorochrome, LLC) in sterile saline was deposit at the site of the nerve cut to enable visualization of injured motor neurons post-injury. Mice were allowed to recover in a clean heated cage and allowed free access to food and water. Pain was managed with buprenorphine injection just prior to surgery, followed by a slow release formulation injected 5 h post-surgery. Neurobehavioral assessments based on a previously published scale$^{37, 38}$ were conducted at days 1, 3, 5 and 7 post injury. Following transcardiac perfusion with saline and 4% paraformaldehyde, tissues were cryopreserved, and subsequently sectioned, then permeabilized and blocked with 0.5% Tween-20 and 5% normal donkey serum (Jackson ImmunoResearch, 017-000-121) for 45 min at room temperature. Primary antibody incubations were performed in 0.3% Tween-20 in PBS overnight at 4°C, followed by an appropriate fluorescently conjugated secondary antibodies against the desired species (Jackson ImmunoResearch). Antibodies used were mouse anti-TDP-43 (1:500, R&D, MAB7778), rabbit anti-G3BP1 (1:4500, Proteintech, 13057-2-AP), and guinea pig anti-FluoroGold (1:750, NM-101 FluGgp, Protos Biotech Corp.). Images were collected using a confocal microscope (SP5; Leica) equipped with LAS AF software (Leica) for acquisition at 63x. Adobe Photoshop CC 2018 was used for quantification of fluorescence intensity. A line was drawn across the longest axis of the neuron and the highest intensity along that line was recorded. This maximum
intensity of a Fluorogold-positive neuron was expressed relative to the maximum intensity mean of 10 Fluorogold-negative neurons. 30 neurons (10 neurons from 3 mice each) were quantified. Importantly, all compared images were acquired with the same microscope settings.

**RNAscope and immunofluorescence on patient autopsy material**

Autopsy tissues from ALS and ALS/FTLD cases were collected in accordance with the local ethics review board at Sunnybrook Health Sciences Centre, Toronto. Orbitofrontal cortex (Broadmann Areas 11 and 47) from four sporadic ALS cases with FTLD, as assessed by presence of TDP-43 pathology (ALS/FTLD-TDP-43), and four sporadic ALS cases without FTLD were used for the study (Supplementary Table II). Formalin-fixed, paraffin-embedded (FFPE) tissue from the orbitofrontal cortex of each case was trimmed and cut into 6 µm sections using a microtome. Slides were dry oven-baked for 20 min at 60°C followed by deparaffinization with fresh xylene and dehydration with sequential ethanol dilutions in water (100%, 95%, 75%, 50%). Endogenous peroxidase was inactivated by RNAscope Hydrogen Peroxide Solution (Advanced Cell Diagnostics, 322335) and antigen target retrieval was achieved by heating the samples in RNAscope Target Retrieval reagent (Advanced Cell Diagnostics, 322000) at 100°C for 30 min before the sections were pretreated with RNAscope Protease Plus (Advanced Cell Diagnostics, 322331) at 40°C for 30 min. Sections were then hybridized with Human G3BP1 RNAscope Probes (Advanced Cell Diagnostics, 567861) at 40°C for 2 h and the resulting signals were amplified and developed with RNAscope 2.5 HD Detection Kit-Red (Advanced Cell Diagnostics, 322360) according to the manufacturer’s instructions. After hybridization, sections were washed with PBS-TX (PBS, 0.15% Triton X-100) and blocked with 4% normal donkey serum in PBS-TX at RT for 1 h. Blocked sections were incubated with TDP-43 antibody (1:1000, Proteintech, 10782-2-AP) at 4°C overnight. After 3 x 5 min washes in PBST, secondary incubation was performed at ambient temperature with donkey α-rabbit 488 Alexa Fluor secondary antibody (1:500, Invitrogen, A-21206). Finally, slides were washed 3 x 5 min with PBS-TX and once with 0.1 M phosphate buffer pH 7.4 (0.08 M sodium phosphate dibasic and 0.02 M sodium phosphate monobasic) prior to coverslipping with ProLong Gold antifade reagent with 4’,6-diamidino-2-phenylindole [DAPI] (Invitrogen, P36931).
Micrograph acquisition and quantification

Fluorescence micrographs of orbitofrontal cortex sections from ALS without FTLD (n=4) and ALSFTLD (n=4) were captured using a Leica DMI6000B microscope with a 100X objective lens on the Volocity Acquisition Suite (v6.3, Perkin Elmer). For each section, a 2-dimensional z-stack projection was produced from 25 micrographs captured at a depth of 0.25 µm through the z-axis. To allow for the quantification of G3BP1-positive mRNA granules (magenta channel) in each neuron with or without TDP-43 pathology (green channel), the green channel intensity was sufficiently enhanced to reveal cell boundaries. The number of G3BP1-positive mRNA granules were quantified from 25 randomly selected neurons demonstrating normal nuclear TDP-43 in the orbitofrontal cortex of ALS without FTLD (n=4) and ALS/FTLD-TDP-43 (n=4) (100 neurons total from each condition). The same sampling strategy was employed resulting in the quantification of G3BP1-positive mRNA granules from an additional 100 neurons demonstrating TDP-43 pathology in ALS/FTLD-TDP-43 cases (n=4). Statistical analysis comparing the mean number of G3BP1-positive mRNA granules in neurons with normal nuclear TDP-43 in ALS without FTLD-TDP-43 pathology and ALS/FTLD-TDP-43 versus neurons with nuclear depletion and cytoplasmic TDP-43 aggregation in ALS/FTLD-TDP-43) was performed using paired one-way ANOVA (p<0.0001).

