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Multiplex SILAC Analysis of a Cellular TDP-43 Proteinopathy Model Reveals Protein Inclusions Associated with SUMOylation and Diverse Polyubiquitin Chains*

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Transactive response (TAR) DNA-binding protein 43 (TDP-43) is a major protein component within ubiquitin-positive inclusions of frontotemporal lobar degeneration and amyotrophic lateral sclerosis. Although TDP-43 is a nuclear DNA/RNA-binding protein, in pathological conditions, TDP-43 has been reported to redistribute to the cytoplasm where it is cleaved and forms insoluble, ubiquitinated, and phosphorylated inclusions. Here we present a cellular model in which full-length human TDP-43 or a splicing isoform (TDP-S6) that lacks the C terminus is overexpressed in a human cell line and mouse primary neurons. Whereas recombinant and endogenous TDP-43 was primarily localized in the nucleus, the shorter TDP-S6 formed highly insoluble cytoplasmic and nuclear inclusions reminiscent of disease-specific pathology. Western blot analysis of detergent-insoluble extracts showed an increase in high molecular weight immunoreactive species for TDP-S6 compared with TDP-43, consistent with ubiquitination or ubiquitin-like modifications. We used a multiplex stable isotope labeling with amino acids in cell culture approach to compare the detergent-insoluble proteome from mock-, TDP-43-, and TDP-S6-transfected cells. TDP-S6 overexpression caused a concomitant increase in both ubiquitin (Ub) and the small Ub-like modifier-2/3 (SUMO-2/3) within the insoluble proteome. Similarly, full-length TDP-43 overexpression also resulted in the elevation of SUMO-2/3. Immunofluorescence showed strong co-localization of endogenous Ub with both cytoplasmic and nuclear TDP-S6 inclusions, whereas SUMO-2/3 was co-localized mainly with the nuclear inclusions. Quantitative mass spectrometry further revealed that mixed Lys-48 and Lys-63 polyUb linkages were associated with the TDP insoluble fractions. Together our data indicate that expression of a TDP-43 splice variant lacking a C terminus recapitulates many of the cellular and biochemical features associated with disease pathology and that the interplay of ubiquitination and SUMOylation may have an important role in TDP-43 regulation. Molecular & Cellular Proteomics 9:705–718, 2010.

Frontotemporal lobar degeneration (FTLD)1 is a progressive neurodegenerative disease characterized by prominent behavioral abnormalities, personality changes, and language dysfunction, and it co-occurs with the development of parkinsonism and motor neuron disease in some patients (1, 2). Like other neurodegenerative diseases, FTLD is characterized by the abnormal accumulation of ubiquitinated intracellular protein aggregates (3, 4). In addition to the aggregation of tau in some forms of FTLD, more than half of cases are marked by ubiquitin-positive inclusions and are subclassified as FTLD-U. Transactive response (TAR) DNA-binding protein 43 (TDP-43) has been identified as a major protein component of inclusions in FTLD-U and amyotrophic lateral sclerosis (ALS) (5). TDP-43 aggregation is also observed in hereditary inclusion body myopathy and Paget disease of the bone with frontotemporal dementia (6) as well as in some cases of Alzheimer and Parkinson diseases (7, 8). This indicates that a common underlying mechanism may broadly define a spectrum of neurodegenerative disorders termed “TDP-43 proteinopathies” (9, 10).

TDP-43 is a 414-amino acid protein that contains two RNA recognition motifs (RRM1 and RRM2) and a glycine-rich C-terminal domain. It is highly conserved in human, mouse, fly, and worm and is expressed in all tissues, including the brain.

1 The abbreviations used are: FTLD, frontotemporal lobar degeneration; TDP, TAR DNA-binding protein; FTLD-U, ubiquitin-positive inclusions of frontotemporal lobar degeneration; ALS, amyotrophic lateral sclerosis; SILAC, stable isotope labeling with amino acids in culture; Ub, ubiquitin; SUMO, small ubiquitin-like modifier; RRM, RNA recognition motif; TAR, transactive response; HEK, human embryonic kidney; HA, hemagglutinin; SENP2, sentrin-specific endopeptidase 2; FDR, false discovery rate; SRM, selective reaction monitoring; STUbL, SUMO-targeted ubiquitin ligase; ARP1, actin-related protein 1.
(11–13). First characterized to bind and repress the promoter activity of TAR DNA in the human immunodeficiency virus 1 long terminal repeat region (14), TDP-43 was later found to regulate splicing of the transcripts of apolipoprotein A-II (15) and cystic fibrosis transmembrane conductance regulator (16). Therefore, TDP-43 can act both as a transcriptional repressor and as a splicing regulator. Although physiological TDP-43 resides mainly in the nucleus, pathology-relevant TDP-43 redistributes from the nucleus to the cytoplasm where it is cleaved and forms phosphorylated and ubiquitinated inclusions (5, 17–19). The degree to which TDP-43 is posttranslationally modified and whether other modifications initiate nuclear to cytoplasmic translocation and aggregation remain unknown.

The primary transcripts of TDP-43 in mouse and human undergo multiple alternative splicing events in which 10 splice variants (S1–S10) have been documented (11). With the exception of full-length TDP-43, all alternatively spliced isoforms are expressed as truncated proteins that lack the C-terminal glycine-rich domain. Isoforms that lack this domain are no longer capable of enhancing the skipping of exon 9 of the cystic fibrosis transmembrane conductance regulator gene via interactions with heterogeneous nuclear ribonucleoprotein A/B (11, 20). At the time, the vast majority of sporadic and familial TARDBP gene variants found in ALS cases (21–25) are reported to have missense mutations resulting in single amino acid substitutions located in the glycine-rich domain. This establishes an intriguing relationship between ALS disease etiology and the function of the TDP-43 C-terminal region. Whether TDP-43 splice variants that lack this C terminus contribute to translocation and aggregation requires further investigation.

