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Neurons selectively targeted in frontotemporal dementia reveal early stage TDP-43 pathobiology

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Abstract

TAR-DNA binding protein 43 (TDP-43) aggregation is the most common pathological hallmark in frontotemporal dementia (FTD) and characterizes nearly all patients with motor neuron disease (MND). The earliest stages of TDP-43 pathobiology are not well characterized, and whether neurodegeneration results from TDP-43 loss-of-function or aggregation remains unclear. In the behavioral variant of FTD (bvFTD), patients undergo selective dropout of von Economo neurons (VENs) and fork cells within the frontoinsular (FI) and anterior cingulate cortices. Here, we examined TDP-43 pathobiology within these vulnerable neurons in the FI across a clinical spectrum including 17 patients with sporadic bvFTD, MND, or both. In an exploratory analysis based on our initial observations, we further assessed 10 patients with *C9orf72*-associated bvFTD/ MND. VENs and fork cells showed early, disproportionate TDP-43 aggregation that correlated with anatomical and clinical severity, including loss of emotional empathy. The presence of a

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A.L.N., M.S., J.L.H., S.E.G., L.L., E.J.K., and Y.P. performed the experiments and cell counting. M.S. performed morphometric analysis. A.L.N., L.P., and I.E.A. analyzed the data. W.W.S., L.T.G., and E.J.H. performed neuropathological diagnoses. D.H.G and G.H.C. performed genetic analysis. J.H.K, K.P.R, G. T, B.L.M., and W.W.S obtained behavioral and clinical data. W.W.S., S.E.G., and A.L.N. conceived and designed the project. W.W.S., and A.L.N. wrote the paper. All authors reviewed the manuscript.

TDP-43 inclusion was associated with striking nuclear and somatodendritic atrophy. An intriguing minority of neurons lacked detectable nuclear TDP-43 despite the apparent absence of a cytoplasmic TDP-43 inclusion. These cells showed neuronal atrophy comparable to inclusion-bearing neurons, suggesting that loss of nuclear TDP-43 function promotes neurodegeneration, even when TDP-43 aggregation is inconspicuous or absent.

Keywords

frontotemporal dementia (FTD); amyotrophic lateral sclerosis (ALS); TAR DNA-binding protein 43 (TDP-43); *C9orf72*; von Economo neuron (VEN); empathy

Introduction

Frontotemporal dementia (FTD) and motor neuron disease (MND) represent the endpoints of a clinicopathological continuum linked to TAR DNA-binding protein 43 (TDP-43) proteinopathy, which is found in more than 50% of patients with bvFTD and almost all patients with FTD-MND and ALS [52]. Within this spectrum, some patients present with FTD, others MND, and still others develop a blended syndrome (FTD-MND) [27]. TDP-43 is a DNA/RNA-binding protein [13] expressed in all healthy neurons; in FTD/MND, neuronal cytoplasmic TDP-43 aggregation is accompanied by a loss of normal nuclear TDP-43. This observation raises the question of which factor drives neurodegeneration: TDP-43 aggregate toxicity, loss of TDP-43 function, or both. Although model-based experiments have begun to inform this debate [5, 32, 77, 82, 84], there remains no consensus yet little work has been done to address this question in human tissues.

In MND, degeneration targets the upper and lower motor neurons within the pyramidal motor system. In FTD, the vulnerable circuits and cell types have just begun to emerge within the past decade. In the behavioral variant (bvFTD), the most common FTD syndrome, early degeneration involves the anterior cingulate (ACC) and ventral anterior insular (i.e., frontoinsular, FI) cortices [8, 66, 87], regardless of the underlying pathological cause [54]. These regions serve as the cortical hubs of a large-scale "salience network" [67] critical for social-emotional-autonomic function and are home to the morphologically distinctive von Economo neurons (VENs) [68, 79] and fork cells [53]. We and others have shown that these large Layer 5 projection neurons undergo early dropout in bvFTD [34, 63-65, 86], but, to date, no bvFTD study has assessed VEN and fork cell pathological changes, such as protein aggregation or neuronal atrophy, occurring prior to neuronal death.

In the present work, we sought to determine whether VEN and fork cell vulnerability in bvFTD due to FTLD-TDP is associated with early TDP-43 aggregation and whether aggregation relates to anatomical and clinical measures of disease severity. To pursue these goals, we studied patients across the bvFTD-MND spectrum, reasoning that patients with pure MND or early bvFTD-MND may represent an early or arrested stage of bvFTD progression. In the course of this work, we recognized an important minority of VENs, fork cells, and other neurons that lack detectable nuclear TDP-43 despite the apparent absence of a cytoplasmic inclusion. These neurons, here termed "nuclear TDP-43-depleted", have been recently noted in FTD and MND by our group and others [6, 7, 78], but they have yet to be

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systematically characterized. Therefore, we also studied the morphology of nuclear TDP-43depleted and TDP-43 inclusion-bearing neurons to determine whether the reduction of nuclear TDP-43 alone is associated with neuronal degeneration.

Materials and methods

Human brain material and subjects

Post-mortem human brain tissue was obtained from the USCF Neurodegenerative Disease Brain Bank. Consent for brain donation was obtained from all subjects or their surrogates in accordance with the Declaration of Helsinki and the research was approved by the University of California, San Francisco Committee on Human Research. Depending on where the autopsy was performed, brains were immersion fixed whole in 10% buffered formalin or cut fresh into 1 cm thick coronal slices. For freshly cut brains, alternate slices were fixed for 48-72 hours in 10% buffered formalin or frozen. Clinical diagnoses of bvFTD were made according to prevailing international consensus criteria at the time of assessment [51, 60]. Clinical diagnoses of ALS were made according to El Escorial Criteria [9, 10]. The presence of MND in patients with bvFTD was assessed by a neurologist based on the neurological examination, usually supported by electrodiagnostic testing, and was recorded prospectively in the Memory and Aging Center database. Neuropathological diagnoses were made following consensus diagnostic criteria [41, 42, 45-47] using previously described histological and immunohistochemical methods [34, 76]. Cases were selected based on clinical and neuropathological diagnoses and genetic analysis (Suppl. Figure 1). Patients were required to have (1) a primary clinical diagnosis of bvFTD, bvFTD-MND, or amyotrophic lateral sclerosis (ALS)/MND, and (2) a neuropathological diagnosis of FTLD-TDP, Type B or U, with or without MND, or ALS-TDP. Within the FTLD-TDP spectrum, Type B, characterized by speckled or dash-like neuronal cytoplasmic TDP-43 inclusions (NCIs), and an unclassifiable "Type U", with insufficient or atypical TDP-43 inclusions, are the most common subtypes in bvFTD \pm MND [41]. We limited the analyses to these types (Suppl. Figure 1) to focus on a more homogeneous pathological substrate, allowing us to make better inferences about stage-wise and clinical symptom relationships. Exclusion criteria were presence of a disease-causing mutation other than a *C9orf72* expansion, Braak neurofibrillary tangle stage > 4, transitional limbic or diffuse neocortical Lewy body disease, and a post-mortem interval > 48h. Control subjects were individuals with no clinical history of neuropsychiatric disorder. Based on these criteria, we studied 39 subjects in total (see Table 1, Table 2, and Suppl. Figure 1 for details). 17 patients across the bvFTD-MND spectrum with no known FTD-related mutation and primary TDP-43 pathology were studied, alongside 10 age- and sex- matched non-neurological controls. To enable blinded characterization of patients with and without the C9orf72 expansion, we included an additional 10 patients (mean age 60, 2 females) with the expansion who met all other inclusion criteria. Two additional patients with C9orf72 expansion were included for morphometric assessment only (Table 2). Anatomical disease stage was assessed using an FTD rating scale [8], as previously described. Anatomical staging of all patients was performed by a single blinded investigator (W.W.S).

Clinical and behavioral assessment

Symptom duration was estimated from the clinical records. Clinical measures were obtained as previously described [34]. Briefly, clinical severity was assessed using the Clinical Dementia Rating (CDR) scale total and Sum of Boxes scores (CDR-SB), using a version of the CDR adapted for FTD [35]. In addition, we chose a validated measure of emotional empathy, the empathic concern subscale of the interpersonal reactivity index (IRI) [17], as completed by the patient's informant. Loss of empathy is a core diagnostic feature of bvFTD, and empathic concern deficits in bvFTD are related to degeneration of anterior insula, anterior cingulate/medial prefrontal, and anterior temporal regions involved in bvFTD [59, 69].

Specimens and tissue processing

Blocks of FI were dissected from ~1 cm thick formalin-fixed coronal slabs (right, left or bilateral, as available, see Table 2 for details) and stored until processed for sectioning and immunohistochemistry. Blocks were removed from storage in formalin or PBS with 0.02% sodium azide (PBS-Az) and cryoprotected in graded sucrose solutions (10, 20 and 30% sucrose in PBS-Az) then sectioned at 50 µm thickness on a freezing microtome. Every 12th section was Nissl-stained with cresyl violet (FD Neurotech) for neuron counts and to determine the anatomical boundaries of the FI, the cortical region of interest.

