RESEARCH ARTICL E

Functional alterations in large-scale resting-state networks of amyotrophic lateral sclerosis: A multi-site study across Canada and the United States

Komal Bharti 1, Simon J. Graham 2, Michael Benatar 3, Hannah Briemberg 4, Sneha Chenji 5, Nicolas Dupré 6, Annie Dionne 6, Richard Frayne 5, Angela Genge 7, Lawrence Korngut 5, Collin Luk 1, Lorne Zinman 2, Sanjay Kalra * , for the Canadian ALS Neuroimaging Consortium (CALSNIC) ¶

1 University of Alberta, Edmonton, Alberta, Canada, 2 University of Toronto, Toronto, Ontario, Canada, 3 University of Miami, Miami, Florida, United States of America, 4 University of British Columbia, Vancouver, British Columbia, Canada, 5 University of Calgary, Calgary, Alberta, Canada, 6 Université Laval, Quebec, Quebec, Canada, 7 McGill University, Montreal, Quebec, Canada

¶ Membership of the Canadian ALS Neuroimaging Consortium (CALSNIC) Group is listed in the Acknowledgments.

* kalra@ualberta.ca

Abstract

Amyotrophic lateral sclerosis (ALS) is a multisystem neurodegenerative disorder characterized by progressive degeneration of upper motor neurons and lower motor neurons, and frontotemporal regions resulting in impaired bulbar, limb, and cognitive function. Magnetic resonance imaging studies have reported cortical and subcortical brain involvement in the pathophysiology of ALS. The present study investigates the functional integrity of resting-state networks (RSNs) and their importance in ALS. Intra- and inter-network resting-state functional connectivity (Rs-FC) was examined using an independent component analysis approach in a large multi-center cohort. A total of 235 subjects (120 ALS patients; 115 healthy controls (HC) were recruited across North America through the Canadian ALS Neuroimaging Consortium (CALSNIC). Intra- and inter-network Rs-FC was evaluated by the FSL-MELODIC and FSLNets software packages. As compared to HC, ALS patients displayed higher intra-network Rs-FC in the sensorimotor, default mode, right and left fronto-parietal, and orbitofrontal RSNs, and in previously undescribed networks including auditory, dorsal attention, basal ganglia, medial temporal, ventral streams, and cerebellum which negatively correlated with disease severity. Furthermore, ALS patients displayed higher inter-network Rs-FC between the orbitofrontal and basal ganglia RSNs which negatively correlated with cognitive impairment. In summary, in ALS there is an increase in intra- and inter-network functional connectivity of RSNs underpinning both motor and cognitive impairment. Moreover, the large multi-center CALSNIC dataset permitted the exploration of RSNs in unprecedented detail, revealing previously undescribed network involvement in ALS.
Introduction

Amyotrophic lateral sclerosis (ALS) is a neurodegenerative disorder characterized by progressive degeneration of both upper motor neurons (UMN) and lower motor neurons [1–3]. ALS affects a wide range of neuronal networks engaged in motor regulation [3]. Extra-motor brain regions are also directly compromised in ALS, and may result in abnormal cognitive and behavioural function [4–6]. Changes in brain function can be investigated in detail using resting-state functional magnetic resonance imaging (Rs-FMRI) [7–9]. Rs-FMRI is a commonly used imaging technique to capture synchronous low-frequency fluctuations in blood oxygen level-dependent (BOLD) signals between functionally connected brain regions [10] in the absence of stimulus-driven tasks [11], and thus can help detect abnormal functioning in pathological conditions where task-based imaging is impractical or logistically challenging.

Previous Rs-FMRI studies have demonstrated the role and importance of several seed-based correlation maps i.e. RSNs in ALS pathophysiology [7, 9, 12–21]. Some of these studies have used an a-priori approach to examine functional alterations in the RSNs associated with the cortical and subcortical brain areas such as the motor cortex, frontal cortex, cerebellum and basal ganglia corresponding to the motor as well as cognitive decline in ALS [7–9, 12, 14, 22, 23]. Alternatively, some Rs-FMRI studies have used a data-driven, independent component analysis (ICA) approach which elucidates whole-brain RSNs without predefined seed regions [13, 15, 17–19, 21]. In ALS, such studies have mainly reported sensorimotor RSN, default mode RSN, and fronto-parietal RSN [13, 16, 18, 21] suggesting the reorganization of motor and extra-motor domains of functional brain networks in of ALS patients. The technique avoids the probability of missing connectivity differences due to seed selection [24–27]. In addition, ICA helps to examine multiple RSNs and thus provides more global information [28, 29] and the potential to probe underlying mechanisms. Keeping in light both approaches, a widespread involvement of not only motor areas, but also extra-motor brain areas actively involved in the underlying pathophysiology of ALS were revealed suggesting not just functional impairment of one or more RSNs, but rather a result of failed communication or interplay of multiple RSNs across the whole brain.

Considering the fact that abnormal whole brain resting state functional connectivity (Rs-FC) plays an important role in the pathophysiology of ALS, we sought to examine whole brain within-network and between-network Rs-FC in a large cohort of ALS patients from multiple centres in Canada and the United States. In addition, we also wished to explore the association of intra-network and inter-network Rs-FC with clinical features relevant to motor and extra-motor dysfunction in ALS patients.

Methods

Canadian ALS Neuroimaging Consortium

Data were acquired for the study across North America through the Canadian ALS Neuroimaging Consortium (CALSNIC) [30]. The main objective of CALSNIC is to develop novel MRI biomarkers for ALS. There are 2 projects in CALSNIC: CALSNIC 1 and CALSNIC 2. The 5 sites of data acquisition for CALSNIC 1 were: University of Alberta, Edmonton; University of Calgary, Calgary; McGill University, Montreal; University of Toronto, Toronto; and University of British Columbia, Vancouver. For CALSNIC 2, data were acquired from the aforementioned sites except Vancouver; and additionally, from University of Miami, Miami, and University of Laval, Québec.
Participants
A total cohort of 235 participants (115 HC; 120 ALS patients) were recruited through CALS-NIC. Patients were diagnosed as possible (25), probable (60), or definite ALS (35) according to revised El Escorial Criteria [31]. Out of 120 ALS patients, 20 patients had bulbar onset and 100 patients had limb onset. The exclusion criteria included the presence of other neurological and psychiatric illness. Excluded were patients with primary lateral sclerosis, progressive muscular atrophy, and frontotemporal dementia. For the control group, 115 age and sex matched HC were recruited with no history of neurological and psychiatric illness. The study was conducted with the approval of the Health Research Ethics Board of each participating site and informed written consent was obtained from the participants.

Clinical assessment
All patients underwent a neurological exam administered by a trained neurologist at each participating site. The ALS functional rating scale revised (ALSFRS-R) was used as a measure of disease severity [32]. Decreased ALSFRS-R corresponds to greater clinical disability. A disease progress rate was subsequently calculated using the ALSFRS-R scores and symptom duration [33]. Severity of upper motor neuron (UMN) degeneration in the limbs and bulbar regions was inferred for patients by calculating an UMN burden score derived from the neurological exam. A UMN burden score was calculated with a maximum score of 12. The presence of spasticity and hyperreflexia in the upper and lower limbs, Babinski sign, and clonus at each ankle were tabulated with a maximum possible score of 6 from each side of the body [34]. Further assessment of motor impairment was measured using finger tapping and foot tapping tasks for the right side and the left side. The Edinburgh Cognitive and Behavioral ALS screen (ECAS) was used to test cognitive performance [35]. In addition to the ECAS “Total” score, the "ALS Specific” and "ALS Non-specific” were also noted. The ECAS and tapping tasks were also performed by each participant in the healthy control group.

Imaging protocol
Structural and functional images were acquired using 3 Tesla MRI systems at all sites. Prior to Rs-FMRI, participants were instructed to remain still, awake, and with their eyes closed, in a fully relaxed condition. The details of data acquisition across the different sites are summarized in Table 1 [30].

Data analysis
Structural and functional data analysis were performed using the FSL (Functional Magnetic Resonance Imaging of the Brain [FMRIB] Software library packages) (http://www.fsl.fmrib.ox.uk/fsl/fslwiki) [36].

Preprocessing steps included data analysis of all the subjects using FSL fMRI expert analysis tool (FEAT, http://www.fsl.fmrib.ox.uk/fsl/fslwiki). Brain extraction was performed on the three dimensional (3D) T1-weighted images using the brain extraction tool (BET) [37]. The Rs-FMRI data were corrected for head motion using the MCFLIRT tool, as well as slice timing correction and spatial smoothing using a Gaussian kernel with a full width at half maximum of 5 mm [38]. The data was then subjected to single-session independent component analysis-based automatic removal of motion artifacts (ICA_AROMA) [39] in order to identify independent components (ICs) representing motion-related artifacts. The FMRIB linear image registration tool (FLIRT) (http://www.fmrib.ox.ac.uk/fsl) was used to register the clean fMRI data from motion related IC’s of each participant with the processed BET images. The output of
Table 1. Details of MRI data acquisition across research centres in Canada and United States.

| Project          | CALSNIC-1 |                     |                     | CALSNIC-2 |                     |                     |                     |
|------------------|-----------|---------------------|---------------------|-----------|---------------------|---------------------|---------------------|
| Site             | University of Alberta | University of British Columbia | University of Calgary | McGill University | University of Toronto | University of Calgary | McGill University |
| Scanner model    | Siemens 3T Prisma      | Philips 3T Intera    | GE 3T Discovery MR 7.50 | Siemens 3T Triotrim | Siemens 3T Discovery MR 7.50 | Siemens 3T Prisma     | Siemens 3T Prisma  |
| Software version | syngo MR E11            | 3.2.31              | DV25.0_EB_1442.a     | syngo MR B17   | DV24.0_R01_1344.a    | syngo MR E11          | syngo MR E11      |

| 3D T<sub>1</sub>-weighted scan, axial acquisition | | | | | | | |
| Acquisition orientation | axial | axial | axial | axial | sagittal | sagittal | sagittal | sagittal |
| Repetition time, ms | 2.300 | 7.9 | 7.4 | 2.300 | 7.4 | 1.700 | 8.1 | 1.700 | 1.800 | 7.1 |
| Echo time, ms | 3.43 | 3.5 | 3.1 | 3.43 | 3.1 | 2.21 | 3.2 | 2.21 | 2.21 | 2.13 | 3.4 |
| Inversion time, ms | 900 | 950 | 400 | 900 | 400 | 880 | 400 | 880 | 880 | 900 | 950 |
| Flip angle, degrees | 9 | 8 | 11 | 11 | 10 | 16 | 10 | 10 | 10 | 10 | 10 |
| Field of view | 256 | 240 | 256 | 256 | 256 | 256 | 256 | 256 | 256 | 256 | 256 |
| Matrix dimension, pixels | 256 x 256 | 240 x 240 | 256 x 256 | 256 x 256 | 232 x 256 | 256 x 256 | 232 x 256 | 256 x 256 | 256 x 256 | 256 x 256 |
| Voxel dimension, mm | 1 x 1 x 1 | 1 x 1 x 1 | 1 x 1 x 1 | 1 x 1 x 1 | 1 x 1 x 1 | 1 x 1 x 1 | 1 x 1 x 1 | 1 x 1 x 1 | 1 x 1 x 1 | 1 x 1 x 1 |
| Slices, n | 167 | 150 | 167 | 167 | 167 | 167 | 167 | 167 | 167 | 167 |
| Acquisition times, min | 0:5:30 | 4:30 | 0:5:30 | 4:30 | 0:3:37 | 0:4:16 | 0:3:37 | 0:3:37 | 0:4:10 | 0:4:08 |

| Resting-state functional MRI | | | | | | | |
| Repetition time, ms | 2.200 | 2.200 | 2.200 | 2.200 | 2.200 | 2.200 | 2.200 | 2.200 | 2.200 | 2.200 |
| Echo time, ms | 30.0 | 30.0 | 30.0 | 30.0 | 30.0 | 30.0 | 30.0 | 30.0 | 30.0 | 30.0 |
| Flip angle, degrees | 70 | 70 | 70 | 70 | 70 | 70 | 70 | 70 | 70 | 70 |
| Field of view | 224 | 224 | 224 | 224 | 224 | 224 | 224 | 224 | 224 | 224 |
| Matrix dimension, pixels | 64 x 64 | 64 x 64 | 64 x 64 | 64 x 64 | 64 x 64 | 64 x 64 | 64 x 64 | 64 x 64 | 64 x 64 | 64 x 64 |
| Voxel dimension, mm | 3.5 x 3.5 x 3.5 | 3.5 x 3.5 x 3.5 | 3.5 x 3.5 x 3.5 | 3.5 x 3.5 x 3.5 | 3.5 x 3.5 x 3.5 | 3.5 x 3.5 x 3.5 | 3.5 x 3.5 x 3.5 | 3.5 x 3.5 x 3.5 | 3.5 x 3.5 x 3.5 | 3.5 x 3.5 x 3.5 |
| Trains, n | 192 | 192 | 192 | 192 | 192 | 250 | 250 | 250 | 250 | 250 |
| Slices, n | 40 | 40 | 40 | 40 | 40 | 40 | 40 | 40 | 40 | 40 |
| Acquisition times, min | 07:11 | –7:00 | 07:11 | –7:00 | 09:18 | 09:18 | 09:18 | 09:18 | 09:17 | 09:23 |