In this study, we overexpressed human TDP-43 and TDP-S6, a splice variant lacking the glycine-rich C terminus, in human embryonic kidney 293 (HEK-293) cells and mouse hippocampal neurons. When compared with TDP-43, TDP-S6 showed both nuclear and cytoplasmic location and extensive aggregation. Biochemical analysis revealed that TDP-S6 was almost completely Sarkosyl-insoluble. Moreover, both insoluble TDP-43 and TDP-S6 displayed varying levels of posttranslational modifications that included ubiquitination and/or ubiquitin-like modifications, phosphorylation, and proteolytic cleavage. To assess protein differences among the detergent-insoluble extracts from mock-, TDP-43-, and TDP-S6-transfected cells, a multiplex stable isotope labeling with amino acids in culture (SILAC) strategy was used in combination with LC-MS/MS (26). This quantitative proteomics approach metabolically introduces a mass difference into proteins by incorporation of heavy isotopic forms of arginine and lysine. We identified and characterized a novel association between the small ubiquitin (Ub)-like modifier-2/3 (SUMO-2/3) and TDP-43. Additional targeted proteomics analysis found multiple polyubiquitin structures in the TDP insoluble fractions. Our data indicate that overexpression of a truncated TDP-43 splice variant recapitulates many of the features associated with disease pathology and that SUMOylation and ubiquitination may play important roles in regulating TDP-43 functions.

EXPERIMENTAL PROCEDURES

DNA Constructs—The full-length TDP-43 coding sequence (nucleotides 1–1245, encoding amino acids 1–414) was amplified by PCR from a TDP-43 expression plasmid (Open Biosystems, Huntsville AL; Clone ID 503890S); the forward and reverse primers were 5′-GGTAG-GAATCCAGCTGCTGATATATCCGGGTAA-3′ and 5′-TTATATGGCC-CCTCACCATTCCCAGCAGA-3′, respectively. The PCR products were digested and cloned into pcDNA3.1-HA expression vector. To investigate downstream of the two HA motifs (peptide YPYDVDYA). The TDP-S6 splice variant (nucleotides 1–888 (11), encoding amino acids 1–295) was also cloned into pcDNA3.1-HA. Both N-terminal HA expression vectors (TDP-43 and TDP-S6) were confirmed by DNA sequencing (Agencourt Bioscience Corp., Beverly MA). The SUMO-2 construct was a kind gift from Dr. Keith Wilkinson (Emory University). The tetracycline-regulated Co., Long Island City, NY) and maintained in Neurobasal medium (1% penicillin/streptomycin) (Cambrex). After plating on Matrigel-coated coverslips, cells were transfected using FuGENE 6 reagent (Roche Applied Science) followed by immunocytochemistry 2 days later as described previously (28). Primary antibodies included rabbit TDP-43 antibody raised against recombinant TDP-43 (ProteinTech Group, Chicago, IL), HA monoclonal antibody (clone 16B12; Covance, Emeryville CA), rabbit SUMO-2/3 antibody (Zymed Laboratories Inc.), rabbit antibody against the neuron-specific marker enolase (Chemicon, Temecula, CA), rabbit Ub antibody (Dako, Carpinteria, CA), and Lys-48 or Lys-63 linkage-specific antibodies, a gift from Dr. V. M. Dixit (29). Briefly, the cells were fixed with 2% paraformaldehyde in PBS, permeabilized and blocked with PBS containing 5% BSA and 0.05% Triton X-100, and incubated in primary antibodies overnight. The cells were then rinsed and incubated with either fluorescein-conjugated or biotinylated secondary antibodies (Jackson ImmunoResearch Laboratories, West Grove, PA) for 60 min. Cells treated with biotinylated secondary antibodies were subsequently incubated with avidin-biotin-peroxidase complex and tyramide-conjugated fluorophores according to the manufacturer’s protocol (PerkinElmer Life Sciences). Following additional rinses, cells were incubated with a Hoechst reagent for nuclear localization, mounted with Vectashield (Vector...
Laboratories, Burlingame, CA), and imaged on a Zeiss LSM 510 confocal microscope (Zeiss, Thornwood, NY). Images were captured with a 1-μm optical thickness for subsequent analysis.

**Subcellular Fractionation**—To examine the solubility of TDP-43 and TDP-S6, sequential extractions were performed as described previously (5) with slight modification. Cells (1 × 10^7) were washed twice, collected with ice-cold PBS buffer, and then lysed in Sarkosyl buffer (10 mM Tris, pH 7.5, 5 mM EDTA, 1 mM DTT, 0.5 mM NaCl, 10% sucrose, 1% N-lauroylsarcosine, 10 mM β-glycerophosphate, 10 mM sodium orthovandate, 10 mM tetrasodium pyrophosphate, 50 mM sodium fluoride). Resulting lysates were spun at 180,000 × g for 30 min at 22 °C to generate the detergent-soluble samples. Insoluble pellets were washed three times with additional Sarkosyl buffer, then extracted with urea buffer (30 mM Tris, pH 8.5, 7 M urea, 2 M thiourea, 2% sodium dodecyl sulfate), and centrifuged at 25,000 × g for 30 min at 22 °C. Protease inhibitors (Roche Applied Science) were added to all buffers prior to use. Protein concentration was determined for Sarkosyl fractions by bicinchoninic acid (BCA) protein assay (Pierce) according to the manufacturer’s instructions. For urea fractions, protein concentration was determined likewise and/or by estimating Coomassie Blue G-250 staining intensity of a small fraction of the urea extracts following electrophoresis in polyacrylamide gels using titrated BSA as a standard (30).

