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Synapse loss in the prefrontal cortex is associated with cognitive decline in amyotrophic lateral sclerosis

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Abstract
In addition to motor neurone degeneration, up to 50% of amyotrophic lateral sclerosis (ALS) patients present with cognitive decline. Understanding the neurobiological changes underlying these cognitive deficits is critical, as cognitively impaired patients exhibit a shorter survival time from symptom onset. Given the pathogenic role of synapse loss in other neurodegenerative diseases in which cognitive decline is apparent, such as Alzheimer’s disease, we aimed to assess synaptic integrity in the ALS brain. Here, we have applied a unique combination of high-resolution imaging of post-mortem tissue with neuropathology, genetic screening and cognitive profiling of ALS cases. Analyses of more than 1 million synapses using two complimentary high-resolution techniques (electron microscopy and array tomography) revealed a loss of synapses from the prefrontal cortex of ALS patients. Importantly, synapse loss was significantly greater in cognitively impaired cases and was not due to cortical atrophy, nor associated with dementia-associated neuropathology. Interestingly, we found a trend between pTDP-43 pathology and synapse loss in the frontal cortex and discovered pTDP-43 puncta at a subset of synapses in the ALS brains. From these data, we postulate that synapse loss in the prefrontal cortex represents an underlying neurobiological substrate of cognitive decline in ALS.

Introduction

Amyotrophic lateral sclerosis (ALS) is the most common form of motor neurone disease (MND), resulting in progressive upper and lower motor neurone dysfunction and loss. This leads to progressively severe muscle wasting and most patients succumb due to respiratory failure, with a mean survival time of 3 years post diagnosis [57]. Due to the striking, and in many cases rapid, deterioration of motor function, ALS was historically considered a homogeneous disease of the motor system. However, both clinical and post-mortem studies increasingly describe ALS as a heterogeneous disorder, with around 50% of patients exhibiting neuropsychological deficits due presumably to breakdown of non-motor areas of the brain [49]. ALS is thought to lie on a disease spectrum with frontotemporal dementia (FTD). These seemingly disparate disorders share both genetic and pathological features [30] with considerable clinical overlap. An estimated 15% of FTD patients present with motor decline meeting the criteria for ALS diagnosis and 15% of ALS cases show
severe cognitive and or behavioral decline, meeting the FTD diagnostic criteria [24, 30, 49]. However, a further 30–40% of ALS patients show milder, more specific cognitive and/or behavior changes (ALS cognitive impairment—ALSci/ALS behavioral impairment ALSbi [49]) that does not meet FTD diagnostic criteria but can still affect daily life [20]. Changes include executive dysfunction, decreased verbal fluency, language impairment and social cognition deficits, which are thought to be mediated by regions of the frontal cortex and are similar to mild FTD-associated impairments [20]. Behavior change is predominantly one of the apathies, while other features of FTD are also seen, including disinhibition and loss of sympathy/empathy [20]. Functional and structural imaging studies support the presence of frontal deficits [11, 34, 47, 56], and specifically relate dysfunction of the dorsolateral prefrontal cortex [including Brodmann Area 9 (BA9)] to the cognitive profile in ALS [1, 2]. It is vitally important that we understand the underlying neuropathology driving these cognitive deficits, as cognitively impaired patients not only endure the personal burden of both motor and cognitive dysfunction, but also exhibit a shorter survival time from symptom onset [15, 58].

Synapse degeneration is a common feature in a large number of neurodegenerative diseases and often plays a role in pathogenesis, rather than simply occurring as a secondary effect of neuron loss [22]. For example, synaptic loss remains the strongest neuropathological correlate of cognitive decline in Alzheimer’s disease (AD [54]). In AD, synaptic accumulation of key pathological forms of tau and beta-amyloid are thought to drive synaptic breakdown [48]. Likewise, studies in ALS mouse models have suggested that synapse defects in motor regions of the cortex are an early/pre-symptomatic feature of the disease [18, 43]. However, to date, few studies have attempted to assess the contribution of synaptic change to pathology of non-motor cortical regions in the brain of human ALS patients and none using direct measurement in patients with defined ALS with cognitive impairment.

By combining the high-resolution imaging techniques of array tomography and transmission electron microscopy (TEM), we have examined synapses in the frontal cortex and motor cortex of cognitively unimpaired and cognitively impaired ALS (ALSci) patients, to define the relationship between synaptic breakdown and cognitive impairment. Cognitive profiling was performed using the Edinburgh Cognitive and Behavioural ALS Screen (ECAS) [3], a short multi-domain assessment designed for people with physical disability which has been validated against extensive neuropsychology [38]. Other post-mortem measurements such as cortical thickness, pathological burden and gliosis were recorded to ensure a comprehensive profiling of each case.

Materials and methods

Participant characteristics

All patients had clinical and electrophysiological evidence of combined upper and lower motor neurone involvement and fulfilled the revised El Escorial criteria for a diagnosis of ALS [10]. Patients were recruited through the Scottish Motor Neurone Disease Register and data collected in the CARE-MND database (Clinical Audit Research and Evaluation). Ethical approval for this register was obtained from Scotland A Research Ethics Committee 10/MRE00/78 and 15/SS/0216. All clinical data were subsequently extracted from the CARE-MND database. Clinical details can be seen in Table 1. There was no documented evidence of dementia in the CARE-MND database.