https://doi.org/10.1371/journal.pone.0269154.t001
linear registration was then non-linearly registered with the Montreal Neurological Institute (MNI) standard space using FMRIB’s non-linear image registration tool (FNIRT) (http://www.fmrib.ox.ac.uk/fsl). To perform higher level group comparisons using the FSL MELODIC software, a temporal concatenation of spatial ICA maps was performed using temporal high pass filter of 100 seconds [40]. Analysis included variance normalization of time courses and an automatic dimensionality estimation using the temporally concatenated ICA technique. For the post stats, ICA maps were thresholded with mean high-resolution 3D T1 weighted images. The group MELODIC output of 20 components generated automatically from 235 subjects was carefully inspected by an experienced researcher (K.B.) The 13 independent components representing the best RSNs were selected according to the previous literature [41], whereas the other 7 components indicating noise were discarded [27]. Subsequently, the 13 selected RSNs listed in the Fig 1 underwent dual regression analysis. This procedure first regressed the group ICA maps onto each participant’s four-dimensional (4D) dataset to give a set of time-courses; and secondly regressed the time-courses into the same 4D dataset to generate participant-specific spatial maps [29]. These participant-specific spatial maps were then considered for intra- and inter- network Rs-FC analysis.

Motion parameter analysis

The Rs-FC data are susceptible to head motion artifacts, potentially confounding the assessment of population differences [42, 43]. Absolute (referenced to the middle time-point) and relative (compared to previous time point) values were recorded from the BOLD time series as the root-mean-square values of three translational axis of X, Y, and Z, and rotational axis of pitch, roll, and yaw. After a careful inspection of the recorded motion correction estimates using FSL MCFLIRT tool, subjects with more than 1.5 mm maximum displacement in any of the axis or 1.5˚ of the angular rotation were excluded from the study [44]. Additionally, any participants with < - 4 minutes of uncorrupted data were excluded from the study [45]. Lastly, the mean absolute, and relative head motion displacement values within 1.5 mm were further...
subjected for group comparisons using a two-sample t test through SPSS software (IBM, Armonk, New York, USA).

**Intra-network resting state functional connectivity**

Intra-network Rs-FC differences were tested by entering participant-specific spatial maps representing the 13 RSNs into a multiple linear regression model implemented in FSL [46]. Age, sex, motion parameters, data acquisition site and MRI system type were included as covariates of no interest in the statistical model. For every participant-specific spatial map representing one of the 13 RSNs (Fig 1), the respective mask was extracted using the ‘fslmath’ command in FSL. The intra-network Rs-FC was then assessed within the mask of the individual RSN using the threshold free cluster enhancement (TFCE) technique in FSL and 5000 nonparametric random permutations [46]. Significant differences in the intra-network Rs-FC were corrected for multiple comparisons using the family wise error (FWE) approach at a significance level of $p<0.05$. The brain areas with statistically significant findings were extracted using the Harvard-Oxford atlas inbuilt in FSL [47].

**Correlation analysis**

In ALS patients, correlation analyses were performed between the intra-network Rs-FC and clinical variables: ALSFRS-R, disease progression rate, symptom duration, finger and foot tapping scores (right and left sides), UMN burden score, ECAS Total, ECAS ALS Specific and ECAS ALS Non-specific. Statistical differences were assessed within the mask of respective RSNs using the TFCE technique involving 5000 nonparametric random permutations, with age, sex, motion parameters, data acquisition site included as covariates of no interest. Moreover, MRI system type was also considered as an additional regressor in the statistical model [48, 49]. Correlations results were corrected for multiple comparisons using the FWE approach at a significance level of $p<0.05$. The brain areas with statistically significant findings were extracted using the Harvard-Oxford atlas inbuilt in FSL [47].

**Inter-network resting state functional connectivity**

Participant-specific spatial maps representing the 13 RSNs were used to investigate inter-network Rs-FC differences using the FSLNets Matlab toolbox (https://fsl.fmrib.ox.ac.uk/fsl/fslwiki/FSLNets). To obtain full and partial correlation matrices, the time series for the 13 RSNs were extracted, normalized and then subjected to correlation analysis. Between-network group comparisons of the normalized time series correlations were performed using a two-sample unpaired t-test with the same covariates of no interest as described above.

**Correlation analysis**

Potential associations between inter-network Rs-FC and clinical variables were assessed using Spearman rank-order correlation in SPSS software. The assessment was performed while regressing out the same covariates of interest and with the same threshold for statistical significance as described above.

**Results**

**Demographic and clinical scores**

Demographic and clinic details of the ALS patients and HC are shown in Table 2. No significant differences were observed in sex between ALS patients and HC ($p>0.05$). However, age differences were observed to be statistically significant between ALS patients and HC.
ALS patients also had lower ECAS scores (Total, Specific and Non-specific) and tapping task scores than the HC group (p < 0.05).

Motion parameter

The mean absolute head displacement values were 0.32 ± 0.22 mm in ALS patients and 0.28 ± 0.19 mm in HC. The mean relative head displacement values were 0.11 ± 0.08 mm in ALS patients and 0.10 ± 0.05 mm in HC. There were no significant differences in motion parameters between the ALS patients and HC (p > 0.05).

Intra-network resting state functional connectivity

Compared to HC, ALS patients displayed higher intra-network Rs-FC in 11 RSNs: sensorimotor, cerebellum, auditory, default mode, right and left fronto-parietal, orbitofrontal, dorsal attention, basal ganglia, medial temporal and ventral stream (Fig 2; Table 3) [FWE; p < 0.05].

Clinical correlations

In ALS, intra-network Rs-FC in the sensorimotor, cerebellum, auditory, default mode, right and left fronto-parietal, orbitofrontal, dorsal attention, basal ganglia, medial temporal, and ventral stream RSNs was negatively correlated with ALSFRS-R. Intra-network Rs-FC in the cerebellum, default mode, left fronto-parietal, basal ganglia, orbitofrontal, and medial temporal RSNs was also negatively correlated with ECAS Total score and its sub scores. Conversely, intra-network Rs-FC in the cerebellum RSN was positively correlated with UMN burden scores, and sensorimotor and right fronto-parietal RSNs were positively correlated with the disease progression rate and symptom duration. Lastly, intra-network Rs-FC within sensorimotor, basal ganglia, and cerebellum RSNs negatively correlated with finger and foot tapping.
scores. Each of these associations is depicted in Fig 3 and summarized in Table 4 (FWE; p < 0.05).

**Inter-network resting state functional connectivity**

Compared to HC, ALS patients displayed higher inter-network Rs-FC between the basal ganglia and orbitofrontal RSNs (Fig 4a) (corrected for multiple comparison approach [FWE; p < 0.05]).

**Clinical correlations**

Higher inter-network Rs-FC between basal ganglia and orbitofrontal RSNs was negatively correlated with the ECAS ALS Specific score (Spearman rank correlation; \(p = 0.01; r_s = -0.21\)) (Fig 4b). No other significant correlations were observed between inter-network Rs-FC and other clinical measures.

**Discussion**

The present study aimed to investigate the role and importance of Rs-FC alterations in different networks in the pathophysiology of ALS. For this purpose, intra- and inter-network Rs-FC was examined in ALS patients and HC using an ICA technique. Compared to HC, ALS patients displayed higher intra-network Rs-FC in 11 out of 13 RSNs: sensorimotor, default mode, right and left fronto-parietal, dorsal attention, orbitofrontal, cerebellum, basal ganglia, auditory, medial temporal, and ventral stream networks. Higher intra-network Rs-FC correlated with greater disability, faster progression rate, and other clinical measures of greater disease severity (UMN scores, tapping scores, and cognitive performance). Additionally, one inter-network Rs-FC alteration was detected in ALS patients: higher functional connectivity between the orbitofrontal and basal ganglia RSNs which associated with the ECAS ALS Specific score. Collectively, these functional alterations within and between multiple RSNs are useful in characterizing and further understanding the pathophysiology of ALS. Notably, the
Table 3. Brain areas exhibiting higher intra-network resting-state functional connectivity (Rs-FC) in resting-state networks (RSNs).

| Group differences | Resting-state networks                  | Brain areas                                                                 | Number of clusters | Clusters voxels | Coordinates X Y Z | Peak—z stat |
|------------------|----------------------------------------|-----------------------------------------------------------------------------|--------------------|-----------------|-------------------|-------------|
| ALS > HC         | Sensorimotor Network                   | Bilateral: Anterior and posterior cingulate gyrus, pre and post central gyrus, precuneus, supplementary motor area, superior frontal gyrus, supramarginal gyrus, superior parietal lobule | 21                 | 5588            | 29 67 60        | 4.7         |
|                  | Cerebellum Network                     | Bilateral: VI, VII a and b, VIII a and b, IX, X, crus I, vermis VIII a Right: Crus II Left: I-IV | 21                 | 1716            | 49 36 19        | 3.7         |
|                  |                                        |                                                                             | 20                 | 133             | 56 47 13        | 4.5         |
|                  | Auditory Network                       | Bilateral: Superior and middle temporal gyrus, insula, planum temporale, planum polare, parietal operculum cortex, heschl’s gyrus | 46                 | 283             | 11 44 43        | 3.4         |
|                  |                                        |                                                                             | 45                 | 276             | 69 50 45        | 3.7         |
|                  |                                        |                                                                             | 44                 | 115             | 78 39 36        | 3.9         |
|                  |                                        |                                                                             | 43                 | 104             | 20 63 39        | 3.6         |
|                  | Default Mode Network                   | Bilateral: Precuneus cortex, cuneal cortex, intracalcarine cortex, supracalcarine cortex, posterior cingulate gyrus, angular gyrus, superior and inferior lateral occipital cortex, posterior supramarginal gyrus, middle temporal gyrus | 20                 | 2316            | 40 31 39        | 4.5         |
|                  |                                        |                                                                             | 19                 | 287             | 17 34 59        | 4.6         |
|                  |                                        |                                                                             | 18                 | 231             | 63 32 58        | 3.5         |
|                  | Right Fronto-Parietal Network          | Right: Superior, middle, and inferior frontal gyrus, precentral gyrus, superior parietal lobule, supramarginal gyrus, angular gyrus, lateral occipital cortex | 38                 | 2437            | 22 78 31        | 4.0         |
|                  |                                        |                                                                             | 37                 | 393             | 22 52 58        | 3.7         |
|                  | Left Fronto-Parietal Network           | Left: Superior, middle, and inferior frontal gyrus, precentral gyrus        | 36                 | 222             | 29 58 35        | 3.9         |
|                  |                                        |                                                                             | 33                 | 726             | 60 79 48        | 4.3         |
|                  | Orbitofrontal Network                 | Bilateral: Frontal pole, frontal orbital and medial cortex, paracingulate gyrus, subcallosal cortex | 32                 | 384             | 73 65 39        | 4.0         |
|                  |                                        |                                                                             | 24                 | 3775            | 44 62 27        | 4.9         |
|                  | Dorsal Attention Network              | Bilateral: Lateral occipital cortex, angular gyrus, precuneus cortex, cuneal cortex, supracalcarine cortex, occipital pole | 17                 | 5632            | 25 26 55        | 4.7         |
|                  | Basal Ganglia Network                 | Bilateral: Thalamus, putamen, pallidium, and insular cortex, precuneus, posterior cingulate gyrus | 27                 | 1605            | 43 51 39        | 4.8         |
|                  |                                        |                                                                             | 26                 | 290             | 64 57 38        | 3.9         |
|                  | Medial Temporal Network               | Bilateral: Temporal fusiform gyrus, parahippocampal gyrus, temporal pole, inferior temporal gyrus, middle temporal gyrus, temporal occipital fusiform cortex | 18                 | 1451            | 29 71 15        | 4.0         |
|                  |                                        |                                                                             | 17                 | 845             | 65 67 16        | 4.4         |
|                  | Ventral Stream Network                | Right: Heschl’s gyrus, planum temporale, superior, middle and inferior temporal gyrus, temporooccipital part, posterior supramarginal gyrus, superior and inferior lateral occipital cortex, angular gyrus | 18                 | 1495            | 30 43 34        | 4.0         |
|                  |                                        |                                                                             | 17                 | 659             | 28 26 45        | 3.9         |

ALS > HC represents higher whole brain intra-network resting-state functional connectivity in Amyotrophic lateral sclerosis (ALS) as compared to healthy controls (HC). Results were corrected for family-wise error at p < 0.05 (FWE corrected; p < 0.05).

