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Heat Shock-induced Phosphorylation of TAR DNA-binding Protein 43 (TDP-43) by MAPK/ERK Kinase Regulates TDP-43 Function

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TAR DNA-binding protein (TDP-43) is a highly conserved and essential DNA- and RNA-binding protein that controls gene expression through RNA processing, in particular, regulation of splicing. Intracellular aggregation of TDP-43 is a hallmark of amyotrophic lateral sclerosis and ubiquitin-positive frontotemporal lobar degeneration. This TDP-43 pathology is also present in other types of neurodegeneration including Alzheimer’s disease. We report here that TDP-43 is a substrate of MEK, a central kinase in the MAPK/ERK signaling pathway. TDP-43 dual phosphorylation by MEK, at threonine 153 and tyrosine 155 (p-T153/Y155), was dramatically increased by the heat shock response (HSR) in human cells. HSR promotes cell survival under proteotoxic conditions by maintaining protein homeostasis and preventing protein misfolding. MEK is activated by HSR and contributes to the regulation of proteome stability. Phosphorylated TDP-43 was not associated with TDP-43 aggregation, and p-T153/Y155 remained soluble under conditions that promote protein misfolding. We found that active MEK significantly alters TDP-43-regulated splicing and that phosphomimetic substitutions at these two residues reduce binding to GU-rich RNA. Cellular imaging using a phospho-specific p-T153/Y155 antibody showed that phosphorylated TDP-43 was specifically recruited to the nucleoli, suggesting that p-T153/Y155 regulates a previously unappreciated function of TDP-43 in the processing of nucleolar-associated RNA. These findings highlight a new mechanism that regulates TDP-43 function and homeostasis through phosphorylation and, therefore, may contribute to the development of strategies to prevent TDP-43 aggregation and to uncover previously unexplored roles of TDP-43 in cell metabolism.

In recent years the role of RNA-binding proteins in neurodegenerative disorders, particularly amyotrophic lateral sclerosis (ALS) and frontotemporal lobar degeneration (FTLD) has become the focus of intense research. The TAR DNA-binding protein (TDP-43) forms cytoplasmic aggregates in almost all ALS and ~50% of FTLD cases (1, 2). TDP-43 pathology is observed in other neurological disorders including Alzheimer’s disease (3, 4). The direct role TDP-43 plays in disease is underscored by >40 ALS/FTLD patient-derived autosomal dominant missense mutations in the TDP-43 gene (TARDBP) (5). TDP-43 inclusions coincide with a dramatic reduction of the normal nuclear TDP-43 detection (2), implying a loss of function upon aggregate accumulation. The link between neurodegeneration and TDP-43 function has not been clearly established and the contribution of TDP-43 aggregation to pathogenesis is poorly understood.

TDP-43 is a highly conserved heterogeneous nuclear ribonucleoprotein (hnRNP) with two canonical RNA recognition motifs (RRMs) of which RRM1 is necessary and sufficient to bind RNA. A low complexity sequence, or prion-like domain at the C terminus mediates protein interactions required for RNA processing (6, 7) and is the main driver of protein aggregation (8). Although TDP-43 shuttles between nucleus and cytoplasm, it is predominantly nuclear (9). In the nucleus, TDP-43 shows a diffuse pattern and recruitment to Cajal bodies and gems (10). Under specific cellular stress conditions, such as oxidative stress, TDP-43 localizes to cytoplasmic stress granules (11), which may contribute to pathological aggregate formation (12).

TDP-43 is essential for development and survival in animal models and cultured cells (13–17). In human cells and mouse brain, TDP-43 binds thousands of transcripts with a strong preference for GU-rich sequences and regulates >600 protein-
coding genes. TDP-43 controls these genes predominantly through splicing regulation (18–20). Although TDP-43 participates in different mechanisms of RNA processing and RNA transport, the regulation of splicing is its best-characterized function (20, 21). In addition, recent findings show that TDP-43 plays a more global role in gene regulation as an inhibitor of cryptic exon inclusion (22).