Immunohistochemistry

Three sections were selected from each block for immunohistochemical staining. Prior to staining, all sections were coded for rater blinding. Free-floating sections were washed (6 \times 10 min) in 0.01 M phosphate buffered saline and underwent antigen retrieval in 10 mM citrate buffer pH 6.0 for 2 hours in an 80 °C water bath. Endogenous peroxidase activity was quenched with 3% hydrogen peroxide in PBS-Az for 30 min. Sections were then washed (3 \times 10 min) in PBS and non-specific staining was blocked by incubating in 5% nonfat milk powder (Carnation) in PBS + 0.25% Triton X-100 (PBT) for 1 h. For TDP-43 immunostaining, sections were incubated with an antibody against TDP-43 (rabbit polyclonal, 1:5,000, Proteintech, RRID: AB 615042) in PBT-Az with gentle agitation (orbital shaker) overnight at room temperature. Following incubation with primary antibody, sections were washed (6×10 min) in PBS, incubated for 1 h in biotinylated secondary antibody (goat anti-rabbit IgG, 1:500 in PBT, Vector), washed in PBS, incubated for 1 h in ABC (Vectastain ABC elite kit, Vector Laboratories, 1:500 each in PBS), washed, then exposed to 0.05% 3,3-diaminobenzidine tetrahydrochloride (Sigma) and 0.01% H_2O_2 to produce a brown reaction product. The sections were washed and mounted onto plus slides. Finally, sections were dehydrated through a graded ethanol series, cleared in xylenes, and coverslipped with Permaslip. TDP-43 stained sections were counterstained with hematoxylin (Fisher).

Immunofluorescence staining and confocal imaging

Multiple immunofluorescence staining was used to further characterize TDP-43 inclusions. Five-color immunofluorescence labeling was performed using fluorescent Nissl to label nuclei, anti-MAP2 (chicken polyclonal, 1:3000, EnCor Biotechnology Inc., RRID:

AB 2138173) to label neuronal somata and proximal dendrites, a pan TDP-43 antibody (mouse monoclonal, 1:30,000, Proteintech, RRID: AB_2200520) to label physiological (nuclear) TDP-43 and TDP-43 inclusions, phosphorylated TDP-43 (rat monoclonal, 1:500, courtesy Manuela Neumann, RRID: AB_2315226), and ubiquitin (rabbit polyclonal, 1:500, DAKO, RRID: AB_2315524) antibodies. Sections were initially photobleached to reduce autofluorescence. Free-floating tissue sections were immersed in PBS-Az, carefully positioned to lay flat, and photobleached for 72 h at 4 °C using a 240 W wide-spectrum light (HTG Supply LED-6B240) with an array of 80 3 W LEDs emitting at 6 bands: 390 nm, 430 nm, 460 nm, 660 nm, 630 nm and 1000 K white/blue light. Sections were then processed for antigen retrieval, blocked, and incubated with the first batch of primary antibodies (anti TDP-43 and anti-MAP2) overnight at room temperature. The next day, sections were washed, incubated with secondary antibodies conjugated to either a fluorescent dye or biotin, washed, then incubated with streptavidin-conjugated pacific orange. The tissue was then sequentially labelled with the second batch of primary antibodies (anti-phosphorylated TDP-43 and anti-ubiquitin), washed, again incubated with the appropriate Alexa Fluorconjugated antibodies, counterstained with Neurotrace blue fluorescent Nissl, mounted onto slides, and dried. Sections were finally incubated with 0.01% Sudan black B to further reduce autofluorescence and coverslipped using Fluoromount G (Southern Biotech). Sections were imaged using a Zeiss LSM 880 confocal laser scanning microscope equipped with 405 (to image blue fluorescent Nissl, and pacific orange), 488 (Alexa Fluor 488), 561 (Alexa Fluor 568), and 633 (Alexa Fluor 647) lasers, and a filterless detection system with customizable emission bandwidths, enabling 5 channels to be imaged without spectral overlap.

ROI definition, image acquisition, and neuron quantification

Layer 5 of area FI was defined as previously described [34]. The extent of the agranular, VEN- and fork cell- containing FI was determined in cresyl-violet stained slides, and FI Layer 5 and adjacent Layers 2-3 were traced using Stereo Investigator (MBF Bioscience; system includes motorized stage and Nikon 80i microscope), mainly using an objective with 10x magnification. To obtain systematic, unbiased neuron counts we used Stereo Investigator's Optical Fractionator probe, in which neurons with nucleoli contained within a three-dimensional counting frame (or crossing the inclusion lines) were counted. Limited tissue availability prevented analysis of the entire FI; thus we did not aim to obtain stereological estimates of the entire population of FI neurons. VEN and fork cell densities and inclusion fractions were assessed in approximately 700 sampling sites, determined by our previous study [34] and pilot data to be sufficient to achieve a coefficient of error of \sim 0.1, from three sections spread out across the entire available FI (see Suppl. Tables 1 and 2 for details). Because VENs and fork cells were counted in the same counting frame, we based the number of sampling sites on the number required to achieve a coefficient of error ~ 0.1 for VENs, and some of the coefficients of error for fork cell density were greater than 0.1. Because neighboring Layer 5 and layer 2-3 neurons are far more abundant, approximately 50-70 frames were counted per subject to assess these neuron densities and inclusion fractions. Nissl-based neuronal density estimates from subjects N1-N3, S1-S4, and C3 have been published previously [34] and are included in the present study for completeness of the neuronal loss data set. VEN, fork cell, and neighboring neuron loss in

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Layer 5 of the FI was assessed in Nissl-stained sections from patients and controls with a 60×, N.A. 1.4 oil immersion objective. We similarly assessed the proportion of each cell type with TDP-43 inclusions among total Layer 5 VENs, forks, and neighboring Layer 5 neurons and Layer 2-3 neurons, but used digitized "virtual" sections for improved workflow and blue/brown color discrimination. To this end, we scanned TDP-43 immunostained sections adjacent to each Nissl-stained sections; for each section, the regions of interest (FI Layer 5 and Layer 2-3) were traced in Stereo Investigator and exported into Zen software (Zeiss). An AxioObserver Z1 microscope (Zeiss) equipped with a motorized stage was used to acquire and montage digital, high magnification (63×, N.A. 1.4 oil immersion objective), image stacks (z-stack interval: 0.4 µm) of the entire FI Layer 5. Stitched montages were saved as Zeiss CZI images, exported as TIFF files, then compressed to create virtual JPX files for viewing in Stereo Investigator. The numbers of VENs, fork cells, neighboring Layer 5 neurons, and Layer 2-3 neurons with and without TDP-43 inclusions were then counted to obtain the percentage of inclusion-bearing neurons (i.e. "inclusion fraction" in each population. While counting, we also noted all neurons that showed loss of nuclear TDP-43 in the absence of a visible cytoplasmic inclusion ("nuclear TDP-43-depleted" neurons).

Cortical thickness measurements

Because apparent local density measurements can be artificially inflated by tissue volume loss, we measured cortical thickness in FI to estimate overall degeneration. Cortical boundaries of the agranular, VEN- and fork cell-containing FI were traced using Stereo Investigator software on cresyl violet-stained sections by following the superficial border of Layer 1 and the deep border of Layer 6 where it meets the white matter. The shortest distance between the traced Layer 1 and Layer 6 borders was measured at five points evenly spaced throughout the traced region of interest. Sulci and obliquely cut regions were excluded.

Interrater and intrarater comparisons

To obtain Layer 5 VEN/total neuron and fork cell/total neuron ratios, and VEN, fork cell and neighboring Layer 5 neuron densities, three blinded raters (E.J.K, L.L. and Y.P) performed counting of Nissl-stained sections after training to predefined competency criteria, as described [34], which are designed to promote inter-rater consistency. Intrarater reliability was assessed independently by each rater every 12 sections. Six sections were recounted by each Nissl rater for intraclass correlation analysis. Intra-rater correlation coefficients across the three raters averaged 0.990 (95% CI 0.923-0.999) for VENs, 0.995 (95% CI 0.959-0.999) for fork cells, and 0.987 (95% CI 0.901-0.999) for neighboring Layer 5 neurons.

To assess TDP-43 inclusions and depletion in VENs, fork cells and neighboring Layer 5 neurons, three blinded raters (S.E.G., A.L.N., and J.H.H.) performed counting of TDP-43 stained sections after training to predefined competency criteria. For TDP-43 inclusion counts, competency was defined as VEN, fork, and neighboring Layer 5 neuron numbers, as well as ratio of affected/total neurons for each cell type, within 10% of values generated for a training section counted by a senior investigator (W.W.S.). Intra-rater reliability was assessed independently by each rater every 6 sections for a total of 6 sections per rater. Intra-

class correlation coefficients for intra-rater reliability of inclusion fraction measurements were high, averaging 0.973 (95% CI 0.905-0.993) for VENs, 0.997 (95% CI 0.988-0.999) for fork cells, 0.978 (95% CI 0.840-0.998) for neighboring Layer 5 neurons, and 0.998 (95% CI 0.985-1.000) for Layer 2-3 neurons across raters.

