Motor cortex transcriptome reveals microglial key events in amyotrophic lateral sclerosis

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Abstract

Objective
To identify transcriptomic changes, neuropathologic correlates, and cellular subpopulations in the motor cortex of sporadic amyotrophic lateral sclerosis (ALS).

Methods
We performed massive RNA sequencing of the motor cortex of patients with ALS (n = 11) and healthy controls (HCs; n = 8) and analyzed gene expression alterations, differential isoform usage, and gene coexpression networks. Furthermore, we used cell type deconvolution algorithms with human single-nucleus RNA sequencing data as reference to identify perturbations in cell type composition associated with ALS. We performed immunohistochemical techniques to evaluate neuropathologic changes in this brain region.

Results
We report extensive RNA expression alterations at gene and isoform levels, characterized by the enrichment of neuroinflammatory and synaptic-related pathways. The assembly of gene coexpression modules confirmed the involvement of these 2 major transcriptomic changes, which also showed opposite directions related to the disease. Cell type deconvolution revealed an overrepresentation of microglial cells in ALS compared with HC. Notably, microgliosis was driven by a subcellular population presenting a gene expression signature overlapping with the recently described disease-associated microglia (DAM). Using immunohistochemistry, we further evidenced that this microglial subpopulation is overrepresented in ALS and that the density of pTDP43 aggregates negatively correlates with the proportion of microglial cells.

Conclusions
DAM has a central role in microglia-related neuroinflammatory changes in the motor cortex of patients with ALS, and these alterations are coupled with a reduced expression of postsynaptic transcripts.
Amyotrophic lateral sclerosis (ALS) is a neurodegenerative disease neuropathologically characterized by the aberrant cytoplasmic aggregation and phosphorylation of the 43-kDa transactive response DNA-binding protein (pTDP43) in the majority of ALS cases at postmortem evaluation, which is known to occur in a sequential manner starting in the motor cortex.1 Although the causes that lead to nongenetic forms of ALS (sporadic ALS) have yet to be fully determined, the presence of neuroinflammation is a consistent feature in the CNS of affected patients.3 Microglia are macrophage-like innate immune cells of the CNS that, as a result of disease conditions, change their gene expression profile to adapt and acquire a reactive state.3 Recently, a study using single-cell RNA sequencing from the CNS of Alzheimer disease and ALS mouse models reported that a specific microglial subpopulation, known as disease-associated microglia (DAM), was the cell type responsible for microgliosis and that triggering receptor expressed on myeloid cells 2 (TREM2) is a major driver of DAM activation.4 In support of this discovery, some chitinase proteins, which are known markers of microglial activation,5,6 have been consistently shown to be increased in the CSF of patients with ALS.7,8

The pivotal role of TDP43 and other genes known to cause ALS (such as FUS, hnRNPA2B1, hnRNPA1, TAF15 or TIA1) in the metabolism of RNA has implicated RNA dyshomeostasis as a crucial event in the pathophysiology of the disease.9 An unbiased resource to obtain a comprehensive signature of gene and isoform expression changes associated with the whole tissue response to disease conditions is total RNA sequencing (RNAseq) in bulk tissue. Furthermore, the advent of novel sequencing tools such as single-nucleus RNAseq (snRNAseq) has opened a new window to better explore the transcriptome at a cell level and has unraveled a complex and huge variety of human cell types with unique expression profiles in the human brain.10-12 These transcriptomic signatures can be applied to perform cell type deconvolution of bulk RNAseq data using novel deconvolution methods. These methods take into account cross-subject and cross-cell variability of gene expression profiles, without relying on preselected markers, thereby providing more realistic cell type proportion estimations and yielding crucial information about the cellular heterogeneity that is associated with a pathologic condition.13

The brain motor cortex is affected at the most early stages of the disease and is one of the most vulnerable regions in ALS. However, studies of this critical region both at the transcriptional and immunohistochemical level are lacking. In the present work, we aim to characterize the motor cortex of sporadic ALS cases through total RNAseq analyses. We also use cell type deconvolution using human snRNAseq data as reference and immunohistochemical analyses to resolve a signature of neuropathologic changes associated with ALS in this particular brain region.