Despite evidence that TDP-43 controls a large number of genes and is dynamically distributed in cells, the cellular stimuli and factors that regulate TDP-43 function through posttranslational modifications are still mostly unknown. We characterized new TDP-43 phosphorylation sites found in non-pathological settings and sought to determine the mechanisms that control them. Our findings show that activation of the cellular heat shock response (HSR) dramatically increases TDP-43 phosphorylation at Thr-153 and Tyr-155 through the MAPK/ERK kinase MEK, in the absence of increased TDP-43 aggregation or protein clearance. Increased Thr-153/Tyr-155 phosphorylation in the presence of active MEK and phosphomimetic substitutions at Thr-153/Tyr-155 show decreased splicing regulatory function. Moreover, phosphomimetic substitutions at Thr-153/Tyr-155 decrease RNA binding affinity, suggesting that phosphorylation at these sites modulates TDP-43 activity.

Results

The Heat Shock Response Controls TDP-43 Phosphorylation in RRM1—TDP-43 phosphorylation at threonine 153 and tyrosine 155, located in RRM1 (Fig. 1, A and B), was identified by large scale phosphoproteome mass spectrometry analysis of human HeLa cells (23). These modifications were determined under non-pathological conditions. To characterize these previously unexplored phosphorylation sites, we developed a novel phospho-specific polyclonal antibody, p-T153/Y155-TDP-43, to recognize single phosphorylation at Tyr-155 (p-Y155-TDP-43). This antibody showed no detection of the p-T153/Y155-TDP-43-associated signal under control or heat shock conditions (supplemental Fig. S2C). Collectively, these results strongly support the specificity of the newly developed p-T153/Y155-TDP-43 antibody for dual TDP-43 phosphorylation at Thr-153 and Tyr-155, and show the regulation of TDP-43 post-translational modification by the HSR.

To confirm our results and to determine whether phosphorylation at Thr-153/Tyr-155 modifies cellular localization of TDP-43, we characterized the p-T153/Y155-TDP-43 antibody by indirect immunofluorescence analysis. In non-treated cells, the p-T153/Y155-TDP-43-associated signal localized in the nucleolar compartment as dense coil-like structures (Fig. 3). This was observed in different human cell lines, including HeLa, SH-SY5Y (Fig. 3, A and B), U2-OS, and HEK-293 cells (not shown). We confirmed antibody specificity for TDP-43 phosphorylation at Thr-153/Tyr-155 in our immunofluorescence assays by RNAi-mediated knockdown of TDP-43 and competition assays with the Thr-153 and Tyr-155 phosphorylated peptide (amino acids 148–161) as above. Detection of the nucleolar signal was blocked in the presence of the phosphorylated peptide, whereas incubation of the antibody with the non-phosphorylated form of the peptide had no effect (supplemental Fig. S3A). Furthermore, knockdown of TDP-43 via shRNA reduced levels of the nucleolar signal compared with control treated cells (supplemental Fig. S3B). These results suggest that p-T153/Y155 is associated with nucleolar localization of TDP-43 under normal conditions. This is in agreement with previous mass spectrometry detection of TDP-43 in the nucleolar domain of HeLa cells (24). The levels of total TDP-43 recruited to nucleoli appear to be considerably lower compared with the rest of the nucleus, based on the detection of phospho-independent TDP-43 (Fig. 3A). Confocal microscopy analyses of HeLa and SH-SY5Y cells suggest that phosphorylated TDP-43 is part of the dense fibrillar component (DFC) of nucleoli as we observed significant colocalization with fibrillarin, a well established DFC marker (Fig. 3B). Localization of p-T153/Y155 in the nucleolus was maintained following heat shock. Quantification of p-T153/Y155 levels throughout the cell
showed an increase in relative fluorescence after heat shock of 50 and 40% in HeLa and HEK-293 cells, respectively (Fig. 3C, supplemental Fig. S4). Further experiments are required to determine whether the increase in phosphorylation during HSR primarily derives from changes in the nucleolar compartment, relative to the rest of the cell.