* Number of clusters represents the exact number of clusters in the fMRI analysis stayed significant after correcting the results for multiple comparisons (FWE corrected; p < 0.05).

* Cluster voxel represents number of voxels in each significant cluster.

* Peak z-stat denotes the maximum statistical value (z-stat) for the peak activity

* Coordinates were extracted from MNI 152 space

https://doi.org/10.1371/journal.pone.0269154.t003
CALSNIC cohort provides the opportunity to characterize effects over participants enrolled from multiple sites using a harmonized MRI and clinical protocol.

**Intra-network functional connectivity**

**Sensorimotor RSN.** ALS patients displayed higher intra-network Rs-FC in the bilateral sensorimotor RSN involving the superior, middle, and inferior frontal gyrus, anterior and posterior cingulate gyrus, pre- and post-central gyrus, supramarginal gyrus, precuneus, supplementary motor area and parietal lobes. These areas play a key role in integrating sensory and motor information [50–52], particularly for processing sensory information into action [53] or for executing motor tasks and/or sensorimotor components [10, 11]. In ALS, functional changes in the sensorimotor RSN have been consistently reported [7, 12–15, 17–19]. Few such studies have reported enhanced Rs-FC within the sensorimotor RSN using an ICA approach [17, 18] whereas others have used an a-priori approach [12, 22] suggesting an underlying mechanism of neurodegeneration in the motor cortex of ALS patients. Enhanced Rs-FC in ALS patients [22] were also reported using network analysis indicating focal reorganisation and remodulation of functional brain networks. Our finding of a Rs-FC increase in the sensorimotor RSN provides further evidence for a breakdown of neuronal circuits in ALS patients [54]. In addition, the present study found that enhanced Rs-FC within the sensorimotor RSN correlated with faster disease progression rate and longer symptom duration. This is in alignment with previous findings [7, 13] suggesting an association between a dynamic pattern of functional alterations in the sensorimotor RSN of ALS patients and an increase in motor impairment [7, 55]. The present study also found a correlation between higher intra-network Rs-FC in the sensorimotor RSN and lower ALSFRS-R. The findings are supportive of an underlying mechanism of increased neuronal connectivity in the motor cortices of ALS patients, potentially acting as a predictor of motor decline and higher clinical disability [56]. Moreover, higher intra-network Rs-FC in the right hemisphere of the sensorimotor RSN

![Significant correlations between higher intra-network resting-state functional connectivity (Rs-FC) and the clinical variables of Amyotrophic lateral sclerosis (ALS) patients.](https://doi.org/10.1371/journal.pone.0269154.g003)
### Table 4. Brain areas showing positive and negative correlations between higher intra-network resting-state functional connectivity (Rs-FC) in resting-state networks (RSNs) and clinical data.

| Clinical parameters and nature of correlation | Resting-state networks | Brain areas | No of clusters | Clusters voxels | Coordinates | Peak-z stat |
|----------------------------------------------|------------------------|-------------|----------------|-----------------|-------------|-------------|
| **ALSFRS-R [Negative correlation]**          |                        |             |                |                 |             |             |
| Sensorimotor Network                         | Bilateral: Anterior and posterior cingulate gyrus, pre- and post- central gyrus, precuneus, supplementary motor area, superior frontal gyrus, supramarginal gyrus, superior parietal lobule. | 20 | 1015 | 28 46 60 | 4.93 |
|                                             |                        | 19 | 232 | 30 77 54 | 4.04 |
|                                             |                        | 18 | 160 | 57 75 53 | 4.13 |
| Cerebellum Network                           | Bilateral: VI, VIII a and b, IX, X, crus I | 12 | 6753 | 36 42 17 | 4.8 |
|                                             |                        | 10 | 3 | 27 39 8 | 2.4 |
| Auditory Network                             | Right: Superior and middle temporal gyrus, insula, planum temporale, planum polare, parietal operculum cortex, heschl’s Gyrus | 11 | 260 | 47 54 49 | 4.54 |
|                                             |                        | 10 | 180 | 72 39 40 | 3.64 |
| Default Mode Network                         | Bilateral: Central operculum cortex | 5 | 1816 | 49 34 36 | 4.41 |
|                                             |                        | 3 | 165 | 20 36 48 | 4.39 |
| Right Fronto-Parietal Network                | Right: Superior, middle, and inferior frontal gyrus, pre-central gyrus, superior parietal lobule, supramarginal gyrus, angular gyrus, lateral occipital cortex | 19 | 1427 | 16 39 44 | 4.49 |
|                                             |                        | 18 | 537 | 20 77 49 | 4.1 |
|                                             |                        | 17 | 101 | 33 95 29 | 3.63 |
| Left Fronto-Parietal Network                 | Left: Superior, middle, and inferior frontal gyrus, pre-central gyrus | 19 | 742 | 67 75 47 | 4.46 |
|                                             |                        | 18 | 684 | 63 37 60 | 4.56 |
| Orbitofrontal Network                        | Bilateral: Frontal pole, frontal orbital and medial cortex, paracingulate gyrus, subcallosal cortex | 5 | 2775 | 47 87 29 | 4.8 |
|                                             |                        | 4 | 8 | 35 65 36 | 3.04 |
| Dorsal Attention Network                     | Bilateral: Lateral occipital cortex, angular gyrus, precuneus cortex, cuneal cortex, supracalcarine cortex, occipital pole, angular gyrus | 3 | 4319 | 22 27 50 | 4.28 |
|                                             |                        | 2 | 466 | 64 40 59 | 3.52 |
| Basal Ganglia Network                        | Bilateral: Thalamus, putamen, pallidum, and insular cortex, precuneus, posterior cingulate gyrus | 3 | 50 | 37 38 47 | 2.75 |
|                                             |                        | 2 | 20 | 38 37 40 | 2.53 |
| Medial Temporal Network                      | Bilateral: temporal fusiform gyrus, parahippocampal gyrus, temporal pole, inferior temporal gyrus, middle temporal gyrus, temporal occipital fusiform cortex | 2 | 1388 | 64 56 19 | 4.02 |
|                                             |                        | 1 | 1223 | 34 59 21 | 4.64 |
| Ventral Stream Network                       | Right: Superior, middle, and inferior temporal gyrus, temporoparietal part, posterior supramarginal gyrus, superior and inferior lateral occipital cortex, angular Gyrus | 5 | 342 | 16 63 27 | 3.66 |
|                                             |                        | 4 | 76 | 32 45 38 | 3.33 |

(Continued)
### Table 4. (Continued)

| Clinical parameters and nature of correlation | Resting-state networks | Brain areas | No of clusters | Clusters voxels | Coordinates | Peak-z stat |
|-----------------------------------------------|------------------------|-------------|----------------|----------------|-------------|-------------|
| **ECAS Total [Negative correlation]**          |                        |             |                |                | X | Y | Z |
| Cerebellum Network                            | Bilateral: VI, VIII a and b, IX, X, crus I | 6 | 1112 | 60 35 19 | 5.65 |
|                                               |                        | 5 | 659 | 28 34 17 | 4.02 |
|                                               |                        | 4 | 166 | 23 27 11 | 3.53 |
| Default Mode Network                          | Bilateral: Precuneus cortex, cuneal cortex, intracalcarine cortex, supracalcarine cortex, posterior cingulate gyrus. | 1 | 2049 | 47 42 45 | 5.17 |
|                                               |                        | 6 | 51 | 65 23 49 | 5.39 |
|                                               |                        | 5 | 7 | 42 24 53 | 2.95 |
|                                               |                        | 4 | 6 | 47 89 27 | 4.99 |
| Left Fronto-Parietal Network                  | Left: Superior, middle, and inferior frontal gyrus, pre-central gyrus | 19 | 742 | 67 75 47 | 4.46 |
| Orbitofrontal Network                         | Bilateral: Frontal pole, frontal orbital and medial cortex, paracingulate gyrus, subcallosal cortex. | 3 | 4143 | 38 88 32 | 5.64 |
|                                               |                        | 2 | 23 | 31 83 37 | 3.95 |
| Basal Ganglia Network                         | Bilateral: Thalamus, putamen, pallidum, and insular cortex | 6 | 1337 | 56 59 43 | 4.55 |
|                                               |                        | 5 | 467 | 42 53 26 | 3.99 |
| Medial Temporal Network                       | Bilateral: Temporal fusiform gyrus, parahippocampal gyrus, temporal pole, inferior and middle temporal gyrus, temporal occipital fusiform cortex | 4 | 1043 | 74 44 28 | 4.92 |
|                                               |                        | 3 | 113 | 30 60 21 | 4.23 |
| **ECAS ALS Specific [Negative correlation]**   |                        |             |                |                | X | Y | Z |
| Orbitofrontal Network                         | Bilateral: Frontal pole, frontal orbital and medial cortex, paracingulate gyrus, subcallosal cortex. | 4 | 1500 | 32 80 31 | 5.60 |
|                                               |                        | 3 | 800 | 28 81 29 | 3.81 |
|                                               |                        | 2 | 200 | 21 72 40 | 2.68 |
|                                               |                        | 1 | 20 | 14 60 27 | 2.58 |
| Medial Temporal Network                       | Bilateral: Temporal fusiform gyrus, parahippocampal gyrus, temporal pole, inferior and middle temporal gyrus, temporal occipital fusiform cortex | 2 | 200 | 59 41 22 | 4.23 |
|                                               |                        | 1 | 150 | 40 52 19 | 3.18 |
| **ECAS ALS Non-Specific [Negative correlation]** |                        |             |                |                | X | Y | Z |
| Default Mode Network                          | Bilateral: Precuneus cortex, cuneal cortex, intracalcarine cortex, supracalcarine cortex, posterior cingulate gyrus. | 15 | 2611 | 42 40 38 | 4.29 |
| **UMN Burden [Positive correlation]**         |                        |             |                |                | X | Y | Z |
| Cerebellum Network                            | Bilateral: VI, VII a and b, VIII a and b, Crus I, vermis VIII a | 3 | 166 | 27 32 16 | 3.54 |
|                                               |                        | 2 | 7 | 34 32 18 | 3.57 |
|                                               |                        | 1 | 6 | 30 24 24 | 3.87 |

(Continued)
Table 4. (Continued)

| Clinical parameters and nature of correlation | Resting-state networks | Brain areas | No of clusters | Clusters voxels | Coordinates | Peak-z stat |
|-----------------------------------------------|------------------------|-------------|----------------|----------------|-------------|------------|
| Disease Progression Rate [Positive correlation] | Sensorimotor Network   | Right: Middle frontal gyrus. | 6              | 453            | 45 70 61    | 6.01       |
|                                                |                        | Bilateral: Anterior cingulate gyrus, supplementary motor area, superior frontal gyrus, precentral gyrus. | 5              | 272            | 33 68 63    | 5.22       |
|                                                |                        | Left: Precuneus          | 4              | 100            | 29 59 65    | 3.99       |
|                                                | Right Fronto-Parietal Network | Right: Superior, middle, and inferior frontal gyrus, pre-central gyrus | 6              | 919            | 28 67 61    | 4.81       |
|                                                |                        |                          | 5              | 192            | 30 92 44    | 3.85       |
|                                                |                        |                          | 4              | 10             | 22 70 63    | 2.89       |
| Symptom Duration [Positive correlation]        | Sensorimotor Network   | Right: Middle frontal gyrus. | 4              | 188            | 33 68 63    | 5.28       |
|                                                |                        | Bilateral: Anterior cingulate gyrus, Supplementary motor area, Superior frontal gyrus, Precentral gyrus. | 3              | 12             | 59 46 68    | 4.11       |
|                                                | Right Fronto-Parietal Network | Right: Superior, middle, and inferior frontal gyrus, pre-central gyrus | 1              | 828            | 28 67 61    | 4.61       |
| Left Foot Tapping [Negative correlation]       | Sensorimotor Network   | Right: Superior, middle, and inferior frontal gyrus, pre-central gyrus | 5              | 13             | 32 63 65    | 2.66       |
| Right Foot Tapping [Negative correlation]      | Cerebellum Network     | Bilateral: VI, VII a and b, VIII a and b, IX, X, crus I, vermis VIII a | 11             | 208            | 28 28 7     | 2.63       |
|                                                |                        |                          | 10             | 100            | 27 39 8     | 2.4        |
| Right Finger Tapping [Negative correlation]    | Basal Ganglia          | Left: Thalamus           | 4              | 16             | 40 50 20    | 3.90       |

Abbreviations: Intra-network resting-state functional connectivity correlation results in Amyotrophic lateral sclerosis (ALS) patients with clinical scale i.e. ALSFRS-R: ALS functional rating scale revised; ECAS: Edinburg cognitive and behavioural ALS screen; UMN; Upper motor neuron burden (FWE corrected; p < 0.05)

* Number of clusters represents the exact number of clusters in the fMRI analysis stayed significant after correcting the results for multiple comparisons (FWE corrected; p < 0.05).
* Cluster voxel represents number of voxels in each significant cluster.
* Peak z-stat denotes the maximum statistical value (z-stat) for the peak activity
* Coordinates were extracted from MNI 152 space

https://doi.org/10.1371/journal.pone.0269154.t004
Moreover, higher intra-network Rs-FC in the right hemisphere of the sensorimotor RSN (superior, middle, inferior frontal gyrus, and pre-central gyrus) significantly correlated with the left foot tapping scores. Tapping is related to UMN functioning [57] which is thought to be affected by degeneration of the primary motor cortex in ALS [54]. Surprisingly, other studies have reported decreased BOLD signals in the sensorimotor RSN using an ICA approach [19]. The contradictory findings between the former and present study are likely due to the differences in patient characteristics and data acquisition parameters. Additionally, previous studies have different analytical methodologies which may also have likely triggered the differences in the findings.