RNA-seq analysis

The RNA-sequencing data from human frontal cortex and cerebellum39 (GEO accession: GSM1642314; SRA study: SRP056477) were downloaded and run in a bioinformatics pipeline using the Compute Canada clusters. Alignment was performed using HiSAT240 against reference genome Hg38. Read counts were obtained with HTSeq-count41 and differential expression analysis were performed with the Bioconductor R package DESeq242. Data were normalized using DESeq2’s median of ratios method. Polyadenylation site usage in human lumbar spinal motor neurons43 (GEO accession: GSE103225; SRA study: SRP116386) was determined with QAPA44.

Statistics

Data were graphed and analyzed using Prism version 6.00 (GraphPad Software). Statistical tests are stated in the figure legends. Data were compared via two-tailed paired and unpaired t
test, Mann-Whitney, one-way and two-way ANOVA with statistical significance established at p<0.05.

Data availability

The data that support the findings of this study are available from the corresponding author, upon reasonable request.

Results

TDP-43 nuclear depletion and disease-associated TDP-43 variants decrease G3BP1 levels

We have previously demonstrated that depletion of TDP-43 results in a downregulation of G3BP1 protein levels[16]. As ALS/FTLD cases predominantly display a cytoplasmic mislocalization of TDP-43 concomitant with its nuclear depletion, we investigated whether cytoplasmic-restricted TDP-43 could rescue G3BP1 protein levels. To this end, we expressed an siRNA-resistant cDNA encoding TDP-43 with an inactivated nuclear localization signal (FLAG-TDP-43ΔNLS) following siRNA-mediated depletion of endogenous TDP-43 in HeLa cells. Immunofluorescence analysis confirmed the cytoplasmic expression of the exogenous construct FLAG-TDP-43ΔNLS in comparison to mock transfected control cells (Supplementary Figure 1A, B), as previously reported[24]. Expression of FLAG-tagged TDP-43ΔNLS in the context of endogenous TDP-43 depletion yielded a ~50% decrease in G3BP1 protein levels (siCTL + mock vs siTDP-43 + TDP-43ΔNLS, P = 0.0029) comparable to G3BP1 levels in TDP-43 siRNA treated cells co-transfected with empty plasmid (Fig. 1A, B). Thus, cytoplasmic-restricted TDP-43 does not rescue G3BP1 protein levels.

Overexpression studies of TDP-43 ALS-associated mutations have reported defects in stress granule disassembly and/or cytoplasmic aggregation[45, 46]. However, this is a phenotype frequently observed in studies involving the overexpression of aggregation prone RBPs. Thus, we opted to evaluate the impact of a familial ALS-causing mutation (N352S) genome edited onto both endogenous TDP-43 alleles in SH-SY5Y cells[47]. Consistent with Melamed et al[47], we noted a 30% decrease in TDP-43 protein levels (P = 0.0334) which correlated with a 30%
reduction in G3BP1 protein levels (Fig. 1C, D; \( P = 0.0393 \)). This reduction was also recapitulated at the mRNA level (Supplementary Figure 1D; \( P = 0.0413 \)). In addition, expression of TDP-35, an N-terminal truncated TDP-43 variant generated via alternative transcriptional start codon usage that is used more often in ALS patients and is predominantly localized in cytoplasmic granules\(^{48}\), also compromised G3BP1 protein levels (Fig. 1E, F, Supplementary Figure 1C; \( P = 0.0065 \)). Here, we also noted that endogenous TDP-43 levels were reduced 40\% (\( P = 0.0489 \)). Thus, expression of two pathological TDP-43 variants, which are associated with decreased endogenous TDP-43 levels, resulted in reduced G3BP1 protein.

**TDP-43 uniquely modulates one of two protein-encoding G3BP1 transcripts**

According to the Genotype-Tissue Expression (GTEx) project, two transcripts encode the same G3BP1 protein (ENT00000394123.7 and ENST00000356245.7), the former of which has only recently been annotated. The two transcripts primarily differ in the size of their 3'UTRs (8,901nt vs. 1,466nt), hereafter called the long and short G3BP1 transcript, with the first 1,466nt being shared between the two isoforms (Fig. 2A). Using primer pairs to the distal 3’ end of the longer transcript, we confirmed the presence of the longer 3’UTR-containing transcript to varying amounts in six different cell lines via qRT-PCR (Fig. 2B).