Morphometry

Cell type-tailored neuronal morphology was assessed in a subset of age- and sex-matched subjects (seven sporadic bvFTD±MND, two C9orf72-ALS, five C9orf72-bvFTD±MND, and 4 controls, see Table 2 for subject details and Suppl. Table 3 for group demographics) selected because their tissue fixation was long-term, ideal for a detailed morphometric study. In studying all subjects with suitable tissue, two subjects were included in the C9orf72 group that had not been included in the blinded C9orf72 cohort for density or inclusion fraction analysis. Control subjects were included to determine whether inclusion-lacking neurons in patients were altered compared with neurons in controls. Using Stereo Investigator and the same digitized slides used for TDP-43 counts, we manually measured apical and basal dendritic diameter and nuclear and somatic area (Suppl. Figure 2) in selected VENs, fork cells, and neighboring Layer 5 neurons. TDP-43 normal cells were selected in a single section by systematic random sampling of Layer 5 to yield a total of 10 neurons per subject. For identification of neurons with TDP-43 neuronal cytoplasmic inclusions (NCIs) or TDP-43 nuclear depletion without NCIs, Layer 5 of a single section of was exhaustively scanned using the meander scan function in Stereo Investigator. If the exhaustive search of Layer 5 of a single section yielded less than 10 neurons of each cell type per group, additional sections were added until at least 10 neurons were identified in each group, or until all three TDP-43 stained virtual sections had been exhaustively searched. If more than 10 neurons per cell type per cell type and group were identified, excess neurons were randomly eliminated. Because neurons with TDP-43 nuclear depletion without an NCI were the rarest group, all identified neurons in this category were included for analysis. Nuclear and somatic areas were obtained by focusing through the cell and tracing the outline of the nucleus or soma at the widest part of the neuron. Apical and basal dendritic widths were measured at the maximum width of the dendrite at the junction between the soma and dendrite. For fork cells, the width of both apical dendrites was measured and the results averaged to yield one apical dendrite measurement per neuron. For pyramidal cells the width of the largest visible basal dendrite was measured.

Statistical analysis

Statistical analysis was performed using R version 3.4.3 in RStudio 1.1.383. Normality was assessed using parametric Q-Q plots and the Shapiro-Wilk normality test. Neuronal loss, inclusion fractions, and morphometry data were assessed using linear mixed-effects models using the R function "lmer" in the lme4 package (http://www.r-project.org). For all models, the only random-effect in the model was subject-specific random intercepts, with all other effects considered as fixed. To examine whether there were significant differences in cell loss between patients with sporadic disease and controls, we used linear mixed-effects models with log transformed neuronal density or fraction of total neurons as outcomes and clinical diagnosis as a primary categorical predictor with cerebral hemisphere and cortical thickness as covariates. To assess differences in TDP-43 inclusion fraction or TDP-43

depletion between cell types, we used a linear mixed-effects model with inclusion fraction as the outcome and cell type as a primary categorical predictor with hemisphere, Broe stage, and clinical diagnosis as covariates. To investigate whether VEN, fork cell, or neighboring Layer 5 neuron density was altered in patients with a C9orf72 mutation we used a linear mixed-effects model with neuronal density as the outcome and clinical diagnosis as a primary categorical predictor with cerebral hemisphere, cortical thickness, and C9orf72 status as covariates. TDP-43 inclusion fraction and TDP-43 depletion in C9orf72 bvFTD/MND were assessed using the linear mixed effect model described above for sporadic bvFTD/MND but with C9orf72 status as an additional fixed effect. The effect of TDP-43 inclusions on cell morphology was tested with logistic mixed-effects models with each morphometric parameter as an outcome, TDP-43 status as the primary categorical predictor, and C9orf72 status and clinical diagnosis as covariates. Normality of residuals was assessed for each model using a Q-Q plot and followed a normal distribution. For models assessing differences in percentages of each neuronal type with TDP-43 inclusions or depletion, residuals following a near normal distribution were accepted without log transforming the data. Normal approximations were used to find p values for mixed effect models. A p < 0.05 (two-tailed) was considered significant. Significant interactions were further explored using the lsmeans package in R; p values for pairwise comparisons were adjusted using the Tukey method for multiple comparisons. Correlations between inclusion fractions and clinical and behavioral measures were assessed by Spearman partial correlation. Because clinical and behavioral measures were taken at a range of intervals prior to death we controlled for time between clinical assessment and death, with the assumption that neuropathological features progress linearly over time.

K-means clustering [39], a nonhierarchical clustering method, was used in this study to obtain class memberships for the measured cells. K-means uses an iterative algorithm to update randomly selected initial cluster centers and to obtain cluster membership for each object, assuming defined boundaries between clusters. Initially, objects are assigned to a cluster on the basis of a minimal distance value. Next, centroids of the clusters are calculated and distance values between the centroids and each of the objects are recalculated. If the closest centroid is not associated with the cluster to which the object currently belongs, the object will switch its cluster membership to the cluster with the closest centroid. The centroid's positions are recalculated every time a component has changed its cluster membership. This process continues until none of the objects has been reassigned using the squared Euclidean distance metric [36, 44]. In this study, measured VENs were classified into three clusters, based on their nucleus and soma areas, and apical and basal dendritic diameters. A setting of 100 iterations and 10 replications was applied. K-means clustering was performed in MATLAB R2017a (https://www.mathworks.com/products/matlab.html) using the kmeans function. The cross-tabulation function in MATLAB was used to determine whether VENs were unevenly distributed across clusters by TDP-43 status.

To explore how each neuronal compartment (i.e. soma, nucleus, or apical or basal dendrite) was affected by TDP-43 aggregation or depletion, morphometric data for each compartment were transformed into z scores referenced to data from that compartment in neurons from control subjects. Z scores for each neuron were then ranked from highest to lowest, with VENs, fork cells, and neighboring layer 5 neurons assessed separately. Chi-square tests were

performed in RStudio to determine whether there was a relationship between TDP-43 inclusion/depletion status and compartment rank for each neuronal type, followed by pairwise comparisons between compartments corrected for multiple comparisons using false discovery rate.

Results

To gain novel insights into TDP-43 pathobiology in humans, we sought to evaluate the earliest clinical stages of sporadic bvFTD, focusing on neuronal populations known to undergo early, selective degeneration. We included patients with ALS having no or few behavioral changes, those with mild bvFTD who developed and died of MND, and those with a full-blown bvFTD syndrome, with or without MND. Although bvFTD can result from the full FTLD pathological spectrum, including FTLD-tau or -FUS, here we included only patients with underlying FTLD-TDP or ALS-TDP.

Cortical atrophy, thinning, and neuronal density in the sporadic bvFTD-MND spectrum

To confirm that our cohort represented a range of stages beginning with limited neuronal loss, we assessed atrophy severity according to a staging system developed for bvFTD [8]. Patients were scored as having no atrophy (stage 0, n = 6), very mild atrophy (stage 1, n = 8), or mild atrophy (stage 2, n = 3). No patient had moderate (stage 3) or severe (stage 4) atrophy. As expected, cortical thickness decreased with progressive Broe stage (Suppl. Figure 3a) and was therefore incorporated into statistical models to adjust for neuronal density increases that result from cortical thinning.

As anticipated, frontoinsular VEN and fork cells showed only mild, statistically nonsignificant reductions in neuronal density (VENs, 32% loss, t = -1.87, p = 0.061; fork cells, 24% loss, t = -1.70, p = 0.089; Suppl. Figure 3b). VEN and fork cell counts as a proportion of total Layer 5 neurons showed similar findings (Suppl. Figure 3c). Neighboring Layer 5 neurons (Suppl. Figure 3b) showed normal densities (t = 1.43, p = 0.15). Laterality had no significant effect on neuronal density. Thus, our diverse patient cohort, ranging in clinical presentation from pure ALS to pure bvFTD, represents an earlier stage of VEN and fork cell degeneration than reported in our previous smaller study of bvFTD due to FTLD-TDP, FTLD-tau, or FTLD-FUS, in which we observed 53% VEN and 68% fork cell loss in FI [34]. Grouping neuron loss by Broe stage (Suppl. Figure 3d) or clinical diagnosis (Suppl. Figure 3e) suggested progressive VEN and fork cell loss with advancing severity, with 40-50% loss by Broe stage 1 (Suppl. Figure 3d). An apparent increase in the ratio of VENs and fork cells to neighboring Layer 5 neurons at Broe stage 2 likely reflected pyramidal neuron dropout as the disease progresses.