**Default mode RSN.** ALS patients displayed higher intra-network Rs-FC in the posterior part of the default mode RSN involving the bilateral precuneus cortex, cuneal cortex, supratemporal and infratemporal cortex, posterior cingulate gyrus, angular gyrus, superior and inferior lateral occipital cortex, supramarginal gyrus, and middle temporal gyrus. The default mode RSN is mainly involved in the regulation of complex cognitive and social functions [58–61]. More specifically, the anterior default mode RSN is involved in self-referential mental thoughts [62] and the posterior default mode RSN is mainly involved in directed attention, such as daydreaming and scene-construction [62]. In ALS patients, several studies have examined the default mode RSN using ICA [13, 17–20] and seed-based approaches [7, 9]. For example, compared to HC, Mohammadi et al (2009) reported decreased Rs-FC in the frontal and temporal areas [18] whereas Agosta et al (2013) reported increased Rs-FC in the left precuneus and decreased Rs-FC in the right inferior orbitofrontal gyrus [13]. When compared to patients with FTD, ALS patients were found to have decreased Rs-FC in the posterior cingulate cortex [20]. This mixed picture of functional response within the default mode RSN may suggest high variability in clinical and cognitive features [13, 17–20, 63]. Because this network is known to be more active when the mind is free to wander and not engaged in any task-driven activity [64], the present findings may indicate that individuals with ALS synchronize more extramotor regions (i.e. temporo-occipital areas) than HC in the posterior default mode RSN. Supporting this argument, atypical patterns of cerebral degeneration have previously been
observed in the temporo-occipital brain regions of ALS patients [65–68]. Increased cortical functional connectivity within the posterior cingulate gyrus and occipital brain regions has also been recorded by resting-state magnetoencephalography (MEG) [69]. Higher Rs-FC is suggestive of higher functional activity at rest [70, 71], in turn reflecting the clinical aspects of cognitive and behavioral impairment in ALS. In addition, higher intra-network Rs-FC in the posterior cingulate gyrus and precuneus of the default mode RSN correlated with lower ALSFRS-R and cognitive tests (ECAS Total and ECAS ALS Non-specific). The ECAS Total score reflects overall cognitive status and ECAS ALS Non-specific captures memory and visuospatial dysfunction [72]. Both the posterior cingulate gyrus and the precuneus have multiple functional roles, and thus damage to these regions results in various cognitive, emotional, and behavioral disturbances [73, 74] and especially executive dysfunction [72]. Since these extra-motor features are more frequently reported in ALS [4, 75], the observed correlation in the present study confirm increased clinical disability beyond muscle weakness i.e. decline of certain cognitive functions, impaired social cognition, and changes in the perception and processing of emotions in ALS patients. This also aligns with the overall characteristics of ALS that as the disease progresses, brain regions beyond those involved in motor control, that are more involved in cognitive processing, become affected and contribute to worsening clinical symptoms [68, 73].

Dorsal attention RSN. Another interesting finding of the present study is that intra-network Rs-FC is higher in the dorsal attention RSN of ALS patients. The dorsal attention RSN, also known as "task-positive RSN" is most active during a stimulus-based task and is typically anti-correlated with engagement of the default mode RSN [64, 76, 77]. Notably, no other ALS neuroimaging studies to date have observed activation of the dorsal attention RSN. However, brain regions within the dorsal attention RSN have been recorded individually in the context of higher-level cognitive processing several task-based fMRI, MEG, and electroencephalography studies of ALS [69, 73, 78, 79]. These studies shed light on functional disturbances in the extra-motor brain networks reflective of cognitive changes in the cortical structure of the cerebral hemisphere. In the present Rs-FMRI study, the dorsal attention RSN was found mainly to encompass bilateral parieto-occipital brain areas such as the precuneus, lateral occipital cortex, angular gyrus, cuneal cortex, and supracalcarine cortex. These brain regions mainly subserve cognitive domains such as attention, executive functions, visuospatial functions, episodic memory, language, and number processing [80–83]. The increased Rs-FC in the dorsal attention RSN could be a compensatory attempt [56] to preserve these cognitive functions and thus maintain attentional resources in the presence of cognitive fatigue [84, 85]. Higher intra-network Rs-FC in the dorsal attention RSN was also found to be negatively correlated with the ALSFRS-R. This supports the idea that cognitive impairment, here specifically subserved by abnormal Rs-FC of the dorsal attention RSN, contributes to the functional impairment in addition to that due to motor weakness.

Right and left fronto-parietal RSN. ALS patients also showed enhanced intra-network Rs-FC mostly in the frontal and parietal brain regions of the right and left fronto-parietal RSN. The involvement of this RSN in ALS pathology has been addressed twice using ICA methods [13, 18]. One study observed increased Rs-FC in the left fronto-parietal RSN of ALS patients correlating with clinical and cognitive deficits, suggesting that network dysfunction between the frontal and parietal brain regions could induce subtle changes in executive functions [13]. The other study did not detect significant group differences [18]. Thus, at present there is no consensus on involvement of the fronto-parietal RSN in ALS pathology. The fronto-parietal RSN tends to consolidate neuronal signals from multiple brain regions and/or diverse brain RSNs, acting as a relay center for executing complex cognitive functions [5, 76, 86–88]. In the present study, enhanced intra-network Rs-FC in the fronto-parietal RSN could be a
dysfunctional outcome of the frontal and parietal brain regions [13] reflecting the inability to maintain the inhibitory/excitatory balance [89] and/or proper functional organization [13]. These effects may potentially impact the complex cognitive circuitry involved in processing higher cognitive and executive functions [90–94].

A negative correlation was also observed between intra-network Rs-FC in the right and left fronto-parietal RSN, and ALSFRS-R score, indicating Rs-FC changes of brain networks involving connections to the frontal and parietal brain regions is due to the progression of clinical dysfunction in ALS. This anti-correlation confirm neurodegeneration in ALS and is commonly reported as the insufficient response of clinical performances in fronto-parietal RSN of ALS leads to attention lapse, poor cognitive task performance, and improper well-being of mental presentation [95]. Moreover, positive correlations with disease progression rate and symptom duration suggest higher neuronal synchronization is linearly associated with an average disease progression rate and severity of clinical symptoms [96, 97]. Last, Rs-FC in the left fronto-parietal RSN was associated with cognitive performance (ECAS Total) indicating that the underlying neural mechanism might be analogous to the dysfunctions of frontal and parietal brain regions [79, 98–101]. Overall, the present findings suggest that functional connectivity in the fronto-parietal network is crucial for instantiating and flexibly modulating cognitive control [102].

**Cerebellum RSN.** ALS patients showed higher intra-network Rs-FC in the cerebellum RSN (left I-IV, right Crus II, and bilateral VI, VII a and b, IX, X, Crus I, Vermis VIII a). These areas regulate numerous motor and extra-motor functions: lobules I-IV and IX regulate tapping and sensorimotor functions; lobule X regulates balance, posture, reflexes, and eye movements; lobule IV, VII, Crus I and II are implicated in higher cognitive functions such as language, working memory, and visuomotor processes [103]. The functional importance of these cerebellar lobules has been reported previously in ALS, associated with motor and cognitive decline [8, 12, 104]. Several studies have reported Rs-FC changes in the cerebellum using a seed-based approach, suggesting a functional response to the disease-related mechanism [8, 12, 105]. Using ICA, another study did not detect any significant Rs-FC differences within the cerebellum RSN [17]. As the cerebellum is involved in regulating both motor and cognitive functions [103, 106–108], the present findings of higher intra-network Rs-FC in the cerebellar lobules potentially reflects the generalized spread of cerebellar functional alterations as part of compensating for motor and cognitive decline [109]. Supporting this argument, higher cerebellar Rs-FC was found to be negatively correlated with ALSFRS-R scores, and with poor cognitive and tapping performance, suggesting that increasing disability is associated with widespread cerebellar dysfunction [105]. Furthermore, a positive association was found between higher intra-network Rs-FC and higher UMN burden scores. Tapping assesses UMN function [110] and provides useful measure of UMN degeneration in the motor cortex [54, 57]. Lower tapping scores indicate worse motor performances and are suggestive of greater UMN degeneration and functional loss [57, 111]. In line with the role of the cerebellum in facilitating highly practiced movement [112], the present findings link well with dysfunctions in balance, posture, coordination, speech, and other smooth and balanced muscular activities in ALS patients.

**Basal ganglia RSN.** Higher intra-network Rs-FC was found bilaterally in the basal ganglia RSN including the thalamus, putamen, pallidum, insula, and precuneus cortex of ALS patients. Alterations of the whole basal ganglia RSN have not been reported previously in the ALS literature. However, the constituent brain regions have been reported very often in the context of motor and extra-motor ALS manifestations [12, 113], suggesting the major role and importance of the basal ganglia and the associated cortico-basal ganglia-thalamo-cortical loops in the pathophysiology of the disease [113–115]. Basal ganglia degeneration has been touted as a
possible imaging biomarker of ALS [113]. The structural and functional relevance of the basal ganglia have also been discussed with respect to widespread dysfunctional connectivity, volumetric atrophy, cortical thinning, and white matter degeneration in ALS [22, 55, 113, 114, 116–118]. While the basal ganglia and its associated pathways are classically known in ALS pathology for preserving the intact motor and cognitive abilities [113, 114], our findings can be interpreted as alterations in the fronto-striatal circuitry is implicated in the dysfunction of motor outcomes, and impaired cognitive, and behavioural flexibility in ALS patients [119–121].

Furthermore, increased Rs-FC of the basal ganglia RSN correlated with lower clinical scores of ALSFRS-R, ECAS Total, and right finger tapping. These correlations align with the overall characteristics of basal ganglia pathology in ALS. The basal ganglia, through their afferent and efferent pathways, not only influence the primary motor cortex but also premotor and prefrontal cortices that are involved in language and cognitive functions [119–121]. Therefore, increasing dysfunction of basal ganglia RSN and its associated pathways is likely contributing to the worsening of clinical symptoms, including greater disability and motor and cognitive decline.

**Auditory, medial temporal, and ventral stream RSNs.** Higher intra-network Rs-FC in the auditory, medial temporal, and ventral stream RSNs was found for ALS patients. These have not previously been implicated in ALS. ALS pathology in the cerebrum is thought to develop first in the motor regions with progressive spread in frontotemporal regions (with the last pathological stage including the medial of the temporal lobe) [9, 122, 123]. A recent Rs-FMRI study has discussed the importance of temporo-occipital brain regions in the pathophysiology of ALS, suggesting their involvement at the later stages of disease [66]. In the present study, therefore, higher intra-network Rs-FC within the auditory, medial temporal, and ventral stream RSNs may suggest a compensatory recruitment of additional resources to overcome early stage dysfunction [124].