Our previous work indicated that siRNA-mediated depletion of TDP-43 decreased G3BP1 mRNA levels\(^{16}\), but these studies were performed using probe sets targeting exonic G3BP1 sequence, and therefore were unable to differentiate between long and short G3BP1 isoforms. To address the question of whether TDP-43 regulates one or both transcripts, we performed single molecule fluorescent in situ hybridization (smFISH), a quantitative method to directly visualize and count transcripts. Specifically, we used smFISH probes against a shared sequence within both G3BP1 mRNA isoforms to target the total pool of G3BP1 transcripts versus probes against a unique region in the 3'UTR of the long G3BP1 transcript. This was performed on Hela cells treated with control, G3BP1, and TDP-43 siRNA (Fig. 2C, Supplementary Figure 2). Notably, the specificity of the smFISH probes was validated by the significant loss of signal in siG3BP1 cells compared to siControl (Fig. 2C, S2A). In siControl, total G3BP1 foci were seen to be significantly more abundant than the long G3BP1 foci (5.7 fold), demonstrating that the long transcript is not the major G3BP1 mRNA isoform (Supplementary Figure 2C).
corroboration, only ~13% of the total G3BP1 foci (magenta) counted colocalized with long G3BP1 foci (green) (Supplementary Figure 2D). Interestingly, we observed a significant reduction (~2.7 fold, P < 0.0001) in the number of total G3BP1 foci in siTDP-43 cells compared to siControl, whereas no significant difference was found for the long G3BP1 foci (Fig. 2C, 2D). Moreover, we observed that ~75% of the long G3BP1 foci (green) colocalized with the total G3BP1 foci (magenta) (Supplementary Figure 2D). We expect that the 25% of non-colocalized foci account for false negatives/positives as well as the orientation of the transcript in the z-plane. Given this data, we can confidently extrapolate that the abundance of the short G3BP1 transcript can be represented by the long foci subtracted from the total foci count. Herein, we show the short G3BP1 transcript specifically to be significantly reduced (~3.7 fold) as a consequence of reduced TDP-43 levels (P < 0.0001, Fig. 2D). We further confirmed this result in several cell lines by separately assessing total and long G3BP1 isoforms via qRT-PCR. Downregulation of TDP-43 using siRNA in HeLa, SK-N-SH and SH-SY5Y lowered total G3BP1 levels (HeLa: P = 0.0222; SK-N-SH: P = 0.0067; SH-SY5Y: P = 0.0271, consistent with our previous results16, 18 (Fig. 2E). However, using a probe set to uniquely identify G3BP1 long transcripts, we found that the longer isoform was not significantly affected by TDP-43 knockdown in all three cell types tested (Fig. 2F-G). Collectively, these data indicate that it is the shorter G3BP1 transcript that is modulated by TDP-43.

TDP-43 stabilizes the short G3BP1 isoform via its 3’UTR

Since TDP-43 depletion is associated with a downregulation of the short G3BP1 transcript, we first sought to determine whether this was mediated by the promotor or the 3’UTR. Using a reporter assay in which the promotor and short 3’UTR were each fused to an optimized Renilla reporter gene (RenSP), siRNA-mediated depletion of TDP-43 reduced steady state luciferase activity of the G3BP1 3’UTR construct by 44% compared to control (GAPDH, Fig. 3A, P = 0.036). In contrast, the luciferase activity of the G3BP1 promotor reporter was not significantly changed compared to controls (Supplementary Figure 3). mRNA abundance is frequently governed by 3’UTR elements linked to mRNA stability. To evaluate whether TDP-43 stabilizes G3BP1 mRNA, we employed a doxycycline-repressible transcript containing His-tagged G3BP1 cDNA fused to the short G3BP1 3’UTR. This approach allowed us to specifically determine exogenous G3BP1 mRNA decay without altering total cellular metabolism (as is observed with Actinomycin D treatment)49. In TDP-43-depleted conditions, the exogenous
G3BP1 transcript was four-fold less stable than in siControl treated cells (t1/2 siCTL: 4.1 ± 0.4 h vs. t1/2 siTDP-43: 1.4 ± 0.1 h, P = 0.0031; Fig. 3B, C). Thus, TDP-43 is required to stabilize the short G3BP1 transcript.

The 3’UTR of G3BP1 harbors a highly conserved UG-rich regulatory element

TDP-43 preferentially binds UG-rich sequences3, 50, which are often associated with mRNA stability51, 52. In order to better characterize TDP-43-dependent stabilization of G3BP1, we analyzed the human short 3’UTR using RBPmap, which predicts RBP binding sites53. A cluster of 13 potential TDP-43 binding sites were identified, located within nt334-358 of the 3’UTR of both transcripts (GRCh38/hg38 chr5: 151,804,220-151,804,244; Fig. 4A). Nucleotide alignment of this sequence across 13 different species indicated an element at 3’UTRnt341-357 (GRCh38/hg38 chr5: 151,804,227-151,804,243) that is highly conserved across all species examined despite overall degeneration of the 3’UTR between species (Supplementary Table III). MEME analysis54 confirms the conservation of UG-repeats in this 16-nt element (Fig. 4B). TDP-43 and its orthologs are structurally conserved throughout evolution, especially within the RBD50. Moreover, G3BP1 is also conserved in C. elegans, with only one protein-encoding transcript, corresponding to the shorter isoform. Consistent with our data in human cell lines, tdp-1 null worms55 exhibit a 2-fold decrease of gtbp-1 (G3BP1 orthologue) mRNA compared to N2 control worms (Fig. 4C; P = 0.0246). Thus, G3BP1 mRNA contains a highly conserved UG-rich cis regulatory element in its 3’UTR that is functionally regulated by TDP-43.