Patients gave pre-mortem consent for brain and spinal cord donation. All tissue retrievals were in line with the Human Tissue (Scotland) Act, and all procedures have been approved by a national ethics committee. Use of human tissue for post-mortem studies has been reviewed and approved by the Sudden Death Brain Bank ethics committee and the ACCORD medical research ethics committee, AMREC (ACCORD is the Academic and Clinical Central Office for Research and Development, a joint office of the University of Edinburgh and NHS Lothian). Further, some brains from human subjects with no cognitive deficits were obtained through the Massachusetts Alzheimer’s Disease Research Centre and Massachusetts General Hospital Neuropathology department. All donor tissue was obtained in accord with local and National Institutional Review Board regulations. Details of all donors used in this study can be found in Supplementary Table 1, with means for each group found below in Table 1.

At post-mortem, the brain was removed as detailed in [46] and cut into coronal slices. Regions of interest were then dissected from each coronal slice. Each region of interest from one hemisphere was then processed for paraffin embedding. Samples from the other hemisphere were dissected into smaller segments then processed for array tomography or transmission electron microscopy (see below for details).

Cognitive profiling

The ECAS is a brief multi-domain assessment which is specifically designed for ALS. It includes the assessment of domains which are typically affected in ALS (Fluency, Executive and Language functions) and which together produce an ALS-Specific Score (max 100). It also includes the assessment of domains which are more typically
Table 1  Summary demographics of all donors

<table>
<thead>
<tr>
<th>Cohort 1-EM</th>
<th>Number</th>
<th>Male/female</th>
<th>Age at diagnosis (median + range)</th>
<th>Age at PM (median + range)</th>
<th>Site of onset</th>
<th>Riluzole (yes/no)</th>
<th>Disease duration (months)</th>
<th>Time between ECAS and death (months)</th>
<th>ECAS total (mean ± SD)</th>
<th>ALS-specific ECAS (mean ± SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>5</td>
<td>4/1</td>
<td></td>
<td>61 (53–77)</td>
<td></td>
<td>4/1</td>
<td>17/9</td>
<td>44 (17–216)</td>
<td>116 ± 4.9</td>
<td>86 ± 3.6</td>
</tr>
<tr>
<td>ALS</td>
<td>20</td>
<td>9/11</td>
<td></td>
<td>63 (40–89)</td>
<td></td>
<td>9/11</td>
<td>44 (17–216)</td>
<td>116 ± 4.9</td>
<td>86 ± 3.6</td>
<td></td>
</tr>
</tbody>
</table>

Cohort 2a-array tomography

| Control     | 14     | 11/3        | 79 (62–95)                        | 4/1            | 17/9 | 44 (17–216) | 116 ± 4.9 | 86 ± 3.6 |
| ALS         | 29     | 18/11       | 60 (32–75)                        | 64 (40–89)     | 20 lim, 7 bulbar | 4/6 | 39 (24–216) | 116 ± 4.9 | 86 ± 3.6 |

Cohort 2b (subset of cohort 2)-cognitive profiling

| Control     | 14     | 11/3        | 79 (62–95)                        | 4/1            | 17/9 | 44 (17–216) | 116 ± 4.9 | 86 ± 3.6 |
| ALS         | 16     | 9/7         | 60 (32–71)                        | 64 (40–83)     | 11 lim, 5 bulbar | 4/6 | 39 (24–216) | 116 ± 4.9 | 86 ± 3.6 |
| ALS-Sci     | 7      | 4/3         | 61 (38–72)                        | 68 (43–75)     | 4 lim, 2 bulbar  | 4/3 | 49 (34–64)  | 116 ± 4.9 | 86 ± 3.6 |

Cohort 3-CSF

| ALS         | 9      | 6/3         | 58 (41–69)                        | 3 lim, 1 bulbar, 5 mixed (bulbar and limb) | 28 (12–84) | 112.6 ± 5.5 |
| ALS-Sci     | 6      | 4/2         | 72 (65–80)                        | 5 lim, 1 bulbar | 15 (1–36) | 62.3 ± 16.6 |

PM post-mortem, ECAS Edinburgh Cognitive and Behavioral ALS Screen, EM electron microscopy, ALS-Sci ALS with cognitive impairment
affected in other disorders such as Alzheimer’s disease (Memory, Visuospatial Functions) and which make up the ALS non-specific score (max 36). Together, the combined ALS-specific and ALS non-specific scores produce an ECAS total Score (max 136). The assessment has been validated against extensive neuropsychology and provides high sensitivity and specificity using published cut-offs with patients classified as ALSci if ALS-specific and/or ECAS total Scores are below threshold [38]. It also shows good convergent validity with other screens not designed for ALS [31, 41].

**Genetics**

A subset of our patient participants actively consented to donate blood and saliva to the Scottish Regenerative Neurology Tissue Bank for genetic analysis. Use of patient samples for genetic profiling has been approved by the Chief Scientist Office Scotland; MREC/98/0/56 1989–2010, 10/MRE00/77 2011–2013, 13/ES/0126 2013–2015, 15/ES/0094 2015-present. C9orf72 genotyping was performed as described in [12] and most cases were genetically analyzed as described in [5]. Some cases were analyzed by next generation sequencing to screen for variants in the coding regions of SOD1, TARDBP, FUS, VCP, UBQLN2, SQSTM1, CHMP2B and VAPB. Library construction was carried out using the Ion AmpliSeq library kit v2.0 (Life Technologies). Emulsion PCR and enrichment was performed using the Ion OneTouch2 instrument (200p template kit) and Ion ES module. Sequencing was carried out on an Ion Torrent Personal Genome Machine (Life Technologies) using the Ion PGM sequencing 200 kit v2 and an Ion 316 chip. All stages followed the manufacturer’s protocols. NextGENe software (Softgenetics) was used with hg19 (GRCh37) human genome as reference to identify variants.