**Biochemical Assays**—To assess TDP-43 phosphorylation, total cell lysate was incubated with 0, 0.15, 0.3, or 0.6 mg/ml bovine alkaline phosphatase Type VII-S (3533 units/mg of protein; Sigma-Aldrich) for 2.5 h at 37 °C followed by Western blot analysis. For deSUMOylation, sentrin-specific endopeptidase 2 (SENPE2; Boston Biochem, Cambridge, MA) was incubated with 10-μg detergent-soluble or 1-μg insoluble fractions from TDP-43-transfected cells in 20-μl reactions (buffer: 50 mM Tris, 100 mM NaCl, 5 mM DTT, pH 8.0; titrated SENPE2 concentration: 0, 1, or 5 μM). All samples were incubated for 3 h at 37 °C and then analyzed by Western blotting.

**Western Blotting**—Immunoblotting was performed according to standard procedures. Briefly, samples in Laemmli sample buffer were separated by SDS-PAGE and transferred overnight to PVDF Immobilon® membranes (Millipore, Billerica, MA). To ensure both equal loading and complete transfer of proteins from the gel, membranes were reversibly stained with Ponceau S (Diasys Europe Ltd., Workingham, UK). Blots were blocked for 1 h at room temperature using Blocking Buffer (USB Corp., Cleveland OH), probed with primary antibody in TBS with 0.1% Tween 20 overnight at 4 °C, and incubated for 1 h at room temperature with secondary antibodies (1:20,000 for secondary antibodies (1:20,000); goat anti-rabbit or goat anti-mouse Igs; Rockland, Gilbertsville PA). Blots were dried, scanned, and quantified with an Odyssey Infrared Imaging System (Li-Cor Biosciences, Lincoln, NE).

**SILAC and LC-MS/MS**—Cells were cultured in Dulbecco’s modified Eagle’s medium (deficient in L-lysine and L-arginine) supplemented with 5% dialyzed fetal calf serum (Invitrogen) as described (31). For stable isotopic labeling, arginine and lysine were added in light (Arg0/Lys0; Sigma), medium (Arg6/Lys4), or heavy forms (Arg10/ Lys8; Cambridge Isotope Laboratories, Andover, MA) to a final concentration of 0.26 μM. Cells were cultured for seven passages to ensure full labeling and then transferred with pcDNAA3.1-HA (mock), HA-TDP-43, and HA-TDP-S6 plasmids, respectively. After 2 days, the cells were harvested, equally mixed, and lysed to prepare the Sarkosyl-insoluble, urea-soluble fraction as described above. The fraction was reduced with 10 mM DTT, alkylated with 50 mM iodoacetamide for 30 min in the dark, and then resolved on a 10% polyacrylamide-SDS gel. After staining with Coomassie Blue, one gel lane was cut into five gel bands, and bands were subjected to in-gel digestion (12.5 μg/ml trypsin). Extracted peptides were loaded onto a C18 column (75-μm inner diameter, 10 cm long, 300 nL/min flow rate, 5-μm resin from Michrom Biosources, Auburn, CA) and eluted during a 10-30% gradient (Buffer A: 0.4% acetic acid, 0.005% heptfluorobutyric acid, 5% ACN; Buffer B: 0.4% acetic acid, 0.005% heptfluorobutyric acid, 95% ACN). The eluted peptides were detected by Orbitrap (350–1500 m/z; 1,000,000 automatic gain control target; 1000-ms maximum ion time; resolution, 60,000 full-width at half-maximum) followed by five data-dependent MS/MS scans in the linear ion trap quadrupole (2 m/z isolation width, 35% collision energy, 5,000 automatic gain control target, 200-ms maximum ion time) on a hybrid mass spectrometer (Thermo Finnigan, San Jose, CA).

Acquired MS/MS spectra were extracted and searched against the human reference database from the National Center for Biotechnology Information (December 13, 2007) using the SEQUEST Sorcerer algorithm (version 2.0, SAGE-N) (32). Searching parameters included mass tolerance of precursor ions (±50 ppm) and product ion (±0.5 m/z); partially tryptic restriction; fixed mass shift for modification of carboxamidomethylated Cys (+57.0215 Da); dynamic mass shifts for oxidized Met (+15.9949), Lys (+4.02511 for 2H2 or +8.01420 for 13C6)^15N4H), and Arg (+6.02013 for 13C6 or +10.00827 for 13C6)^15N4H); five maximal modification sites; and three maximal missed cleavages. Only b and y ions were considered during the database match. To compare medium and heavy labeled samples, searches were performed with static modifications +4.02511 on Lys and +6.02013 on Arg and dynamic modifications of +3.98909 on Lys and +3.98814 on Arg to account for the mass difference between medium and heavy labeled peptides. To evaluate the false discovery rate (FDR), all original protein sequences were reversed to generate a decoy database that was concatenated to the original database (a total of 53,830 protein entries) (33, 34). The FDR was estimated by the number of decoy matches (nD) and total number of assigned matches (nA). FDR = 2 × nD/nA, assuming that mismatches in the original database were the same as in the decoy database. To remove false positive matches, assigned peptides were grouped by a combination of trypticity (fully, partial, and non-tryptic) and precursor ion charge state (1+, 2+, 3+, and 4+). Each group was first filtered by mass accuracy (10 ppm for high resolution MS) and by dynamically increasing XCorr (minimal 1.8) and ΔCn (minimal 0.05) values to reduce protein FDR to less than 0.2%. The identified proteins/peptides are listed in the supplemental data (supplemental Tables S1–S6 and S8) with accession number, matched peptide number, sequencing coverage, mass shift, and matching scores. Every peptide in the tables was linked to assigned MS/MS spectra, precursor ion mass, and charge state. If peptides were matched to multiple members of a protein family, the matched members were clustered into a single group (supplemental Tables S7 and S9).