Methods

Human samples

The study included human motor cortex (Brodmann area 4) samples provided by the Neurological Tissue Bank (NTB) of the Biobanc-Hospital Clinic-IDIBAPS. None of the brain tissues presented any infarcts in the motor cortex. Diagnosis of ALS complied with the El Escorial criteria during life,14 and none of the patients had a family history of ALS or dementia, nor did show any sign of cognitive impairment. All patients presented pTDP43 inclusions in the motor cortex at postmortem examination. None of the samples carried the C9orf72 hexanucleotide repeat expansion or mutations in the TBKI gene, the most common genetic causes related to adult-onset ALS in Spain.15,16

RNA extraction and sequencing

Using a mortar and liquid nitrogen, the tissue (60 mg) was grinded to powder and transferred to a solution of 600 μL of TRIzol reagent (Thermo Fisher Scientific, Waltham, MA). We used standard recommendations and procedures to extract RNA with TRIzol, the RNeasy Mini Kit (Qiagen, Hilden, Germany), and the Rnase-Free Dnase Set (Qiagen). Qubit was used to measure RNA concentration, whereas RNA integrity (RIN) was verified on an Agilent 2100 Bioanalyzer (Agilent Technologies, Santa Clara, CA). Only samples with a quantity threshold of 5 μg and RIN ≥6.5 were used for total RNAseq. The final study group included 11 ALS cases and 8 healthy controls (HCs) (table 1). Paired-end sequencing libraries were prepared using the TruSeq Stranded Total RNA Library Preparation kit (Illumina, San Diego, CA) and sequenced on the Illumina HiSeq 2500 platform by the Centro Nacional de Análisis Genómicos (CNAG; Barcelona, Spain) with 101-bp paired-end reads to achieve at least 100 million reads.
paired-end reads for each sample (ranging from 103,447.000 to 138,990.000 paired-end reads).

Data processing
We aligned FASTQ files to the Grch38.p12 genome assembly using STAR v2.6.1a.17 We then used GATK Best Practices18 for variant calling and FeatureCounts (within the Subread v.1.6.2 package19) to assign fragments to each gene feature included in the Grch38v94 gene transfer format file. For differential isoform usage, we used the STAR v.2.6.1 package in quantMode TranscriptomeSAM mode and used Salmon to obtain isoform expression values.20 For differential gene expression and differential isoform usage analyses, we used DESeq2 v1.24,21 which applies the Benjamini-Hochberg for p value adjustment. We decided to use RPL13 as a housekeeping gene as it showed the smallest coefficient of variation (<3.5%) compared with the 2 other putative housekeeping genes that we assessed (RPL0 and GAPDH).

Table 1 Demographic, clinical, and quality control data

| Sample ID | Group | Age at death, y | Sex | Age at onset, y | Duration, mo | Site of onset | RIN | PMI, h |
|-----------|-------|----------------|-----|----------------|-------------|---------------|-----|-------|
| AF5214    | ALS   | 64             | Male | 62             | 33          | Limb          | 8.2 | 7     |
| AF5215    | ALS   | 50             | Male | 46             | 58          | Limb          | 8.8 | 13    |
| AF5216    | ALS   | 60             | Female | 54            | 72          | Limb          | 8.3 | 6     |
| AF5218    | ALS   | 53             | Female | 50            | 32          | Limb          | 8.4 | 9     |
| AF5220    | ALS   | 82             | Male | 81             | 1           | Bulbar        | 7.7 | 16    |
| AF5222    | ALS   | 54             | Female | 53            | 18          | Limb          | 9   | 7     |
| AF5224    | ALS   | 70             | Female | 68            | 24          | Limb          | 8.4 | 14    |
| AF5227    | ALS   | 78             | Male | 76             | 24          | Limb          | 8.1 | 12    |
| AF5228    | ALS   | 77             | Female | 75            | 31          | Bulbar        | 7   | 16    |
| AF5229    | ALS   | 57             | Female | 54            | 24          | Limb          | 6.8 | 5     |
| AF5232    | ALS   | 52             | Male | 51             | 12          | Limb          | 7.4 | 18    |
| AF5219    | HC    | 64             | Male | —              | —           | —             | 8.2 | 10    |
| AF5221    | HC    | 78             | Male | —              | —           | —             | 7.9 | 6     |
| AF5226    | HC    | 81             | Female | —            | —           | —             | 7.5 | 23    |
| AF5231    | HC    | 83             | Female | —            | —           | —             | 7.3 | 7     |
| AF5234    | HC    | 58             | Male | —              | —           | —             | 7.5 | 4     |
| AF5235    | HC    | 76             | Male | —              | —           | —             | 7.5 | 11    |
| AF5236    | HC    | 83             | Female | —            | —           | —             | 7.7 | 7     |
| AF5237    | HC    | 68             | Female | —            | —           | —             | 6.5 | 13    |

