Deep learning predictions of survival based on MRI in amyotrophic lateral sclerosis

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Abstract

Amyotrophic lateral sclerosis (ALS) is a progressive neuromuscular disease, with large variation in survival between patients. Currently, it remains rather difficult to predict survival based on clinical parameters alone. Here, we set out to use clinical characteristics in combination with MRI data to predict survival of ALS patients using deep learning, a machine learning technique highly effective in a broad range of big-data analyses. A group of 135 ALS patients was included from whom high-resolution diffusion-weighted and T1-weighted images were acquired at the first visit to the outpatient clinic. Next, each of the patients was monitored carefully and survival time to death was recorded. Patients were labeled as short, medium or long survivors, based on their recorded time to death as measured from the time of disease onset. In the deep learning procedure, the total group of 135 patients was split into a training set for deep learning (n = 83 patients), a validation set (n = 20) and an independent evaluation set (n = 32) to evaluate the performance of the obtained deep learning networks. Deep learning based on clinical characteristics predicted survival category correctly in 68.8% of the cases. Deep learning based on MRI predicted 62.5% correctly using structural connectivity and 62.5% using brain morphology data. Notably, when we combined the three sources of information, deep learning prediction accuracy increased to 84.4%. Taken together, our findings show the added value of MRI with respect to predicting survival in ALS, demonstrating the advantage of deep learning in disease prognostication.

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predicting disease course, i.e. going beyond the establishment of patient-control identification, is presumably a more difficult problem, and arguably clinically more relevant. In the present study we therefore set out to explore the use of clinical characteristics in combination with MRI-based metrics of connectivity and brain morphology. White matter brain connectivity was derived from diffusion-weighted connectome imaging and brain morphology (i.e. cortical thickness and subcortical volume), was extracted from T1-weighted images. We used deep learning, a powerful technique shown to be of great value in many classification problems (Dean et al., 2012; Hinton et al., 2012; Krizhevsky et al., 2012; Mohamed et al., 2012; Wu et al., 2015), with Google’s visual search algorithm and Google’s AlphaGo program as well-known examples of applications of deep learning (Dean et al., 2012; Krizhevsky et al., 2012; Silver et al., 2016). Deep learning methods have shown high value in the fields of image classification (Krizhevsky et al., 2012; Wu et al., 2015), speech recognition (Hinton et al., 2012; Mohamed et al., 2012) as well as in elucidating complex relationships in MRI data (Plis et al., 2014).

Focusing on testing the predictive power of deep learning in differentiating between three survival duration subgroups (i.e. short, medium and long survivors), we assess the prediction accuracy of four deep learning networks which are based on 1. clinical data, 2. structural connectivity MRI data, 3. morphology MRI data and 4. a combined approach, in which clinical and imaging data are combined using layered deep learning. We show that the prediction accuracy of future survival time of ALS patients can be as high as 84% on the basis of combined clinical and neuroimaging data.

2. Materials and methods

2.1. Patients

A total dataset of 135 patients with sporadic ALS was included in this study (Table 1). Patients were diagnosed according to the EL Escorial criteria (Brooks et al., 2009) and recruited from the outpatient clinic for motor neuron diseases of the University Medical Center Utrecht. Parts of this dataset have been described in earlier publications in the context of examining patient-control group effects (Schmidt et al., 2014; Verstraete et al., 2011) (demographics given in Table 1). The included set involved data of patients that were either deceased (n = 122) or still alive with a disease duration of over 50 months (n = 13) at the time of analysis, providing the opportunity to test survival predictions using information from the first clinical MRI appointment.

The included clinical characteristics (Table 1) consisted of eight metrics. These comprised 1. site of disease onset, 2. age at disease onset and 3. time to diagnosis (i.e. the time from disease onset until the diagnosis of ALS was given). In addition, 4. the ALSFRS slope was included to provide an indication of disease progression based on the revised ALS Functional Rating Scale (ALSFsR-R) (Cedarbaum et al., 1999) and time T (in months) between symptom onset and first examination: slope = (48 − ALSFRS-R) / T (Kimura et al., 2006; Kollewe et al., 2008; Qureshi et al., 2006). Other clinical variables taken into account were 5. the forced vital capacity (FVC), 6. C9orf72 phenotype status, 7. frontotemporal dementia (FTD) status as derived from the (revised) Neary criteria (Neary et al., 1998; Rascovsky et al., 2011) and 8. EL Escorial criteria diagnostic category.