Moreover, higher intra-network Rs-FC in the auditory, medial temporal, and ventral stream RSNs negatively correlated with ALSFRS-R, indicating that physical impairment is associated with the altered neural connectivity accompanying the progression of loss in functional outcomes [105, 125]. Higher intra-network Rs-FC in the medial temporal RSN also correlated negatively with ECAS Total and ECAS ALS Specific scores, indicating that greater medial temporal connectivity is associated with worse cognitive performance specifically involving language, verbal fluency, and executive functions [95, 98, 126]. The abnormal Rs-FC involving the mesial temporal lobe is in line with frontotemporal lobar degeneration. Because the significance of hyper-connectivity is poorly understood in ALS in the context of cognition, these correlation results shed additional light on the medial temporal involvement and the nature of cognitive decline in ALS patients consistent with the characteristics of later-stage disease pathology.

**Orbitofrontal RSN.** ALS patients had higher intra-network Rs-FC bilaterally in the frontal pole, frontal orbital, frontal medial, paracingulate gyrus, and subcallosal cortex of the orbitofrontal RSN. This particular RSN occupies the prefrontal cortex of the frontal lobe [127] and is involved in the processing of memory, executive functions, reward, decision making, goal-directed behavior, and control of inhibitory functions [128–137]. Regions of the orbitofrontal RSN are also engaged in processing emotional responses and social behavior such as empathy, external environmental stimuli, predicting future events, and thoughts about the self and others [138]. The orbitofrontal RSN has been previously discussed in ALS in the context of stimulus-driven tasks [139] given the importance of these networked regions in cognition and in executive processing. Using graph-theory methods, enhanced resting-state fluctuations in the right orbital frontal and prefrontal cortex were proposed as a means to assess disease severity
in the later stages of ALS [63]. A decreased functional response in the right inferior orbitofrontal gyrus of the default mode RSN has also been reported to suggest cognitive impairment [13].

As discussed above, motor-related Rs-FC changes in the motor areas of the frontal cortex are pronounced at stage one of ALS pathology. Extra-motor regions including the prefrontal cortex and orbitofrontal cortex occur at stage three [54, 122, 140]. This suggests that the findings in the present study of ALS Rs-FC changes within the orbitofrontal RSN reflect dysfunction in the extra-motor system caused by progressive neurodegeneration at the later stages of the disease [63, 122] as also observed in other disorders such as Parkinson’s disease gait [141–144]. From the literature already discussed, it is speculated that the higher intra-network Rs-FC is a compensatory mechanism [56] within the orbitofrontal RSN to maintain cognitive, affective, and executive functions. This notion is further supported by the negative correlation of higher orbitofrontal Rs-FC with lower ALSFRS-R and the negative correlations with ECAS Total and ECAS ALS Specific scores.

Inter-network resting-state functional connectivity

**Basal ganglia and orbitofrontal RSNs.** ALS patients displayed higher inter-network Rs-FC between the basal ganglia RSN and the orbitofrontal RSN. The inter-functional relationship between these RSNs has not been assessed previously in the ALS literature, but the structural and functional relevance of basal ganglia and orbitofrontal brain regions in ALS pathology have been studied extensively as described above, indicating that the disease can produce widespread neurodegeneration [13, 63, 113, 115]. Studies have reported involvement of basal ganglia grey matter pathology in terms of the degree of volumetric changes, shape, and density in ALS [113, 115] suggesting an association of the basal ganglia with widespread areas of the neocortex [115] and/or more specific dysregulation of the fronto-striatal network [113]. Moreover, dysfunction in the projections of the basal ganglia and orbitofrontal cortex may lead to altered fronto-striatal connectivity patterns [145]. Cortico-basal ganglia circuits are involved in monitoring the motor and cognitive aspects of altered cortical processes [145–147] and thus influence sensorimotor, limbic and cognitive functions [121, 148, 149]. These circuits are known to play a key role in ALS pathology [150] and act as an underlying substrate in the control of motor and cognitive capabilities of the prefrontal cortex [149]. The orbitofrontal cortex also has reciprocal anatomical connections with the basal ganglia controlling multiple functions, such as decision-making [151, 152].

Based on this literature, the present findings of increased Rs-FC between the basal ganglia RSN and orbitofrontal RSN likely are a consequence of compensatory processes attempting to preserve motor and cognitive functions in the presence of ALS [56]. Furthermore, a negative correlation of higher Rs-FC of these RSNS with lower ECAS ALS Specific score is consistent with the idea that damage to the fronto-striatal circuits (as can occur in ALS) may lead to the impaired cognitive and executive functions [145, 147]. Overall, the present findings suggest that ALS pathology significantly alters the interplay between the basal ganglia and orbitofrontal RSNs.

**Conclusion**

This study highlights the role and importance of multiple RSNs in the pathophysiology of ALS, and sheds light on functional disturbances within and between RSNs. Both motor and extra-motor RSNs were found to be altered in ALS, and Rs-FC changes were associated with clinical measures of motor, cognitive, and general disease status. ALS is a multi-system disorder, confirmed by the extent of diffuse involvement of RSNs [5].
Previous neuroimaging literature has demonstrated increased Rs-FC in ALS patients [17, 69, 105] this methodology holding promise as a neuroimaging biomarker [143, 153]. The present study findings are supportive of prior works that higher Rs-FC represents a pathobiological feature related to key clinical aspects of the disease [71, 89, 154]. Higher inter-network Rs-FC could also indicate an early compensatory or adaptive response mechanism to overcome motor and cognitive decline, mediated by functional reorganization and plasticity-related changes [56]. The association between higher neuronal synchronization and disease-related modulatory response in ALS is also suggestive of such changes [155].

From a methodological perspective, the present work also highlights the utility of the ICA approach to explore whole brain RSN functional connectivity in ALS. The multi-center CALSNIC Rs-FMRI data facilitated exploration of RSNs in a large population sample in unprecedented detail, revealing the involvement of specific networks in ALS not previously described. Discrepancies with previous literature could be due to sample size and/or methodological differences, as well as patient variability and recruitment bias.

**Limitations**

There are limitations to our study. First, the multicenter approach introduces site-dependent bias and variability in clinical and imaging data. However, a harmonized imaging and clinical protocol was adopted across all sites and the Rs-FMRI data were carefully processed in a stringent and standardized manner to extract Rs-FC information. Before performing the statistical analysis, the data were carefully de-noised to suppress confounding effects including head motion [39] followed by additional motion parameter analysis to remove bias from motion artifact [43]. The statistical analysis also included a multiple linear regression model accounting for multiple co-variates of no interest (age, sex, motion parameters, site, etc.) to increase the likelihood that the observed Rs-FC changes related to disease state (ALS, HC) and clinical variables.

Second, the association of network connectivity with clinical disease staging was not explored. This could be explored in the future by subgrouping patients according to the King’s or Mitos staging systems.

Third, associations between functional and structural changes were not explored. Such association are important to investigate, to assess the extent that the observed increases in Rs-FC reflect grey matter or white matter pathology [14]. An association between functional alterations in large-scale RSNs and structural damage could be addressed accordingly in future studies.

**Acknowledgments**

We thank the patients and healthy subjects for their participation in CALSNIC.

**Membership of author group:**

Komal Bharti, Simon J. Graham, Michael Benatar, Hannah Briemberg, Sneha Chenji, Nicolas Dupré, Annie Dionne, Richard Frayne, Angela Genge, Lawrence Korngut, Collin Luk, Lorne Zinman, Sanjay Kalra.

**Author Contributions**

**Conceptualization:** Komal Bharti, Sanjay Kalra.

**Formal analysis:** Komal Bharti.

**Investigation:** Komal Bharti.
Methodology: Komal Bharti.

Supervision: Sanjay Kalra.

Validation: Komal Bharti.

Visualization: Komal Bharti, Hannah Briemberg, Nicolas Dupré, Annie Dionne, Richard Frayne, Angela Genge, Lawrence Korngut, Collin Luk, Lorne Zinman.

Writing – original draft: Komal Bharti.

Writing – review & editing: Komal Bharti, Simon J. Graham, Michael Benatar, Sneha Chenji, Sanjay Kalra.

References

1. Beghi E, Logroscino G, Chiò A, Hardiman O, Mitchell D, Swingle R, et al. The epidemiology of ALS and the role of population-based registries. Biochim Biophys Acta. 2006; 1762: 1150–1157. https://doi.org/10.1016/j.bbadis.2006.09.008 PMID: 17071069

2. Hardiman O, van den Berg LH, Kiernan MC. Clinical diagnosis and management of amyotrophic lateral sclerosis. Nat Rev Neurol. 2011; 7: 639–649. https://doi.org/10.1038/nrneurol.2011.153 PMID: 21989247

3. Rowland LP, Shneider NA. Amyotrophic lateral sclerosis. N Engl J Med. 2001; 344: 1688–1700. https://doi.org/10.1056/NEJM20010531344207 PMID: 11386269

4. Goldstein LH, Abrahams S. Changes in cognition and behaviour in amyotrophic lateral sclerosis: nature of impairment and implications for assessment. Lancet Neurol. 2013; 12: 368–380. https://doi.org/10.1016/S1474-4422(13)70026-7 PMID: 23518330

5. Phukan J, Pender NP, Hardiman O. Cognitive impairment in amyotrophic lateral sclerosis. Lancet Neurol. 2007; 6: 994–1003. https://doi.org/10.1016/S1474-4422(07)70265-X PMID: 17945153

6. Phukan J, Elamin M, Bede P, Jordan N, Gallagher L, Byrne S, et al. The syndrome of cognitive impairment in amyotrophic lateral sclerosis: a population-based study. J Neurol Neurosurg Psychiatry. 2012; 83: 102–108. https://doi.org/10.1136/jnnp-2011-300188 PMID: 21836033

7. Chenji S, Jha S, Lee D, Brown M, Seres P, Mah D, et al. Investigating Default Mode and Sensorimotor Network Connectivity in Amyotrophic Lateral Sclerosis. PLoS ONE. 2016; 11. https://doi.org/10.1371/journal.pone.0157443 PMID: 27322194

8. Gius T, Zhang Y, Tang X, Liu X, Wang Y, Zhou C, et al. Precentral degeneration and cerebellar compensation in amyotrophic lateral sclerosis: A multimodal MRI analysis. Hum Brain Mapp. 2019; 40: 3464–3474. https://doi.org/10.1002/hbm.24609 PMID: 31020731

9. Schultheiss I, Gorges M, Müller H-P, Lülé D, Del Tredici K, Ludolph AC, et al. Functional connectivity changes resemble patterns of pTDP-43 pathology in amyotrophic lateral sclerosis. Sci Rep. 2016; 6: 38391. https://doi.org/10.1038/srep38391 PMID: 27929102

10. Biswal B, Yetkin FZ, Haughton VM, Hyde JS. Functional connectivity in the motor cortex of resting human brain using echo-planar MRI. Magn Reson Med. 1995; 34: 537–541. https://doi.org/10.1002/mrm.1910340409 PMID: 8524021

11. Smith SM, Fox PT, Miller KL, Glahn DC, Fox PM, Mackay CE, et al. Correspondence of the brain’s functional architecture during activation and rest. Proc Natl Acad Sci U S A. 2009; 106: 13040–13045. https://doi.org/10.1073/pnas.0905267106 PMID: 19620724

12. Agosta F, Valsasina P, Absinta M, Riva N, Sala S, Prelle A, et al. Sensorimotor Function Connectivity Changes in Amyotrophic Lateral Sclerosis. Cereb Cortex. 2011; 21: 2291–2298. https://doi.org/10.1093/cercor/bhr002 PMID: 21368084

13. Agosta F, Canu E, Valsasina P, Riva N, Prelle A, Corni G, et al. Divergent brain network connectivity in amyotrophic lateral sclerosis. Neurobiol Aging. 2013; 34: 419–427. https://doi.org/10.1016/j.neurobiolaging.2012.04.015 PMID: 22608240

14. Douaud G, Filippini N, Knight S, Talbot K, Turner MR. Integration of structural and functional magnetic resonance imaging in amyotrophic lateral sclerosis. Brain J Neurol. 2011; 134: 3470–3479. https://doi.org/10.1093/brain/awr279 PMID: 22075069

15. Fang X, Zhang Y, Wang Y, Zhang Y, Hu J, Wang J, et al. Disrupted effective connectivity of the sensorimotor network in amyotrophic lateral sclerosis. J Neurol. 2016; 263: 508–516. https://doi.org/10.1007/s00415-015-8013-z PMID: 26743627
16. Ma X, Zhang J, Zhang Y, Chen H, Li R, Wang J, et al. Altered cortical hubs in functional brain networks in amyotrophic lateral sclerosis. Neuror Sci. 2015; 36: 2097–2104. https://doi.org/10.1007/s10072-015-2319-6 PMID: 26198762

