Estimated connectivity networks outperform observed connectivity networks when classifying people with multiple sclerosis into disability groups

Ceren Tozlu a, Keith Jamison a, Zijin Gu b, Susan A. Gauthier a, c, d, Amy Kuceyeski a, d, *

a Department of Radiology, Weill Cornell Medicine, New York, NY, USA
b Electrical and Computer Engineering Department, Cornell University, Ithaca 14850, USA
c Department of Neurology, Weill Cornell Medicine, New York, NY, USA
d Brain and Mind Research Institute, Weill Cornell Medicine, New York, NY, USA

A B S T R A C T

Background: Multiple Sclerosis (MS), a neurodegenerative and neuroinflammatory disease, causing lesions that disrupt the brain’s anatomical and physiological connectivity networks, resulting in cognitive, visual and/or motor disabilities. Advanced imaging techniques like diffusion and functional MRI allow measurement of the brain’s structural connectivity (SC) and functional connectivity (FC) networks, and can enable a better understanding of how their disruptions cause disability in people with MS (pwMS). However, advanced MRI techniques are used mainly for research purposes as they are expensive, time-consuming and require high-level expertise to acquire and process. As an alternative, the Network Modification (NeMo) Tool can be used to estimate SC and FC using lesion masks derived from pwMS and a reference set of controls’ connectivity networks.

Objective: Here, we test the hypothesis that estimated SC and FC (eSC and eFC) from the NeMo Tool, based on an individual’s lesion masks, can be used to classify pwMS into disability categories just as well as SC and FC extracted from advanced MRI directly in pwMS. We also aim to find the connections most important for differentiating between no disability vs evidence of disability groups.

Materials and Methods: One hundred pwMS (age: 45.5 ± 11.4 years, 66% female, disease duration: 12.97 ± 8.07 years) were included in this study. Expanded Disability Status Scale (EDSS) was used to assess disability, 67 pwMS had no disability (EDSS < 2). Observed SC and FC were extracted from diffusion and functional MRI directly in pwMS, respectively. The NeMo Tool was used to estimate the remaining structural connectome (eSC), by removing streamlines in a reference set of tractograms that intersected the lesion mask. The NeMo Tool’s eSC was used then as input to a deep neural network to estimate the corresponding FC (eFC). Logistic regression with ridge regularization was used to classify pwMS into disability categories (no disability vs evidence of disability), based on demographics/clinical information (sex, age, race, disease duration, clinical phenotype, and spinal lesion burden) and either pairwise entries or regional summaries from one of the following matrices: SC, FC, eSC, and eFC. The area under the ROC curve (AUC) was used to assess the classification performance. Both univariate statistics and parameter coefficients from the classification models were used to identify features important to differentiating between the groups.

Results: The regional eSC and eFC models outperformed their observed FC and SC counterparts (p-value < 0.05), while the pairwise eSC and SC performed similarly (p = 0.10). Regional eSC and eFC models had higher AUC (0.66–0.68) than the pairwise models (0.60–0.65), with regional eFC having highest classification accuracy across all models. Ridge regression coefficients for the regional eFC and regional observed FC models were significantly correlated (Pearson’s r = 0.52, p-value < 10e-7). Decreased estimated SC node strength in default mode and ventral attention networks and increased eFC node strength in visual networks was associated with evidence of disability.

Abbreviations: AUC, area under curve; EDSS, extended disability status score; eFC, estimated functional connectivity; eSC, estimated structural connectivity; FC, functional connectivity; GM, gray matter; HC, healthy control; QSM, quantitative susceptibility imaging; LST, lesion segmentation tool; MS, multiple sclerosis; NeMo, network modification; pwMS, people with multiple sclerosis; SC, structural connectivity; WM, white matter.

E-mail address: amk2012@med.cornell.edu (A. Kuceyeski).