**Neuropathology**

Fresh post-mortem tissue blocks were fixed in 10% formalin for a minimum of 24 h. Tissue was dehydrated in an ascending alcohol series (70–100%), followed by three xylene washes, all for 4 h each. Next, three paraffin waxing stages (5 h each) were performed to ensure full penetration of the embedding wax and the blocks were cooled. Tissue sections were cut on a Leica microtome at 4 µm and collected on glass slides. All sections were dried at 40 °C for at least 24 h before staining. Immunohistochemistry was performed using standard protocols, enhanced using the Novolink Polymer detection system and visualized using DAB as chromogen. See Table 2 for antibody information. Slides were finally counterstained with hematoxylin for 30 s to stain cell nuclei. For each staining run, positive and negative control tissue for each pathological marker were used to ensure specificity and success of each experiment. All staining was performed by experimenters blind to clinical diagnosis.

For pTDP-43 and pTau analysis, cases were described as being positive or negative depending on the presence of clearly stained protein aggregates.

**Burden quantification**

All sections stained for β-amyloid, CD68 or GFAP was assessed for plaque, microglia and astrocytic burden, respectively, using Stereo Investigator. Cortical gray matter was outlined in each section and immune-positive objects identified using an automated color-based thresholding algorithm. The area of immuno-positive cortex was expressed as a percentage of total cortex in each brain region. All imaging and analysis were performed by experimenters blind to clinical diagnosis.

### Table 2 Primary antibody information

#### Histopathology

<table>
<thead>
<tr>
<th>Antibody</th>
<th>Company</th>
<th>Code</th>
<th>Dilution</th>
<th>Pre-treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Beta amyloid (BA4)</td>
<td>Dako</td>
<td>M087201-2</td>
<td>1:100</td>
<td>98% Formic acid 5 min</td>
</tr>
<tr>
<td>pTDP-43</td>
<td>2B Scientific</td>
<td>CAC-TIP-PTD-MO1</td>
<td>1:4000</td>
<td>Pressure cooker/citric acid</td>
</tr>
<tr>
<td>pTau (AT8)</td>
<td>Thermo</td>
<td>MN1020</td>
<td>1:2500</td>
<td>None</td>
</tr>
<tr>
<td>GFAP</td>
<td>Dako</td>
<td>Z0334</td>
<td>1:800</td>
<td>None</td>
</tr>
<tr>
<td>CD68</td>
<td>Dako</td>
<td>M0876</td>
<td>1:100</td>
<td>Pressure cooker/citric acid</td>
</tr>
</tbody>
</table>

#### Array tomography

<table>
<thead>
<tr>
<th>Antibody</th>
<th>Company</th>
<th>Code</th>
<th>Dilution</th>
<th>Secondary antibody</th>
</tr>
</thead>
<tbody>
<tr>
<td>Synaptophysin</td>
<td>Abcam</td>
<td>Ab8049</td>
<td>1:50</td>
<td>Donkey α Mouse—AF594 or AF488</td>
</tr>
<tr>
<td>PSD95</td>
<td>Synaptic Systems</td>
<td>124,014</td>
<td>1:50</td>
<td>Donkey α Guinea Pig—AF647</td>
</tr>
<tr>
<td>pTDP-43</td>
<td>Proteintech</td>
<td>22309-1-AP</td>
<td>1:50</td>
<td>Donkey α Rabbit—AF594 or AF488</td>
</tr>
</tbody>
</table>
Cortical thickness

Gray matter thickness was calculated at ten randomly selected points throughout each section. Thickness was measured from the pial surface to the border of grey/white matter, below cortical layer 6. All ten measurements were averaged to give a cortical thickness for each brain region. All imaging and analysis were performed by experimenters blind to clinical diagnosis.

Neuron counts

Paraffin-embedded sections from the frontal cortex of 10 ALS cases were used to assess neuronal density as previously described [21]. The five ALS cases with the highest and the five with the lowest synapse density as measured by array tomography were analyzed. All imaging and analysis were performed by experimenters blind to clinical diagnosis.

Neurofilament light-chain CSF ELISA

Fifteen patients provided CSF for this study. Patients were seen at the Department of Neurology in Ulm. The study, whose overall aim was to find biomarkers for neurological diseases, was approved by the local ethics committee in Ulm according to institutional guidelines and patients provided written informed consent to participate (approval number 20/10). A summary of patient group characteristics is given in Supplementary Table 1. The physical function of ALS patients’ status was classified with the revised ALS functional rating scale (ALSFRS-R) with a maximum of 48 points, where lower values represent a more severe disease stage.