Abbreviations: ALS = amyotrophic lateral sclerosis; HC = healthy control; PMI = postmortem interval; RIN = RNA integrity number.

Validation of transcriptome changes by quantitative real-time PCR
We performed quantitative real-time PCR (qPCR) on the same brain-derived RNA samples used for total RNAseq (11 sALS cases and 8 controls). A total of 500 ng of RNA was used to generate cDNA with the RevertAid First Strand cDNA Synthesis Kit (Thermo Fisher Scientific), as per the manufacturer’s instructions. All qPCRs were conducted using Fast SYBR Green Master Mix (Thermo Fisher Scientific) and run on an ABI Prism 7900HT Fast Real-Time PCR System (Applied Biosystems, Foster City, CA). All primer pairs used are listed in table e-1, links.lww.com/NXI/A280. For relative quantification, we applied the DD_Ct method, using RPL13 as an endogenous control to normalize the data. We decided to use RPL13 as a housekeeping gene as it showed the smallest coefficient of variation (<3.5%) compared with the 2 other putative housekeeping genes that we assessed (RPL0 and GAPDH).

Cell type deconvolution
We used the recently developed Multi-Subject Single Cell deconvolution (MuSiC)13 method to perform cell type deconvolution of our bulk RNAseq data. As a reference, we
used the most comprehensive human snRNAseq data set available. This data set includes the RNAseq from 70,634 cells in the frontal cortex (Brodmann area 10) of 24 Alzheimer disease cases and 24 HCs from the Religious Orders Study and Memory and Aging Project (ROSPMAP),12 grouped into 8 major cellular populations (excitatory neurons, inhibitory neurons, microglia, astrocytes, oligodendrocytes, oligoden-
drocyte precursor cells, endothelial cells, and pericytes). To check consistency and validate our results, we used 2 in-
dependent snRNAseq repositories as a reference to estimate major cell type proportions. To this aim, we downloaded the medial temporal gyrus snRNAseq data derived from the Allen Brain Atlas, which includes 15,928 cells from 8 human tissue donors. We selected this data set as it contains double the number of nuclei than the primary visual cortex and the an-
terior cingulate cortex, also contained in the database.10 The third data set comprised 10,319 cells derived from the frontal cortex of 4 individuals.11 Cell type deconvolution of cellular subpopulations was performed using the ROSMAP data as reference.

Weighted gene coexpression network analyses
We accounted for genes with more than 10 counts across all individuals. DESeq2 was then used to normalize the input data to construct coexpression networks using the weighted gene coexpression network analyses (WGCNA) package.24 We then constructed the signed weighted correlation network, which takes into account both negative and positive correlations, using the manual function in WGCNA, applying the biweight midcorrelation, selecting a power of 7 (the lowest possible power term where topology fits a scale free network), and run in a single block analysis. We defined all modules by the hybrid treecutting option with deepsplit pa-
rarameter = 2, applying a minimal module size of 30 genes and merging modules with a cutHeight ≤0.25. Module eigengenes (first principal component as a summary of each module) summarized the modules and correlations performed using spearman correlation for continuous traits and Pearson cor-
relation for binary traits (bicor and corValueStudent, in-
spearman correlation for continuous traits and Pearson cor-
relations, using the manual function in WGCNA, applying