**SILAC Quantification and Bioinformatics Analysis**—Quantitative pairwise comparisons of control, TDP-S6-, and TDP-43-transfected cells were carried out according to reported methods (35, 36) with slight modification. (i) For ion extraction from MS scans, the ion currents for identified peptides were extracted in MS survey scans of high resolution (60,000) based on the isotopic ion selected for MS/MS sequencing. A number of parameters were defined, including precursor m/z, charge state, retention time, ion peak width, height, area, and noise level. The noise level was derived by averaging signal intensity of all ions in the MS scan after removing outliers that were at least two S.D. away from the mean. The intensity of ions was presented by the peak height and normalized according to the noise intensity under the assumption that the noise level of MS scans reflects, at least partially, variable ionization efficiency. The peaks used in the analysis had a minimal intensity of 2 signal-to-noise ratio. (ii) Ion matching among light, medium, and heavy isotopes was allowed with a tolerance of 10 ppm. If a sequenced peptide could not be matched, we estimated that the maximum ion current for undetected signal was equal to the noise level and used it to derive the peptide ratio. (iii) For data
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integration, the ratio of every peptide was transformed into logarithmic (log₂) values that were averaged over all peptides of a particular protein to determine the protein ratio. If a protein was quantified by both matched peptide ratios and unmatched peptide ratios, only the matched data were averaged. The results are shown in supplemental Table S7 with the number of peptides quantified for every protein. (iv) For data normalization, according to the null hypothesis, the histogram of all protein log₂ ratios was fitted to a Gaussian distribution to evaluate systematic bias (according to the mean) and experimental variation (based on S.D.). The data were then normalized by subtracting the mean in every protein to The S.D. values in all SILAC comparisons were less than 0.4. (v) For data filtering, we selected the cutoff of log₂ ratios that were outside a 95% confidence interval (~2 S.D.) from the mean of the Gaussian distribution. Finally, the quantified proteins were manually examined with respect to MS/MS assignment, peak matching, and ion intensity.

Semiquantitative Proteomics by Spectral Counts—To compare differences between the detergent-soluble and -insoluble proteome, we compared the identified proteins in both samples based on spectral counts. The spectral counts were first normalized to ensure that average spectral counts per protein was the same in the two data sets (37). A G test was used to judge statistical significance of protein abundance difference (38). Briefly, the G value of each protein was calculated as shown in Equation 1.

\[
G = 2 \times (S_{tc} \times \ln[S_{tc} + ((S_{tc} + S_{si}) \times 2)] + S_{si} \times \ln[S_{si} + ((S_{tc} + S_{si}) \times 2)]) \quad (\text{Eq. 1})
\]

where \(S_{tc}\) and \(S_{si}\) are the detected spectral counts of a given protein in the total cell lysate and in the Sarkosyl-insoluble fraction, respectively, and “ln” is the natural logarithm. Although theoretical distribution of the \(G\) values is complex, these values approximately fit to the \(\chi^2\) distribution (1 degree of freedom), allowing the calculation of related \(p\) values.

Quantitative Analysis of Polyubiquitin Chains and Other Targeted Proteins by LC/Selective Reaction Monitoring (SRM)—The analysis of polyUb linkages, ubiquitin E1 enzyme, and proteasome subunit Rpn2 was performed with metabolically heavy labeled cells as internal standards (+8.01420 for Lys and +10.00827 for Arg) using a previously reported protocol (36, 39). The labeled cells were spiked into transfected cells followed by protein extraction. Sarkosyl-insoluble fractions were resolved on a one-dimensional SDS gel. The gel regions above 80 kDa, which contained the vast majority of polyUb species, were used for in-gel trypsin digestion that produced a pair of light and heavy GG-linked ubiquitin peptides corresponding to every polyUb linkage. Digested peptides were analyzed by the same LC system as above in which peptide ion pairs of interest were selected for fragmentation and quantified by related product ion pairs, a process termed SRM or multiple reaction monitoring. The detailed LC/SRM parameters are shown in supplemental Table S10.

**RESULTS**

Expression, Localization, and Biochemical Properties of Recombinant TDP-43 and TDP-S6—We cloned and overexpressed HA-tagged (at the N terminus) human TDP-43 and TDP-S6 in HEK-293 cells (Fig. 1). Human TDP-S6 is generated via an additional splicing event within exon 6 and encodes a 295-amino acid protein in which the first 277 amino acids are identical to those of TDP-43 (Fig. 1A). Although mouse TDP-S6 has been shown to have altered nuclear structure in mammalian cells (40), the human TDP-S6 transcript, identified in HEK-293 cells (11), has not yet been characterized. Western blotting of total cell lysate prepared from the transfected cells indicated that both recombinant proteins were expressed and recognized by the HA antibody. TDP-43 also displayed two higher molecular mass species at ~50 kDa (Fig. 1, A and C, asterisk). After treatment with an increasing amount of alkaline phosphatase, the upper band disappeared, and the middle band became significantly weaker, whereas the intensity of unmodified TDP-43 remained stable. This supports that TDP-43 is preferentially phosphorylated on the C terminus on at least two different residues (Fig. 1C), consistent with current identification of two phosphorylation sites near the C terminus of TDP-43 in diseased tissues (41, 42).