TDP-43 stabilizes G3BP1 mRNA via a conserved 3’UTR regulatory element

To investigate the relevance of the identified conserved sequence, we revisited the luciferase reporter assay using a construct where a region including the regulatory element was deleted (G3BP1 3’UTRΔ319–372; Fig. 4D). Unlike the intact short G3BP1 3’UTR, or the positive control HDAC6, siRNA-mediated depletion of TDP-43 had no effect on the steady state luciferase activity of G3BP1 3’UTRΔ319–372 (Fig. 4E). Moreover, using a doxycycline-repressible version
of this construct with only the 16-nt conserved element deleted (G3BP1 3’UTR^341-357, Fig. 4D), we determined that the half-life of this exogenous transcript was comparable in the presence and absence of TDP-43 (4.6 ± 1.7 h vs 3.9 ± 0.8 h, P = 0.58; Fig. 4F, G). The data obtained here indicate that the identified 16-nt UG-rich regulatory element is required for TDP-43-mediated stabilization of the short G3BP1 transcript.

TDP-43 directly binds the conserved element with high affinity

To determine if TDP-43 mediates stabilization of G3BP1 mRNA via binding to the transcript, we performed immunoprecipitation of endogenous TDP-43 followed by RT-PCR of associated mRNAs. G3BP1 transcripts, including the long transcript, were recovered from TDP-43 immunoprecipitates, as was the positive control CAMKII^56 suggesting that other factors such as other RNA-binding proteins or a structural feature may also influence the regulation. HSPA1A served as a negative control^56 (Fig. 5A). Next, in order to determine if the stabilizing effect of TDP-43 on the short G3BP1 transcript was due to binding to the identified conserved element, we performed an RNA pulldown. Using a biotinylated probe (G3BP1 3’UTR^nt319-372) and HeLa whole cell extracts, we retrieved TDP-43 from the lysate, similar to a probe derived from GRN which has been previously reported to be bound by TDP-43^57 (Fig. 4B). The binding was considered specific since an AC-rich probe [(AC)_{12}] was not bound by endogenous TDP-43 but was by hnRNP L, an RBP with a preference for AC-rich sequences^58. To determine if the interaction between the 3’UTR regulatory element of G3BP1 and TDP-43 was direct, the RNA pulldown was repeated using commercially-produced recombinant TDP-43 protein. Both the G3BP1 and GRN probes, but not the (AC)_{12} probe, were bound by recombinant TDP-43 (rTDP-43) (Fig. 5B). Thus, TDP-43 directly binds the region containing the conserved regulatory element in the 3’UTR of G3BP1.

To establish that the binding was via the RBD of TDP-43 and to determine the affinity of the interaction of TDP-43 for the 3’UTR element, we performed an in vitro electrophoretic mobility shift assay (EMSA) using highly-purified components. The RBD of TDP-43 (TDP-43^102-269) efficiently bound a 32-nt RNA encompassing the G3BP1 3’UTR conserved regulatory sequence (G3BP1-RNA^32) and corresponding to G3BP1 3’UTR^nt340-378). This binding was accompanied by formation of two main shifted bands on the gel, as observed in previous binding studies with (TG)_{12} DNA oligomers^59. Given that the G3BP1-RNA^32 contains two GUGUGU sequences that have the potential to serve as independent binding sites, we
speculate that these two bands represent the formation of RNA-protein complexes containing either one or two molecules of TDP-43_{102-269}. Interestingly, the wild-type TDP-43_{102-269} binds with very high affinity to G3BP1-RNA_{32}. In contrast, there is no evidence of RNA binding for the TDP-43_{102-269} protein fragment containing mutations that inactivate the RNA binding pocket (F147L/F149L) at concentrations up to 1 µM (Fig. 5C, D). EMSA experiments performed in multiple replicates with the wild-type TDP-43_{102-269} indicate an average $K_d$ value of 3.1 ± 0.4 nM. Thus, taken together, these data demonstrate a high affinity of TDP-43 for the identified conserved element in the short 3`UTR of G3BP1.

**Injury-induced TDP-43 nuclear efflux correlates with reduced G3BP1 in motor neurons in vivo**

TDP-43 nuclear efflux has been reported in motor neurons of mice subjected to axotomy or permanent ligation\(^{37, 60}\). Thus, as a way to evaluate whether TDP-43 nuclear depletion can impact G3BP1 in vivo, we axotomized C57Bl/6 mice by severing the sciatic nerve just past the sciatic notch, as it exits the pelvic bone (Fig. 6A). The behavioural evaluation of all injured mice was performed using the previously published NBA scoring system\(^{37, 38}\). At day 1, axotomized mice exhibited notable paralysis of the foot with dragging, knuckle walking, and no toe extension on the injured side (Fig. 6B). This phenotype attenuated in the following days post-injury as shown by the significant decrease in the NBA score at day 7 post-injury (Fig. 6B). We confirmed the presence of axotomized motor neurons on the ipsilateral, but not the contralateral, side of the ventral spinal cord using FluoroGold retrograde labelling (Fig. 6C). Investigation of TDP-43 expression/localization in injured neurons labelled with FluoroGold revealed 80% depletion of nuclear TDP-43 at day 7 post-injury consistent with previous reports\(^{37, 60}\). This was accompanied by 70% reduction in G3BP1 expression in these motor neurons compared to the contralateral side (Fig. 6C, D; $P < 0.0001$). To verify the specificity of these observations, we also immunolabelled for FUS (Supplementary Figure 4). After injury, FUS remained localized to the nucleus even at day 7, consistent with previous work indicating that FUS is not regulated by TDP-43\(^{16, 18}\). Taken together, these results demonstrate an in vivo correlation between nuclear TDP-43 depletion and G3BP1 reduction in adult motor neurons.
Short G3BP1 transcript predominates in affected cell types and is reduced in ALS and ALS/FTD neurons