**TDP-43 Phosphorylation Is Not Associated with Aggregation, Stress Granules, or Protein Clearance**—We analyzed TDP-43 aggregation following heat shock to determine whether p-T153/Y155 was associated with changes in protein solubility, and to compare p-T153/Y155 with the two previously determined TDP-43 phosphorylation sites Ser-403/404, Ser-409/410, which are tightly linked to aggregation and pathology (25, 26). Heat shock significantly induced misfolding and accumulation of total TDP-43 in the insoluble fraction following protein extraction from cellular lysate (CL) into RIPA (R) and urea (U) soluble fractions. Immunoblot analysis showed a great reduction of total TDP-43 in the RIPA soluble fraction and a corresponding increase in the urea soluble sample after heat shock (Fig. 4A). In contrast, p-T153/Y155 was present exclusively in the RIPA soluble fraction. Recovery from heat shock by incubation at 37 °C reduced the levels of total TDP-43 in the urea fraction, indicating clearance of misfolded TDP-43 and/or chaperone-assisted refolding of the protein. At the same time,
the levels of p-T153/Y155 greatly decreased during recovery. These results are consistent with imaging analysis, which showed no accumulation of p-T153/Y155 into visible aggregates following heat shock (Fig. 3C). Even though total TDP-43 accumulated in the urea fraction following heat shock, we did not observe accumulation of total TDP-43 into cytoplasmic aggregates (Fig. 3C), suggesting that TDP-43 present in the urea fraction after heat shock represents misfolded species that may precede formation of larger, visible aggregates.

To determine whether phosphorylation of Thr-153 and Tyr-155 is associated with pathological inclusions in disease conditions, we analyzed p-T153/Y155 localization in FTLD cases. As positive control for the presence of TDP-43 pathology, we used an antibody recognizing phosphorylated Ser-409/410, which is a well established marker of TDP-43 disease-associated inclusions (25). Immunohistochemical analysis of frontotemporal dementia brain sections showed no significant accumulation of p-T153/Y155 in aggregates (supplemental Fig. S5). These initial observations suggest that, unlike previously identified phosphorylation events (i.e. Ser(P)-403/404 and Ser(P)-409/410), p-T153/Y155 is not linked to the formation of aggregates in FTLD. This is in agreement with our cell-based data suggesting that p-T153/Y155 regulates TDP-43 function and localization and that it is linked to soluble protein under conditions that promote TDP-43 misfolding (Fig. 4A). However, at this time, we cannot exclude the possibility that increased p-T153/Y155 levels, or specific patterns of detection may be observed in different types of TDP-43 proteinopathies.

TDP-43 recruitment to stress granules (SGs) has been observed upon exposure to oxidative and proteotoxic stress (11). To investigate whether phosphorylation mediates recruitment of TDP-43 to SGs upon heat shock, we analyzed the colocalization of p-T153/Y155 and the SG marker fibrillarin in HeLa and SH-SYSY cells as seen by confocal microscopy. C, detection of p-T153/Y155 in HeLa cells upon heat shock compared with control-treated cells. Bars, 10 μm.
phospho-independent TDP-43 to SGs, in agreement with previous reports (27). In summary, our findings indicate that phosphorylation at Thr-153/Tyr-155 is not linked to aggregation or stress granule recruitment of TDP-43. Instead, we find that in conditions that trigger TDP-43 misfolding, such as heat shock, p-T153/Y155 is associated with a soluble conformation of the protein. However, at this time, we cannot exclude the possibility that increased TDP-43 aggregation leads to structures that make the Thr-153/Tyr-155 inaccessible to phosphorylation.