Subcellular localization of full-length TDP-43 and TDP-S6 was assessed by immunofluorescence confocal microscopy. Overexpressed TDP-43 in HEK-293 cells had primarily nuclear localization (Fig. 2A, upper panel), consistent with that of endogenous TDP-43 in untransfected cells (Fig. 2A, upper panel, left bottom corner). In contrast, TDP-S6 formed aggregates that were mainly localized to the cytoplasm with some inclusions in the nucleus (Fig. 2A, middle and bottom panels).
Observed puncta were of variable size, ranging from fine granular deposits to large, aggresome-like inclusions. To assess whether the TDP-S6 phenotype observed in HEK-293 cells could be recapitulated in neurons, we transfected primary mouse hippocampal neurons with TDP-43 or TDP-S6 constructs (Fig. 2B). Consistent with results in HEK-293 cells, TDP-43 in neurons was nuclear, whereas the majority of TDP-S6 formed granular deposits in the cytoplasm of the soma and neurites (Fig. 2, B and C). Enolase was stained as a neuron-specific marker.

To evaluate the solubility of overexpressed TDP-43 or TDP-S6, HEK-293 cells were transfected with either construct and
then mixed equally to eliminate experimental variability in subsequent processing. The mixed cells were sequentially extracted with Sarkosyl-containing buffer and urea and then analyzed by immunoblotting. The differences in molecular mass between HA-TDP-43 (47 kDa), native TDP-43 (43 kDa), and HA-TDP-S6 (37 kDa) allowed analysis of all three proteins in a single lane (Fig. 3A). In addition, phosphorylated TDP-43 isoforms were also resolved. Although almost all of HA-TDP-S6 was insoluble, 60 ± 7% of unmodified HA-TDP-43 and virtually none of the native unmodified TDP-43 were present in the insoluble fraction (Fig. 3, A and B). The degree of TDP-S6 and TDP-43 enrichment in the insoluble fraction was also repeatable using a second preparation of cells (data not shown). Based on these findings, biochemical insolubility of TDP-S6 is consistent with the inclusions observed by immunofluorescence (Fig. 2). The overexpression of HA-TDP-43 may account for the differences in solubility between the recombinant and native proteins as has been recently reported (43). Interestingly, the phosphorylated species of recombinant HA-TDP-43 (44 ± 1%) and endogenous TDP-43 (44 ± 1%) were more enriched in the insoluble fraction than the unmodified forms, respectively (60 ± 7 and <5%), supporting previous findings linking TDP-43 phosphorylation with aggregation and insolubility (5). To ensure that overexpression of TDP-43 or TDP-S6 did not dramatically bias the global distribution of proteins to the insoluble fraction, the percentage of total protein in each fraction (detergent-soluble and -insoluble) was calculated from mock-, HA-TDP-43-, and HA-
TDP-S6-transfected cells. Approximately 96% of all cellular proteins were detergent-soluble irrespective of the recombinant protein expressed, indicating that overexpression of our target proteins did not cause a gross increase in the total amount of protein within the insoluble fraction.

To further characterize proteins enriched in the Sarkosyl-insoluble fraction, we identified proteins in the total cell lysate and in the Sarkosyl-insoluble sample using a label-free LC-MS/MS approach. The relative abundance of proteins in the two samples was compared by spectral counts, and the difference was statistically evaluated by \( G \) test (38). Among 1265 proteins profiled (supplemental Table S9), the level of 267 proteins was significantly altered with corresponding \( p \) values below 0.01. Listed in Fig. 3C are proteins with large changes, including components associated with the proteasome, caveolae, aggresomes, and a number of nuclear complexes. Specifically, flotillin-1 and -2 are markers for caveolae/lipid rafts that are highly enriched in cholesterol and are thus more resistant to detergent extraction (44). Vimentin is a protein marker of aggresomes that is often associated with proteasome proteins (45). Finally, particular DNA/RNA-interacting proteins were found to be intrinsically detergent-insoluble. Therefore, the protocol of Sarkosyl-based differential extraction is capable of enriching both aggregated proteins and other detergent-insoluble complexes.

Further analysis of detergent-soluble and insoluble fractions from transfected cells revealed that high molecular mass TDP-43-immunoreactive species were more abundant in fractions corresponding to TDP-S6, rather than TDP-43, overexpression. This observation is consistent with ubiquitination, ubiquitin-like modifications, and/or protein polymerization typically associated with protein insolubility (Fig. 4). To this end, immunoblotting of Sarkosyl-insoluble fractions with a ubiquitin-specific antibody demonstrated an increase in global protein ubiquitination with TDP-43 (\( \sim \)250 kDa) and more dramatically with TDP-S6 overexpression (Fig. 4). TDP-S6 also displayed two short TDP-43-immunoreactive species (\( \sim \)30 and \( \sim \)20 kDa) that were not detected by an HA antibody. In addition, a third fragment (\( \sim \)37 kDa) was identified in TDP-S6 urea fractions by both TDP-43 and HA antibodies. Although TDP-S6 lacks the extreme C terminus, the proteolytic fragmentation observed here is in agreement with reported C-terminal TDP-43 fragmentation in FTLD-U cases (19). Full-length TDP-43 was also cleaved but at a much lower level (observed on overexposed images; data not shown). These results show that an increase in phosphorylation, ubiquitination, and proteolysis is related to the insolubility of TDP proteins. Taken together, the microscopic and biochemical data for TDP-S6 in cell culture are highly consistent with that of pathologic TDP-43 observed in disease tissue (5, 9, 18, 19).