Among the group of 135 patients, there was no history of brain injury, psychiatric illness, epilepsy, or neurodegenerative diseases other than ALS. The Ethical Committee for human research of the University Medical Center Utrecht approved the study protocols and informed written consent according to the Declaration of Helsinki was obtained from each patient.

2.2. Short, medium, long survivors

Each of the 135 patients was categorized according to the true survival time (i.e. time between disease onset and death): short survivors...
with survival up to 25 months after disease onset, medium survivors with survival between 25 and 50 months after disease onset, and long survivors living over 50 months after disease onset (Elamin et al., 2015). The group of long survivors consisted of patients who either died after a disease duration of at least 50 months or were still alive and had a disease duration of at least 50 months at time of analysis.

2.3. Image acquisition and preprocessing

T1 and diffusion-weighted scans were acquired from all patients using a 3 Tesla Philips Achieva Medical Scanner with a SENSE receiver head-coil, described in detail by (Verstraete et al., 2011) and (Schmidt et al., 2014). A high-resolution T1-weighted image was acquired for anatomical reference by a 3D fast field echo using parallel imaging (TR/TE = 10/4.6 ms, flip-angle 8°, slice orientation: sagittal, voxel size = 0.80 × 0.75 × 0.75 mm, field of view = 176 × 240 × 240 mm covering the whole brain). For each subject, two sets of 30 weighted diffusion scans and 5 unweighted B0 scans were acquired with opposite k-space readouts (Andersson et al., 2003) using the following settings: parallel imaging SENSE p-reduction 3, high angular gradient set of 30 point on the cortical mantle (Fischl and Dale, 2000). Volumes of the regions, cortical thickness was measured by computing the distances sphere), 14 subcortical areas, and the brainstem). Of the 68 cortical regions, were parcellated using FreeSurfer (V5.1.0) according to the Desikan-Killiany atlas (Desikan et al., 2006) dividing the segmented gray matter into 83 distinct brain regions (68 cortical regions (34 for each hemisphere), 14 subcortical areas, and the brainstem). Of the 68 cortical regions, cortical thickness was measured by computing the distances between the gray/white matter boundary and pial surface at each point on the cortical mantle (Fischl and Dale, 2000); Volumes of the 14 subcortical areas and the brainstem were computed with FreeSurfer’s automated procedure for volumetric measurements (Fischl et al., 2002). Preprocessing of diffusion-weighted images included corrections for susceptibility and eddy-current distortions (Andersson and Skare, 2002; Andersson et al., 2003). Next, a tensor was fitted to the diffusion signals in each voxel and diffusion tensor imaging metrics, such as fractional anisotropy (FA) (Alexander et al., 2007), were derived. White matter tracts were reconstructed using Fiber Assignment by Continuous Tracking (FACT) (Mori et al., 1999; Mori et al., 2002; Mori and van Zijl, 2002); tracking was initiated by 8 seeds per white matter voxel and stopped using conditions as detailed in previous work (Schmidt et al., 2014). For each subject, an individual brain network was reconstructed by selecting the interconnecting tracts from the total cloud of reconstructed streamlines for each pair of regions included in the used cortical atlas (van den Heuvel and Sporns, 2011; Verstraete et al., 2011). We focused on white matter connectivity strength measured in terms of FA (from here on referred to as connection weight). In support of using this metric as a marker for disease effects, previous studies have extensively shown FA changes in ALS patients (Agosta et al., 2011; Ciccarelli et al., 2006; Menke et al., 2012; Schmidt et al., 2014; Senda et al., 2011; Turner et al., 2011; Verstraete et al., 2010; Verstraete et al., 2014). Moreover, the extent to which FA alterations are observed has been suggested to reflect distinct sequential ALS disease stages (Kassubek et al., 2014; Müller et al., 2016; Schmidt et al., 2015), and has been noted to mirror the pattern of phosphorylated 43 kDa TAR DNA-binding protein (pTDP-43, Brettschneider et al., 2013) aggregation. FA values of tracts interconnecting brain regions were stored in a weighted connectivity matrix.