Next, we asked whether increased TDP-43 phosphorylation during HSR occurs upon activation of two pathways that mediate TDP-43 clearance: the ubiquitin proteasome system (UPS) and macroautophagy (autophagy) (28–30). Cells treated with the UPS inhibitor MG132, which induces the accumulation of TDP-43 destined for UPS degradation (28), did not show changes p-T153/Y155 levels (Fig. 4B). Immunofluorescence analysis showed the accumulation of cytoplasmic aggregates of total TDP-43 following MG132 treatment (Fig. 4D), as previously shown (29). However, we observed no colocalization of p-T153/Y155 with these aggregates. We then asked whether p-T153/Y155 mediates autophagy-associated degradation of TDP-43. Cells were exposed to conditions that either increase autophagy (trehalose, serum starvation), or prevent autophagosome-lysosome fusion (thapsigargin). The levels of p-T153/Y155 were unaffected under these conditions (Fig. 4B). Moreover, we observed no colocalization of p-T153/Y155 with microtubule-associated protein light chain 3 (LC-3), a marker of autophagy vesicles (Fig. 4E). These results indicate that phosphorylation at Thr-153/Tyr-155 is not linked to known pathways of TDP-43 clearance.

**MEK Regulates TDP-43 Phosphorylation at Thr-153/Tyr-155**—The dual specificity mitogen-activated protein/ERK kinase (MEK) (isoforms 1/2, MAP2K1/2) is a prime candidate to phosphorylate at Thr-153/Tyr-155 based on prediction analysis of consensus sequence. Amino acid sequence alignment of the Thr-153/Tyr-155 region with the dual phosphorylation consensus sequences of other kinases in the MAPK family. The glutamic acid residue in the motif is critical for efficient ERK1/2 phosphorylation by MEK (31). Heat shock treatment dramatically increased activation and phosphorylation of MEK (Fig. 5B), in agreement with previous results (32). This triggered phosphorylation and activation of ERK as well as phosphorylation of a downstream substrate MSK1 (Fig. 5B). We investigated MEK phosphorylation of TDP-43 by use of specific kinase inhibitors, PD184352 and PD98059. Inhibitor treatment prior to heat shock prior to heat shock blocked Thr-153/Tyr-155 phosphorylation (Fig. 5C). However, inhibition of the downstream kinase ERK with FR182045 did not decrease Thr-153/Tyr-155 phosphorylation (Fig. 5C). Addition of okadaic acid, which blocks MEK inactivation by inhibiting protein phosphatase PP1/2A, increased p-T153/Y155 levels under control and heat shock conditions (Fig. 5D). This effect was blocked by addition of the MEK inhibitor PD184352. Furthermore, overexpression of a constitutively active mutant of MEK1, GFP-MEK1_DD (S218D/S222D), robustly increased TDP-43 phosphorylation at Thr-153/Tyr-155 in non-treated cells (Fig. 5E). We also observed increased
phosphorylation of a hemagglutinin (HA)-tagged TDP-43 expressed in the presence of the constitutively active MEK compared with control (supplemental Fig. S6). Collectively, these findings strongly suggest that TDP-43 is a novel substrate of MEK and that heat shock is not a prerequisite for Thr-153/Tyr-155 phosphorylation by constitutively active MEK.