**Quantitative Analysis of Insoluble TDP-43 and TDP-S6 Proteome Using Multiplex SILAC—**As both recombinant TDP-43 and TDP-S6 are highly enriched in detergent-insoluble extracts, we used a multiplex SILAC approach to evaluate proteins that co-enrich in the detergent-insoluble proteome of cells overexpressing TDP-43 or TDP-S6 (46) (Fig. 5A). Human HEK-293 cells were fully labeled with light, medium, or heavy arginine and lysine amino acids. The three labeled cell populations were transfected with HA-TDP-43, HA-TDP-S6, or a mock plasmid. After incubation for 2 days, the cells were harvested, equally mixed, and subjected to sequential protein extraction. Although pooling the labeled cells minimizes experimental variation during protein extraction and subsequent steps in the SILAC analysis, it is possible for biological variation to occur during cell labeling and DNA transfection. Therefore, we performed a second independent analysis as a
biological replicate with the labeled isotopes “swapped” in HEK-293 cells prior to transfection (Fig. 5A). The detergent-insoluble extracts were then resolved on an SDS gel, excised into gel bands, digested with trypsin, and analyzed by LC-MS/MS on a high resolution Orbitrap mass spectrometer (Fig. 5B). After database searching and stringent filtering mainly by mass accuracy (≤10 ppm) and SEQUEST scores (XCorr and ΔCn), a total of 2670 proteins that were clustered into 1117 groups were identified and quantified (supplemental Tables S1–S7). The corresponding false discovery rate was calculated to be less than 0.2% according to the target-decoy strategy (33, 34).

To evaluate the quality of the quantitative data, we analyzed the results based on the null hypothesis as described previously (35). The protein ratios in all pairwise comparisons were converted into log2 values, and the resulting histogram of all values was fitted to a Gaussian distribution (Fig. 5C). The majority of proteins fit the curve well, indicating that they did not change under the conditions analyzed. Although the fitted mean (−0.01) suggested little systematic bias introduced by sample handling (e.g. a slightly different amount of starting cells), the S.D. (0.31) provided a good measure of the magnitude of variations in the analysis. In all six pairwise comparisons of light, medium, and heavy labeling in the two experiments, the value of the S.D. was stable (0.28 ± 0.03; Fig. 5D), indicating that the variations in our SILAC analyses were consistent. Proteins were considered changed if their values fell outside more than 2 S.D. (0.6; 95% confidence interval) and showed consistency in experiments 1 and 2. A large number of proteins were removed when comparing results from experiments 1 and 2, suggesting that the biological replicate is essential to reduce false positives in the SILAC assay. The filtered proteins were further validated by manual examination of paired ion currents in raw files, resulting in the acceptance of six proteins with altered expression (Table I).

Validation and Subcellular Co-localization of SUMO-2/3 and Ubiquitin—The list of proteins with altered levels in the Sarkosyl-insoluble fraction included overexpressed TDP proteins (TDP-43 or TDP-S6), ubiquitin, and SUMO-2/3 (Table I). In gel band 1 (>180 kDa; Fig. 5B), the MS results (Fig. 6, A and B) indicated that TDP-S6-expressing cells had more TDP in the insoluble proteome compared with TDP-43-expressing cells, consistent with the high molecular mass immunoreactivity in Western blotting (Fig. 4B). As expected, native TDP-43 from mock-transfected cells was not observed at this high molecular mass range. SILAC also confirmed our Western blot results for ubiquitin (Fig. 4B), which increased concomitantly with TDP levels within this molecular mass range. However, although TDP-S6 showed significant ubiquitin enrichment compared with TDP-43-expressing cells, consistent with the high molecular mass immunoreactivity in Western blotting (Fig. 4B), no enrichment was observed for ubiquitin in the TDP-43 insoluble fraction (Table I). In contrast, the significant increase in total ubiquitin enrichment for TDP-S6 may be a reflection of its aggregation observed in cell culture. Interestingly, SUMO-2/3 was identified as a novel component enriched in both the TDP-43 and TDP-S6 insoluble proteome (summed over all molecular mass regions), suggesting the presence of both polyubiquitination
and SUMOylation in the insoluble (Fig. 6C) proteome. The elevation of SUMO-2/3 modifications was correlated to the degree of TDP protein insolubility (Fig. 3), suggesting that insoluble TDP proteins are direct targets of SUMOylation.

To assess whether TDP-43 is covalently modified by SUMO-2/3, we performed a deSUMOylation assay using SENP2 (47). An increasing amount of SENP2 was added to either detergent-soluble or -insoluble samples from cells overexpressing TDP-43. A Western blot with anti-SUMO-2/3 antibodies. Because unmodified HA-TDP-43 has a molecular mass of ~45 kDa and SUMO-2/3 is ~10 kDa in size, these bands likely represent mono-, di-, and triSUMOylated forms of TDP-43. The tri- and diSUMOylated HA-TDP-43 showed decreasing signals when incubated with titrated SENP2. In contrast, the monoSUMOylated band was increased at 1 μM SENP2, possibly due to processing of tri- and diSUMOylated bands. This increase was appropriately reduced with increased SENP2 concentration (5 μM). Thus, HA-TDP-43 is directly modified by SUMO-2/3 in detergent-insoluble fractions.

To assess whether SUMO-2/3 co-localizes with TDP-43 or TDP-S6 in HEK-293 cells, we performed immunofluorescence confocal microscopy (Fig. 7A). Endogenous SUMO-2/3 in untransfected cells was diffusely expressed mainly in the nucleus with some accumulation in nuclear bodies as described previously (48). In cells expressing full-length TDP-43, the majority of endogenous SUMO-2/3 was diffusely localized throughout the nucleus, although co-localization was also observed in certain SUMO-2/3 bodies when TDP-43 expression levels were higher (Fig. 7A, arrowheads). In TDP-S6-expressing cells, SUMO-2/3 was sequestered within TDP-S6 nuclear inclusions. Notably, SUMO-2/3 did not co-localize with cytoplasmic aggregates of TDP-S6. Similar results were obtained in cells co-expressing green fluorescent protein-tagged SUMO-2 and TDP constructs (data not shown). These data strongly support that SUMOylation is associated with TDP protein insolubility and localization in the nuclei but not in the cytoplasm.