Given that only one of the two G3BP1 transcripts is sensitive to TDP-43 levels, it was imperative to determine the relative abundance of each transcript in cells/tissue of relevance to disease. Analysis of published transcriptomics data from two different brain regions of healthy individuals\textsuperscript{39} revealed a paucity of reads corresponding to the extended 3'UTR region of the long G3BP1 transcript in the frontal cortex, compared to the cerebellum (Fig. 7A), suggesting that the short G3BP1 transcript predominates in the frontal cortex. Analysis of global expression of G3BP1 (i.e. both transcripts) revealed a modest reduction in G3BP1 expression in the frontal cortex compared to the cerebellum, which demonstrated high variability across individuals (Fig. 7B; log\textsubscript{2} FC = 0.30; \(P = 0.04\)). We also examined polyadenylation site usage in published transcriptomics data generated from lumbar spinal motor neurons isolated by laser capture microdissection from healthy individuals\textsuperscript{43}. This analysis clearly indicated the preferential utilization of the proximal polyadenylation site, reflecting that the short G3BP1 transcript is more abundant than the longer isoform in the most ALS-vulnerable neurons (Fig. 7C; \(P = 0.0004\)). Both of these aforementioned datasets also included ALS and/or ALS/FTD cases. However, we were unable to detect a significant difference between G3BP1 3’UTR usage between controls and individuals with ALS or ALS/FTD, presumably due to the relatively low overall abundance of G3BP1 transcripts (data not shown). (Note, this dataset had an average depth of only \textasciitilde28 million reads per sample, and neither study implemented a strategy to enrich for low-abundance transcripts.) Thus, we assessed G3BP1 transcript levels in human neurons differentiated from induced pluripotent stem cells (iPSCs) via Ngn1-2 expression to yield a near-homogenous population of iPSC-derived neurons with properties consistent with glutamatergic, excitatory forebrain-like neurons (i\textsuperscript{3}Neurons)\textsuperscript{28, 61, 62}. These i\textsuperscript{3}Neurons were modified to express a single copy of wild type or mutant (M337V) TDP-43. Via qRT-PCR, we observed that G3BP1 total transcript levels, but not the long transcript, were reduced 49\% in TDP-43\textsuperscript{M337V} expressing i\textsuperscript{3}Neurons compared to TDP-43\textsuperscript{WT} i\textsuperscript{3}Neurons (Fig. 7D, \(P = 0.0272\)). In agreement with our data, this result suggests that it is the shorter G3BP1 transcript impacted by the TDP-43 mutation.

Finally, to directly quantify total G3BP1 transcripts in the context of TDP-43 pathology, we performed quantitative in situ hybridization using RNAscope probes to G3BP1 coupled with immunofluorescence labeling using TDP-43 antibody on orbitofrontal cortices of ALS/FTLD
cases with cortical TDP-43 pathology in comparison with ALS cases without FTLD and no cortical TDP-43 pathology ($n = 4$ each, **Supplementary Table 2**). The number of $G3BP1$ transcripts detected using RNAscope was equivalent in cortical neurons with normal nuclear TDP-43 localization (**Fig. 7E**) in both ALS/FTLD-TDP-43 cases and ALS/no FTLD cases ($14.1 \pm 3$ and $14.4 \pm 4$, respectively, $n = 100$ neurons per group, **Fig. 7F**). In contrast, $G3BP1$ transcript levels were reduced 60% in ALS/FTLD neurons with TDP-43 pathology ($5.9 \pm 2.5$ puncta per neuron; $n = 100$ neurons, $P = 2.5 \times 10^{-35}$; **Fig. 7E, F**), as evidenced by nuclear depletion and cytoplasmic mislocalization of TDP-43 (**Fig. 7E**, indicated with arrows). Taken together, these data are consistent with the concept that $G3BP1$ mRNA is destabilized in the absence of nuclear TDP-43 in disease affected neurons in ALS/FTLD.