Quantification analysis
We obtained full-section immunohistochemistry (IHC) images with Pannoramic MIDI II (3DHistech). Blinded to the neuropathologic diagnoses, we delimited 6 gray matter areas of the motor cortex from each case, and for each gray matter area, we randomly generated 6 regions of interest (ROIs) with the same size using the multicrop.m script. We used the Iba1.m script to quantify densities for Iba1 staining in each ROI, which represents a total of 36 images per in-
dividual. This algorithm binarizes the random images and compute densities of protein expression to quantify the number of immunoreactive objects. Both in-house algo-
rithms can be freely accessed at github.com/Memo-
ryUnitSantPau. We used MATLAB R2017b software to develop the algorithm (The MathWorks Inc., Natick, MA). For pTDP43 quantification, 2 different researchers blinded to neuropathologic diagnoses quantified immunoreactive positive objects manually. We evaluated immunofluores-
cence using FIJI imaging software. We estimated an auto-
mated threshold to create binary images. Following the noise removal with the despeckle and the remove outliers’ filters, we determined the number of Iba1+ microglial cells expressing MHC class II markers using the analyze particles function.
Statistical analysis
We used the Shapiro-Wilk test to test for deviation from a standard distribution. For correlation analyses, we determined the Pearson or Spearman correlation coefficients using the gssscatter function within the ggpubr package in R. We performed mean comparison analyses through the Student t test or Student test with Welch correction (using t.test function in R), depending on data distribution. Statistical significance for all tests was set at 5% (α = 0.05), and all statistical tests were 2 sided. We used R version 3.6.2 to perform all statistical analyses.

Standard protocol approvals, registrations, and patient consents
The Sant Pau Hospital Ethics Committee approved the study with all experiments with human tissue performed in accordance with the Declaration of Helsinki. The NTB of the Biobanc-Hospital Clinic-IDIBAPS supplied all tissue following approval from their Scientific Advisory Committee.

Data availability statement
We have deposited all raw sequencing data (FASTQ files) at the European Genome-phenome Archive (EGA), which is hosted by the EBI and the CRG under accession number EGAS00001004286.

Results
Demographics, clinical features, and sample composition
We performed RNAseq analysis of postmortem motor cortex samples from 11 ALS cases (6 females) and 8 individuals without neurologic disease (4 females). Postmortem interval and RIN did not differ between groups (p = 0.68 and p = 0.089, respectively). The mean age at death was lower for ALS cases compared with HC (63.4 years; range: 54–82 years in ALS and 73.9 years; range: 58–83 years in HC, p = 0.04). Patients had an average disease onset of 60.9 years (range: 46–81 years) and a disease duration of 29.9 months (range: 1–72 months). Nine of the 11 cases (81.8%) presented a limb disease onset (table 1). We used RNAseq data to rule out the presence of other ALS-disease causing mutations, confirming the presumably sporadic nature of our ALS group of cases.

Gene expression alterations and differential isoform usage
Differential gene expression between patients with ALS and HCs disclosed a total of 108 upregulated and 16 downregulated genes (adjusted p < 0.05) (figure 1A and table e-2, links.lww.com/NXI/A280). We validated 9 of the 10 genes selected by qPCR, as we did not confirm the CHI3L2 gene overexpression in patients with ALS (p = 0.085) (figure 1B). Correlation analyses between counts derived from total RNAseq data and qPCR expression values indicated a high and significant concordance between both expression measures in these 10 genes (table e-3, links.lww.com/NXI/A280). GO and KEGG pathway analyses revealed an enriched involvement of the inflammatory response in ALS (figure 1C). The assessment of changes at the isoform expression level resulted in 167 upregulated and 40 downregulated isoforms, of which 181 were from unique genes (table e-4, links.lww.com/NXI/A280). Enrichment analyses revealed the involvement of postsynaptic density and immune response as dominant pathways (figure e-1, links.lww.com/NXI/A279). To further characterize and confirm the synaptic involvement, we used the SynGO database, which provides an expert-curated resource for synaptic function and gene enrichment analysis.23 This analysis confirmed the enrichment of postsynaptic components in our list of differentially expressed isoforms (table e-5, links.lww.com/NXI/A280). Among the list of altered genes identified by the differential isoform expression and gene expression models, 32 of them overlapped between both approaches (table e-6, links.lww.com/NXI/A280). The involvement of synaptic-related changes was confirmed from the differential isoform usage analysis (figure e-2, links.lww.com/NXI/A279), whereas the enrichment of inflammatory markers was evidenced in the gene model approach (figure e-3, links.lww.com/NXI/A279).