Von Economo neurons and fork cells show early, disproportionate TDP-43 aggregation in the sporadic bvFTD-MND spectrum

Having established that our study represents a range of neurodegeneration severities starting from no or minimal neuronal loss, we asked whether and at what stage VENs and fork cells undergo TDP-43 inclusion formation. Visual review of the materials suggested a striking propensity of VENs and fork cells to form TDP-43 neuronal cytoplasmic inclusions in

comparison with neighboring Layer 5 neurons. An exhaustive map of an FI section from a representative patient with bvFTD-ALS at Broe stage 1 (Fig. 1a) reveals the distribution of normal and TDP-43 inclusion-bearing VENs and fork cells, which were distributed throughout Layer 5 (Figs. 1a, 1b). VENs and fork cells most frequently presented with punctate/diffuse "speckled" or dash-like TDP-43 neuronal cytoplasmic inclusions typical of FTLD-TDP, Type B (Fig. 1c). To characterize the post-translational modifications of the characteristic "speckled" TDP-43 inclusions in VENs and fork cells, we performed multiple immunofluorescence labelling for TDP-43, phosphorylated TDP-43, and ubiquitin (Fig. 1d). In patients with ALS, the VEN and fork cell TDP-43 inclusions adopted dash-like or reticular, mesh-like structures in the cytoplasm; these deposits were almost always phosphorylated, at least in part, with sparse, punctate ubiquitin labelling across the TDP-43 mesh-like structure. In patients with bvFTD, VEN inclusions were also reticular and phosphorylated but contained larger, more globular areas of ubiquitination not part of the mesh-like TDP-43 inclusion structure. We did not observe VENs with phosphorylated TDP-43-positive inclusions that were stained negatively using the pan-TDP-43 antibody. To our initial surprise, but in keeping with recent studies from our group [78] and others [6, 7]. we also observed neurons lacking normal nuclear TDP-43 despite the absence of any visible cytoplasmic inclusion on brightfield imaging (Fig. 1c). Although no TDP-43 inclusions were identified in these neurons, they may contain small amounts of cytoplasmic TDP-43 below the detection threshold of our antibody, but the degree of nuclear TDP-43 depletion was unambiguous in comparison to other neurons in the same patient or in controls.

Systematic, unbiased counting confirmed that, despite the mild neuronal loss, VENs (t = 8.20, p < 0.0001) and fork cells (t = 8.98, p < 0.0001) were three- to five-fold more prone to TDP-43 inclusion formation than were neighboring Layer 5 neurons (Fig. 2a). This effect was observed in both the left and right FI (inclusion fractions, left: VENs, 25%; fork cells, 28%; neighboring Layer 5 neurons, 6%; inclusion fractions, right: VENs, 26%; fork cells, 26%; neighboring Layer 5 neurons, 8%) (Figs. 2a and Suppl. Figure 4a). Nuclear TDP-43-depleted neurons showed the same selectivity pattern across cell types (Fig. 2b), with significantly higher rates of nuclear TDP-43 depletion found in VENs (t = 4.14, p < 0.0001) and fork cells (t = 4.48, p < 0.0001) than in neighboring Layer 5 neurons, although rates of nuclear TDP-43 depletion were generally low and varied between subjects (Suppl. Figure 4a). Consistent with our current and previous [34] studies of FI VEN and fork cell loss, which showed no distinct hemispheric pattern, we saw no consistent difference in inclusion fraction between left and right FI (t = -1.20, p = 0.23) in the ten patients for whom bilateral FI was available. Some showed right-predominant, others left-predominant, and the remainder symmetrical TDP-43 inclusion formation rates (Suppl. Figure 4b).

Superficial cortical layers show prominent neuropil degeneration [11, 12] and inclusion burden [14, 40, 62] in FTLD-TDP. We therefore compared inclusion burden in FI Layer 2-3 neurons to that seen in FI Layer 5 neurons. Layer 2-3 neurons were significantly more prone to TDP-43 inclusions (left, 21%; right, 22%, t = 6.42, p < 0.0001) compared with Layer 5 non-VEN, non-fork neurons (i.e. neighboring Layer 5 neurons), yet showed a slightly lower inclusion fraction than VENs and fork cells in absolute terms (Fig. 2a). Nuclear TDP-43depleted neuron fractions, however, were significantly higher in VENs (t = 2.64, p = 0.0083) and fork cells (t = 2.98, p = 0.0029) than in Layer 2-3 neurons (Fig. 2b). Nuclear TDP-43-

depleted neuron fractions did not differ significantly between Layer 2-3 and neighboring Layer 5 neurons (t = -1.45, p = 0.15).

Despite the absence of gross atrophy, cortical thinning, and significant overall neuronal loss at Broe stage 0, 18% of FI VENs and 20% of fork cells, but only 5% of neighboring Layer 5 neurons, showed TDP-43 aggregation (Fig. 2c). Layer 2-3 neurons (13% inclusion fraction) fell midway between VENs/fork cells and neighboring Layer 5 neurons. Patients with more advanced disease (stages 1-2) showed a trend toward a higher proportion of neurons with TDP-43 inclusions across cell types, with the gap between VENs/fork cells and Layer 2-3 neurons narrowing at stage 1 and closing by stage 2 (Fig. 2c). In contrast, the rate of nuclear TDP-43 depletion was similar in VENs, fork cells, and Layer 2-3 neurons at Broe stage 0, although nuclear TDP-43 depletion was more prominent in VENs and fork cells than neighboring Layer 5 neurons and Layer 2-3 neurons at Broe stages 1 and 2 (Fig. 2d). Due to substantial intersubject variability, however, the interaction between Broe stage and neuron type was not significant for TDP-43 inclusions (F = 1.49, p > 0.1) or nuclear depletion (F = 1.54, p > 0.1).

VEN and fork cell TDP-43 inclusion fraction relates to clinical diagnosis, overall symptom severity, and loss of emotional empathy in the sporadic bvFTD/MND spectrum

Patients with bvFTD develop progressive social-emotional deficits that undermine day-today work and family life. Here, we found a significant interaction (F = 8.49, p < 0.01) between neuron type and clinical diagnosis, such that VEN, fork cell and Layer 2-3 neuron TDP-43 inclusions in right FI were more frequent among patients with behavioral symptoms sufficient to warrant a bvFTD diagnosis (ALS vs bvFTD-MND or bvFTD, p < 0.005) (Fig. 2e). Although patients with clinical ALS showed only rare TDP-43 inclusions in VENs (inclusion fraction 2.6%) and fork cells (inclusion fraction 2.4%), these VEN/fork cell inclusions were 3-4 times more frequent than TDP-43 inclusions in the other right FI cell types we studied (neighboring Layer 5 neurons: 0.7%, Layer 2-3 neurons: 0.6%). Nuclear TDP-43-depleted VENs and fork cells were more frequent in ALS than in bvFTD (Fig. 2f and Suppl. Figure 4a), suggesting that loss of nuclear TDP-43, through whatever mechanism, may precede TDP-43 aggregation. To assess whether selective VEN/fork pathobiology can be detected even in patients with pure ALS, we combined TDP-43 depletion and aggregation measures in this group and compared this composite measure across cells types (Suppl. Figure 4a. S1-S4). The results show that 11.4% (left) and 11.5% (right) of VENs and 11.0% (left) and 8.9% (right) of fork cells show TDP-43 pathobiology in pure ALS, representing a four- to five-fold increase over neighboring Layer 5 neurons (VENs: t = 9.75, p < 0.0001; fork cells: t = 10.67, p < 0.0001) and a two- to three-fold increase over Layer 2-3 neurons (VENs: t = 2.59, p = 0.30; fork cells: t = 3.49, p = 0.034).

In contrast to a previous study of later stage bvFTD, in which greater functional impairment (CDR sum of boxes) correlated with right hemisphere VEN and fork cell dropout [34], in the present study it was the proportion of right hemisphere TDP-43 inclusion-bearing neurons that predicted greater impairment (VENs + fork cells: r = 0.83, p = 0.00048, Fig. 2g; neighboring Layer 5 neurons: r = 0.82, p = 0.00062; Layer 2-3 neurons: r = 0.63, p = 0.025). Using a more specific measure of social-emotional dysfunction, the Interpersonal

Reactivity Index (IRI), we found that the fraction of right hemisphere Layer 5 neurons bearing TDP-43 inclusions predicted deficits in emotional empathy (IRI, empathic concern subscale; VENs + fork cells: r = -0.69, Fig. 2h), p = 0.038; neighboring Layer 5 neurons: r = -0.69, p = 0.039). Layer 2-3 neuron inclusion fractions did not correlate with measures of empathic concern (r = -0.30, p = 0.47). Right FI inclusion fractions in VENs, fork cells, and neighboring Layer 5 neurons showed high multicollinearity (spearman correlation coefficients 0.75 - 0.96; variance inflation factor = 3.56 - 5.51), precluding mediation analysis to determine which cell type was most likely driving the behavioral change. CDR (VENs: r = 0.073, p = 0.81; fork cells: r = -0.18, p = 0.55; neighboring Layer 5 neurons: r = 0.34, p = 0.25) and empathic concern (VENs: r = -0.010, p = 0.98; fork cells: r = -0.12, p = 0.75; neighboring Layer 5 neurons: r = 0.023, p = 0.95) showed no correlation with neuronal densities. Collectively, these findings suggest that, in early stages of bvFTD, TDP-43 inclusion formation may lead to targeted neuronal dysfunction that contributes to clinical deficits despite limited neuronal loss.