**Discussion**

In this study, we demonstrate the mechanism by which the loss of TDP-43 leads to reduced levels of $G3BP1$ mRNA and protein\textsuperscript{16-18}. We found that TDP-43 binds to and regulates the $G3BP1$ short transcript via its 3′UTR, similar to that described for HDAC6, ADD2 and MAPT\textsuperscript{51, 52, 57, 63}. TDP-43 depletion reduces the stability of the most abundant $G3BP1$ transcript, explaining the observed decrease in G3BP1 protein level in TDP-43-depleted cells. In general, cellular stress and pathology are associated with an increase in the proportion of shorter transcripts, suggesting that this TDP-43 regulated transcript may become elevated in adverse conditions\textsuperscript{64-66} and thus outcompete the longer transcript for translation. Moreover, short and long 3′UTRs can be differentially regulated with long 3′UTR transcripts generally being longer-lived\textsuperscript{67}. Several factors including RNA structure, RNA modifications and competitive/collaborative binding with other RBPs or miRNAs can impact translatability and localization of a given transcript. As TDP-43 binds both transcripts, it is possible that these other factors are key elements to differential $G3BP1$ transcript stability and warrant further investigation.

We propose that the mechanism by which TDP-43 stabilizes the short $G3BP1$ transcript is by binding the nascent transcript in the nucleus prior to its export across the nuclear membrane and protecting it from cytoplasmic mRNA decay. This is akin to HuR-mediated mRNA stability, which binds transcripts in the nucleus and protects them from TTP-mediated decay\textsuperscript{68}. Our smFISH data suggests that the long $G3BP1$ mRNA has a higher ratio of nuclear transcripts than the short $G3BP1$ mRNA (**Supplementary Figure 2E**). Thus, one possibility is that the
long transcript is not susceptible to siTDP-43 mediated instability as less of the transcript is present in the cytoplasm, where mRNA decay mainly occurs, compared to the short G3BP1 transcript. However, as mentioned above, further study is required to fully evaluate the mechanisms that could account for differential G3BP1 transcript stability.

In the context of ALS/FTD, TDP-43 is typically found to be cytoplasmic, and thus in theory should be able to stabilize G3BP1 transcripts in the cytoplasm. However, consistent with our data, it is plausible that nuclear TDP-43 depletion as observed in ALS patients means that TDP-43 is not available to nascent G3BP1 transcripts in the nucleus as they are transcribed, and thus these transcripts are not protected from mRNA decay upon transfer to cytoplasm. Another consideration is that cytoplasmic TDP-43 as detected in post-mortem samples is typically in an aggregated state, phosphorylated and insoluble – i.e. it is in a conformational state that precludes its normal physiological function. This is supported by Mann et al[69] who used optogenetics to demonstrate that light-induced TDP-43 aggregates are phosphorylated, insoluble and can be reversed by RNA binding. Moreover, RNA is absent from these TDP-43 aggregates. Perhaps most compelling is that the authors also showed that RNA is not detected in phospho-TDP-43 inclusions in ALS spinal cord neurons or FTLD hippocampal neurons[69]. Taken together, this supports the model that cytoplasmic aggregated TDP-43 is not in a conformational state that can bind mRNA and thus would not be expected to colocalize with or stabilize G3BP1 mRNA in this compartment.

TDP-43 stabilizes G3BP1 transcripts via an evolutionary conserved cis regulatory sequence located within the 3'UTR. In alignment with recent findings of the ENCODE project, the identified regulatory element is UG-rich but not a pure UG cluster[70]. TDP-43 directly binds the regulatory element with a $K_d$ that is 6-23x higher than that reported for small UG-rich RNAs (20-70 nM)[50], but is similar to that reported for a 16-nt (2.8 nM)$^2$ and a 30-nt RNA (5 nM)$^2$ consisting entirely of UG repeats. While the entire G3BP1 3'UTR sequence is poorly conserved amongst the 13 species examined (and 10 additional species, data not shown), the 16-nt element we identified is very highly conserved and functional, as supported by our data in C. elegans where the sequence is 67% conserved. Together, these findings indicate that the regulatory element is necessary and sufficient for TDP-43 mediated stabilization of G3BP1 mRNA. Interestingly, the conserved element we have identified lies within a region that harbors very few SNPs (UCSC genome browser, VarSome), which demonstrates a conservation of the sequence and possibly indicates that deviation from this sequence is poorly tolerated. The few SNPs observed are predicted to be pathogenic and all increase or decrease
the number of UG in the sequence (VarSome). Further work will be needed to define whether any of these SNPs can be exploited to inform on disease onset or progression. Taken together, our data reinforce the evolutionary importance of TDP-43-mediated regulation of G3BP1, and by extension, its cellular function.

Transcriptomics data indicate that G3BP1 is expressed at relatively low levels in neurons (GTEx) and this could suggest that it is subject to very fine/tight regulation. However, our experience with mouse spinal cord shows that motor neurons are the major cell type in which G3BP1 signal is detected (Fig. 6 and unpublished data). We show that the short 3’UTR G3BP1 transcript is the predominant one expressed in disease-vulnerable frontal cortex and lumbar spinal motor neurons and G3BP1 mRNA levels were reduced in ALS/FTLD neurons bearing TDP-43 cytoplasmic inclusions/nuclear clearance. Our data in axotomized motor neurons, where loss of nuclear TDP-43 positively correlated with a sharp decrease in total G3BP1 levels is supportive of this possibility. Finally, near-physiological expression of ALS-associated TDP-43 mutations had the same impact on G3BP1 levels as nuclear TDP-43 depletion suggesting that TDP-43 mutations can act through a loss of function mechanism.