Cell type deconvolution
To characterize the variability of cell types and compare cellular proportions between groups, we applied the recently developed MuSiC algorithm. As reference, we used data from the first and most comprehensive human snRNAseq study performed in brain tissue of individuals with a neurodegenerative disease, which includes 24 patients with Alzheimer disease and 24 HCs.12 Cell type deconvolution revealed an overrepresentation of microglial cells in the ALS motor cortex compared with HC (p = 0.025, fold change = 1.65) and a reduction in excitatory neurons (p = 0.101, fold change = -0.85) (figure 2). We confirmed microglial upregulation using 2 additional independent reference data sets.10,11 In both cases, we found the same pattern of downregulation of excitatory neurons (figures e-4 and e-5, links.lww.com/NXI/A279). To further gain insight into which microglial subpopulation might be driving this effect, we estimated the proportion of each subpopulation identified in the ROSMAP study. Our results showed that a specific microglial subpopulation (Mic1, as named in the study performed by Mathys and collaborators and recognized as the human DAM12) is disproportionately presented in the ALS motor cortex (p = 6.6 × 10⁻³, fold change = 4.33) (figure 2). Among the 77 marker genes, which characterize the Mic1 subcluster as DAM (Mic1 unique genes or those overlapping with DAM),12 22 of them (28.6%) are genes differentially expressed in our bulk RNAseq analysis (unadjusted p < 0.05; (table e-7, links.lww.com/NXI/A280). Of interest, the proportion of Mic1 cells showed a positive correlation with the expression of TREM2 (p = 1.6 × 10⁻³; R = 0.67; figure e-6, links.lww.com/NXI/A279), a receptor required for the transition from the homeostatic microglia to DAM.
Gene coexpression network analyses

To identify gene clusters with varying coexpression patterns that could behave differently between patients and healthy individuals, and elucidate possible biological mechanisms driving pathologic processes, we performed a gene coexpression network analyses using the WGCNA package. The analysis disclosed a total of 8 modules. Among them, 3 modules associated with disease status (MEblack, \( p = 0.003; R = 0.64 \); MEyellow, \( p = 0.025; R = 0.57 \); and MEpink, \( p = 0.011; R = -0.51 \); figure 3). Moreover, MEblack and MEyellow were enriched for inflammatory responses, whereas MEpink had an overrepresentation of genes involved in synaptic and neuronal functions (table e-8, links.lww.com/NXI/A280). The SynGO database confirmed the postsynaptic signature of this enrichment in MEpink (table e-9, links.lww.com/NXI/A280). Of interest, the 2 inflammatory gene coexpression modules (MEblack and MEyellow) showed a strong negative correlation with the postsynaptic module (MEpink) (\( p = 8 \times 10^{-3}; R = -0.78 \) and \( p = 3 \times 10^{-4}; R = -0.74 \), respectively) (figure 3), suggesting that both inflammation and synaptic alterations are interconnected in ALS pathologic processes. Furthermore, the proportion of Mic1 cells showed a strong direct correlation with the 2 inflammatory gene coexpression modules associated with ALS (MEblack; \( p = 2.7 \times 10^{-8}; R = 0.92 \); and MEyellow; \( p = 1.4 \times 10^{-5}; R = 0.68 \)) and a negative correlation with the synaptic module (MEpink; \( p = 2.4 \times 10^{-4}; R = -0.75 \)) (figure 3). Overall, our results from differential gene and isoform expression, gene coexpression modules, and cell type deconvolution comparisons strongly indicate that microglia-related inflammatory changes, mainly driven by the Mic1 subpopulation (also known as DAM), are central in ALS pathophysiology and that these inflammatory processes are closely related to the synaptic disturbances that are present in the motor cortex.

Immunohistochemistry

To further investigate the inflammatory response in postmortem brain tissue, we performed IHC analyses using Iba1, a protein whose expression is primarily restricted to homeostatic
microglia, to examine the broad population of microglial cells, and MHC class II, a marker responsible for antigen recognition and the activation of the adaptive immune system, which is expressed in the Mic1 cell subpopulation. Positive immunostaining was increased in the ALS motor cortex for Iba1; however, differences between patients and controls did not reach statistical significance (p = 0.106; figure e-7, links.lww.com/NXI/A279). Notwithstanding, co-immunofluorescence of Iba1 and MHC class II markers revealed an increase in the number of microglial cells expressing MHCII markers in ALS compared with HC (70.1% vs 10%, respectively; p = 0.021) (figure 4 and figure e-8, links.lww.com/NXI/A279), strengthening our results and suggesting that Mic1 is driving neuroinflammation in the ALS motor cortex.