**MEK Phosphorylation at Thr-153 and Tyr-155 Reduces TDP-43 regulation of Splicing**—To explore the effect of Thr-153/Tyr-155 phosphorylation on TDP-43 function, we used a reporter of splicing in cells. The well established cystic fibrosis transmembrane conductance regulator (CFTR) exon 9 mini-gene reporter (6, 33) was used in combination with a Thr-153/Tyr-155 phosphomimetic TDP-43 variant (T153E/Y155E). TDP-43 prevents exon 9 splicing through binding to a GU-rich sequence upstream of the 3' splice site (Fig. 6A) (34). Direct binding of TDP-43 to this sequence recruits additional hnRNPs, in particular hnRNP A2, preventing recognition of the 3' splice site by steric hindrance (6). Under control conditions, exon 9 showed 26% inclusion and, as expected, siRNA-mediated knockdown of TDP-43 greatly increased exon inclusion (supplemental Fig. S7A) (30). Splicing inhibition was re-established upon expression of siRNA-resistant wild-type TDP-43 (WTsiRes) (6, 33). The double mutant Phe-147 and Phe-149 to Leu (F147L/P149L) was used as control in our assays because of its inability to regulate splicing due to greatly reduced RNA binding (35, 36). As expected, F147L/F149L showed ~50% loss in activity compared with wild-type (Fig. 6B). We found that the phosphomimetic T153E/Y155E substitutions caused a 30% loss in splicing inhibition compared with wild-type and T153E/Y155A, where the phosphosites were mutated to alanine to prevent phosphorylation (Fig. 6B). Similar assays were carried out to further analyze the effect of Thr-153/Tyr-155 and MEK regulation on TDP-43 activity. Fig. 6C shows CFTR exon 9 splicing activity of control and siRNA-treated cells as a function of MEK_DD expression. The relative TDP-43 activity shown in Fig. 6C is a measure of CFTR exon 9 inclusion levels in each condition compared with WTsiRes cells. In control cells, MEK_DD reduced inhibition of CFTR exon 9 splicing by ~30% and this effect was lost upon TDP-43 knockdown. These results suggest that MEK_DD phosphorylation of endogenous TDP-43 decreases splicing regulatory activity, as seen with the phosphomimetic mutation (Fig. 6B). Accordingly, the effect of MEK_DD expression on reporter exon splicing was re-established in the presence of WTsiRes TDP-43, where MEK_DD reduced splicing regulation by ~50%. In contrast, this effect of MEK_DD was significantly diminished upon expression of the siRes, non-phosphorylatable, T153A/Y155A TDP-43 mutant. Collectively, our observations suggest that MEK phosphorylation at Thr-153/Tyr-155 inhibits the TDP-43-mediated regulation of splicing.

To elucidate whether the reduced splicing regulatory activity seen with T153E/Y155E may be caused by changes in RNA binding affinity, we generated recombinant TDP-43 using a bacterial expression system. We expressed and purified TDP-43 fused to a His-SUMO tag at the N terminus to improve yield and protein solubility. Next, we developed a fluorescence-based RNA binding assay to measure the apparent dissociation constant ($K_{d(app)}$) of TDP-43-RNA interactions as an alternative to the electromobility shift assay (EMSA). This new method was based on our observations that the intrinsic fluorescence of TDP-43 dramatically decreases upon RNA binding.
(approximately 50%). Using this assay we obtained \( K_{d(app)} \approx 2.3 \pm 0.7 \) nM for wild-type TDP-43 binding to A(GU)\(_6\) with a 1:1 stoichiometry (Fig. 7A, Table 1), in close agreement with values obtained by EMSA (37). As expected, mutations of Phe-147 and Phe-149 (F147L/P149L), previously shown to make direct contacts with GU repeats (35, 36), showed a great reduction in binding affinity (Table 1). We investigated the contribution of tryptophan residues to the changes in TDP-43 intrinsic fluorescence associated with RNA binding to further validate our assays. Removal of the C-terminal (ΔC) or both N- and C-terminal domains (fragment 102–269) did not significantly alter affinity or decrease the change in fluorescence compared with full-length TDP-43 (Table 1). These results indicate that Trp residues in the N and C terminus did not contribute to the fluorescence change linked to RNA binding. Moreover, they are in agreement with previous work showing that the carboxyl- and amino-terminal domains of TDP-43 do not contribute to RNA binding (35). We then substituted the remaining tryptophans at positions 113 and 172, found in RRM1. Substitution of Trp-113 to Ala (W113A) results in the loss of the fluorescence signal change as a function of RNA, whereas W113F reduced the fluorescence change by 50%, but had a modest effect on the \( K_{d(app)} \) (Table 1). Mutations of Trp-172 to either phenylalanine or alanine did not affect binding affinity or fluorescence change in our assays (Table 1). Our findings strongly suggest that...
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changes surrounding the Trp-113 region are principally responsible for the observed decrease in intrinsic fluorescence upon RNA binding. Trp-113 makes sequence-specific contacts with GU-rich RNA and substitutions of this residue reduce binding affinity (36, 37). These interactions, according to our data, result in dramatic changes in intrinsic fluorescence that may be exploited to accurately measure RNA binding affinity.