VENs and fork cells show increased TDP-43 aggregation and nuclear depletion in the *C9orf72*-bvFTD/MND spectrum

The C9orf72 hexanucleotide repeat expansion is the major genetic cause of inherited FTD/ALS [18, 61] and is almost always accompanied by TDP-43 proteinopathy [43, 49]. In addition, C9orf72 expansions are associated with several additional pathological features, including repeat RNA foci [18] and dipeptide repeat protein inclusions [4, 48], Therefore, for this study we elected to separate the *C9orf72* and sporadic patients in our cohort. We retained and randomized 10 patients with C9orf72-bvFTD/MND to neuron quantification runs so that raters would remain blind to genetic status, enabling a future blinded study comparing sporadic to C9orf72-bvFTD/MND. An interim analysis of TDP-43 inclusions in VENs, fork cells, and neighboring Layer 5 neurons in the C9orf72 group, however, provided data critical to the understanding of our sporadic bvFTD/MND findings, so we elected to present this analysis here. We observed no significant neuronal loss in patients with C9orf72 FTD/MND, although there was a trend toward VEN and fork cell loss (Suppl. Figure 3f) that mirrored the degree of neuronal loss observed in the sporadic FTD/MND group (Suppl. Figure 3a). Although the overall proportion of neurons forming TDP-43 inclusions was much lower in *C9orf72*-byFTD than in sporadic byFTD, VENs (t = 3.82, p = 0.00013) and forks cells (t = 4.36, p < 0.0001) were again four times more vulnerable to TDP-43 inclusions than were neighboring Layer 5 neurons (Fig. 3b). Interestingly, in C9orf72bvFTD/MND, neurons lacking nuclear TDP-43 without a TDP-43 inclusion (nuclear TDP-43 depleted) were as common as neurons with TDP-43 inclusions overall (Figs. 3a, 3b), although these rates varied between subjects (Fig. 3c). In one C9orf72-ALS patient, over 20% of VENs showed nuclear TDP-43 depletion while fewer than 5% showed TDP-43 aggregation (Fig. 3c).

VENs and fork cells with nuclear TDP-43 depletion, with or without aggregation, show nuclear and somatodendritic atrophy

Having shown that VENs and fork cells are more prone to TDP-43 aggregation and nuclear depletion, we next asked how these specific phenomena relate to the morphological integrity of individual neurons. To our knowledge, no human study has quantified the relationship

between TDP-43 pathobiology and neuronal morphology, perhaps because the enormous variation in cortical neuron size and shape would undermine implementation of such efforts. VENs and fork cells, being relatively homogeneous, provided an opportunity to overcome these issues. Indeed, upon visual inspection, VENs and fork cells with TDP-43 inclusions or depletion exhibited a shrunken, degenerate morphology (Fig. 1 and Suppl. Figure 5a). We further noted that, in patient tissues, inclusion-lacking VENs and fork cells with normal nuclear TDP-43 were often unusually large. Consistent with these observations, morphometric analysis confirmed that TDP-43 aggregation is associated with atrophy across multiple measures (Figs. 4a, 4b). VEN somatic (t = -13.37, p < 0.00001) and nuclear area (t = -10.90, p < 0.00001) and apical (t = -18.91, p < 0.00001) and basal (t = -17.48, p = 0.00001) dendritic diameter were smaller in cells with cytoplasmic TDP-43 aggregation than unaffected neurons (Fig. 4a). The pattern in fork cells (Fig. 4b) mirrored that seen in VENs: somatic (t = -6.33, p < 0.00001) and nuclear area (t = -7.16, p < 0.00001) and apical (t = -15.90, p < 0.00001) and basal (t = -11.92, p < 0.00001) dendritic diameters were smaller in fork cells with cytoplasmic TDP-43 inclusions than fork cells with normal nuclear TDP-43. Inclusion-bearing pyramidal neurons showed reduced apical (t = -5.86, p = 4.64×10^{-9}) and basal dendritic diameter (t = -2.10, p = 0.036), but no change in nuclear (t = 0.63, p = 0.52) or somatic (t = 1.15, p = 0.24) area (Suppl. Figure 5b). The findings were not influenced by C9orf72 status overall (genotype as a fixed effect was not significant).

TDP-43 inclusion formation is universally accompanied by a loss of nuclear TDP-43. In the course of the present work, our recognition of the nuclear TDP-43-depleted neurons indicated that the reverse is not necessarily true (Fig. 1c and Suppl. Figure 4a). This dissociation suggested an avenue for informing the ongoing controversy about whether neuronal degeneration in TDP-43 proteinopathy results from inclusion toxicity or a loss of TDP-43 function. In our data, nuclear TDP-43-depleted VENs and fork cells showed somatodendritic and nuclear atrophy (Figs. 4a, 4b; VENs: somatic area, t = -11.80, p < 0.00001; nuclear area, t = -10.22, p < 0.00001; apical dendritic diameter, t = -16.57, p < 0.00001; basal dendritic diameter, t = -16.01, p < 0.00001; fork cells: somatic area, t =-6.43, p < 0.00001; nuclear area, t = -6.64, p < 0.00001; apical dendrite diameter, t = -12.01, p < 0.00001; basal dendritic diameter, t = -9.86, p < 0.00001) just as severe as that seen in inclusion-bearing neurons, suggesting that loss of nuclear TDP-43 function alone proves deleterious. We observed no significant differences between inclusion-bearing and nuclear TDP-43-depleted VENs (t = -1.38 - -0.61, p = 0.17 - 0.54) or fork cells (t = -1.76 - 0.54) -0.33, p = 0.079 - 0.74) for any neuronal compartment. In layer 5 pyramidal neurons, somata (t = -2.17, p = 0.03) but not nuclei or dendrites (t = -1.37 - -0.52, p = 0.17 - 0.60), were significantly smaller in nuclear TDP-43-depleted cells than those with TDP-43 inclusions. Remarkably, for reasons that remain uncertain, VENs with normal TDP-43 localization showed larger nuclear (VENs: t = 2.00, p = 0.045) and somatic (VENs: t = 2.28, p = 0.022) areas in patients than in controls.

Our morphometric data suggested that, in patients, VENs and fork cells might each be divided into 2-3 neuronal classes or "states". To pursue this idea, we used K-means clustering, an iterative data-partitioning algorithm that assigns a set of observations to a predefined number of clusters, to classify VENs into three clusters (Fig. 4c) based on their somatic and nuclear area but without using information about the source (patient vs. control)

or TDP-43 status (inclusion-bearing, depleted, or unaffected). Although assigned to clusters without regard to TDP-43 status, VENs were unevenly distributed across clusters according to TDP-43 status (Suppl. Table 4, X^2 = 135.48, p < 0.0001). TDP-43 inclusion-bearing and depleted neurons clustered together (Cluster 3). Neurons with normal TDP-43 were spread over two clusters. Cluster 1 was composed of predominantly normal cells from patients, whereas Cluster 2 had normal cells from healthy controls along with some TDP-43 inclusion-bearing or nuclear depleted neurons from patients. Overall, these findings further support the view that neurons with TDP-43 inclusions or nuclear depletion are characterized by similar atrophy profiles, while normal neurons from patients are larger than normal neurons in controls.

To further characterize how TDP-43 pathobiological states influenced the pattern of atrophy across the four neuronal compartments (i.e., soma, nucleus, apical dendrite, and basal dendrite; Fig. 4d and Suppl. Tables 5 and 6), we ranked each TDP-43 inclusion-bearing or TDP-43 nuclear-depleted neuron in terms of which compartment was most to least affected compared to control neurons of the same type. Across the three neuron types and two TDP-43 pathobiological states, we found a relationship between atrophy severity rankings and neuronal compartment (Suppl. Table 5). In TDP-43 inclusion-bearing and nuclear TDP-43 depleted VENs and fork cells, the apical dendrite was most affected, followed by the basal dendrite, nucleus, then soma (Fig. 4d and Suppl. Table 6). The apical dendrite was most affected compartments in pyramidal neurons with nuclear TDP-43 depletion (Fig. 4d and Suppl. Table 6).

Discussion

In many neurodegenerative disorders, early neuronal loss targets a specialized pool of regionally concentrated and morphologically unique neurons [28, 31]. In bvFTD, this pool is composed of the VENs and fork cells of the anterior cingulate and frontoinsular cortices. Here, we leveraged this anatomical framework to provide new insights into selective vulnerability and TDP-43 pathobiology. Our study identifies VENs and fork cells as an early target of TDP-43 pathobiology in bvFTD and shows that VEN and fork cell TDP-43 aggregation (1) correlates with regional atrophy and behavioral symptoms, (2) occurs in both sporadic and C9orf72 bvFTD/MND, and (3) is linked to nuclear and somatodendritic atrophy at the level of individual neurons. In addition, we show that loss of nuclear TDP-43 can occur in the absence of, or perhaps prior to, overt TDP-43 aggregation and may be sufficient to induce neurodegeneration. These novel human findings have major implications for the pathogenesis and future treatment of FTD, ALS, and other disorders associated with TDP-43 inclusion pathology, suggesting that loss of nuclear TDP-43 function may be the earliest and most consequential aspect of TDP-43 pathobiology. At the very least, the findings show that mature TDP-43 inclusions, though they may be deleterious, are not required for neuronal degeneration in FTD/MND.