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1. Introduction

Multiple sclerosis (MS) is a disease that is largely characterized by brain lesions, detected using clinical MRI, that can cause disability. The number and size of lesions, however, is not always proportional to an individual’s disability ( Barkhof, 2002); recent work has demonstrated that lesions’ disruption to structural and physiological connectivity networks may be more important for accurate lesion-dysfunction mapping (Kuceyeski et al., 2018; A. F. Kuceyeski et al., 2015b). Connectomics, or brain connectivity network analysis, provides a promising tool to directly measure the effect of MS-related pathology and to capture the reorganization mechanisms in response to pathology. Having a more accurate lesion-dysfunction map, particularly a lesion-future dysfunction map, would allow development of more accurate prognoses and could be used to personalize treatment plans to improve clinical care of people with MS (pwMS).

Advanced neuroimaging techniques such as diffusion magnetic resonance imaging (dMRI) and functional MRI (fMRI) are commonly used to quantify the structural (white matter) connectome (SC) and the functional (co-activation) connectome (FC). Previous studies have shown that SC metrics can successfully differentiate healthy controls (HC) from pwMS, classify pwMS who showed different clinical phenotypes, and be used to assess its relationship with disability (Li et al., 2013; Muthuraman et al., 2016; Shu et al., 2011). However, tractography can be challenging in areas of complex fiber orientations and can produce false-positive and false-negative connections. Lesions, pathologies in normal-appearing white matter, inflammation, demyelination and axonal loss may decrease SNR in dMRI and make existing tractography issues even more pervasive in pwMS (Kuceyeski et al., 2011; Lipp et al., 2020; Pagani et al., 2007). In terms of measuring FC in pwMS, inflammation induces alterations in synaptic transmission (Rossi et al., 2012; Stampanoni Bassi et al., 2017) and thus may affect the BOLD signal in non-trivial ways (Bassi et al., 2017; Tomassini et al., 2016). FC changes in pwMS relative to controls may also be driven by compensation (Tommasini et al., 2018), maladaptation (Cruz-Gomez et al., 2014) or both (Tona et al., 2014), making the mapping of FC to dysfunction even more complicated. Therefore, observed SC and FC extracted from advanced MRI in pwMS may not be accurately representing underlying brain networks and/or be informative of disability.

In addition to the above outlined challenges using advanced MRI in pwMS, the imaging techniques themselves are expensive, time-consuming and require a high level of expertise. Therefore, alternative techniques that use only conventional, clinically-acquired MRI may allow more widespread use of SC and FC measures for understanding the mechanisms and connectome-behavior mapping in pwMS (Poulon et al., 2018; Griffis et al., 2021). As an alternative to performing tractography in pwMS, lesion masks extracted from conventional MRI can be used to calculate estimated structural connectivity (eSC) via the Network Modification (NeMo) Tool (Kuceyeski et al., 2013). The recently updated NeMo Tool version 2.0 (https://kuceyeski-wcm-web.s3.us-east-1.amazonaws.com/upload.html) uses tractography results from 420 unrelated Human Connectome Project controls as a reference; a lesion mask from a pwMS is used to remove “damaged” streamlines that pass through the lesions from the reference tractographies and the remaining streamlines are used to obtain eSC. A recent work introduced a deep neural network trained to predict FC from SC (Sarwar et al., 2021). Using their framework, we created a neural network trained on the NeMo Tool’s 420 control reference subjects to predict FC from SC. This neural network was then integrated into the NeMo Tool so that it could take its eSC and produce the corresponding estimated FC (eFC), a functionality which as far as we know is not available in other tools that estimate structural dysconnectivity (Foulon et al., 2018; Griffis et al., 2021). To summarize, the NeMo Tool version 2.0 uses only an individual’s lesion mask to produce estimates of their SC and FC networks.

Observed connectivity from advanced imaging and estimated disruptions in connectivity based on conventional imaging have been independently associated with disability and cognitive impairment in pwMS (Has Silemek et al., 2020; Kuceyeski et al., 2018; Pagani et al., 2019; Tommasin et al., 2018). A recent study (Salvalaggio et al., 2020) showed structural disconnectivity due to stroke lesions is a better predictor than indirect functional connectivity, estimated using symptom-lesion network mapping (Boes et al., 2015), in predicting multiple functional deficits in stroke. Moreover, the same study showed observed FC from fMRI was superior than indirect functional disconnectivity in the subset of the stroke patients. However, no study has yet compared the ability of observed and estimated SC/FC metrics in classifying pwMS into disability categories.