Analysis of T153E/Y155E using the fluorescence-based assay showed a 5-fold decrease in A(GU)_6 binding affinity in the presence of the phosphomimetic substitutions compared with wild-type, whereas the corresponding alanine mutations did not affect binding (Table 1). These results are consistent with the reduced splicing regulatory function seen in the presence of the phosphomimetic substitutions and strongly suggest that phosphorylation modulates TDP-43 activity.

Discussion

TDP-43 regulates hundreds of genes through different RNA processing mechanisms, in particular through the regulation of splicing (18, 19, 22). TDP-43 cellular localization is highly dynamic and defects in nuclear import are linked to aggregation and neurotoxicity (9, 38, 39). How these various TDP-43 functions are controlled in cells is largely unknown. Previously determined TDP-43 phosphorylation sites at Ser-403/404 and Ser-409/410 are poorly associated with the control of TDP-43 function and are mostly markers of pathological aggregates (25, 26). Here we report dual phosphorylation of TDP-43 at Thr-153/Tyr-155 and provide strong evidence that it is specifically controlled by the MAPK/ERK kinase MEK. This phosphorylation is dramatically up-regulated by the HSR in human cells. The increase in p-T153/Y155 is not associated with greater aggregation or protein clearance under the conditions tested. We find, instead, that p-T153/Y155 remains soluble under conditions that promote TDP-43 misfolding. In support of a functional role of p-T153/Y155, our results show that increased MEK activity significantly alters TDP-43-mediated control of exon splicing in cells. Our observations suggest that the effect of MEK is mediated by phosphorylation at Thr-153/Tyr-155. The decrease in TDP-43 activity may be at least partly explained by a reduction in RNA binding as we find that T153E/Y155E binds GU-rich RNA with 5-fold lower affinity. Future studies should determine whether phosphorylation changes RNA sequence specificity, or whether it affects protein interactions required for TDP-43 regulation of splicing, such as binding to hnRNP A2 (6).

The dramatic up-regulation of TDP-43 phosphorylation by heat shock, along with recent findings on the reduction of TDP-43 aggregation upon heat shock factor 1 (HSF1) activation (40, 41), point to a previously unappreciated role of HSR in TDP-43 homeostasis and posttranslational regulation. In addition to hyperthermic stress, HSR is activated by other types of insults that compromise protein homeostasis (42, 43), and defects in this response are commonly found in human disease including neurodegeneration. Therapies targeting HSR pathways, such as those showing promising effects in ALS (44), may be designed to specifically target TDP-43. We posit that p-T153/Y155 might mediate interactions with chaperones/heat shock proteins that prevent TDP-43 misfolding and aggregation, such as DNAJB2a and Hsp70 (40). Alternatively, it may be associated with an HSR-mediated clearance pathway, different from UPS and autophagy, which prevents aggregate accumulation. We also propose the alternative, tantalizing scenario that TDP-43 might regulate the expression of the HSR target genes whose levels are controlled by alternative splicing during the proteotoxic stress response (45). Based on our findings, this HSR-specific function of TDP-43 may be controlled by phosphorylation at Thr-153/Tyr-155.

MEK is activated upon heat shock and was recently shown to maintain protein homeostasis, at least in part, by phosphorylating and activating HSF1 (32). Previously, the only other recognized MEK substrate was ERK. Now, we provide evidence that TDP-43 is a novel MEK substrate in the presence and absence of heat shock. MEK may not be the only kinase to phosphorylate TDP-43 at Thr-153/Tyr-155, however, we observed no significant reduction of HSR-induced p-T153/Y155 levels upon specific inhibition of MAPK-related dual specificity kinases, namely JNK and p38 kinase (supplemental Fig. S8). Our evidence suggests that cellular processes associated with active MEK, such as proliferation and differentiation, may also induce p-T153/Y155 and regulate TDP-43 function. Based on our findings that p-T153/Y155 is associated with TDP-43 solubility in conditions that trigger protein misfolding, future work should determine whether MEK activation prevents TDP-43 aggregation and blocks TDP-43-associated neurotoxicity.