VENs and fork cells are selectively prone to TDP-43 aggregation at the earliest stages of sporadic bvFTD

Several studies have demonstrated ACC VEN loss as a general feature of the bvFTD syndrome, including patients with underlying FTLD-TDP, -tau or -FUS [63-65, 86]. Only one previous study examined FI [34], also across the FTLD spectrum, and found 53% VEN and 68% fork cell loss in bvFTD and bvFTD-MND. No previous study has assessed VEN or fork cell disease protein aggregation in bvFTD. In the previous study of FI, VEN and fork cell dropout was greatest in patients with FTLD-tau, FTLD-FUS, and FTLD-TDP without MND [24], likely because MND often truncates the disease course when it accompanies bvFTD. We therefore focused the present study on TDP-43, the most common bvFTDrelated disease protein, and on patients representing points along the bvFTD-MND spectrum, hoping to capture stages prior to severe VEN dropout during which the anatomical correlates of bvFTD symptoms are just beginning to emerge. Due to these efforts, the present sample showed only a 32% reduction in FI VEN density and a 24% reduction in fork cell density, making this the mildest byFTD sample studied to date and allowing us to probe TDP-43 pathobiology in surviving neurons. We included patients with ALS because we hypothesized that, just as some patients with pure bvFTD show TDP-43 aggregation in scattered motor neurons, some patients with pure ALS might show TDP-43 inclusions in VENs and fork cells. Confirming our hypothesis, and in line with a qualitative study identifying VEN TDP-43 depletion in pure ALS [6], we found that the proportion of VENs and fork cells with TDP-43 inclusions in pure ALS was low but higher than seen in the other cell types we studied. Separating patients by clinical diagnosis and atrophy severity suggested that VENs and fork cells are affected during the earliest identifiable stages, at a time when Layer 2-3 neurons are just beginning to form TDP-43 inclusions and when neighboring Layer 5 neurons are still largely spared. In patients with more advanced clinical-anatomical deficits, Layer 2-3 neurons become nearly as affected as VENs and fork cells, whereas inclusions in Layer 5 non-VEN/fork neurons rise only slightly. Nonetheless, the strong link between TDP-43 aggregation and clinical measures suggests that TDP-43 pathobiology is sufficient to induce neuronal dysfunction, leading to behavioral deficits, even while neuronal loss remains mild or absent. The characteristic "speckled" dot- and dash-like TDP-43 inclusions in VENs and fork cells have been described by some as earlystage "pre-inclusions", which often show only partial ubiquitination, as observed here (Fig. 1d) [25, 29, 71].

VEN and fork cell involvement in C9orf72-bvFTD/MND

As in sporadic bvFTD/MND, [6]in *C9orf72*-bvFTD/MND VENs and fork cells were more prone to TDP-43 inclusion formation compared to neighboring Layer 5 neurons. Interestingly, although overall rates of TDP-43 inclusion formation were lower in *C9orf72*bvFTD/MND, VEN and fork cell density reductions were nearly identical to those seen here in sporadic bvFTD/MND and consistent with a previous *C9orf72*-bvFTD study of VEN dropout in the anterior cingulate cortex [86]. The low TDP-43 inclusion fractions might suggest that *C9orf72*-specific disease mechanisms, such as repeat RNA foci and dipeptide repeat protein inclusions [4, 61], contribute to neuronal dysfunction and death in *C9orf72*bvFTD and ALS or that other brain regions contribute to bvFTD-related network dysfunction [37, 86]. Alternatively, *C9orf72*-specific pathological features may sensitize

vulnerable neurons to TDP-43 pathobiology, such that early stage TDP-43 mislocalization and aggregation lead to accelerated neuronal dropout. Although some human *C9orf72*-FTD/MND studies have suggested a strong correlation between regional TDP-43 aggregation and neurodegeneration [43], our previous work has shown that TDP-43 aggregation can be scarce or even absent in patients with clinical *C9orf72*-bvFTD [78]. Future studies should examine the frequency of other *C9orf72*-specific pathological phenomena in VENs and fork cells and the relationship of these features to TDP-43 nuclear depletion and cytoplasmic aggregation.

Our data, however, suggest a third possibility: that VEN and fork cell dysfunction in *C9orf72*-bvFTD/MND may relate to depletion of nuclear TDP-43 without, or prior to, inclusion formation. This "nuclear TDP-43-depleted" state was also seen, albeit less often, in the sporadic patients, where nuclear TDP-43 depletion without NCI formation was particularly prominent in pure ALS. Only three previous studies have reported nuclear TDP-43-depleted neurons in ALS or FTD [6, 7, 78]. Depletion of nuclear TDP-43 may represent a separate pathway toward neurodegeneration, perhaps related to nuclear import/ export dysfunction [58], or a transitional state en route to TDP-43 aggregation. A previous report from our group characterized temporal lobe resection (for epilepsy) and autopsy tissue from the same *C9orf72*-FTD patient and found nuclear TDP-43 depletion without aggregation in the resection specimen taken five years prior to symptom onset but prominent TDP-43 inclusion pathology at autopsy 13 years later [78]. These finding, together with the present study, suggest the possibility that nuclear TDP-43 depletion may ultimately transition toward aggregate formation within the same neurons.

Association of TDP-43 aggregation and nuclear depletion with neuronal atrophy

Microscopic examination of patient tissue suggested striking size differences between inclusion-bearing neurons and inclusion-lacking neurons with normal nuclear TDP-43, leading us to undertake a detailed morphometric study. In sporadic and C9orf72-bvFTD/ MND, VEN and fork cell apical and basal dendritic diameter and nuclear and somatic area were significantly shrunken in neurons with TDP-43 inclusions compared to those without. Many studies in model systems suggest that TDP-43 overexpression and aggregation lead to cell dysfunction and death [5, 77, 84, 85], but no previous human neuropathologic study has linked TDP-43 pathobiology to degeneration at the level of individual neurons. Moreover, it has remained unclear whether TDP-43 aggregate toxicity or loss of normal nuclear TDP-43 function is the key driver of pathogenesis in patients. Our data show that nuclear TDP-43depleted neurons develop somatodendritic and nuclear atrophy similar to that seen in inclusion-bearing neurons. This crucial observation suggests that loss of nuclear TDP-43 function may be sufficient to undermine neuronal integrity in vivo. Our data also suggest that TDP-43 aggregation-at least the type visible using conventional methods-may not be necessary for neurodegeneration. Despite the lack of a visible TDP-43 NCI in these nuclear TDP-43-depleted neurons, it is possible that low levels of a toxic non-aggregated form of cytoplasmic TDP-43, below the detection threshold of our staining protocol, are contributing to the morphometric deficits seen in these neurons, in keeping with model system findings in which elevated but non-aggregated cytoplasmic TDP-43 precipitates cell death [5]. In addition, it is possible that undetected low levels of nuclear TDP-43 remain in these neurons

and are sufficient to support normal TDP-43 function. Despite these caveats, our data suggest that loss of nuclear TDP-43 function deserves major consideration as a contributor to pathogenesis in FTLD-TDP and ALS-TDP.

In sporadic ALS, upper motor neurons and VENs have also been reported to show clearing of nuclear TDP-43 without cytoplasmic TDP-43 aggregation [6, 7]. Identification of the nuclear TDP-43-depleted VENs and fork cells provides an opportunity for future studies to explore neurodegeneration mechanisms associated with TDP-43 loss-of-function. Interestingly, conditional mouse models overexpressing a human form of TDP-43 with a mutated nuclear localization signal develop behavioral and motor changes with little TDP-43 aggregation [1, 32], relating to suppression of mouse TDP-43 expression. Likewise, mice expressing a common human mutation in TDP-43 showed no TDP-43 aggregation but frequent nuclear TDP-43 depletion, accompanied by progressive neuromuscular weakness and death [83]. Recently, novel hypotheses have emerged about how loss of nuclear TDP-43 function may lead to neuronal dysfunction and degeneration. In mouse embryonic stem cell, HeLa cell, and conditional TDP-43 knockout mouse models, TDP-43 mutations, depletion, or sequestration into inclusions lead to aberrant splicing [3, 38, 56, 75] and altered expression of numerous proteins [57, 70], including those involved in nucleocytoplasmic transport, RNA processing, and DNA repair. Loss of nuclear TDP-43 results in, for example, incorporation of cryptic exons into mRNAs [33, 38, 74, 75]; often such exons introduce frameshifts or premature stop codons that lead the transcript to undergo nonsense mediated decay. The selective vulnerability framework and experimental platform developed through this study can now be used to explore these potential mechanisms directly in human tissues.