Ub is attached to protein substrates in monomeric form or as polymers (polyUb) assembled through the N-terminal amino group and the side chains of all seven lysine residues (Lys-6, Lys-11, Lys-27, Lys-29, Lys-33, Lys-48, and Lys-63). These diverse polyUb structures may regulate downstream signaling specificity and determine the consequence of ubiquitination (36, 47). Whereas the Lys-63 polyUb linkage is proposed to mediate nonproteolytic events, including protein trafficking and inclusion formation (49), the Lys-48 linkage and other atypical polyUb linkages mediate proteasomal degradation (36). To elucidate the role of different polyUb linkages with TDP protein aggregates, we examined transfected cells with Lys-48 or Lys-63 polyUb linkage-specific antibodies (29) compared with untransfected cells and found that both linkages were clearly present in the cytoplasmic and nuclear TDP inclusions (Fig. 7B and C). These results indicate that mixed polyUb chains (Lys-48 and Lys-63) are specifically associated with TDP protein aggregates.

To confirm the presence of Lys-48 and Lys-63 polyUb linkages observed in TDP-S6 cytoplasmic aggregates, we analyzed the levels of polyUb linkages in the Sarkosyl-insoluble proteome by targeted mass spectrometry (36, 39). Metabolically heavy labeled proteins were used as internal standards for their unlabeled counterparts, allowing the quantification of native proteins in each sample. Trypsin cleaves endogenous polyUb chains and their corresponding internal standards to generate linkage-specific peptides tagged with two Gly residues (51). During the LC/SRM analysis, heavy and light peptides co-eluted and were fragmented to produce product ion pairs (Fig. 8, A and B), which were used as surrogates to provide sensitive measurements of relative protein abundance. Among all polyUb linkages, Lys-63 increased ~3.3-fold in the insoluble fraction of TDP-S6-expressing cells over that of mock-transfected cells. In addition, Lys-48 link-
ages (~2.0-fold), Lys-11 linkages (~1.5-fold), and Lys-29 linkages (~1.3-fold) were also measured in this sample (Fig. 8C), whereas the other four potential linkages mediated by three lysines (Lys-6, Lys-27, and Lys-33) and the N terminus were not detected. In the TDP-S6 insoluble fraction, all measured linkages other than Lys-29 were significantly increased. In comparison, within the TDP-43 insoluble fraction, Lys-63 was the only polyUb linkage significantly increased (~2.2-fold) over the level in mock-transfected cells. This is in good agreement with the localization of Lys-63-linked polyUb and nuclear TDP-43 bodies by immunofluorescence in HEK-293 cells expressing a high level of TDP-43 (Fig. 7C). These data further support the involvement of multiple polyUb chains in TDP protein aggregation and indicate that the Lys-63 polyUb linkage may be the most significant form associated with TDP-43 or TDP-S6 overexpression.

**DISCUSSION**

The abnormal accumulation of phosphorylated and ubiquitinated species in protein inclusions is observed in a wide variety of neurodegenerative diseases (3, 4). In Alzheimer and Parkinson diseases, these intracellular inclusions are called neurofibrillary tangles and Lewy bodies and are composed of pathologically altered forms of tau and α-synuclein, respectively (3, 5). In FTLD-U and ALS, TDP-43 has been identified as the major component of tau- and α-synuclein-negative inclusions where it is both phosphorylated and ubiquitinated (5). Although there are 10 known splice variants of human TDP-43, their roles in abnormal TDP-43 translocation and pathologic aggregation have not yet been characterized. In this study, we expressed human recombinant TDP-43 and a truncated splice variant, TDP-S6, in HEK-293 cells and pri-
mary neurons. The full-length protein was expressed almost exclusively in the nucleus where it co-localized with endogenous TDP-43. In sharp contrast, the shorter TDP-S6 formed highly insoluble cytoplasmic and nuclear inclusions reminiscent of disease-specific pathology. Moreover, overexpression of both TDP-43 and TDP-S6 resulted in the enrichment of specific post-translational modifications within their respective insoluble protein extracts. These post-translational modifications included ubiquitination, phosphorylation, and TDP-43-specific C-terminal fragmentation. Based on our multiplex SILAC proteomics approach, we identified the significant enrichment of SUMO-2/3 and Ub within the Sarkosyl-insoluble, urea-soluble extracts.

In our cellular model, endogenous Ub showed strong association with the TDP aggregates in both biochemical analysis and immunostaining, supporting the hypothesis that aggregation-prone TDP-43 is a major ubiquitinated species.

Although TDP-S6-expressing cells displayed inclusions of various sizes, only a fraction of recombinant TDP-43-expressing cells formed small nuclear aggregates dependent on high level expression (Fig. 7). Quantitative MS analysis revealed that several main polyUb chains were present in the insoluble extracts from these cells with Lys-63 linkages being the most up-regulated. These findings are reminiscent of previous characterization of tau- and SOD1-positive inclusions (49) and are consistent with the idea that Lys-63-linked chains may facilitate the formation of inclusions and direct subsequent clearance by the autophagy pathway.