In this paper, models based on eSC and eFC derived from clinical MRI-derived lesion masks and the NeMo Tool were compared with models based on SC and FC extracted directly from advanced MRI in their ability to classify pwMS into disability groups. Specifically, the classification performance of models based on pairwise and regional SC, FC, eSC, and eFC metrics were compared. We hypothesized that models based on eSC and eFC metrics would perform at least as well as the SC and FC metrics from advanced imaging. Second, the most important pairwise and region-wise connections in the classification task were quantified.

2. Materials and methods

2.1. Subjects

One hundred pwMS (age: 45.5 ± 11.4, 66% females) with a diagnosis of clinically isolated syndrome (CIS) or MS (7 CIS, 88 relapsing-remitting, 5 primary or secondary progressive MS) were enrolled in our study. Possible participants were excluded if they had contraindications to MRI. Demographic data was collected (age, sex, race, clinical phenotype, and disease duration) and subjects underwent an MRI scan. Extended Disability Status Score (EDSS) was used to quantify disability, where an EDSS < 2 was considered no disability and EDSS ≥ 2 considered evidence of disability. This threshold was defined based on the fact that pwMS having EDSS values of 0–1.5 present some abnormal signs/symptoms but do not have functional disability. The spinal cord lesion number was included in the patients’ clinical radiology report and this was used to estimate spinal cord burden in patients. All studies were approved by an ethical standards committee on human experimentation, and written informed consent was obtained from all patients.

2.2. Image acquisition, processing, and connectome extraction

MRI data were acquired on a 3 T Siemens Skyra scanner (Siemens, Erlangen, Germany) with a 20-channel head-neck coil and a 32-channel spine-array coil. Anatomical MRI (T1/T2/T2-FLAIR, 1 mm3 iso-voxel), resting-state fMRI (6 min, TR = 2.3 s, 3.75 × 3.75 × 4 mm voxels) and diffusion MRI (55 directions HARDI, b = 800, 1.8 × 1.8 × 2.5 mm voxels) acquisitions were performed. Sagittal STIR images were
acquired for identification of spinal lesions (TR = 3.5 s, TI = 220 ms, TE = 45 ms, in-plane resolution 0.43 mm, FOV = 22 mm, slice thickness 3 mm). Multi-echo 2D GRE field maps were collected for use with both fMRI and diffusion MRI (0.75 × 0.75 × 2 mm voxels, TE1 = 6.69 ms, ΔTE = 4.06 ms, number of TEI = 6).

### 2.2.1. Structural and functional connectivity extraction

White matter (WM) and gray matter (GM) were segmented and GM further parcellated into 86 regions of interest (68 cortical and 18 subcortical/cerebellar) using FreeSurfer (Fischl and Dale, 2000). The white and gray matter surfaces were checked for each subject on FreeSurfer and hand-edited with control points and reconstruction editing if necessary. As described elsewhere (Kuceyeski et al., 2016), fMRI pre-processing included simultaneous nuisance regression and removal of WM and cerebrospinal fluid (CSF) effects (Hallquist et al., 2013), followed by band-pass filtering (0.008–0.09 Hz) using the CONN v18b toolbox (Whitfield-Gabrieli and Nieto-Castanon, 2012) and SPM12 in Matlab. Nuisance regressors included 24 motion parameters (6 rotation and translation, temporal derivatives, and squared version of each) and the top 5 eigenvectors from eroded masks of both WM and CSF. The mean fMRI signal over all voxels in a region was calculated and the mean regional time series correlated (Pearson’s correlation) between every pair of regions to obtain pairwise FC matrices. Regional FC node strengths were calculated by taking the sum of the columns in the FC matrix after removing the negative entries.