2.4. Training, validation and test set

The total set of 135 datasets (i.e. 135 patients) was randomly divided into a training set (n = 83), a validation set (n = 20), and an independent evaluation set (n = 32), by first randomly selecting an evaluation set of the total dataset, and next by dividing the remaining subset of the data into a training set and a validation set, with the proportional splits between 70/30 and 80/20 (Crowther and Cox, 2005; Shahin et al., 2004). The training, validation and evaluation sets had similar survival class distributions (Fig. 1) and were not significantly different for each of the eight clinical characteristics (Table 1).

2.5. Deep learning neural network

A deep learning approach on the basis of an artificial neural network was applied. In brief (see for details on the applied deep learning procedures below), the procedure included deep learning on the training set, with the validation set used to stop the training process on time to prevent overfitting of the classifier to the training set (Duda et al., 2001). The evaluation set was then used to assess the final performance of the trained neural network on an independent sample.

For clarity, we note that the term ‘neural network’ here is taken from the field of supervised learning (Bishop, 1995; Duda et al., 2001; Hinton et al., 2012; Larochelle et al., 2009) and does not refer to the concept of brain network as used in the field of connectomics (Bullmore and Sporns, 2009; van den Heuvel et al., 2012; van den Heuvel et al., 2008). Deep learning networks refer to the subclass of neural networks comprising multiple hidden layers (Hinton et al., 2012) that allow for a detailed input-to-output mapping. Due to the inclusion of multiple hidden layers, deep learning networks are particularly useful for modeling high-level abstractions from data (Larochelle et al., 2009). In total, four deep learning networks were constructed (Fig. 2). These were based on 1. clinical data, 2. structural connectivity MRI data, 3. morphology MRI data and 4. a combination of the previous three information sources. The input vectors or features of these networks included the normalized clinical characteristics (8 in total), normalized cortical thickness and subcortical volumes derived from T1-weighted MRI and/or the connection weights as stored in the connectivity matrices. Normalization was performed using a min-max feature scaling in order to accelerate training (Priddy and Keller, 2005). Average values of features were imputed for missing values (i.e. either unknown clinical characteristics or connections that could not be detected in patients), with missing values being accounted for in an additional binary input vector. The output vectors, i.e. the classes that the deep learning

Fig. 1. Survival class distributions of the 135 subjects across the training, validation, and evaluation sets. The three survival categories (short, medium and long) are displayed for each data subset. The training set contained 83 patients and the validation set 20 patients. The evaluation set consisted of 32 patients. There are no significant differences in known survival class distributions and demographics between the three datasets (all p > 0.05, see Table 1 for more details).
network had to classify, represented the predefined classes of short, medium and long survivors. In what follows, we will describe the construction of the four deep learning networks, their training and their evaluation.

2.6. Deep learning network construction

1. Clinical deep learning
First, a deep learning network was constructed for the clinical characteristics (Fig. 2). The input vector of this network represented the eight clinical characteristics, including site of onset, age at onset, time to diagnosis, ALSFRS slope, FVC, C9orf72 phenotype status, FTD status and El Escorial criteria diagnostic category. Each layer in the deep learning network consisted of nodes and was connected to other layers using weighted edges (Duda et al., 2001). The number of input nodes was based on the number of clinical characteristics and the number of output nodes was based on the number of classes (short, medium, long). The number of hidden layers was set to two, in order to balance between the benefit of discovering more complex relations, the risk of overfitting, and training time and complexity (Karsoliya, 2012). The number of hidden nodes was set by means of a fine neuron grid search during the training phase (described below), with the number of hidden nodes in both layers varying from 1 to 500.

2. Structural connectivity deep learning
Second, a deep learning network was constructed for the structural connectivity MRI metric (Fig. 2). This second network employed connection weight of each reconstructed connection as an input node for the deep learning (2285 features in total: 83 brain regions × 82 / 2 = 3403 possible connections of which 2285 are existing connections). The number of output nodes was set to three survival classes. Two hidden layers were used and the sizes of these layers were found using a fine neuron grid search in the same search domain as previously described for structural connectivity deep learning.

3. Morphology deep learning
Next, a deep learning network was constructed based on the cortical thickness and subcortical volume measurements (Fig. 2). The input vector of the morphology network included 68 cortical thickness values and volume values of 14 subcortical regions and the brainstem, resulting in a total of 83 input nodes. The number of output nodes was set to three survival classes and two hidden layers were used. The size of these layers was set by means of a fine neuron grid search in the same range as for the other networks during the training phase. Output nodes (three nodes) represented survival class.