VENs and fork cells with normal nuclear TDP-43 were larger in patients with bvFTD than in control subjects and much larger than the atrophied inclusion-bearing and nuclear TDP-43-depleted neurons seen within the same patients. This unexpected observation could reflect compensation for other degenerating neurons in the circuit or a stage of disease just prior to nuclear TDP-43 depletion and aggregation. Strikingly, TDP-43 targets another population of large, long-range projection neurons—the upper motor neurons lost in ALS [2, 52]. We hypothesize that there are common features among VENs, fork cells, and upper motor neurons that render them particularly vulnerable to TDP-43-related degeneration. Like corticospinal motor neurons, VENs and fork cells are large Layer 5 projection neurons with likely subcerebral targets [15]. In a mutant SOD1 mouse model of ALS, vulnerable populations of spinal motor neurons showed an initial increase in soma cross-sectional area, followed by atrophy later in disease [21]. This alteration in cell size was predicted to result in altered neuronal excitability [21]. Hyperexcitability is an established early feature of ALS, seen across humans [22, 80], mouse models [55] and induced pluripotent stem cell-derived neurons studies [19, 81], including motor neurons from patients with C9orf72-ALS [19]. Cortical hyperexcitability is described in early stage sporadic [80] and C9orf72-associated ALS [26]. In VENs and fork cells, similar hyperexcitability mechanisms could increase neuronal size before giving way to TDP-43 pathobiology and neuronal atrophy as neurons progress toward death.

VENs and fork cells appear to be more prone to nuclear TDP-43 depletion and aggregation (present data) but also more vulnerable to death [34, 65] in the face of TDP-43 pathobiology.

It will be important to determine whether the increased rates of VEN and fork cell death result only from their increased proneness to TDP-43 pathobiology or whether they are also more likely than other cells to die once TDP-43 pathobiology emerges. Post-mortem studies cannot trace a neuron's course from nuclear TDP-43 depletion to aggregation to degeneration to ultimate cell dropout, and commonly used laboratory mammals lack VENs and fork cells. For these reasons, a natural next step in pursuing this work would be to develop human cell-based *in vitro* assays, akin to human iPSC-derived motor neuron assays used in ALS research. Such an approach would enable observation of longitudinal VEN/fork cell trajectories and experimental manipulations that can determine causal influence. On the other hand, still much more can be done with human tissue, following the model provided here, to determine which cellular events associated with TDP-43 pathobiology are most strongly linked to neuronal degeneration.

The impact of VEN and fork cell degeneration on large-scale network function has yet to be elucidated. The ACC and FI, key regions affected early in bvFTD [66, 87, 88] are involved in social and emotional function [16, 73]. The FI plays an important role in autonomic function, representing the internal milieu [16, 50], and connects with limbic and autonomic brain regions, including ACC, amygdala, ventral striatum, and brainstem. Intriguingly, the present study links VEN and fork cell inclusion formation with deficits in empathic concern for others. Although we have previously shown that VEN and fork cell loss correlates with behavioral symptom severity in bvFTD [34], the present study is the first to correlate inclusion formation in VENs and fork cells, or any other neuron population, with a specific social-emotional deficit. Most likely, the relationship between these cell types and emotional empathy is not one-to-one but mediated by the large-scale networks anchored by these and other FI neurons. Future studies could investigate links between the cellular, network, and behavioral levels. VENs and fork cells are thought to have subcerebral projections [15], although their precise connections are still under study in the macaque [23]. VENs and fork cells express the alpha 1a adrenergic receptor (ADRA1A), the GABA-A receptor theta subunit (GABRQ), and the vesicular monoamine transporter (VMAT2), suggesting a role for VENs and fork cells in autonomic control circuits [20]. Thus, autonomic dysfunction in bvFTD, which we have strongly linked to ACC and FI degeneration and dysfunction [30, 72], may be initiated by FI VEN and fork cell dysfunction and degeneration before spreading throughout the system to undermine some of our most specialized socialemotional capacities.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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Fig. 1. Distribution of normal and inclusion bearing neurons and patterns of TDP-43 aggregation in VENs, fork cells, and neighboring Layer 5 neurons.

(a, b) Mapping of all VENs and fork cells located in a section of the FI (outlined in a) in subject S5 shows similar distribution of normal and inclusion-bearing VENs and fork cells across Layer 5 (b). B shows an expanded region of the FI indicated by the dashed rectangle in a. (c) Punctate TDP-43 neuronal cytoplasmic inclusions (NCIs) were the most common inclusion type in VENs, fork cells, and neighboring Layer 5 neurons (L5 NNs). Neurons without TDP-43 NCIs but lacking normal nuclear TDP-43 ("nuclear TDP-43-depleted") were observed for all cell types in the majority of patients. All images in (c) are from subject S5. (d) Images of VENs from a single confocal z slice with TDP-43 (red), phospho-TDP-43 (pTDP-43, green) and ubiquitin (cyan) immunostaining, and Nissl-stained nuclei (blue). MAP2 immunostaining (not shown) was used to trace VEN outlines. Normal VENs in

control subjects (a VEN from subject N4 is shown) and patients with bvFTD/MND showed robust nuclear TDP-43 staining and no TDP-43 aggregation, phosphorylated TDP-43, or ubiquitin staining. In subjects with ALS (subject S4 is shown), TDP-43 inclusions were phosphorylated, with punctate, scattered ubiquitin staining dispersed around the TDP-43 inclusions. A few characteristic ubiquitin puncta are indicated by arrows on the single z slice from the patient with ALS, and are more clearly seen on a maximum intensity projection of 15 z slices through the VEN soma. In severely affected subjects (subject S15 is shown), TDP-43 inclusions were phosphorylated and foci of ubiquitination were larger and more numerous. The merge of red (TDP-43) and green (phospho-TDP-43) is shown as yellow. Brightness and contrast for each color channel in D were adjusted separately. Scale bars in (c) and (d) represent 10 μ m. dAI: dorsal anterior insula; FI: frontoinsular cortex; MIP: maximum intensity projection; Pu: putamen; TP: temporal pole.





(a) Layer 5 (L5) VENs and fork cells, and Layer 2-3 (L2-3) neurons were significantly more prone to TDP-43 neuronal cytoplasmic inclusions (NCIs) than were neighboring Layer 5 non-VEN/fork neurons (L5 NNs). (b) L5 VENs and fork cells showed significantly more nuclear TDP-43-depletion than L5 NNs or L2-3 neurons. * p < 0.01. The patterns of TDP-43 inclusion formation and nuclear TDP-43 depletion were similar in left and right FI (a,b). (c) VEN and fork cell TDP-43 aggregation emerged in the absence of gross atrophy (Broe stage 0), while neighboring Layer 5 neurons showed low inclusion fractions. TDP-43 inclusion formation rates in Layer 2-3 neurons appeared to more closely follow VENs and fork cells.

(d) At Broe stages 0-2, depletion of nuclear TDP-43, without formation of overt cytoplasmic inclusions, (nTDP-43 depletion) was observed in a small percentage of VENs, fork cells, and Layer 2-3 neurons. (e) VEN, fork cell, and L2-3 neuron inclusions were infrequent in ALS but were present by behavioral symptom onset (p < 0.005). (f) In contrast, nuclear TDP-43 depletion without NCI formation was more prominent in VENs and fork cells than Layer 2-3 neurons and Layer 5 neighboring neurons in ALS cases. Data in c-f are shown as group mean \pm SEM from right FI. Data for Stage 2 in c-d are shown as mean and range (n = 2 Broe stage 2 subjects with right FI). (g, h) The proportion of VENs and fork cells with TDP-43 inclusions correlated with Clinical Dementia Rating (CDR) (g) and inversely correlated with empathic concern (h) in the right hemisphere. For clarity, only VENs + fork cells (combined) are shown in g and h.





(a) Nuclear TDP-43-depleted neurons were more frequent in *C9orf72*-bvFTD/MND (arrow: VEN, arrowhead: pyramidal neuron). Scale bar in a represents 10 μ m. (b) In *C9orf72*-bvFTD/MND, VENs and fork cells were also more prone to TDP-43 neuronal cytoplasmic inclusions (NCIs) and nuclear TDP-43 depletion without NCIs (nTDP-43 depleted) than neighboring Layer 5 neurons (L5 NNs) although TDP-43 inclusion formation in VENs and fork cells occurred at a lower rate in *C9orf72*-bvFTD/MND than sporadic bvFTD/MND. Data in (b) is from the right hemisphere and is shown as mean ± SEM. * p < 0.02 compared with L5 NNs. (c) Percentage of affected (TDP-43 NCI and nuclear TDP-43-depleted) neurons in Layer 5 of the FI in each patient with *C9orf72*-bvFTD/MND. The fraction of VENs, fork cells, and neighboring Layer 5 neurons with TDP-43 depletion was as high as the fraction of neurons with TDP-43 inclusions in six of the *C9orf72*-bvFTD/MND cases.

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Fig. 4. TDP-43 aggregation and TDP-43 depletion are linked to VEN and fork cell atrophy in both sporadic and *C9orf72*-bvFTD/MND.