We finally assessed the frequency of pTDP43 immunoreactive structures in the motor cortex. As expected, patients with ALS showed a higher density of pTDP43 inclusions compared with HCs (p = 0.017) (figure e-9A, links.lww.com/NXI/A279). Of interest, a high degree of variability on the number of pTDP43 aggregates was noted in ALS and was not explained by age at death, age at onset, or disease duration. The burden of pTDP43 inclusions inversely correlated with the proportion of microglial cells in our group of patients with ALS (p = 0.0026; R = −0.81) (figure e-9B, links.lww.com/NXI/A279), and the same pattern was found with the microglial subpopulation Mic1 (p = 0.019; R = −0.69).

**Discussion**

The motor cortex is one of the major vulnerable and early affected regions in ALS and represents a target region to disentangle key pathologic processes in this neurodegenerative disorder. To date, few studies have performed a whole transcriptomic assessment of the CNS in ALS. Through
Figure 3 WGCNA visualization and correlation map constructed with WGCNA module eigengenes and cell type proportions

Network diagrams showing the 50 most connected genes for each of the 3 modules associated with ALS (MEblack, MEyellow, and MEpink) are depicted on the top of the figure. Node darkness and size are proportional to the number of connections within the module. Correlation matrix showing the correlation coefficients between cell type proportions is depicted for the 6 major cell types and the Mic1 (DAM) cell subtype. The 3 significant modules (MEblack, MEyellow, and MEpink), each of one sharing unique groups of coexpressed genes that are differentially expressed between patients with ALS and controls, are also included in the correlogram. Only significant correlations are depicted with a colored circle (blue for direct and red for inverse correlations). The size of the circle is proportional to the correlation significance. The plot indicates the high correlation between the 3 modules and how these modules are related to the proportion of specific cell types. ALS = amyotrophic lateral sclerosis; DAM = disease-associated microglia; HC = healthy control; WGGNA = weighted gene coexpression correlation network analyses.

| Mic1 subtype | MEblack (inflammatory) | MEyellow (inflammatory) | MEpink (postsynaptic) |
|--------------|------------------------|-------------------------|-----------------------|
| MEblack      | -0.78                  | -0.74                   | -0.79                 |
| MEyellow     | -0.74                  | -0.79                   | -0.75                 |
| MEpink       | -0.92                  | 0.68                    |                       |
| Excitatory neutrons | 0.52               | -0.42                  | -0.48                 |
| Oligodendrocytes | -0.31              | 0.01                    | -0.10                 |
| Inhibitory neurons | -0.02               | -0.06                  | 0.22                  |
| Microglia    | -0.57                  | 0.89                    | 0.72                  |
| Astrocytes   | 0.29                   | 0.05                    | 0.25                  |
| Endothelial  | -0.57                  | 0.49                    | 0.71                  |
| Pericytes    | -0.18                  | 0.04                    | 0.33                  |
an unbiased transcriptomic analysis using high-throughput RNA sequencing, we have elucidated gene and isoform expression alterations, gene coexpression networks, and cell type proportions associated with ALS. For the first time in the ALS field, we have performed cell type deconvolution using human brain single-nucleus RNA sequencing data from 3 independent sources as reference, thus providing highly reliable results that have been reinforced through our immunohistochemical analyses.

We report a list of 124 genes differentially expressed in the ALS motor cortex, which reflect the RNA expression changes in this brain region. Among them, chitinase-related genes CHI3L1 and CHI3L2 presented 2 of the most prominent shifts in gene expression, whereas CHIT1 showed a nominal significance. Of interest, these chitinases are neuroinflammatory biomarkers, which have been shown to be consistently increased in the CSF of patients with ALS compared with neurologically healthy individuals.7,8 These results indicate that among the unbiased signature of RNA alterations provided herein, some of them might lead to the discovery of novel promising biomarkers.