2.7. Network training
All four deep learning networks were trained using the following procedures. Each hidden node was assigned a non-linear (logistic) activation function (Hjelm et al., 2014). Feeding each training data point through the network produced the output vector of weights. This output vector was compared to the target values, with any difference in the predicted outcome and the real outcome (i.e. short, medium, long survivors) defined as error using the cross-entropy error function (Murphy, 2012; Rubinstein and Kroese, 2004). After presenting all examples of the training set to the network, the network weights were updated by backpropagation learning (Rumelhart et al., 1986) using the scaled conjugate gradient algorithm (Møller, 1993) to correct the weights in a direction that reduced the error of the network. During this training phase, overfitting of the network was prevented by the use of L2-regularization technique (i.e. penalizing by adding the sum of the squared values of the weights to the error function) with parameter $\lambda = 0.1$ (Ng, 2004) and performance comparison against the validation set. The validation data was presented to the network after
each training iteration to obtain a non-training performance error. Training was stopped when the validation error ceased to decrease (Sarle, 1995). This training procedure was repeated for all networks constructed in the neuron grid search; performance of these networks was evaluated using the measures described below, and the optimal deep learning network size was selected.

2.8. Network evaluation

After the training stage, performance of the obtained neural network was assessed in the evaluation phase by means of the evaluation dataset. The input features (i.e. clinical characteristics (network 1), connectivity matrices (network 2), morphology values (network 3) or all three information sources (network 4)) of the subjects in the evaluation set (n = 32) were presented to the trained networks. The softmax activation function (Bishop, 1995) was used for the output nodes, resulting in a vector of values varying between 0 and 1 that add up to 1 and the output node with the highest probability was selected as the predicted class label using a winner-take-all approach (Duan et al., 2003; Hinton, 2002; Hinton et al., 2012; Lefebvre et al., 2013). To determine whether the predicted class was correct, the network output label was compared to the true class label (i.e. true survival class of a patient). Good classifications were marked by equal labels for the true and predicted class, incorrect classifications were marked by a mismatch between prediction and truth. Next, a mosaic plot of the trained network (Fig. 3) was computed to visualize the distribution of datasets over the different classes after prediction (Hartigan and Kleiner, 1981), including the positive predictive values (PPV) of the network, defined as the percentage of patients with a predicted label that coincided with the true class label (Altman and Bland, 1994; Fletcher and Fletcher, 2005). For each class a PPV score was computed as $PPV_i = \frac{N_{correct}}{N_{label}}$ (i ∈ {short, medium, long}), with $N_{correct}$ being the number of patients that correctly received class label i and $N_{label}$ the total number of patients predicted to have class label i. PPVs were computed to give an impression of the discriminative power and thus predictive value of the network.

In addition, the overall performance of a network was assessed as the overall accuracy of the predictions, denoting the percentage of patients for whom the network predicted the correct class label and was calculated using the formula $N_{correct} / N$. Here, $N_{correct}$ denoted the number of patients for whom the prediction by the deep learning network was equal to the true survival class (i.e. highlighted diagonal elements in mosaic plot), and $N$ denoted the total number of patients included in the set.

3. Results

3.1. Clinical deep learning

The clinical network was trained on eight clinical characteristics (site of onset, age at onset, time to diagnosis, ALSFRS slope, FVC, C9orf72 phenotype status, FTD status and El Escorial criteria diagnostic category). Neuron grid search resulted in a deep learning network with

![Fig. 3. The distribution of the prediction results shown in mosaic plots. The columns represent the known survival class of patients, where the width of columns is relative to the number of subjects in that column. The colors orange, gray and blue represent the predicted survival classes short, medium and long, respectively. The highlighted diagonal cells (bottom left to top right) denote the number of patients that were correctly classified and the off-diagonal cells the number of patients that were mispredicted (opaque cells), i.e. the wrong class was predicted by the network. The positive predictive value (Fletcher and Fletcher, 2005) for each predicted class can be derived by dividing the correctly predicted subjects by the total number of predictions of that class (i.e. highlighted cell/sum of same colored cells). The overall accuracy is computed by summing the highlighted diagonal cells and dividing this number by the total population. For the evaluation set (n = 32), the clinical deep learning network, the structural connectivity deep learning network, morphology deep learning network and the clinical-MRI combined deep learning network obtained overall accuracies of 68.8%, 62.5%, 62.5% and 78.1%, respectively. (Box) A perfect classification and a random classification mosaic plot are displayed for comparison of the results obtained on the evaluation set. In the perfect classification, all subjects are correctly classified and therefore an accuracy of 100% is achieved. In a random classification, three class labels are randomly distributed over the subjects, only predicting a third of the total subject population correctly. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)](image-url)
8 input nodes, 158 nodes in the first hidden layer, 448 nodes in the second hidden layer, and 3 output nodes. The predicted short survivors had a PPV of 72.7% on the evaluation set; medium survivors had a PPV of 64.3% and long survivors 71.4%. The highest PPV was obtained for the predicted short survivors, indicating that a predicted short survival on the basis of clinical metrics was more often correct than a predicted medium or long survival. The optimal network based on clinical characteristics gave an evaluation accuracy of 68.8% (Fig. S1), a training accuracy of 78.3% (Fig. S1) and a validation accuracy of 70.0% (Fig. S2).