G3BP1 is essential for neuronal homeostasis and survival\textsuperscript{71-74}. Indeed, G3BP1 genetic deletion in 129/sv mice results in widespread neuronal loss, while the same deletion on a Balb/c background yields synaptic and locomotor deficits, with 75% of homozygotes displaying paralysis of at least one limb\textsuperscript{72, 73}. In addition, these mice have impairments in neuronal plasticity and in calcium/glutamate signaling\textsuperscript{72, 75}. G3BP1 is also a known target of certain enteroviruses, such as poliovirus and coxsackievirus B3, which each encode a protease that cleaves G3BP1 and effectively cripples the stress granule response in order to facilitate virus replication in motor neurons\textsuperscript{76-78}. It is noteworthy that poliovirus exhibits tropism for ALS-vulnerable regions (\textit{i.e.} it infects the anterior horns of the spinal cord, brainstem and the motor cortex but not the oculomotor nerve)\textsuperscript{77, 79}. As lytic viruses, they cause selective motor neuron death and, in the case of poliovirus, induce a progressive paralysis (post-poliomyelitis syndrome) similar to that experienced by ALS patients\textsuperscript{80}.

Collectively, our data support that under normal conditions, TDP-43 functions to stabilize G3BP1 transcripts via a highly conserved \textit{cis} 3’UTR regulatory element. In addition, we observed reduced G3BP1 mRNA levels in neurons bearing the TDP-43 pathological signature of ALS/FTLD. The loss of G3BP1 mRNA, and thus the encoded protein that is essential to stress granule assembly, would preclude the launch of the protective neuronal response known
as stress granule formation and draws into question the model that TDP-43 inclusions derive from defective stress granule disassembly. As we have previously published that deficient stress granule assembly contributes to enhanced neuronal vulnerability\textsuperscript{16-18}, our data are consistent with the idea that stabilization of \textit{G3BP1} mRNA and maintenance of stress granule dynamics could be a valid therapeutic avenue.

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**Competing interests**

The authors report no competing interests.
Supplementary material

Supplementary material is available at *Brain* online.

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Figure legends

Figure 1. TDP-43 nuclear depletion and ALS-related species induce G3BP1 down-regulation. (A) HeLa cells were transfected with siTDP-43 or siControl (siCTL), then transfected with empty plasmid (Mock) or Flag-TDP-43ΔNLS and immunoblotted. (B) Quantification via densitometry of G3BP1 protein levels normalized to actin. (C) Protein from SH-SY5Y WT and TDP-43N352S/N352S cells were extracted, and immunoblotted. Quantification via densitometry of (D) TDP-43 and (E) G3BP1 protein levels normalized to actin. (F) HeLa cells were transfected with empty plasmid (Mock) or Flag-tagged TDP-35 and immunoblotted. Quantification via densitometry of (G) TDP-43 and (H) G3BP1 protein levels normalized to actin. Data from 3-5 independent experiments are expressed as the mean fold change ± SEM; Unpaired t test *P < 0.05; **P < 0.01.

Figure 2. TDP-43 depletion only affects the short G3BP1 transcript. (A) Schematic of Genome Reference Consortium Human GRCh38.p12 G3BP1 transcripts. Long transcript has 8,901nt 3’UTR while the shorter version has a 1,466nt 3’UTR. (B) qRT-PCR for total G3BP1, long G3BP1 transcripts and GAPDH of extracted RNA from HeLa, HEK293, SK-N-SH, SH-SY5Y, U2OS and Mo3.13 cells, normalized with GAPDH and 18S, n = 3-4. (C) Representative images of smFISH in HeLa cells treated with siControl and siTDP-43, using probes against total G3BP1 (magenta) and long G3BP1 (green) mRNA. Cell nuclei were counterstained with Hoechst. Yellow foci show co-localization of total and long signal. (D) Total G3BP1 and long G3BP1 were quantified and short G3BP1 was calculated as the number of long foci subtracted from total magenta foci for each cell. Data is expressed as the mean ± SEM, n = 3, N = 45. Unpaired t test, **** P <0.0001, scale bar 25 μm. (E-F) qRT-PCR for total G3BP1 and long G3BP1 transcripts extracted RNA from HeLa, SK-N-SH and SH-SY5Y cells transfected with siControl or siTDP-43 and normalized with GAPDH and 18S. (G) Ratio of the levels of long G3BP1 transcripts:total G3BP1, mean ± SEM, n = 3-5, Unpaired t test *P < 0.05, **P < 0.01.

Figure 3. TDP-43 regulates and stabilizes G3BP1 via its 3’UTR. (A) HeLa cells were transfected with siTDP-43 or siCTL, then co-transfected with the indicated reporter plasmids. Luciferase activity of G3BP1 3’UTR is expressed relative to siCTL cells. GADPH and R03 (random sequences) are used as controls, mean ± SEM, n = 4, Unpaired t test **P < 0.01, ***P
< 0.001. (B) qRT-PCR for His-G3BP1 transcripts, normalized to 18S following doxycycline treatment for 6 h to shut off expression, n = 3. (C) Half-lives of His-G3BP1 transcripts in HeLa-Tet-off cells transfected with siCTL compared to cells transfected with siTDP-43, n = 3, Unpaired t test **P < 0.01.