Diffusion MRI was interpolated to isotropic 1.8 mm voxels, and then corrected for eddy current, motion and EPI-distortion with the eddy command from FSL 5.0.11 (Andersson and Sotiropoulos, 2016) using the outlier detection and replacement option (Andersson et al., 2016). MRtrix3Tissue (https://3Tissue.github.io), a fork of MRtrix3 (Tournier et al., 2019) was used to estimate a voxel-wise single-shell, 3-tissue constrained spherical deconvolution model (SSST-3D) and then compute whole-brain tractography for each subject. The tractography seed mask was generated using MRtrix to estimate gray matter and white-matter volume fractions from local diffusion properties, and thresholding to include all voxels with a gray + white volume fraction ≤ 0.05, essentially including all non-CSF voxels within the brain volume, including lesion voxels. We seeded 92 streamline randomly within in each mask voxel, and performed deterministic (sd-stream) tractography with MRtrix3 (Tournier et al., 2010; Tournier et al., 2012) with a fiber-orientation distribution (FOD) amplitude cutoff of 0.05, leading to approximately 5 million streamlines per subject. We then applied the SIFT2 global filtering algorithm (Smith et al., 2015) to account for bias that exists in greedy, locally-optimal streamline reconstruction algorithms. The SC matrix was constructed by taking the sum of the SIFT2 weights of streamlines connecting pairs of regions and then dividing by the sum of the two regions’ volumes. In addition to the pairwise SC measures, regional SC node strength was quantified by taking the sum of each of the columns in the SC matrix.

#### 2.2.2. Lesion mask creation

The WM hyperintensity lesion masks were created by running the T2 FLAIR images through the Lesion Segmentation Tool (LST) and were further hand-edited as needed. T2 FLAIR-based lesion masks were transformed to the individual’s T1 native space using the inverse of the T1 to GRE transform and nearest-neighbor interpolation. Individual T1 images were then normalized to MNI space using FSL’s linear (FLIRT) and non-linear (FMRIT) transformation tools (http://www.fmrib.ox.ac.uk/fsl/index.html); transformations with nearest-neighbor interpolation were then applied to transform the native anatomical space T2FLAIR lesion masks to MNI space. The transformations were concatenated (T2FLAIR to T1 to MNI) to minimize interpolation effects. Lesions were manually inspected after the transformation to MNI space to verify the accuracy of coregistration.

#### 2.2.3. Network Modification Tool

The NeMo Tool (Kuceyeski et al., 2013) used each subject’s MNI-space T2 FLAIR lesion mask to calculate eSC and eFC using a reference database of SCs and FCs from 420 unrelated controls from the Human Connectome Project (see Fig. 1 for the workflow and supplementary material for details). The NeMo Tool’s original functionality was to estimate SC (or structural disconnectivity) due to a lesion mask by identifying streamlines from each healthy control tractography set that pass through the lesion mask, and then recording the gray matter regions they connect. The eSC matrix is then the sum of the SIFT2 weights of the remaining streamlines connecting pairs of regions, divided by the sum of the two regions’ volumes averaged over the 420 individuals in the reference set. Regional eSC node degree was taken to be the sum of the columns in the eSC matrix.

In addition, we extended the NeMo Tool’s functionality to create eFC from lesion masks by first creating a deep neural network that can predict FC from SC using a previously published approach (Sarwar et al., 2021). We used the NeMo Tool’s 420 subjects to train, validate and test the model, which was optimized to minimize the difference between predicted and observed FC but also minimize the l1 distance between inter-subject similarity of the predicted FC and observed FC across the individuals in the training set. The model, which takes in the upper triangular part of the SC matrix, is a feed-forward fully connected neural network comprised 8 hidden layers with 1024 neurons and a dropout rate of 0.5 for each layer. We used leaky rectified linear unit (ReLU, leakage parameter set to 0.2) and hyperbolic tangent (tanh) as the activation functions and initialized the weights using the Xavier algorithm. Adam was used as our optimizer. The hyperparameters in the loss function include λ, the regularization parameter, and γ, the parameter that modeled averaged the inter-subject correlation of the observed FC of each individual in the training set, were determined by grid search in the interval of [0.1, 0.7] with steps of 0.1 for γ, and across values of [0.0001, 0.0005, 0.001, 0.005, 0.01, 0.05, 0.1, 0.5, 1] for λ. The best value for these parameters was found to be γ = 0.2 and λ = 0.01. We trained the model using 340 individuals’ FCs and SCs (out of 420 total) for 20,000 epochs with batch size n = 10 and learning rate α = 1e-5. The final model was tested on the remaining 80 individuals, see Supplemental Fig. 1 for a violin plot showing the correlation between predicted and true FC and correlations between inter-subject predicted FC, which were similar to the original publication’s results (Sarwar et al., 2021). Once trained and validated on the NeMo Tool’s controls, the neural network was then applied to the eSCs from each of the individual lesion masks to produce an eFC matrix for each pwMS. Regional eFC node strengths were calculated by taking the sum of each of the columns in the eFC matrix after removing the negative entries.