3.2. Structural connectivity deep learning

The structural connectivity MRI network, based on connection weights, consisted of 2285 input nodes (the total number of reconstructed tracts), and was fitted 134 and 313 nodes in the first and second hidden layer respectively, and comprised 3 output nodes. The PPV scores for the predicted short, medium and long survivors of the evaluation set were 62.5%, 57.1% and 100.0%, respectively. The highest PPV in this network was obtained for the long survivor class, indicating that this structural connectivity-based deep learning network was highly reliable when it gave predictions for the class of long survivors. The structural connectivity deep learning network reached an evaluation prediction accuracy of 62.5% (Fig. 3), a training accuracy of 79.5% (Fig. S1) and a validation accuracy of 60.0% (Fig. S2). With average simulated PPV chance levels of 38.6% (short), 38.6% (medium) and 23.0% (long) when assigning random class labels to patients, these findings show that objective connectivity values alone can provide valuable information on disease survival.

3.3. Morphology deep learning

The morphology MRI deep learning network consisted of 83 input nodes (68 cortical thickness values and 15 subcortical volumes), and was fitted to 181 and 178 nodes in the first and second hidden layer respectively, and contained included the three survival classes as output nodes. The PPV scores of the evaluations set were 64.3% (short), 61.5% (medium) and 60.0% (long survival), respectively. The highest PPV in this network was obtained for the short survivor class, indicating that the morphology network, similar to the clinical network, was more reliable when it predicted a short survivor. The morphology deep learning network reached an evaluation prediction accuracy of 62.5% (Fig. 3), a training accuracy of 80.7% (Fig. S1) and a validation accuracy of 60.0% (Fig. S2). These findings support the relation between morphology and disease progression, and thus disease survival.

3.4. Clinical-MRI combined deep learning

Next, the prediction probabilities from the clinical deep learning network and the prediction probabilities from the two MRI networks were presented as input for a combined deep learning network. Grid search during network training resulted in a network configuration with 9 input nodes, 171 nodes in the first hidden layer, 108 nodes in the second hidden layer, and 3 output nodes. PPV scores of the combined network on the evaluation set included 90.9%, 83.3% and 77.8% for the predicted short, medium and long survivor classes, respectively. The combined network reached an evaluation accuracy of 84.4% (Fig. 3), a training accuracy of 88.0% (Fig. S1) and a validation accuracy of 80.0% (Fig. S2). Statistical testing indicated that survival prediction was significantly improved (p < 0.001) due to the addition of structural connectivity data and morphology MRI findings to the clinical characteristics (Fig. 4, see Supplementary materials for details).

4. Discussion

We evaluated the use of deep learning to predict survival time of ALS patients on the basis of clinical characteristics and advanced MRI metrics. Our findings show that MRI data alone (i.e. structural connectivity and brain morphology data, consisting of cortical thickness and subcortical volumes) can provide valuable predictions of survival time. Furthermore, combining clinical characteristics and MRI data into a layered deep learning approach can further improve predictions about whether a patient will have a short, medium or long survival time.

Previous studies have used proportional Cox hazard classification (Cox, 1972) employing clinical characteristics such as site of onset, executive dysfunction and diagnostic delay (Elamin et al., 2015; Scotton et al., 2012) to develop prognostic models for survival. These models already showed the predictive power of clinical data, with PPVs and overall accuracies lower than or similar to results in our study, indicating a potentially better predictive power of survival classes using deep learning. It should however be noted that PPV scores are dependent on the prevalence of a subtype in the total population and might also influence the differences in scores.