**Figure 4. Identification of a conserved element in the G3BP1 3’UTR.** (A) Schematic representation of the conserved element identified in the human G3BP1 3’UTR between nt319-372 (GRCh38/hg38 chr5: 151,183,766-151,183,816). The blue lines represent TDP-43 binding sites suggested by RNAmap. (B) Consensus sequence of the conserved sequence obtained from 13 species using the consensus sequence generator MEME. (C) qRT-PCR for gtpb-1 of extracted RNA from N2 (Wild-type control) and tdp-1(ok803) null worms. n = 3, Unpaired t test *P < 0.05. (D) Schematic representation of the G3BP1 3’UTR constructs used. The red rectangle represents the conserved element. (E) HeLa cells were transfected with siTDP-43 or siCTL, then co-transfected with the indicated reporter plasmids. GADPH and R03 (random sequences) are negative controls and HDAC6 is a positive control, mean ± SEM, n = 3-8, Unpaired t test *P < 0.05, **P < 0.01. (F) qRT-PCR for His-G3BP1Δ341-357 transcript, normalized to 18S following doxycycline treatment for 6 hours to shut off expression. (G) Half-life of His-G3BP1Δ341-357 transcript in HeLa-Tet-off cells transfected with siCTL compared to cells transfected with siTDP-43, n = 3.

**Figure 5. TDP-43 directly binds the conserved element.** (A) TDP-43 protein and its associated transcripts were co-immunoprecipitated from HeLa cells homogenates. mRNAs binding to TDP-43 were extracted, reverse transcribed and amplified for G3BP1 transcripts, CaMKII (positive control) and HSPA1A (negative control), showing that TDP-43 binds total G3BP1 transcripts and the longer transcripts. Flag-IP serves as a control to demonstrate specificity of TDP-43 immunoprecipitation. (B) Proteins from whole HeLa cell extracts or TDP-43 recombinant protein (rTDP-43) were pulled down using biotinylated RNA probes containing the TDP-43 binding sequence in GRN (positive control), G3BP1 3’UTRnt319-372, or (4C)12 (negative control). hnRNP L served as a positive control for AC-repeat binding. (C) Typical EMSA performed with 10 pM 5’-[32P]-labeled RNA and increasing concentrations of TDP-43102-269. (D) Typical binding curve using WT TDP-43102-269 (black line) and the F147L/F149L mutant of TDP-43102-269 (red line).

**Figure 6. Changes in TDP-43 and G3BP1 expression in injured motor neurons following medial axotomy.** (A) Schematic of the medial site of axotomy. (B) NBA score of the
axotomized mice at days 1, 3, 5 and 7. Data is expressed as the mean± SEM; Unpaired t test
****P < 0.0001. (C) Representative images of control non-axotomized (ipsilateral) and
axotomized mice (contralateral) spinal cord tissues at day 7, showing injured spinal motor
neurons of the ventral horn using mouse TDP-43, rabbit G3BP1 and guinea pig Fluorogold
antibody. The grey line shows the cell boundary. (D) Fluorescence intensity of nuclear TDP-43
and total G3BP1 for the control non-injured and injured motor neurons at day 7, n = 3 mice
and N = 10 cells per mouse. Data is expressed as the mean ± SEM; Two-way ANOVA ****P
< 0.0001. Scale bar, 10 µm.

Figure 7. G3BP1 short mRNA is destabilized and reduced in patients with TDP-43-
mediated neurodegeneration. (A) Sashimi plots of GEO dataset GSE67196 for human
G3BP1 in cerebellum and frontal cortex of control case, showing reduced reads for the long
G3BP1 3’UTR in the frontal cortex. (B) Comparison of G3BP1 levels (both transcripts) in
human cerebellum and frontal cortex in healthy controls. *P < 0.05. (C) Quantification of
polyadenylation site usage in G3BP1 transcripts in lumbar spinal motor neurons isolated by
laser capture microdissection (GEO dataset GSE103225). Data is expressed as the mean ±
SEM; Unpaired t test ***P < 0.001. (D) qRT-PCR for total G3BP1 and long G3BP1 transcripts
extracted RNA from I3 neurons derived from TDP-43^{M337V} and isogenic control fibroblast,
mean ± SEM, n = 3, Paired t test *P < 0.05. Normalized to GAPDH and 18S. (E) Neurons in
the orbitofrontal cortex of ALS and ALS/FTLD cases showing normal nuclear localization of
TDP-43 (top row, green) or nuclear TDP-43 depletion/cytoplasmic TDP-43 accumulation
(bottom row, green). G3BP1 mRNA (magenta dots) is labelled with RNAscope probes, nuclei
are marked with DAPI (blue). (F) Quantification of G3BP1 mRNA signals in neurons of the
orbitofrontal cortex of ALS and ALS/FTLD with or without TDP-43 pathology (defined as
obvious cytoplasmic accumulations or reduced nuclear TDP-43 levels). Data is expressed as
the mean± SEM; One-way Anova ****P < 0.0001. Scale bar, 10 µm.
Figure 1, Sidibe et al.

190x254mm (300 x 300 DPI)
Figure 2, Sidibé et al.
Figure 4, Sidibé et al.
Figure 5, Sidibé et al.

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