An exacerbated innate immune response with microgliosis has been recently described in the ALS motor cortex.28 A recent study that has investigated the whole transcriptome of the ALS motor cortex suggested that among ALS cases, a subgroup of them is characterized by a molecular signature related to glial activation and inflammation.27 Our results clearly reinforce the idea of these inflammatory-related changes in the human motor cortex and emphasize the role of DAM as a key factor in this process. First, differential gene and isoform expression data point toward an inflammatory response as a major event that is strongly intensified in the ALS motor cortex. Second, the assembly of weighted gene coexpression networks resulted in 2 significant modules associated with ALS (MEblack and MEyellow), both highly enriched with genes related to inflammatory functions. Third, cell type deconvolution of the bulk RNAseq data demonstrated an increased proportion of microglial cells compared with the motor cortex of healthy cases. In this context, studies in mice have recently evidenced DAM as the microglial subpopulation with a more prominent role in ALS.4 Our results show that Mic1, the human microglial subpopulation that harbors the majority of markers found in the previously described DAM transcriptomic signature,12 is the main microglial subpopulation that drives microgliosis in ALS. In fact, almost a third of Mic1 (DAM) marker genes are deregulated in the motor cortex of our group of patients with ALS. Of note, Mic1 proportion highly correlated with the expression of TREM2, which is required to enhance the proinflammatory stage of DAM.4

Our immunohistochemical analyses did not show a relevant increased density of Iba1 (a widely used marker of microglial processes) in ALS-related brain tissue, nor was the expression...
of its coding gene (AIF1) altered in our RNASeq data set. Previous studies have provided contradictory results when using this marker to assess microgliosis. This somewhat unexpected result could be explained by the fact that some marker genes might downregulate on microglial activation. Also, whereas cell type deconvolution is performed using a complete catalogue of gene expression profiles obtained from brain-derived snRNAseq, immunohistochemistry only uses a single marker and does not reflect the complexity and heterogeneity of this cell type, making it an underpowered method to detect differences in the proportion of cell subpopulations. That being said, we did validate the increased proportion of Mic1 cells in the ALS motor cortex through co-immunofluorescence of 2 markers (Iba1 and MHC class II) previously shown to characterize this microglial subpopulation. This finding further establishes Mic1 (the human disease-associated microglia) as the microglial subpopulation that drives microgliosis in the ALS motor cortex.

We observed a high degree of variability in the density of pTDP43 inclusions across patients, which could not be explained by any of the demographic or clinical features available in this study. Notably, we found a striking inverse correlation between the proportion of microglial cells and the amount of pTDP43 aggregates. These results are in line with recent in vivo and in vitro studies, suggesting that although microgliosis arises as a phagocytic response to pTDP43 aggregates, at some point, these cells lose their ability to clear these neuropathologic insults and are downregulated. Whether biofluid levels of microglial markers, such as TREM2, could be used as a proxy of pTDP43 density in the motor cortex is an avenue worth pursuing and would be a valuable addition to the biomarker arsenal for use in designing clinical trials and assessing therapeutic efficacy.

Synaptic dysfunction is an early pathogenic event in ALS. Our data point toward an underrepresentation of post-synaptic markers and a decrease of excitatory neurons in patients with ALS. Together, these results are consistent with upper motor neuron degeneration occurring in this brain region. A limitation of our approach is the lack of an available motor cortex snRNAseq data set, precluding any firm and more detailed conclusion related to the alteration of Betz cells, a unique class of motor neurons expressed in this specific brain area. Recent studies in other neurodegenerative diseases have suggested that microglia are a key and early mediator of synapse loss through phagocytosis induced by the complement cascade. Our gene expression data show an increased expression of some key complement cascade-related genes and the most significant enriched pathway corresponds to the complement and coagulation cascades, reinforcing this hypothesis. Furthermore, our results indicate that the 2 neuroinflammatory-related modules of gene expression (MEblack and MEyellow) and the proportion of Mic1 cells inversely correlate with the presence of synaptic markers.

Overall, our study strongly suggests that DAM plays a key role in driving neuroinflammatory changes and synapse loss in the ALS motor cortex. The identification of specific microglial populations with well-defined transcriptional signatures will contribute to disentangle new mechanisms and novel therapeutic targets to fight against this devastating disorder.

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**Disclosure**

The authors report no disclosures relevant to the manuscript. Go to Neurology.org/NN for full disclosures.

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