CSF was obtained by lumbar puncture, centrifuged, aliquoted and stored within 2 h at − 80 °C until analysis. Neurofilament light-chain levels in the CSF were examined using a commercial 96-well ELISA kit from UmanDiagnostics (Umea, Sweden). CSF samples were diluted 1:20 in sample diluent before analysis and the ELISA was run according to the manufacturer’s instructions, with all CSF samples and kit standards run in duplicate.

Electron microscopy

Fresh post-mortem samples, stored in 0.1 M PB were trimmed into small cortical blocks and fixed in 4% paraformaldehyde and 2.5% glutaraldehyde in 0.1 M PB for 3 h. Fixed blocks were washed twice in 0.1 M PB before being cut at 70 μm with a vibratome. Sections were then treated with osmium tetroxide (1% in 0.1 M PB) for 30 min (protected from light) and dehydrated in an ascending series of ethanol and propylene oxide, before embedding in Durcupan resin. During dehydration, the sections were treated with uranyl acetate (1% in 70% ethanol) for 40 min in the dark. Durcupan resin was polymerized for 48 h at 56 °C. Small regions of interest were cut from the Durcupan-embedded sections and glued onto Durcupan blocks, before cutting into 70 nm ribbons using an ultracut microtome (Leica) with a Jumbo Histo Diamond Knife (Diatome, Hatfield, PA). Ribbons were collected on grids and stained with lead citrate before imaging on a Philips CM12 transmission electron microscope equipped with a Gatan digital camera. Qualitative and quantitative synapse analyses were performed by an investigator who was blinded to the disease status of each sample. Individual synaptic profiles were identified based on the presence of clear pre- and post-synaptic terminals, containing synaptic vesicles and post-synaptic density, respectively. Degenerating synapses were classified as those with an electron-dense cytoplasm, as previously described [19, 26].

Array tomography

Brain samples were prepared for array tomography as outlined previously [26]. Briefly, fresh post-mortem tissue was trimmed into small cortical blocks and fixed in 4% paraformaldehyde and 2.5% sucrose in PBS for 2–3 h. Samples were then dehydrated through ascending ethanol washes and into LR White resin (Electron Microscopy Sciences) overnight. Blocks were placed individually into capsules containing LR White and polymerized overnight at 53 °C. Tissue blocks were cut into ribbons of 70 nm serial sections using an ultracut microtome (Leica) with a Histo Jumbo Diamond Knife (Diatome, Hatfield, PA) and collected on gelatin-coated glass coverslips. For synaptic density analysis, ribbons were immunostained as described previously [21], with a primary antibody against synaptophysin overnight (see Table 2 for antibody details) and no primary negative control was included to rule out non-specific binding of the secondary antibodies. The following day, the staining was developed with a fluorescently labeled secondary antibody (1:50 dilution) and finally mounted onto slides with Immumount mounting media. To obtain improved staining with the pTDP-43 antibody, array ribbons underwent epitope retrieval. Ribbons were pressure-cooked for 2 min in citrate buffer (pH 6), and then allowed to cool to room temperature. Ribbons were washed with TBS (2 × 5 mins) and then underwent the same staining protocol as described above. Three antibodies were used in combination (see Table 2 for details), to analyze the presynapse (synaptophysin), postsynapse (PSD95) and pTDP43 co-localization. Once the ribbons had been stained and mounted, images were obtained at the same position on each section along the ribbon, using a Zeiss axio Imager Z2 epifluorescent microscope equipped CoolSnap digital camera and AxioImager software with array tomography macros (Carl Zeiss, Ltd, Cambridge, 2017).
UK). First, a tile scan of the entire ribbon was obtained at 10× magnification; then regions of interest were chosen on the tilescan and a position list generated. High-resolution images were obtained from the position list with a 63× 1.4 NA Plan Apochromat objective. At least two image stacks were captured from at least two tissue blocks per region, per case. This meant at least four image stacks were captured per region for each case.

Stacks were aligned using the MultiStackReg ImageJ plugin [55]. Regions of interest (ROI) were automatically chosen in the neuropil of each aligned image stack using an in-house MATLAB cropping script and coded for blind analysis, before thresholding with an automated algorithm in ImageJ. To remove false background staining which was only found on single sections, we used the Watershed script [35] which also provides 3-dimensional (3D) object counts. These 3D objects (synaptic puncta) were used to generate synapse densities (synaptic puncta per mm³) based on the ROI volume. All individual ROI synapse densities (between 100 and 300 ROIs per region, per case) from each region, from each case were combined and a mean density value generated. All imaging and analysis were performed by experimenters blind to clinical diagnosis.

Statistics

GraphPad Prism 7.02 was used to test data for normality (D’Agostino and Pearson normality test). A mean or median for each subject was calculated for each measurement, following which parametric (t test or ANOVA) or non-parametric (Mann–Whitney or Kruskal–Wallis) tests were performed as required. Significance was reported when \( p < 0.05 \). Data are usually presented as mean ± standard deviation, unless otherwise stated.

Data sharing

Data used in this manuscript including custom image analysis scripts for ImageJ will be freely shared on the University of Edinburgh Data Repository. Analysis macros can be found at the following link: https://dx.doi.org/10.7488/ds/2268. Original raw image files can be found at the following link: http://dx.doi.org/10.7488/ds/2269.