(a, b) Morphometric analyses revealed that VEN (a) and fork cell (b) somatic and nuclear area, and apical and basal dendritic diameter were significantly smaller (20-39%) in cells with TDP-43 neuronal cytoplasmic inclusions (NCIs) than those without in bvFTD/MND patients. Somatodendritic atrophy was also evident in nuclear TDP-43-depleted neurons without NCIs (nTDP-43-depleted). Intriguingly, VENs (a) with normal nuclear TDP-43 in bvFTD patients showed higher nuclear (16%) and somatic area (18%) compared to VENs in controls. Normal VENs and fork cells neurons in bvFTD had a significantly larger nucleus and soma than TDP-43 inclusion-bearing and nuclear TDP-43-depleted neurons. Data in a and b are shown as mean \pm SEM, with the mean and range shown for nuclear TDP-43-depleted neurons in sporadic cases (nuclear TDP-43-depleted VENs and fork cells were identified in two of the sporadic cases that underwent morphometric analysis). * p < 0.05 compared with normal controls. (c) Clustering analysis of VEN nucleus and soma area, and

apical and basal dendritic diameter identified three clusters, primarily corresponding to normal cells from patients (cluster 1) and inclusion bearing/nuclear TDP-43-depleted VENs (cluster 3), with normal cells from control subjects and some TDP-43 inclusion/depleted neurons combined in cluster 2. Only two (nuclear and soma area) of the four parameters used to define clusters are shown for clarity. (d) Apical dendrites, followed by basal dendrites, were the most affected compartments in TDP-43 inclusion-bearing and nuclear TDP-43 depleted VENs and fork cells, with the highest mean rank, corresponding to highest z score.

Table 1.

Subject demographics

	Controls	Sporadic bvFTD/MND	C9orf72-bvFTD/MND				
Ν	10	17	10				
Clinical diagnoses	N/A	3 ALS, 1 MND, 2 bvFTD-ALS, 5 bvFTD-MND, 6 bvFTD	2 ALS, 5 bvFTD-MND, 3 bvFTD				
Neuropathological diagnoses	N/A	4 ALS-TDP; 2 FTLD-TDP-B, ALS; 8 FTLD-TDP-B, MND; 3 FTLD-TDP-B	1 ALS-TDP; 5 FTLD-TDP-B, MND; 3 FTLD-TDP-U, MND; 1 FTLD- TDP-U				
Gender (M:F)	6:4	10:7	8:2				
Age at death (years)	63.6 (9.8)	61.8 (7.4)	59.8 (3.4)				
Symptom duration (years)	N/A	5.2 (2.8)	7.1 (6.9)				
PMI (hours)	43.1 (38.2)	13.0 (10.2)	13.3 (6.3)				
Broe stage; median (range)	N/A	1 (0-2)	1 (0-2)				
Brain weight (g)	1244 (151)	1284 (138)	1257 (172)				
CDR total	N/A	1.8 (1.1)	1.3 (0.9)				
CDR Sum of boxes	N/A	8.9 (5.2)	7.5 (5.0)				
Worst NPI, sum of apathy, disinhibition, motor, eating	N/A	41.9 (25.1)	44.3 (27.5)				
IRI, empathic concern subscale	N/A	20.0 (8.0)	17.6 (7.3)				

Unless otherwise noted, numbers in parentheses are SD of preceding mean.

Table 2.

Detailed subject characteristics.

Case no.	Age	Sex	Clinical diagnosis	Primary neuropathological diagnosis ^a	Side studied	Sx dur (years)	Broe stage	Thal phase	Braak stage	CERAD score	ADNC	CDR, sum of boxes	IRI – Empathic concern
N1	46	F	NC	N/A	R & L	N/A	ND	ND	ND	ND	ND	ND	ND
$N2^{b}$	57	М	NC	N/A	R & L	N/A	ND	ND	ND	ND	ND	ND	ND
N3	59	F	NC	N/A	R & L	N/A	ND	ND	ND	ND	ND	ND	ND
N4	76	М	NC	N/A	R & L	N/A	ND	1	2	Sparse	Low	ND	ND
N5	73	М	NC	N/A	R & L	N/A	ND	0	1	Absent	Not	ND	ND
N6 ^b	55	F	NC	N/A	R & L	N/A	ND	0	0	Absent	Not	ND	ND
N7	72	F	NC	N/A	R & L	N/A	ND	4	4	Frequent	Int.	ND	ND
N8	73	М	NC	N/A	R & L	N/A	ND	2	2	Absent	Low	ND	ND
$N9^{b}$	64	М	NC	N/A	R & L	N/A	ND	0	1	Absent	Not	ND	ND
N10 ^b	61	М	NC	N/A	R & L	N/A	ND	1	1	Absent	Low	ND	ND
S 1	49	М	ALS	ALS-TDP	R & L	3	0	1	0	Absent	Low	3	ND
S2	64	М	ALS	ALS-TDP	R & L	9	1	0	0	Absent	Not	ND	ND
S 3	52	М	MND	ALS-TDP	R & L	7	0	0	1	Absent	Not	0	31
S4	67	F	ALS	ALS-TDP	R & L	2	1	4	2	Frequent	Low	0	ND
s5 ^b	51	F	bvFTD-MND	FTLD-TDP-B; MND	R & L	3	1	2	0	Absent	Low	11	31
S6	58	F	bvFTD-MND	FTLD-TDP-B; MND	R & L	3	1	1	0	Absent	Low	11	23
S 7	66	М	bvFTD-MND	FTLD-TDP-B, MND	L	3	2	2	2	Absent	Low	9	20
S8	72	М	bvFTD-MND	FTLD-TDP-B	R & L	1	0	0	4	Absent	Not	18	ND
89 ^b	60	М	bvFTD-ALS	FTLD-TDP-B, ALS	R	7	1	3	1	Frequent	Low	10	23
S10 ^b	65	М	bvFTD-MND	FTLD-TDP-B, MND	R & L	5	0	1	1	Sparse	Low	10	9
S11 ^b	64	М	bvFTD-ALS	FTLD-TDP-B, MND	R	9	0	1	1	Absent	Low	5.5	20
S12	53	F	bvFTD	FTLD-TDP-B	R & L	6	1	0	0	Absent	Not	13	ND
S13	75	F	bvFTD	FTLD-TDP-B; MND	R & L	3	0	3	4	Absent	Int.	9	19
S14 ^b	66	М	bvFTD	FTLD-TDP-B, MND	R	8	2	2	2	Sparse	Low	5.5	ND
S15	65	F	bvFTD	FTLD-TDP-B, MND	R	9	1	1	2	Absent	Low	15	12
S16 ^b	57	F	bvFTD	FTLD-TDP-B, ALS	R	3	2	0	0	Absent	Not	7	25
S17 ^b	66	М	bvFTD	FTLD-TDP-B	R	8	1	2	2	Moderate	Low	15	7
C1 ^{<i>b</i>}	62	F	ALS	FTLD-TDP-B, MND	R & L	2	0	0	1	Absent	Not	4	30
$C2^{b}$	58	М	ALS	ALS-TDP	R	2	1	1	2	Absent	Low	0	28
C3	59	М	bvFTD-MND	FTLD-TDP-U, MND	L	7	0	2	2	Absent	Low	5	18
C4	57	М	bvFTD-MND	FTLD-TDP-B; MND	R & L	1	0	1-2	0	Absent	Low	7	19
C5 ^{<i>b</i>}	59	М	bvFTD-MND	FTLD-TDP-B; MND	L	9	2	0	0	Absent	Not	6	14
C6	58	М	bvFTD-MND	FTLD-TDP-U, MND	R	3	1	2	1	Sparse	Low	3.5	ND

Case no.	Age	Sex	Clinical diagnosis	Primary neuropathological diagnosis ^a	Side studied	Sx dur (years)	Broe stage	Thal phase	Braak stage	CERAD score	ADNC	CDR, sum of boxes	IRI – Empathic concern
C7 ^b	56	М	bvFTD-MND	FTLD-TDP-U, MND	R	14	2	1	1	Absent	Low	8	15
C8 ^b	67	М	bvFTD	FTLD-TDP-B; MND	R	23	1	3	3	Sparse	Int.	16	15
C9	65	F	bvFTD	FTLD-TDP-U	R	8	1	1	3	Absent	Low	15	9
C10	57	М	bvFTD	FTLD-TDP-B, MND	L	2	0	1	0	Absent	Low	10	10
C11 ^c	59	М	bvFTD-MND	ALS-TDP	R	12	ND	1	1	Absent	Low	8	ND
C12 ^C	60	F	bvFTD-MND	FTLD-TDP-B, MND	R	17	1	1	4	Absent	Low	10	ND

^aThe diagnosis thought most likely to have caused the clinical syndrome; Sx dur: symptom duration; ADNC: Alzheimer's disease neuropathologic change

 ${}^{b}\!\!\!\mathrm{Subject}$ included in inclusion fraction and morphometry studies

^cSubject included in morphometry study only

NC: neurologically unaffected control; S: sporadic disease; C: C9orf72-associated disease; N/A: not applicable; R: right; L: left; ND: not done