17. Menke RAL, Proudfoot M, Wu J, Andersen PM, Talbot K, Benatar M, et al. Increased functional connectivity common to symptomatic amyotrophic lateral sclerosis and those at genetic risk. J Neuror Neuropsych Psychiatry. 2016; 87: 580–588. https://doi.org/10.1136/jnp-2015-311945 PMID: 26733601

18. Mohammadi B, Kollewe K, Samii A, Krampf K, Dengler R, Münte TF. Changes of resting state brain networks in amyotrophic lateral sclerosis. Exp Neurol. 2009; 217: 147–153. https://doi.org/10.1016/j.expneurol.2009.01.025 PMID: 19416664

19. Tedeschi G, Trojci F, Testi S, Cerbo D, Sagnelli A, Paccone A, et al. Interaction between aging and neurodegeneration in amyotrophic lateral sclerosis. Neurobiol Aging. 2012; 33: 886–898. https://doi.org/10.1016/j.neurobiolaging.2010.07.011 PMID: 20739098

20. Trojci F, Sorrentino P, Sorrentino G, Tedeschi G. Neurodegeneration of brain networks in the amyotrophic lateral sclerosis-frontotemporal lobar degeneration (ALS-FTLD) continuum: evidence from MRI and MEG studies. CNS Spectr. 2018; 23: 378–387. https://doi.org/10.1017/S109285291700075X PMID: 29076800

21. Welsh RC, Jelsone-Swain LM, Foerster BR. The utility of independent component analysis and machine learning in the identification of the amyotrophic lateral sclerosis diseased brain. Front Hum Neurosci. 2013; 7: 251. https://doi.org/10.3389/fnhum.2013.00251 PMID: 23772210

22. Basaia S, Agosta F, Cividini C, Trojci F, Riva N, Spinelli EG, et al. Structural and functional brain connectome in motor neuron diseases: A multicenter MRI study. Neurology. 2020; 95: e2552–e2564. https://doi.org/10.1212/WNL.0000000000010731 PMID: 32913015

23. Jelsone-Swain LM, Fling BW, Seidler RD, Hovatter R, Gruis K, Welsh RC. Reduced Interhemispheric Functional Connectivity in the Motor Cortex during Rest in Limb-Onset Amyotrophic Lateral Sclerosis. Front Syst Neurosci. 2010; 4: 158. https://doi.org/10.3389/fnsys.2010.00158 PMID: 21228916

24. Lee MH, Smyser CD, Shimony JS. Resting-state fMRI: a review of methods and clinical applications. AJNR Am J Neuroradiol. 2013; 34: 1866–1872. https://doi.org/10.3174/ajnr.A3263 PMID: 22936095

25. Rosazza C, Minati L. Resting-state brain networks: literature review and clinical applications. Neurolog Sci Off J Ital Neurol Soc Ital Soc Clin Neurophysiol. 2011; 32: 773–785. https://doi.org/10.1007/s10072-011-0636-y PMID: 21667095

26. Kolra S, Khan ML, Barlow L, Beaulieu C, Benatar M, Brimmberg H, et al. The Canadian ALS Neuroimaging Consortium (CALSNIC)—a multicentre platform for standardized imaging and clinical studies in ALS. medRxiv. 2020; 2020.07.10.20142679. https://doi.org/10.1101/2020.07.10.20142679

27. Brooks BR. The El Escorial World Federation of Neurology criteria for the diagnosis of amyotrophic lateral sclerosis. Subcommittee on Motor neuron Diseases/Amyotrophic Lateral Sclerosis of the World Federation of Neurology Research Group on Neuromuscular Diseases and the El Escorial “Clinical limits of amyotrophic lateral sclerosis” workshop contributors. J Neuror Sci. 1994; 124 Suppl: 96–107. https://doi.org/10.1016/0022-510x(94)90191-0

28. Cederbaum JM, Stambler N, Malta E, Fuller C, Hilt D, Thurmound B, et al. The ALSFRS-R: a revised ALS functional rating scale that incorporates assessments of respiratory function. BDNF ALS Study Group (Phase III). J Neuror Sci. 1999; 169: 13–21. https://doi.org/10.1016/s0022-510x(99)00210-5 PMID: 10540002

29. Kimura F, Fujimura C, Ishida S, Nakajima H, Furutama D, Uehara H, et al. Progression rate of ALSFRS-R at time of diagnosis predicts survival time in ALS. Neurology. 2006; 66: 265–267. https://doi.org/10.1212/01.wnl.0000194316.91908.8a PMID: 16434671
34. Ishaque A, Mah D, Seres P, Luk C, Johnston W, Chenji S, et al. Corticospinal tract degeneration in ALS unmasked in T1-weighted images using texture analysis. Hum Brain Mapp. 2019; 40: 1174–1183. https://doi.org/10.1002/hbm.24437 PMID: 30367724

35. Niven E, Newton J, Foley J, Colville S, Swingler R, Chandran S, et al. Validation of the Edinburgh Cognitive and Behavioural Amyotrophic Lateral Sclerosis Screen (ECAS): A cognitive tool for motor disorders. Amyotroph Lateral Scler Front Degener. 2015; 16: 172–179. https://doi.org/10.3109/21678421.2015.1030430 PMID: 25967542

36. Smith SM, Jenkinson M, Woolrich MW, Beckmann CF, Johansen-Berg H, et al. Advances in functional and structural MR image analysis and implementation as FSL. Neuroimage. 2004; 23 Suppl 1: S208–219. https://doi.org/10.1016/j.neuroimage.2004.07.051 PMID: 15501092

37. Smith SM. Fast robust automated brain extraction. Hum Brain Mapp. 2002; 17: 143–155. https://doi.org/10.1002/hbm.10062 PMID: 12391568

38. Beckmann CF, Smith SM. Tensorial extensions of independent component analysis for multisubject FMRI analysis. NeuroImage. 2005; 25: 294–311. https://doi.org/10.1016/j.neuroimage.2004.10.043 PMID: 15734364

39. Pruim RHR, Mennes M, Buitelaar JK, Beckmann CF. Evaluation of ICA-AROMA and alternative strategies for motion artifact removal in resting state fMRI. NeuroImage. 2015; 112: 278–287. https://doi.org/10.1016/j.neuroimage.2015.02.063 PMID: 25770990

40. Beckmann CF, Smith SM. Probabilistic independent component analysis for functional magnetic resonance imaging. IEEE Trans Med Imaging. 2004; 23: S208–219. https://doi.org/10.1109/TMI.2003.822821 PMID: 14964560

41. Beckmann CF, DeLuca M, Devlin JT, Smith SM. Investigations into resting-state connectivity using independent component analysis. Philos Trans R Soc Lond B Biol Sci. 2005; 360: 1001–1013. https://doi.org/10.1098/rstb.2005.1634 PMID: 16087444

42. Maknojia S, Churchill NW, Schweizer TA, Graham SJ. Resting State fMRI: Going Through the Motions. Front Neurosci. 2019; 13. https://doi.org/10.3389/fnins.2019.00825 PMID: 31465656

43. Van Dijk KRA, Sabuncu MR, Buckner RL. The Influence of Head Motion on Intrinsic Functional Connectivity MRI. NeuroImage. 2012; 59: 431–438. https://doi.org/10.1016/j.neuroimage.2011.07.044 PMID: 21810475

44. Siegel JS, Power JD, Dubis JW, Vogel AC, Church JA, Schlaggar BL, et al. Statistical improvements in functional magnetic resonance imaging analyses produced by censoring high-motion data points. Hum Brain Mapp. 2013; 35: 1981–1996. https://doi.org/10.1002/hbm.22307 PMID: 23861343

45. Parkes L, Fulcher B, Yücel M, Fornito A. An evaluation of the efficacy, reliability, and sensitivity of motion correction strategies for resting-state functional MRI. NeuroImage. 2018; 171: 415–436. https://doi.org/10.1016/j.neuroimage.2017.12.073 PMID: 29278773

46. Nichols TE, Holmes AP. Nonparametric permutation tests for functional neuroimaging: a primer with examples. Hum Brain Mapp. 2002; 15: 1–25. https://doi.org/10.1002/hbm.1058 PMID: 11747097

47. Desikan RS, Ségonne F, Fischl B, Quinn BT, Dickerson BC, Blacker D, et al. An automated labeling system for subdividing the human cerebral cortex on MRI scans into gyral based regions of interest. NeuroImage. 2006; 31: 968–980. https://doi.org/10.1016/j.neuroimage.2006.01.021 PMID: 16530430

48. Turner JA, Damaraju E, van Erp TGM, Mathalon DH, Ford JM, Voyvodic J, et al. A multi-site resting state fMRI study on the amplitude of low frequency fluctuations in schizophrenia. Front Neurosci. 2013; 7: 137. https://doi.org/10.3389/fnins.2013.00137 PMID: 23964193

49. Valk SL, Di Martino A, Milham MP, Bernhardt BC. Multicenter mapping of structural network alterations in autism. Hum Brain Mapp. 2015; 36: 2364–2373. https://doi.org/10.1002/hbm.22776 PMID: 25727958

50. Chung SJ, Im J-H, Lee J-H, Lee MC. Stuttering and gait disturbance after supplementary motor area seizure. Mov Disord Off J Mov Disord Soc. 2004; 19: 1106–1109. https://doi.org/10.1002/mds.20136 PMID: 15372607

51. Shum M, Shiller DM, Baum SR, Gracco VL. Sensorimotor integration for speech motor learning involves the inferior parietal cortex. Eur J Neurosci. 2011; 34: 1817–1822. https://doi.org/10.1111/j.1460-9568.2011.07889.x PMID: 22098364

52. Wolpert DM, Kawato M. Multiple paired forward and inverse models for motor control. Neural Netw Off J Int Neural Netw Soc. 1998; 11: 1317–1329. https://doi.org/10.1016/s0893-6080(98)00066-5 PMID: 12662752

53. Vahdat S, Darainy M, Milner TE, Ostry DJ. Functionally Specific Changes in Resting-State Sensorimotor Networks after Motor Learning. J Neurosci. 2011; 31: 16907–16915. https://doi.org/10.1523/JNEUROSCI.2737-11.2011 PMID: 22114261
54. Ragagnin AMG, Shadfar S, Vidal M, Jamali MS, Atkin JD. Motor Neuron Susceptibility in ALS/FTD. Front Neurosci. 2019; 13. https://doi.org/10.3389/fnins.2019.00532 PMID: 31316328

55. Verstraete E, van den Heuvel MP, Veldink JH, Blanken N, Mandl RC, Hulshoff Pol HE, et al. Motor network degeneration in amyotrophic lateral sclerosis: a structural and functional connectivity study. PloS One. 2010; 5: e13664. https://doi.org/10.3389/fnins.2010.001366 PMID: 21060689

56. Hillary FG, Roman CA, Venkatasesan U, Rajtmajer SM, Bajo R, Castellanos ND. Hyperconnectivity is a fundamental response to neurological disruption. Neuropsychology. 2015; 29: 59–75. https://doi.org/10.1037/neu0000110 PMID: 24933491

57. Huynh W, Simon NG, Grosskretz J, Turner MR, Vucic S, Kiernan MC. Assessment of the upper motor neuron in amyotrophic lateral sclerosis. Clin Neurophysiol. 2016; 127: 2643–2660. https://doi.org/10.1016/j.clinph.2016.04.025 PMID: 27291884

58. Laird AR, Fox PM, Eickhoff SB, Turner JA, Ray KL, McKay DR, et al. Behavioral Interpretations of Intrinsic Connectivity Networks. J Cogn Neurosci. 2011; 23: 4022–4037. https://doi.org/10.1162/jocn_a_00077 PMID: 21671731

59. Li W, Mai X, Liu C. The default mode network and social understanding of others: what do brain connectivity studies tell us. Front Hum Neurosci. 2014; 8. https://doi.org/10.3389/fnhum.2014.00074 PMID: 24605094

60. Raichle ME. The brain’s default mode network. Annu Rev Neurosci. 2015; 38: 433–447. https://doi.org/10.1146/annurev-neuro-071013-014030 PMID: 25938726

61. Schilbach L, Eickhoff SB, Rotarska-Jagiela A, Fink GR, Vogeley K. Minds at rest? Social cognition as the default mode of cognizing and its putative relationship to the “default system” of the brain. Conscious Cogn. 2008; 17: 457–467. https://doi.org/10.1016/j.concog.2008.03.013 PMID: 18434197

62. Xu X, Yuan H, Lei X. Activation and Connectivity within the Default Mode Network Contribute Independently to Future-Oriented Thought. Sci Rep. 2016; 6. https://doi.org/10.1038/srep21001 PMID: 26867499