Results

Transmission electron microscopy (TEM) analysis of the prefrontal cortex [Brodmann Area 9 (BA9); ~150 synapses per case] from 5 control and 20 ALS cases (Table 1; cohort 1) revealed a significant decrease in synaptic density in the ALS cohort (Fig. 1a–c). In addition, there was a significant increase in the number of electron-dense, degenerating pre-synaptic terminals in ALS cases compared to controls (Fig. 1b, d) [19].

To expand our synaptic analyses, we next examined a largely independent cohort (Table 1; cohort 2a) of subjects using higher-throughput array tomography. Analysis of over 1 million synapses (~36,000 synapses per case) from 14 controls and 29 ALS brains using array tomography confirmed our initial TEM findings revealing synapse loss in BA9 of ALS patients (2-tailed unpaired \( t \) test; \( p = 0.013 \); Fig. 1e–g). Post-mortem interval (PMI) did not influence synapse density (Pearson \( r = –0.21, p = 0.17 \)). Importantly, the loss of synaptic density was not accompanied by any detectable gross atrophy of BA9 (2-tailed unpaired \( t \) test; \( p = 0.68 \); Fig. 1h). Furthermore, stereological estimates of neuron density counts from 5 cases with the highest and lowest synapse densities were not significantly different (Low: \( n = 5, 23,370 ± 2756 \) neurons/mm³; High: \( n = 5, 28068 ± 2094 \) neurons/mm³). Unpaired 2-tailed \( t \) test; \( p = 0.22 \), suggesting that synapse loss was occurring in the absence of concomitant, overt neuronal loss.

We then performed some exploratory analyses to assess the influence of numerous genetic and pathological factors that may associate with increased synapse loss. Genetic screening revealed that 6 ALS cases had a hexanucleotide repeat expansion in C9orf72, 3 cases had SOD1 variations and 2 had NEK1 variations (Supplementary Table 1). However, we did not observe any significant association between lower synapse density in BA9 and distinct genetic mutations in ALS with this sample size (Fig. 1g; Supp. Fig. 1).

Pathological changes in disease-associated proteins are strongly implicated in driving synapse degeneration in dementias [48]. Thus, we next assessed the presence of disease-associated pathological aggregates in formalin-fixed paraffin-embedded (FFPE) sections from BA9 and their association with synapse loss in array tissue from the same case. We found no association between the presence of beta-amyloid or pTau pathology in BA9 and synapse loss (Supp. Fig. 2). We also found no association between the presence of pTDP-43 aggregates in BA9 and synapse loss (Supp. Fig. 3). However, it is known that ALS patients with SOD1 mutations do not express pTDP-43 pathology [32]. When we remove our three SOD1 + ve cases from the analysis, we find a significantly lower synapse density in ALS cases with pTDP-43 pathology (Supp. Fig. 3). Due to this potential association and to the literature from Alzheimer’s disease in particular, indicating that pathological protein accumulation at synapses can drive synapse degeneration [48], we used array tomography to assess the presence of pTDP-43 at the synapse. We observed pTDP-43 pathology in all of the 9 ALS cases screened, often exhibiting as large cytoplasmic aggregates or neuritic threads (Fig. 1i). Interestingly, we also
noticed small pTDP-43 puncta in the neuropil of our 9 cases, which in a subset of synapses co-localized with synaptic markers (Fig. 1i, j).

Twenty-three patients from whom we collected tissue for array tomography had previously been cognitively profiled using the Edinburgh Cognitive and Behavioural ALS Screen (ECAS). These individuals (Table 1; cohort 2b) were classified as unimpaired (ALS) or ALS with cognitive impairment (ALSci) according to established threshold scores for ECAS Total or ALS-specific ECAS results [3]. The scores revealed 16 unimpaired cases and 7 impaired cases [70% unimpaired, 30% impaired (Fig. 2a)]. Comparing ECAS data with synaptic data, we found a significant difference between our three groups in synaptic