Deep learning on diffusion-weighted imaging data led to a prediction accuracy of 62.5%. Deep learning on T1-weighted images data resulted in a prediction accuracy of 62.5%. A combination of these imaging metrics yielded an improved prediction accuracy of 78.1% (see Supplementary materials for more details), indicating the predictive power of combining imaging metrics in deep learning. Previous studies have used MRI data for the prediction of diagnosis; that is, they used structural connectivity MRI data to differentiate between ALS patients and healthy controls, resulting in prediction accuracies between 70 and 80% (Fekete et al., 2013; Welsh et al., 2013). Other studies used cortical thickness measurements to discriminate between patients and controls in various diseases, such as Alzheimer’s disease (Lerch et al., 2008), childhood onset schizophrenia (Greenstein et al., 2012), and major depression (Foland-Ross et al., 2015) with prediction accuracies ranging from 60 to 75%. In our study we examined a presumably more difficult task of predicting survival time within the group of patients, with a priori chance levels (i.e. true positive rate) here equal to 33.3% for the three survival classes, rather than 50% for patient/control status. The potential of MRI in patient classification and prognostication was previously also shown for the prediction of disease status in Alzheimer’s disease, where machine learning differentiated between two subtypes of dementia based on T1-weighted images with accuracies of 89%.

![Fig. 4. Fitted normal curves on accuracy distributions of the four networks. Normal curves are fitted on the accuracy distributions of the clinical deep learning network (blue), the structural connectivity deep learning network (orange), morphology deep learning network (purple) and the clinical-MRI combined deep learning network (gray), based on 16 randomly selected subjects (repeated 10,000 times) from the first evaluation set (n = 32). The mean accuracies (dashed lines) of these distributions were 68.7%, 62.5%, 62.4% and 84.4% for the clinical deep learning network, structural connectivity deep learning network, morphology deep learning network and clinical-MRI combined deep learning network, respectively. A paired t-test showed significant differences between accuracies of each pair of networks (all p < 0.001). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)](image-url)
In this study we included all reconstructed connections instead of focusing on connections between specific brain regions, for example from motor regions to other brain areas (Schmidt et al., 2014). By considering all connections, a deep learning method is allowed to identify combinations of affected connections that are most valuable for survival prediction. As such, the deep learning network may detect relevant patterns in connections that are only slightly affected, thereby adding valuable information for prediction. The ability of deep learning to distill complex relationships from large datasets makes it a promising tool for disease prognostication.

The predictions of the network based on clinical parameters and the two MRI networks were often predicted correctly by the combination network using prediction information from the other two networks. By utilizing the predicting probabilities of the survival classes instead of the survival class label, the uncertainty of predictions was taken into account. The prediction probabilities of the clinical, structural connectivity MRI and morphometry MRI networks contributed equally to the combined prediction.

Deep learning shows promising results for the prediction of survival categories for individual ALS patients, but several points have to be taken into account. Large training and evaluation sets are preferred to ensure convergence of prediction accuracies and to prevent overfitting (Ng, 2004). In addition, external validation is crucial for the development of a reliable prognostic tool and should be incorporated in future examinations. Second, while deep learning can effectively make predictions on datasets with complex relationships, dependency among input variables and between input and output variables cannot be easily deduced. In future research, it would therefore be worthwhile to investigate possibilities to reveal these dependencies and gain more insight into the mechanisms underlying disease progression. Third, prediction may also be improved using additional deep learning networks based on fMRI scans, as used in previous studies investigating disease diagnosis (Fekete et al., 2013; Welsh et al., 2013). Finally, additional clinical characteristics or diffusion tensor imaging metrics may also improve prediction. For example, radial diffusivity differences have been shown between ALS patients and healthy controls (Agosta et al., 2010; Metwalli et al., 2010) and therefore might also be of value in survival prognostication.

Deep learning is a powerful approach with successful applications in many real world issues. Here, we show that deep learning can also be of benefit to medical problems. Our findings show that deep learning can contribute to early prognostication of survival in ALS by combining clinical characteristics and brain imaging data. Our study provides promising results and may contribute to developing an automated prognostic tool for the estimation of survival in individual patients. Supplementary data to this article can be found online at http://dx.doi.org/10.1016/j.nicl.2016.10.008.

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