#### 2.3. Mass univariate analysis

First, demographics and clinical variables were tested for differences between the no disability vs evidence of disability groups using the Chi-square test for qualitative variables and Wilcoxon rank-sum test for quantitative variables. Second, the pairwise and regional SC, FC, eSC, and eFC were tested for differences between the two disability groups using the Wilcoxon rank-sum test. Differences were considered significant when p < 0.05 after Bonferroni-Hochberg (BH) correction for multiple comparisons (Benjamini and Hochberg, 1995). Entries in the eSC and SC were only considered in the univariate tests if they were non-zero for a majority of pwMS. All statistical analyses were performed and graphs created using R version 3.4.4 and Matlab version R2020a.

#### 2.4. Classification analysis

Logistic regression with ridge regularization was used to classify pwMS into disability categories based on demographics, clinical information (sex, age, race, disease duration, clinical phenotype, and spinal lesion burden) and one of the pairwise or regional connectivity metrics
derived from advanced MRI (SC and FC) or estimated from clinical imaging and the NeMo Tool (eSC and eFC). All pairwise connections and regional information were included in the pairwise and regional models, respectively. The number of the variables was 3655 (upper-triangle of the 86x86 connectivity matrix) for the pairwise models and 86 for the regional models.

The models were trained with outer and inner loops of k-fold cross-validation (k = 5) to optimize the hyperparameters and test model performance. The folds for both inner and outer loops were stratified to ensure that each fold contained the same proportion of subjects in the two classes as the original dataset. The inner loop (repeated over 5 different partitions of the training dataset only) optimized the set of hyperparameters that maximized validation AUC. A model was then fitted using the entire training dataset and those optimal hyperparameters, which was then assessed on the hold-out test set from the outer loop. The outer loop was repeated for 100 different random partitions of the data (See Fig. 4 in the supplementary document). The median of AUC (overall 5 folds × 100 iterations = 500 test sets) was calculated to assess the performance of the models. In addition to the AUC values; confusion matrices, brier scores, sensitivity, specificity, and balanced accuracy are provided. The classification performances of different models were compared using permutation test (David, 2008) of 1000 permutations. P-values were the number of permutations that had a difference in means bigger than the original difference and were considered significant when p < 0.05 after BH correction for multiple comparisons (Benjamini and Hochberg, 1995).

The important variables were identified using both univariate statistics (t-test or Wilcoxon rank sum test based on the connectivity type) and the mean feature weights (the beta parameter coefficients) over all 500 models (100 partitions of the data into 5 folds) (Tian et al., 2021). Regional and region-pair feature weights were also summarized at a functional network level by assigning each of the 68 cortical regions to one of seven canonical functional networks (Yeo et al., 2011); subcortex and cerebellum were also added as networks. Pearson’s correlation coefficient was used to assess similarity of the feature weights from the four estimated and observed connectivity model pairs. Finally, as a sensitivity analysis, we repeated the classification analysis with EDSS of 3 as the threshold for defining disability groups (88 pwMS who had not disability vs 12 pwMS).