Fig. 1 Synapse loss in ALS frontal cortex. Representative electron micrographs showing two excitatory synapses in the frontal cortex of a healthy control (a) and an ALS case (b). Edges of the post-synaptic density (PSD) are marked by red arrowheads and presynaptic terminals by black asterisks. The white asterisk in b highlights an electron dense, degenerating presynaptic terminal. Scale bars 100 nm. Decrease in synaptic density in the frontal cortex [(c) 2-tailed unpaired t test; \( p = 0.016 \)], and increased numbers of degenerating synapses [(d) 2-tailed Mann–Whitney test; \( p = 0.014 \)] in ALS patients (\( n = 20 \)) compared to controls (\( n = 5 \)), measured using TEM. Each data point represents the mean total synapse (c) or degenerating synapse (d) count per 100 \( \mu m^2 \) for each individual. Three-dimensional reconstructions of fifteen 70 nm array tomography sections from a control frontal cortex (e) and an ALS frontal cortex (f), stained for synaptophysin (green) and DAPI (white). Scale bar is 10 \( \mu m \). g Decrease in synaptic puncta (2-tailed unpaired t test; \( p = 0.017 \)) in ALS patients compared to controls, measured using array tomography. Each data point represents the mean synapse count per mm\(^3\) of frontal cortex, for each individual (control \( n = 14 \); ALS \( n = 29 \)). Blue icons = C9orf72 expansions, Red icons = SOD1 variant and Green icons = NEK1 variant. h Cortical thickness comparisons between control (\( n = 7 \)) and ALS (\( n = 34 \)) revealed no difference (2-tailed unpaired t test; \( p = 0.76 \)). Each data point represents the mean of ten measurements per cortical block. All histogram bars plot the mean ± SD. i 3D reconstruction of 25 array tomography sections (each 70 nm thick), stained for synaptophysin (red), PSD95 (blue), pTDP-43 (green) and DAPI (white). A large perinuclear pTDP-43 aggregate (white arrow), numerous pTDP-43 + ve neuritic threads (white arrowheads) and small pTDP-43 puncta (open arrows) are clearly visible. Scale bar 10 \( \mu m \). By cropping the image in i, compressing 3 consecutive sections and then splitting the channels, it is evident in panel (j) that pTDP-43 aggregates are found both in some pre (synaptophysin, red) and post (PSD95, blue) synaptic compartments. Scale bar 1 \( \mu m \)
density and that the cognitively impaired group had a significantly lower synapse density in BA9 than controls (Fig. 2b). We found no significant difference between the ALS unimpaired cohort and either controls or ALSci, most likely due to two cases with very low densities in the unimpaired group (Fig. 2b). When we plotted ECAS score versus synapse density, we found no significant correlation between these data (Pearson $r = 0.31$, $p = 0.15$), again due to the influence of the two unimpaired cases with low synapse density mentioned above (Supp. Fig. 4). One potential confounding factor is the length of time between cognitive testing and post-mortem synapse analysis. However, we found no correlation between synapse density at post-mortem and time since cognitive testing (Pearson $r = 0.043$, $p = 0.85$). Although there was no association between genetic status and synapse density or cognitive impairment, it was interesting to note that 3 of our 4 genetically screened cognitively impaired cases were positive for C9orf72 repeat expansions (Fig. 2b), warranting further study of the effects of C9orf72 expansions, as more cases prepared for these techniques become available. Cortical thickness did not vary between the groups (ALS, ALSci, Controls) (Fig. 2c), suggesting that atrophy and overt neuron loss were not driving cognitive decline. We found no association between age and ECAS score (Supp. Fig. 5) or cortical thickness and ECAS score (Supp. Fig. 5). Furthermore, the presence of beta-amyloid or pTDP-43 pathology in BA9 did not associate with lower ECAS score (Fig. 2d, f). Interestingly, although the presence of pTau pathology did not associate with a lower mean ECAS score (2-tailed unpaired t tests; $p > 0.05$), pTau +ve cases were predominantly ALSci (5 of the 7 pTau +ve cases were ALSci), whereas pTau-ve cases were predominately unimpaired ALS cases (12 of the 14 pTau-ve cases were unimpaired ALS; Fisher’s exact test:
p = 0.005). This suggests that pTau pathology may associate with cognitive impairment and warrants further study once more cases become available.

Eleven of the ECAS-screened cases also underwent the ECAS behavior interview, which was completed by an informant/carer in vivo. Six cases were unimpaired and five were classified as ALS with behavioral impairment (ALSbi), with changes including apathy, lack of empathy and behavioral disinhibition. Analysis revealed no difference in BA9 synapse densities between the three groups of controls, ALS and ALSbi (One-way ANOVA; F = 1.93; p = 0.17).

Together, these data suggest that synapse density is a robust indicator of ECAS score and cognitive performance in ALS.

To address whether synapse breakdown, in the absence of cortical atrophy, is a global feature of pathology in the ALS brain, or a specific event occurring in the prefrontal cortex, we employed an array tomography approach to analyze ~750,000 synapses from the primary motor cortex. Comparing synapse density in 7 controls versus 25 ALS patients, we discovered no difference in synapse density (Fig. 3a–c). Grouping the BA4 synapse values based on cognitive status did not reveal any difference in synapse density between the groups (Fig. 3d). However, there was significant cortical thinning of the motor cortex in ALS patients compared to controls (Fig. 3e). When the ALS group was divided into unimpaired ALS and ALSci, it was found that both groups exhibited significant cortical atrophy compared to controls (Fig. 3f). Furthermore, the assessment of neurofilament light-chain in the cerebrospinal fluid (CSF) of another cohort of ALS and ALSci patients (Table 1; cohort 3) revealed comparable levels (ALS = 7950 ± 2030 pg/ml, ALSci = 11,068 ± 3073 pg/ml. 2-tailed unpaired t test; p = 0.39), suggesting the presence of similar neuronal breakdown across the brain in both groups (Supp. Fig. 6). When comparing synapse density in the frontal cortex to the motor cortex within the healthy control brains, we discovered a lower density in the motor cortex (Supp. Fig. 7; 2-tailed paired t test p = 0.036). This difference in regional synapse density was also observed in the larger group of cognitively unimpaired ALS cases (Supp. Fig. 7; 2-tailed paired t test p = 0.005), but was lost in the ALSci cases (Supp. Fig. 7; 2-tailed paired t test p = 0.75). This may suggest that synapse densities in the frontal cortex of ALSci brains had dropped to levels similar to those found in the motor cortex. We found no association between beta-amyloid, pTau or pTDP-43 pathology in BA4 and synapse density (Fig. 3g–i). Further, we found no association between beta-amyloid (Fig. 3j), pTau (Fig. 3k) or pTDP-43 (Fig. 3l) pathology in the motor cortex and cognitive score.