Our findings showing the specific localization of phosphorylated TDP-43 suggest that nucleolar TDP-43 recruitment is regulated by phosphorylation and may provide new insight into a previously unrecognized role of TDP-43 in cellular function linked to nucleolar RNA processing. In the C9ORF72 hexanucleotide repeat expansion, which is the most common cause of familial ALS and FTD (46, 47), the nucleolar compartment is disrupted by the generation of arginine-rich (GR and PR) dipeptide repeats (48–50). GR and PR are toxic, accumulate in nucleoli, and bind to proteins containing low complexity sequence domains including TDP-43 (51, 52). These findings strongly suggest that one of the principal pathogenic mechanisms in ALS/FTD is abnormal nucleolar function. Future experiments will determine TDP-43 function, particularly the role of p-T513/Y155 in nucleolar-specific RNA processing, and the effect of Arg-rich dipeptide repeats on this function. This may elucidate a convergent process resulting from TDP-43 and C9ORF72-associated dysfunction in ALS/FTD pathogenesis.

Experimental Procedures

Materials—All reagents are from Sigma unless otherwise specified. The human cell lines: neuroblastoma SH-SY5Y, epithelial HeLa, and embryonic kidney HEK-293 cells were purchased from ATCC.

Plasmid Construction—FLAG-tagged wild-type TDP-43, siRNA-resistant construct for mammalian expression was previously described (7). This served as template for site-directed mutagenesis using the QuikChange site-directed mutagenesis kit (Agilent Technologies) to generate TDP-43 mutants for mammalian expression. MEK1-DD-GFP (S218D/S222D) was generated by site-directed mutagenesis using the wild-type MEK1-GFP construct (Addgene, 14746) as template. pEFP-
LC3 vector was purchased from Addgene (24920). Constructs for bacterial TDP-43 expression (SUMO-TDP43) were made by cloning TDP-43 in pET-28b/His-SUMO (53) using pQTD-PBam_FW and SacTDPend_RV. This construct was the template to generate TDP-43 mutants for bacterial expression through site-directed mutagenesis as above. pQTDPPBam_FW and Sac_269_RV; Bam_102_FW and Sac_269_RV were used to clone ΔC-TDP-43 and fragment 102–269, respectively. Oligonucleotides used for mutagenesis and cloning are described in supplemental Table S1. The pLKO vector for TDP-43 down-regulation was generated by ligation of shT2_FW and shT2_RV annealing product into pLKO.1-puro vector digested with AgeI and EcoRI.

**Cell Culture, Fractionation, and Splicing Assays—**Human cell lines were grown in growth media, Dulbecco’s modified Eagle’s medium, 4,500 mg/liter of glucose, L-glutamine, and sodium bicarbonate, and supplemented with filtered fetal bovine serum at 10%. For the soluble and insoluble fractions cell pellets were resuspended in RIPA buffer (150 mM NaCl, 50 mM Tris, pH 8.0, 1% Nonidet P-40, 5 mM EDTA, 0.5% sodium deoxycholate, 0.1% SDS, 1× protease inhibitor mixture, 1× Phostop Phosphatase inhibitor mixture (Roche Applied Science)). Lysates were sonicated using a Bioruptor Pico (Diagenode) using 10, 30-s on/30-s off, cycles at 4 °C and centrifuged at 40,000 × g for 30 min at 4 °C. The pellet was rinsed with RIPA buffer and the final pellet was resuspended in 30 mM urea, 2 mM thiourea, 4% CHAPS, and 30 mM Tris, pH 8.5. Stable HEK-293 cells expressing HA-tagged TDP-43 under tetracycline induction (7) were used for immunoprecipitation experiments as previously described (54). RNAi-mediated down-regulation of TDP-43 was carried out as previously described (33) and siCONTROL Nontargeting siRNA#1 (Dharmacon) was used as control. shRNA-mediated down-regulation of TDP-43 was carried out by generating lentiviral particles in HEK-293T cells using a pLKO vector system (pCMV8.2AR, pCMV-VSV-G, generous gifts of Susana Gonzalo’s lab, St. Louis University). shRNA luciferase lentiviral particles were generated for control transduction. HEK-293T cells were transfected with Lipofectamine 2000 (Life Technologies) in Opti-MEM (Life Technologies) according to manufacturer’s protocols. After 6 h, media was replaced with Reduced Serum Media (Life Technologies) according to man-
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Described (53). Ulp1 was generously provided by Sergey Korolev’s group. All steps were carried out at 4 °C.