Recent studies have suggested that ubiquitinated inclusions in FTLD-U brains are primarily composed of C-terminal fragments of TDP-43 and that these fragments may comprise the primary pathologic species of TDP-43 aggregates by serving as a seed for aggregation (53). Moreover, caspase-3 activation has been implicated in the proteolytic processing of TDP-43 into three C-terminal fragments similar to those seen...
SUMOylation assay, we provide experimental evidence that partial SUMO-targeted lysines (56). Using an residue, amino acids 135–138 of TDP-43) among its potential components in the formation of protein inclusions. SUMO-2/3 was found to be preferentially enriched within the TDP-43 and TDP-S6-expressing cells, but further studies using targeted MS analysis will be necessary to validate enzyme-specific TDP-43 cleavage.

Currently, it remains unknown whether aberrant splicing of the TDP-43 transcript results in the production of pathologic TDP isoforms in ALS or FTLD-U. However, two TDP-43 splicing isoforms have been identified previously in human brain and spinal cord tissue (55). One of these isoforms, which closely resembles TDP-S6, was identified in two ALS cases. This novel transcript lacked all of exon 3 and a significant portion of exon 6, resulting in expression of a protein product (~28 kDa) without a glycine-rich C terminus (55). In this study, TDP-S6, a C-terminal truncated splice variant previously identified in HEK-293 cells (11), was found predominately localized to the cytoplasm and mimicked the aggregation observed in FTLD-U and ALS. It should be noted that although the human TDP-S6 variant formed both cytoplasmic and nuclear inclusions, overexpression of the mouse TDP-S6 in HEK-293 cells resulted only in the formation of nuclear specific speckle-like structures, termed TDP bodies (40). Thus, despite sharing 96% amino acid identity, inconsistency between mouse and human TDP-S6 localization may result from differences in species-specific nuclear/cytoplasmic translocation or different experimental conditions.

SILAC analysis of this cellular model revealed novel components in the formation of protein inclusions. SUMO-2/3 was found to be preferentially enriched within the TDP-43 and TDP-S6 insoluble proteomes and was subsequently validated by Western blotting. Interestingly, immunofluorescence confocal microscopy indicated that SUMO-2/3 co-localized mainly with TDP-S6 nuclear inclusions, thereby implicating a unique nuclear association between SUMO-2/3 and TDP-S6. This nuclear association is not completely surprising because TDP-43 has been previously found to associate with promyelocytic leukemia bodies (40), nuclear structures known to co-localize with SUMO (27, 48). Moreover, sequence analysis of TDP-43 revealed a canonical SUMO conjugation site motif (ΨKX(D/E); where Ψ indicates hydrophobic and χ indicate any residue, amino acids 135–138 of TDP-43) among its potential SUMO-targeted lysines (56). Using an in vitro deSUMOylation assay, we provide experimental evidence that TDP-43 itself is directly SUMOylated within the insoluble fraction. This is also supported by a recent large scale study wherein TDP-43 was identified as a SUMO-2 conjugate that was accumulated over 7-fold in response to heat shock (57). Notably, protein SUMOylation has also been linked to aggregated proteins in Alzheimer disease, Parkinson disease, and poly(Q) disorders (58). In addition, SUMO-1 has been shown to specifically co-localize with ubiquitinated nuclear neuronal inclusions in FTLD-U tissues (59). Although protein SUMOylation functions in regulating transcription and nuclear transport (60), it has also been shown to cross-talk with ubiquitination because the SUMO tag on proteins may serve as a recognition signal for subsequent ubiquitination and proteasome-mediated degradation (61–63). More specifically, the recent identification of SUMO-targeted ubiquitin ligases (STUbLs) (64) raises the intriguing possibility that upon SUMOylation misfolded nuclear TDP-S6 is subjected to sequestration in nuclear aggregates and STUbL-dependent ubiquitination and degradation (65). Interestingly, a STUbL-mediated mechanism for the cellular regulation of insoluble TDP-S6 is supported by the immunochromeproton of polyubiquitin chains with both nuclear and cytoplasmic TDP-S6 inclusions (Fig. 7). The mechanism of TDP-43 SUMOylation underlying disease-specific protein translocation, degradation, or aggregation is worth further investigation.

In addition to Ub and SUMO-2/3, actin-related protein 1 (ARP1) was also found to be enriched in the Sarkosyl-insoluble fractions of TDP-43- and TDP-S6-transfected cells. ARP1 is a subunit of dynactin, a macromolecular complex that interacts with both microtubules and cytoplasmic dynein. ARP1 is also involved in protein transport and vesicular trafficking (66). Interestingly, gene mutation in another subunit of dynactin, p150Glued, has been reported in ALS cases (67), and the related mouse model developed motor neuron disease (68). In addition, overexpression of TDP-S6 reduces N-ethylmaleimide sensitive factor, an ATPase involved in intracellular trafficking, and dynamin 3, a microtubule-associated protein involved in vesicle budding (69). These results suggest that protein transportation may be dysregulated in neurodegenerative diseases. Furthermore, these findings, coupled with an increase in Lys-63 polyUb linkages, further support an association between TDP-43 and protein sorting pathways.

In sum, our data show that the C-terminal domain is necessary for TDP-43 nuclear localization and that without this region TDP-43 becomes primarily sequestered within cytoplasmic aggregates, consistent with a previous report showing that the C terminus is essential for the solubility and cellular localization of TDP-43 (50). Thus, it might be plausible that the genetic mutations in TDP-43 found within the C-terminal region (21–25) somehow disrupt critical amino acid residues needed for TDP-43 nuclear localization and proper function. Although further studies are needed to evaluate the presence of TDP-S6 or other similar short isoforms in disease tissue, our data raise the intriguing possibility that dysregula-
tation of human TDP-43 alternative splicing or preferential pro-
teolytic processing to produce TDP N-terminal fragments can
contribute to the pathology of FTLD-U and ALS.

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