Glial cells are thought to play a prominent role in ALS pathogenesis [40], so we analyzed GFAP and CD68 burdens as a measure of astrocyte and microglial activity, respectively, in both BA9 and BA4. We found no correlation between GFAP burden and synapse density (Supp. Fig. 8) in BA9 (Pearson r = 0.127, p = 0.51) or in BA4 (Pearson r = 0.16, p = 0.44). There was also no correlation between CD68 burden and synapse density (Supp. Fig. 8) in BA9 (Pearson r = −0.012, p = 0.95) or BA4 (Pearson r = 0.036, p = 0.86). Furthermore, there was no difference in GFAP or CD68 burden in BA9 (Supp. Fig. 8) in ALS patients compared to ALSci (2-tailed unpaired t tests. GFAP; p = 0.09; CD68: p = 0.59). This suggests that synapse loss in the ALSci patients is not associated with altered gliosis.

Taken together, these data suggest that synapse degeneration, occurring specifically within frontal (and not motor) cortex of ALS patients with cognitive deficits, is one underlying neurobiological contributor to cognitive decline. Of all the pathological changes measured, synapse degeneration in BA9 was the strongest predictor of cognitive impairment.

**Discussion**

We have optimized a combination of cognitive and genetic profiling with post-mortem tissue collection, neuropathology and high-resolution subcellular analyses. Our human tissue collection and processing protocol [26] robustly preserves synaptic structures for both electron microscopy and array tomography analysis. To the best of our knowledge, this is the only ALS cohort in the world that has been extensively characterized to this level of detail.

A few previous studies attempted to assess synaptic changes in human post-mortem tissue using less direct measurements such as optical density of immunohistochemical staining or western blotting [25, 45], but failed to observe any difference between ALS patients and controls. However, one potential confound is that we do not know the cognitive status of the patients in these studies [25, 45] and taken together our study highlights the importance of using highly precise, high-resolution approaches (such as array tomography and TEM) when studying synaptic changes in post-mortem material. One very recent study has detected a loss of synaptic terminals in the putamen and caudate of ALS patients, using immunohistochemical approaches [44]. Patients with severe cognitive decline meeting the diagnosis of FTD or FTD-MND had significant synaptic loss, whereas the ALS group had a more variable phenotype. Unfortunately, the ALS group was not stratified by cognitive status, so we cannot discern whether the synapse loss is related to cognitive decline in the ALS cases [44].

Synapse loss is the strongest correlate of cognitive decline in Alzheimer’s disease [54] and a recent meta-analysis highlighted pre-synaptic terminals as the major site of early pathogenesis in the disease [14]. Therefore, the loss of
Patients with cognitive decline, compared to cognitively
expression of TDP-43 aggregates in the frontal cortex of
synapse loss in the frontal cortex of ALSci patients is not
in the frontal cortex and lower ECAS score (Fig. 2). This
in pathology progression through the central nervous
system has been described [8]. It is believed that TDP-
pathology may progress through synaptically connected
brain regions [6, 16], in a similar manner to other
neurodegenerative diseases [7], thus placing synapses at a
crucial point in ALS pathogenesis. Furthermore, in a human TDP-
(HTDP43) overexpressing mouse model, synaptophysin
to note widespread brain homogenates [33] suggesting
up to TDP-43 function may drive synapse loss.
Intriguingly, four studies have described what appears to be
"synaptic" labeling of pTDP-43 in the neuropil of human
post-mortem cortex from ALS and FTD patients [4, 37, 44,
A recent study described pTDP-43 puncta in very close
proximity to synaptophysin puncta in human ALS brain
[53], with the suggestion that small pTDP-43 aggregates
may be accumulating in dendritic spines. However, another
recent study suggests pTDP-43 puncta co-localize with syn-
aptophysin [44], thus suggesting these small aggregates are
in fact presynaptic. In support of this, we also noted punctate
labeling of pTDP-43 in the neuropil of some of our ALS
cases and found pTDP-43 puncta in a number of synapses in
a subset of ALS cases (Fig. 1i, j). Given that synaptic accu-
mulation of pathological proteins in Alzheimer’s disease and
FTLD-tau animal models drives synapse loss [27–29, 51,
52], it is possible that a similar build-up of pTDP-43 in the
ALS synapse may initiate synapse breakdown. An alterna-
tive explanation is the phagocytosis of synaptic components
by pathologically active glial cells, as recently described
in Alzheimer’s disease [23]. In support of this, we recently
observed that when TDP-43 is specifically knocked out in
microglial cells, they adopt a hyper-phagocytic phenotype.
The microglial cells actively phagocytosed synapses, result-
ing in a significantly lower synaptic density in the cortex
of these transgenic mice compared to controls [39]. Fur-
thermore, in a human cohort of ALS patients, pTDP-43
positive cases had significantly higher expression of the
phagocytic microglial marker CD-68 [39]. Interestingly, in
our current study, we found the same result in the frontal
cortex of ALS patients. Those with pTDP-43 in the frontal
cortex had a higher CD-68 burden than ALS cases with-
out pTDP-43 (pTDP-43-positive n = 14, pTDP-43-negative
n = 20; 2-tailed unpaired t test; p = 0.028). However, CD68
expression did not correlate with synaptic density (Pearson
r = −0.012; p = 0.99) and thus further work is required to
understand the role of phagocytic microglia in synapse loss.