63. Zhou C, Hu X, Li J, Li J, Yin N, Chen L, et al. Altered Brain Network in Amyotrophic Lateral Sclerosis: A Resting Graph Theory-Based Network Study at Voxel-Wise Level. PLOS ONE. 2016; 10. https://doi.org/10.3389/fnins.2016.00204 PMID: 27242409

64. Fox MD, Snyder AZ, Vincent JL, Corbetta M, Van Essen DC, Raichle ME. The human brain is intrinsically organized into dynamic, anticorrelated functional networks. Proc Natl Acad Sci U S A. 2005; 102: 9673–9678. https://doi.org/10.1073/pnas.0504136102 PMID: 15976020

65. Abrahams S, Goldstein LH, Simmons A, Brammer M, Williams SCR, Giampietro V, et al. Word retrieval in amyotrophic lateral sclerosis: a functional magnetic resonance imaging study. Brain. 2004; 127: 1507–1517. https://doi.org/10.1093/brain/awh170 PMID: 15163610

66. Loewe K, Machts J, Kaufmann J, Petri S, Heinze H-J, Borgelt C, et al. Widespread temporo-occipital lobe dysfunction in amyotrophic lateral sclerosis. Sci Rep. 2017; 7. https://doi.org/10.1038/srep40252 PMID: 28067298

67. Ogura A, Watanabe H, Kawabata K, Ohdake R, Tanaka Y, Masuda M, et al. Semantic deficits in ALS related to right lingual/fusiform gyrus network involvement. EBioMedicine. 2019; 47: 506–517. https://doi.org/10.1016/j.ebiom.2018.09.022 PMID: 31492562

68. Sudharshan N, Hanstock C, Hui B, Pyra T, Johnston W, Kalra S. Degeneration of the Mid-Cingulate Cortex in Amyotrophic Lateral Sclerosis Detected In Vivo with MR Spectroscopy. Am J Neuroradiol. 2011; 32: 403–407. https://doi.org/10.3174/ajnr.A2289 PMID: 21087934

69. Fomina T, Weichwald S, Synofzik M, Just J, Schöls L, Schölkopf B, et al. Absence of EEG correlates of self-referential processing depth in ALS. PLOS ONE. 2017; 12: e0180136. https://doi.org/10.1212/WNL.0000000000005333 PMID: 29661904

70. Deco G, Jirsa VK, McIntosh AR. Emerging concepts for the dynamical organization of resting-state activity in the brain. Nat Rev Neurosci. 2011; 12: 43–56. https://doi.org/10.1038/nrn2961 PMID: 21170073

71. Menon P, Kiernan MC, Vucic S. Cortical hyperexcitability precedes lower motor neuron dysfunction in ALS. Clin Neurophysiol Off J Int Fed Clin Neurophysiol. 2015; 126: 803–809. https://doi.org/10.1016/j.clinph.2014.04.023 PMID: 25227219

72. Abrahams S, Newton J, Niven E, Foley J, Bak TH. Screening for cognition and behaviour changes in ALS. Amyotroph Lateral Scler Front Degener. 2014; 15: 9–14. https://doi.org/10.3109/21678421.2013.805784 PMID: 23781974

73. Fomina T, Weichwald S, Synofzik M, Just J, Schöls L, Schölkopf B, et al. Absence of EEG correlates of self-referential processing depth in ALS. PLOS ONE. 2017; 12: e0180136. https://doi.org/10.1371/journal.pone.0180136 PMID: 28662161
74. Leech R, Sharp DJ. The role of the posterior cingulate cortex in cognition and disease. Brain. 2014; 137: 12–32. https://doi.org/10.1093/brain/awt162 PMID: 23869106

75. McCombe PA, Wray NR, Henderson RD. Extra-motor abnormalities in amyotrophic lateral sclerosis: another layer of heterogeneity. Expert Rev Neurother. 2017; 17: 561–577. https://doi.org/10.1080/14737157.2017.1273772 PMID: 27983884

76. Corbetta M, Shulman GL. Control of goal-directed and stimulus-driven attention in the brain. Nat Rev Neurosci. 2002; 3: 201–215. https://doi.org/10.1038/nrn755 PMID: 11994752

77. Vossel S, Geng-JJ, Fink GR. Dorsal and ventral attention systems: distinct neural circuits but collaborative roles. Neurosci Rev. 2014; 20: 150–159. https://doi.org/10.1177/1073858413494269 PMID: 23835449

78. Thorns J, Jansma H, Peschel T, Grosskreutz J, Mohammad B, Dengler R, et al. Extent of cortical involvement in amyotrophic lateral sclerosis—an analysis based on cortical thickness. BMC Neurol. 2013; 13: 148. https://doi.org/10.1186/1471-2377-13-148 PMID: 24138960

79. Witiuk K, Fernandez-Ruiz J, Mc Kee R, Alahyane N, Coe BC, Melanson M, et al. Cognitive Deterioration and Functional Compensation in ALS Measured with fMRI Using an Inhibitory Task. J Neurol. 2017; 264: 88–101. https://doi.org/10.1007/s00415-016-8309-7 PMID: 27778161

80. Gordon EM, Breeden AL, Bean SE, Vaidya CJ. Working memory-related changes in functional connectivity persist beyond task disengagement. Hum Brain Mapp. 2014; 35: 1004–1017. https://doi.org/10.1002/hbm.22230 PMID: 23281202

81. Grill-Spector K, Kourtzi Z, Kanwisher N. The lateral occipital complex and its role in object recognition. Vision Res. 2001; 41: 1409–1422. https://doi.org/10.1016/s0042-6989(01)00073-6 PMID: 11322983

82. Spadone S, Penna SD, Sestieri C, Betti V, Tosoni A, Perrucci MG, et al. Dynamic reorganization of human resting-state networks during visuospatial attention. Proc Natl Acad Sci. 2015; 112: 8112–8117. https://doi.org/10.1073/pnas.1415439112 PMID: 26080395

83. Babu Henry Samuel I, Wang C, Burke SE, Kluger B, Ding M. Compensatory Neural Responses to Cognitive Fatigue in Young and Older Adults. Front Neural Circuits. 2019; 13. https://doi.org/10.3389/fncir.2019.00012 PMID: 30853901

84. Holtzer R, Shuman M, Mahoney JR, Lipton R, Verghese J. Cognitive Fatigue Defined in the Context of Attention Networks. Neuropsychol Dev Cogn B Aging Neuropsychol Cogn. 2011; 18: 108–128. https://doi.org/10.1080/13825585.2010.517826 PMID: 21128132

85. Dixon ML, Vega ADL, Mills C, Andrews-Hanna J, Spreng RN, Cole MW, et al. Heterogeneity within the frontoparietal control network and its relationship to the default and dorsal attention networks. Proc Natl Acad Sci. 2018; 115: E1598–E1607. https://doi.org/10.1073/pnas.1715766115 PMID: 29382744

86. Marek S, Dosenbach NUF. The Frontoparietal Attention Network of the Human Brain: Action, Saliency, and a Priority Map of the Environment. The Neuroscientist. 2012; 18: 502–515. https://doi.org/10.1177/1073858411409051 PMID: 21636849

87. Clark RM, Brizuela M, Blizzard CA, Dickson TC. Reduced Excitability and Increased Neurite Complexity of Cortical Interneurons in a Familial Mouse Model of Amyotrophic Lateral Sclerosis. Front Cell Neurosci. 2018; 12. https://doi.org/10.3389/fncel.2018.00328 PMID: 3023744

88. Hensch TK. Critical period plasticity in local cortical circuits. Nat Rev Neurosci. 2005; 6: 877–888. https://doi.org/10.1038/nn1787 PMID: 16261181

89. Kelsom C, Lu W. Development and specification of GABAergic cortical interneurons. Cell Biosci. 2013; 3: 19. https://doi.org/10.1186/2045-3701-3-19 PMID: 23618463

90. Owens DF, Kriegstein AR. Is there more to GABA than synaptic inhibition? Nat Rev Neurosci. 2002; 3: 715–727. https://doi.org/10.1038/nrm919 PMID: 12209120

91. Wang X-J, Tegné r J, Constantinidis C, Goldman-Rakic PS. Division of labor among distinct subtypes of inhibitory neurons in a cortical microcircuit of working memory. Proc Natl Acad Sci U S A. 2004; 101: 1368–1373. https://doi.org/10.1073/pnas.0305337101 PMID: 14742867

92. Whittington MA, Traub RD. Interneuron diversity series: inhibitory interneurons and network oscillations in vitro. Trends Neurosci. 2003; 26: 675–682. https://doi.org/10.1016/j.tins.2003.08.016 PMID: 14624852
95. Benbrika S, Desgranges B, Eustache F, Viader F. Cognitive, Emotional and Psychological Manifestations in Amyotrophic Lateral Sclerosis at Baseline and Overtime: A Review. Front Neurosci. 2019; 13: 951. https://doi.org/10.3389/fnins.2019.00951 PMID: 31551700

96. Kellmeyer P, Grosse-Wentrup M, Schulze-Bonhage A, Ziemann U, Ball T. Electrophysiological correlates of neurodegeneration in motor and non-motor brain regions in amyotrophic lateral sclerosis—implications for brain–computer interfacing. J Neural Eng. 2018; 15: 041003. https://doi.org/10.1088/1741-2552/aabfa5 PMID: 29676287

97. Woo JH, Wang S, Melhem ER, Gee JC, Cucchiara A, McCluskey L, et al. Linear Associations between Clinically Assessed Upper Motor Neuron Disease and Diffusion Tensor Imaging Metrics in Amyotrophic Lateral Sclerosis. PLOS ONE. 2014; 9: e105753. https://doi.org/10.1371/journal.pone.0105753 PMID: 25144708

98. De Marchi F, Carrarini C, De Martino A, Diamanti L, Fasano A, Lupica A, et al. Cognitive dysfunction in amyotrophic lateral sclerosis: can we predict it? Neuro Sci. 2021; 42: 2211–2222. https://doi.org/10.1007/s12022-021-01588-0 PMID: 33772353

99. McMackin R, Dukic S, Broderick M, Iyer PM, Pinto-G grau M, Mohr K, et al. Dysfunction of attention switching networks in amyotrophic lateral sclerosis. NeuroImage Clin. 2019; 22: 101707. https://doi.org/10.1016/j.nicl.2019.101707 PMID: 30735860

100. Meoded A, Kwan JY, Peters TL, Huey ED, Danielian LE, Wiggs E, et al. Imaging Findings Associated with Cognitive Performance in Primary Lateral Sclerosis and Amyotrophic Lateral Sclerosis. Dement Geriatr Cogn Disord Extra. 2013; 3: 233–250. https://doi.org/10.1159/000353456 PMID: 24052798

101. Sheffield JM, Repovs G, Harms MP, Carter CS, Gold JM, MacDonald AW, et al. Evidence for Accelerated Decline of Functional Brain Network Efficiency in Schizophrenia. Schizophr Bull. 2016; 42: 753–761. https://doi.org/10.1093/schbul/sbv148 PMID: 26472685

102. Stoodley CJ, Valera EM, Schmahmann JD. Functional topography of the cerebellum for motor and cognitive tasks: an fMRI study. NeuroImage. 2012; 59: 1560–1570. https://doi.org/10.1016/j.neuroimage.2011.08.065 PMID: 21907811

103. Chiò A, Paganini M, Agosta F, Calvo A, Cistaro A, Filippi M. Neuroimaging in amyotrophic lateral sclerosis: insights into structural and functional changes. Lancet Neurol. 2014; 13: 1228–1240. https://doi.org/10.1016/S1474-4422(14)70167-X PMID: 25453462

104. Nijssen J, Comley LH, Hedlund E. Motor neuron vulnerability and resistance in amyotrophic lateral sclerosis. Acta Neuropa thol (Berl). 2017; 133: 863–885. https://doi.org/10.1007/s00401-017-1708-8 PMID: 28409262

105. Grieve SM, Menon P, Korgaonkar MS, Gomes L, Foster S, Kiernan MC, et al. Potential structural and functional biomarkers of upper motor neuron dysfunction in ALS. Amyotrophic Lateral Scler Front Degener. 2015; 17: 85–92. https://doi.org/10.3109/21678421.2015.1074707 PMID: 26458122

106. Manto M, Bower JM, Conforto AB, Delgado-Garcı ´ a JM, da Guarda SNF, Gerwig M, et al. Consensus Paper: Roles of the Cerebellum in Motor Control—The Diversity of Ideas on Cerebellar Involvement in Movement. Cerebellum Lond Engl. 2012; 11: 457–487. https://doi.org/10.1007/s12311-011-0331-9 PMID: 23889583