Fluorescence Titration RNA Binding Assays—TDP-43 intrinsic fluorescence was measured upon titration with increasing concentrations of A(GU)_6 RNA on a Fluoromax-4 spectrofluorometer (Horiba Scientific). Fluorescence readings were performed in a 1-ml cuvette at four different recombinant TDP-43 concentrations in the range of 10–100 nM in the presence of 200 mM NaCl, 20 mM Tris, pH 8.0, 0.1% PEG-8000, 2% glycerol, at 25 °C. Protein was excited at 280 nm (slits 5/10, exposure time of 0.02 s), and emission was recorded at 340 nm. We observed no significant photobleaching under these conditions. Data were analyzed with Origin (version 8.1, OriginLab) according to Equation 1,

\[ F = F_0 + \Delta F_{\text{max}} \theta \]

(Eq. 1)

where \( F \) is fluorescence, \( F_0 \) is \( F \) in the absence of RNA, \( \Delta F_{\text{max}} \) is the total change in \( F \) at saturation.

\[ \theta = \frac{K_{d(app)} + N_T + R_4 - \sqrt{(K_{d(app)} + N_T + R_4)^2 - 4N_TR_4}}{2N_T} \]

(Eq. 2)

Equation 2 describes the fractional saturation in terms of the apparent equilibrium dissociation constant \( K_{d(app)} \), the total concentration of TDP-43 (\( T_4 \)), total RNA concentration (\( R_4 \)), and TDP-43:RNA binding stoichiometry, \( N \). Four TDP-43:RNA binding curves were analyzed simultaneously to obtain \( K_{d(app)} \) and \( N \).

Antibodies—Immunoblots and indirect immunofluorescence were performed with: rabbit polyclonal and mouse monoclonal anti-TDP-43 (ProteinTech 10782-2-AP, Abcam, ab109535), anti-TIAR (BD Biosciences, 610352), anti-fibrillarin (Abcam, ab4566), mouse anti-tubulin (Andres Muro, Trieste, Italy), anti-GAPDH (Abcam, ab181602), anti-MEK (Abcam, ab178767), anti-phospho-MEK1/2 (Cell Signaling, number 9154), anti-ERK1/2 (Cell Signaling, number 4695), anti-phospho-ERK1/2 (Cell Signaling, number 4370), anti-MSK1 (Cell Signaling, number 3489), and anti-phospho-MSK1 (Cell Signaling, number 9595). The rabbit polyclonal antibodies recognizing p-T153/Y155 and p-Y155-TDP-43 were produced by 21st Century Biochemicals, MA. Peptide including TDP-43 amino acids 148–161 phosphorylated at either Thr-153/Tyr-155 or Tyr-155 served as epitope for antibody production. Antibody specificity was enhanced by affinity purification to select for binding to phosphorylated Thr-153/Tyr-155 or Tyr-155, but not the corresponding non-phosphorylated peptides.

**Author Contributions**—W. L., A. N. R., B. L., P. S., E. E. F., C. R. K., R. L. F., E. H. B., and Y. M. A. designed the research, performed the experiments and analyzed the data. Y. M. A. wrote the manuscript.

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**References**


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