We found no direct association between the presence
of the dementia-associated pathologies, beta-amyloid and pTau
in the frontal cortex and lower ECAS score (Fig. 2). This
is in agreement with recent studies showing no association
between Alzheimer’s pathology and cognition in ALS [9,
42]. However, it is interesting to note that 5 out of the 7
ALSci cases were positive for pTau and 5 of the 7 pTau + ve
cases were ALSci. This finding aligns with the proposed
model that certain forms of pTau in the frontal cortex may
associate with cognitive decline in ALS [50, 59, 60] and
warrants further study once more cases become available to

Synapses from the frontal cortex of ALS patients described
here fits well with the features of other disorders where cog-
nitive decline is observed. ALMS mouse models, including the hSOD1G93A
and FUSR521C models, have revealed pre-symptomatic spine
loss in the motor cortex [18] and sensorimotor cortex [43],
respectively. Intriguingly, in the hSOD1G93A model, early
pre-symptomatic loss of spines on pyramidal cells in the
medial prefrontal cortex preceded changes in dendritic
arborization [17]. The earliest onset of anatomical change
occurred in the motor cortex, with such severe neuronal
breakdown that cortical thinning was detected at pre-sym-
tomatic time-points [17]. Thinning of the medial prefrontal
cortex was only detected in adult diseased mice, by which
time spine loss and dendritic changes were significant [17].
This temporal pattern of cortical breakdown observed in
mouse models fits with our observations in human post-
mortem tissue. We observed synapse loss in the prefrontal
cortex without neuron loss, yet found significant cortical
thinning in the ALS motor cortex. Furthermore, our CSF
results suggest that similar levels of neuronal breakdown
occurred across the brain of ALS and ALSci groups (Supp.
Fig. 5). This is most likely due to the cortical thinning
observed in the motor cortex and supports our finding that
synapse loss in the frontal cortex of ALSci patients is not
due to neuronal loss.

Two human post-mortem studies have found greater
expression of TDP-43 aggregates in the frontal cortex of
patients with cognitive decline, compared to cognitively
unimpaired ALS patients [9, 13]. TDP-43 aggregates are
often found in the brain of ALS patients, and a sequential
staging of pathology progression through the central nervous
system has been described [8]. It is believed that TDP-
43 pathology may progress through synaptically connected

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\text{Fig. 3 Synapse density in the ALS motor cortex does not associate with cognitive decline. Three-dimensional reconstruction of twenty-four 70 nm array tomography sections from a control motor cortex (a) and an ALS motor cortex (b), stained for synaptophysin (green) and DAPI (white). Scale bar is } 10 \mu \text{m. e Histogram showing no change in synaptic puncta within the motor cortex (2-tailed unpaired } t \text{ test; } p = 0.73) \text{ between control and ALS. Each data point represents the mean synapse count per } \text{mm}^3 \text{ of motor cortex, for each individual (control } n = 7; \text{ ALS } n = 25). \text{ d When split by cognitive status, there is no difference in synapse density within the motor cortex (One-Way ANOVA; } F = 0.06, \text{ p = 0.94). e Histogram showing a significant decrease in the thickness of ALS motor cortex compared to controls (2-tailed unpaired } t \text{ test; } p = 0.0002). \text{ f Both ALS and ALSci motor cortices are significantly thinner than control (One-Way ANOVA; } F = 9.98, \text{ p = 0.0006; Tukey post hoc test; } p < 0.05 \text{ and } p < 0.01, \text{ respectively). The presence of beta-amyloid (g), pTau (h) and pTDP-43 (i) in the motor cortex, had no effect on synapse density (2-tailed unpaired } t \text{ tests; } p > 0.05). \text{ The presence of beta-amyloid (j), pTau (k) and pTDP-43 (l) in the motor cortex, had no effect on cognitive score (2-tailed unpaired } t \text{ tests; } p > 0.05). \text{ Red icons = ALSci. All histogram bars plot the mean ± SD.}
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us. One of the limitations of this study is the small number of cognitively impaired patients who have donated brain tissue for these synaptic analyses. As more people donate tissue, we may be able to further dissect the contributions of pathological protein accumulation within synapses and genetic contributors to synaptic loss and cognitive decline. Another potential confound in the study is that the control group prepared for array tomography imaging was significantly older than the ALS group (Table 1). However, since subtle age-related changes in synapse loss have been reported [36], we predict that this age discrepancy would make the detection of synapse loss in ALS more difficult and thus are confident that this age difference did not lessen the main finding of synapse loss in frontal cortex of ALS cases, which is worse with cognitive impairment. In support of this idea, we did not detect any correlation between age and synapse density in our data (Pearson $r = 0.23$, $p = 0.23$).

In summary, we have combined two high-resolution imaging techniques to reveal synapse loss in the frontal cortex of ALS patients. We show that ALS patients with cognitive impairment have lower synaptic densities than controls. Synapse density did not differ in the primary motor cortex between ALS and controls, most likely due to the significant cortical thinning we observed in the ALS motor cortex. These findings significantly advance our understanding of ALS-associated neuropathology and identify the underlying neurobiological substrate for cognitive decline in ALS. Further, these results suggest potential synaptic biomarkers for predicting which ALS patients will experience cognitive changes, and point to the potential of synapse-directed therapeutics to combat both the cognitive and potentially also the motor symptoms of ALS.

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