107. Bede P, Elamin M, Byrne S, McLaughlin RL, Kenna K, Vajda A, et al. Basal ganglia involvement in amyotrophic lateral sclerosis. Neurology. 2013; 81: 2107–2115. https://doi.org/10.1212/01.wnl.0000437313.80913.2c PMID: 24212388
114. Bede P, Omer T, Finegan E, Chipika RH, Iyer PM, Doherty MA, et al. Connectivity-based characterisation of subcortical grey matter pathology in frontotemporal dementia and ALS: a multimodal neuroimaging study. Brain Imaging Behav. 2018; 12: 1696–1707. https://doi.org/10.1007/s11682-018-9837-9 PMID: 29423814

115. Machts J, Loewe K, Kaufmann J, Jakubiczka S, Abdulla S, Petri S, et al. Basal ganglia pathology in ALS is associated with neuropsychological deficits. Neurology. 2015; 85: 1301–1309. https://doi.org/10.1212/WNL.0000000000002017 PMID: 26385880

116. Li W, Zhang J, Zhou C, Hou W, Hu J, Feng H, et al. Abnormal Functional Connectivity Density in Amyotrophic Lateral Sclerosis. Front Aging Neurosci. 2018; 10. https://doi.org/10.3389/fnagi.2018.00215 PMID: 30065647

117. Radakovic R, Puthusserryppady V, Flanagan E, Kienan MC, Mioshi E, Hornberger M. Frontostriatal grey matter atrophy in amyotrophic lateral sclerosis: a visual rating study. Dement Neurocogn Disord. 2018; 12: 388–393. https://doi.org/10.1590/1980-57642018dr12-040008 PMID: 30546849

118. Sharma KR, Sheriff S, Maudsley A, Govind V. Diffusion Tensor Imaging of Basal Ganglia and Thalamus in Amyotrophic Lateral Sclerosis. J Neuroimag Imaging Off J Am Soc Neuroimaging. 2013; 23: 368–374. https://doi.org/10.1011/j.1552-6569.2011.00679.x PMID: 22273090

119. DeLong MR, Wichmann T. Circuits and circuit disorders of the basal ganglia. Arch Neurol. 2007; 64: 20–24. https://doi.org/10.1001/archneur.64.1.20 PMID: 17210805

120. DeLong M, Wichmann T. Changing Views of Basal Ganglia Circuits and Circuit Disorders. Clin EEG Neurosci. 2010; 41: 61–67. https://doi.org/10.1177/155005941004100204 PMID: 20521487

121. Middleton FA, Strick PL. Basal ganglia and cerebellar loops: motor and cognitive circuits. Brain Res Rev. 2000; 31: 236–250. https://doi.org/10.1016/S0165-0173(99)00040-5 PMID: 10719151

122. Braak H, Brettschneider J, Ludolph AC, Lee VM, Trojanowski JQ, Del Tredici K. Amyotrophic lateral sclerosis—a model of corticofugal axonal spread. Nat Rev Neurosci. 2013; 9: 708–714. https://doi.org/10.1038/nrn3842 PMID: 24217521

123. Brettschneider J, Del Tredici K, Toledo JB, Robinson JL, Inwijn DJ, Grossman M, et al. Stages of pTDP-43 pathology in amyotrophic lateral sclerosis. Ann Neurol. 2013; 74: 20–38. https://doi.org/10.1002/ana.23937 PMID: 23686809

124. Cosottini M, Pesaresi I, Piazza S, Diciotti S, Cecchi P, Fabbri S, et al. Structural and functional evaluation of cortical motor areas in Amyotrophic Lateral Sclerosis. Exp Neurol. 2012; 234: 169–180. https://doi.org/10.1016/j.expneurol.2011.12.024 PMID: 22226599

125. Wirth AM, Khomenko A, Baldarano D, Kobor I, Hsam O, Grimm T, et al. Combinatorial Biomarker Use of Cortical Thickness, MUNIX, and ALSFRS-R at Baseline and in Longitudinal Courses of Individual Patients With Amyotrophic Lateral Sclerosis. Front Neurol. 2018; 9. https://doi.org/10.3389/fneur.2018.00614 PMID: 30104996

126. Crockford C, Newton J, Lonergan K, Chiwera T, Booth T, Chandran S, et al. ALS-specific cognitive and behavior changes associated with advancing disease stage in ALS. Neurology. 2018; 91: e1370–e1380. https://doi.org/10.1212/WNL.0000000000002017 PMID: 30209236

127. Goldman-Rakic PS. Circuitry of Primate Prefrontal Cortex and Regulation of Behavior by Representative Memory. Comprehensive Physiology. American Cancer Society; 2011. pp. 373–417.

128. Bechara A, Damasio H, Damasio AR. Emotion, Decision Making and the Orbitofrontal Cortex. Cereb Cortex. 2000; 10: 295–307. https://doi.org/10.1093/cercor/10.3.295 PMID: 10731224

129. Damasio AR, Tranel D, Damasio H. Individuals with sociopathic behavior caused by frontal damage fail to respond autonomically to social stimuli. Behav Brain Res. 1990; 41: 81–94. https://doi.org/10.1016/0166-4328(90)90144-4 PMID: 2288668

130. Gottfried JA, O’Doherty J, Dolan RJ. Appetitive and aversive olfactory learning in humans studied using event-related functional magnetic resonance imaging. J Neurosci Off J Soc Neurosci. 2002; 22: 10829–10837. https://doi.org/10.1523/JNEUROSCI.22-24-10829.2002 PMID: 12486176

131. Ikeda A, Lüders HO, Collura TF, Burgess RC, Morris HH, Hamano T, et al. Subdural potentials at orbitofrontal and mesial prefrontal areas accompanying anticipation and decision making in humans: a comparison with Bereitschaftspotential. Electroencephalogr Clin Neurophysiol. 1996; 98: 206–212. https://doi.org/10.1016/0013-4694(95)00239-1 PMID: 8631280

132. Izquierdo A, Suda RK, Murray EA. Bilateral orbital prefrontal cortex lesions in rhesus monkeys disrupt choices guided by both reward value and reward contingency. J Neurosci Off J Soc Neurosci. 2004; 24: 7540–7548. https://doi.org/10.1523/JNEUROSCI.1921-04.2004 PMID: 15329401

133. Nobre AC, Coull JT, Frith CD, Mesulam MM. Orbitofrontal cortex is activated during breaches of expectation in tasks of visual attention. Nat Neurosci. 1999; 2: 11–12. https://doi.org/10.1038/4513 PMID: 10195173
134. O’Doherty JP, Deichmann R, Critchley HD, Dolan RJ. Neural responses during anticipation of a primary taste reward. Neuron. 2002; 33: 815–826. https://doi.org/10.1016/s0896-6273(02)00603-7 PMID: 11879657

135. Roesch MR, Olson CR. Neuronal activity related to reward value and motivation in primate frontal cortex. Science. 2004; 304: 307–310. https://doi.org/10.1126/science.1093223 PMID: 15073380

136. Roesch MR, Olson CR. Neuronal activity in primate orbitofrontal cortex reflects the value of time. J Neurophysiol. 2005; 94: 2457–2471. https://doi.org/10.1152/jn.00373.2005 PMID: 15958600

137. Schoenbaum G, Takahashi Y, Liu T-L, McDannald MA. Does the orbitofrontal cortex signal value? Ann N Y Acad Sci. 2011; 1239: 87–99. https://doi.org/10.1111/j.1749-6632.2011.06210.x PMID: 22145878

138. Stone VE, Baron-Cohen S, Knight RT. Frontal lobe contributions to theory of mind. J Cogn Neurosci. 1998; 10: 640–656. https://doi.org/10.1162/089892998562942 PMID: 9802997

139. Stoppel CM, Vielhaber S, Eckart C, Machts J, Kaufmann J, Heinze H-J, et al. Structural and functional hallmarks of amyotrophic lateral sclerosis progression in motor- and memory-related brain regions. NeuroImage Clin. 2014; 5: 277–290. https://doi.org/10.1016/j.nicl.2014.07.007 PMID: 25161894

140. Walhout R, Verstraete E, van den Heuvel MP, Veldink JH, van den Berg LH. Patterns of symptom development in patients with motor neuron disease. Amyotroph Lateral Scler Front Degener. 2018; 19: 21–28. https://doi.org/10.1080/21678421.2017.1386688 PMID: 29037065

141. Bharti K, Suppa A, Pietracupa S, Upadhyay N, Giannì C, Leodori G, et al. Abnormal Cerebellar Connectivity Patterns in Patients with Parkinson’s Disease and Freezing of Gait. The Cerebellum. 2019; 18: 298–308. https://doi.org/10.1007/s12311-018-0998-4 PMID: 30392037

142. Cole DM, Beckmann CF, Searle GE, Plisson C, Tziortzi AC, Nichols TE, et al. Orbitofrontal Connectivity with Resting-State Networks Is Associated with Midbrain Dopamine D3 Receptor Availability. Cereb Cortex. 2012; 22: 2784–2793. https://doi.org/10.1093/cercor/bhr354 PMID: 22186675

143. Hohenfeld C, Werner CJ, Reetz K. Resting-state connectivity in neurodegenerative disorders: Is there potential for an imaging biomarker? NeuroImage Clin. 2018; 18: 849–870. https://doi.org/10.1016/j.nicl.2018.03.013 PMID: 29876270

144. Skidmore FM, Yang M, Baxter L, von Deneen K, Collingwood J, He G, et al. Apathy, depression, and motor symptoms have distinct and separable resting activity patterns in idiopathic Parkinson disease. NeuroImage. 2013; 81: 484–495. https://doi.org/10.1016/j.neuroimage.2011.07.012 PMID: 21782030

145. Morris LS, Kundu P, Dowell N, Mechelmans DJ, Favre P, Irvine MA, et al. Fronto-striatal organization: Defining functional and microstructural substrates of behavioural flexibility. Cortex J Devoted Study Ner Syst Behav. 2016; 74: 118–133. https://doi.org/10.1016/j.cortex.2015.11.004 PMID: 26673945

146. Alexander GE, DeLong MR, Strick PL. Parallel organization of functionally segregated circuits linking basal ganglia and cortex. Annu Rev Neurosci. 1986; 9: 357–381. https://doi.org/10.1146/annurev.ne.09.030186.002041 PMID: 3085570

147. Draganski B, Kherif F, Klöppel S, Cook PA, Alexander DC, Parker GJM, et al. Evidence for segregated and integrative connectivity patterns in the human Basal Ganglia. J Neurosci Off J Soc Neurosci. 2008; 28: 7143–7152. https://doi.org/10.1523/JNEUROSCI.1486-08.2008 PMID: 18614684

148. Leisman G, Braun-Benjamin O, Melillo R. Cognitive-motor interactions of the basal ganglia in development. Front Syst Neurosci. 2014; 8. https://doi.org/10.3389/fnsys.2014.00016 PMID: 24592214

149. Middleton FA, Strick PL. Cerebellar Projections to the Prefrontal Cortex of the Primate. J Neurosci. 2001; 21: 700–712. https://doi.org/10.1523/JNEUROSCI.21-02-0070.2001 PMID: 11160449

150. Shepherd GMG. Corticostriatal connectivity and its role in disease. Nat Rev Neurosci. 2013; 14: 278–291. https://doi.org/10.1038/nrn3468 PMID: 23511908

151. Frank MJ, Claus ED. Anatomy of a decision: striato-orbitofrontal interactions in reinforcement learning, decision making, and reversal. Psychol Rev. 2006; 113: 300–326. https://doi.org/10.1037/0033-295X.113.2.300 PMID: 16637783

152. Kringlebach ML, Rolls ET. The functional neuroanatomy of the human orbitofrontal cortex: evidence from neuroimaging and neuropsychology. Prog Neurobiol. 2004; 72: 341–372. https://doi.org/10.1016/j.pneurobi.2004.03.006 PMID: 15157726

153. Filippi M, Spinelli EG, Cividini C, Agosta F. Resting State Dynamic Functional Connectivity in Neurodegenerative Conditions: A Review of Magnetic Resonance Imaging Findings. Front Neurol. 2019; 13. https://doi.org/10.3389/fneuro.2019.00657 PMID: 31281241

154. Shibuya K, Park SB, Geevasinga N, Menon P, Howells J, Simon NG, et al. Motor cortical function determines prognosis in sporadic ALS. Neurology. 2016; 87: 513–520. https://doi.org/10.1212/WNL.0000000000002912 PMID: 27402995

155. Keller J, Böhm S, Aho-Özhan HEA, Loose M, Gorges M, Kassubek J, et al. Functional reorganization during cognitive function tasks in patients with amyotrophic lateral sclerosis. Brain Imaging Behav. 2018; 12: 771–784. https://doi.org/10.1007/s11682-017-9738-3 PMID: 28600740