Fig. 1. Workflow of the study. (A) Observed SC and FC were extracted from dMRI and dMRI directly in pwMS. The eSC and eFC matrices were computed using the lesion masks extracted from T2 FLAIR images and the Network Modification (NeMo) Tool. The eSC matrix was estimated by overlaying the lesion mask on a reference set of tractography results and removing streamlines that passed through the lesion mask. A deep neural network trained to predict FC from SC using the NeMo Tool’s control data (see tan box) was used to predict eFC from eSC.
who had evidence of disability). The demographics, clinical and imaging variables were given in the supplementary document (please see Supplementary Table 2) for the groups created with EDSS 3.

When the data contains class imbalance, models tend to favor the majority class. Due to the class imbalance in our data (67 vs 33 pwMS with no disability vs evidence of disability), the over-sampling approach Synthetic Majority Over-sampling Technique (SMOTE) (Chawla et al., 2002) was used to obtain a balanced training dataset during cross-validation. SMOTE compensates for imbalanced classes by creating synthetic examples using nearest neighbor information and has been shown to be among the most robust and accurate methods with which to control for imbalanced data (Santos et al., 2018).

**Data/code availability statement:** The codes that were used in this study are publicly available. Please see

1) https://github.com/cerent/MS-Estimated-Observed-SC-FC for the classification and statistical analyses as well as the violin plots,
2) https://github.com/zijin-gu/sc2fc for the codes of the deep neural network framework for structural and functional connectivity mapping, and
3) https://github.com/kjamison/connviewer for the circle and glass brain plots.

The Network Modification tool that can be used to upload the lesion masks to compute the estimated structural connectivity can be reached via: https://kuceyeski-wcm-web.s3.us-east-1.amazonaws.com/upload.html

The deidentified data (the connectivity matrices) that support the findings of this study are available upon reasonable request from the corresponding author. Data sharing agreement and approval from our ethics committee are needed to share the connectivity matrices.

**Ethics statement:** All studies were approved by an ethical standards committee on human experimentation, and written informed consent was obtained from all patients.

### 3. Results

#### 3.1. Patient characteristics

The 67 pwMS who did not have disability were significantly younger than those 33 pwMS having disability (corrected p < 0.05) and had a trend for shorter disease duration (corrected p < 0.10) (See Table 1). The pwMS with progressive diagnoses were in the disability group, while those with CIS were in the no disability group. The population average lesion mask showed that lesions tended to occur most commonly in periventricular white matter, see Supplementary Fig. 2.

#### 3.2. Comparison of connectivity between disability groups

There was no significant difference in the pairwise or regional SC, FC, eSC or eFC, or between the regional eFC between the no disability vs evidence of disability groups. However, the pairwise eFC between the right rostral anterior cingulate and right temporal pole was significantly greater in pwMS who had no disability compared to those with evidence of disability (corrected p-value = 0.04). Figs. 2 and 3 illustrate the p-values (before correction) of group differences in pairwise and regional connectivity metrics between two disability groups, respectively. FC between ventral attention and frontoparietal and between cerebellum and subcortex, and eFC between ventral attention and visual were the connections that showed the biggest difference between pwMS who had evidence of disability vs those who had no disability. Regional node strength differences are shown in Fig. 3; weaker eSC in all networks (except the cerebellum) and stronger eFC in all networks was observed in those pwMS who had evidence of disability. Observed FC was stronger in most networks (except subcortex, cerebellum and ventral attention) while observed SC was weaker in cerebellum and subcortex and stronger in limbic networks in pwMS who had evidence of disability.

#### 3.3. Classification results

Fig. 4 shows that models based on pairwise SC and eSC had similar AUC (p = 0.21, BH corrected), while pairwise eFC significantly outperformed FC (p-value < 10e-4) and had the highest AUC of all the pairwise models. Both models based on estimated regional node strength outperformed observed regional node strength in classifying pwMS into disability groups (corrected p-value < 0.01 for both regional SC vs eSC and regional FC vs eFC). The highest AUC over all 8 models tested was the one based on regional eFC, with a median of 0.681. Regional eSC and eFC had higher AUC results than pairwise eSC and eFC, and, while pairwise eFC outperformed pairwise eSC (corrected p-value < 10e-4), regional eSC and eFC were not significantly different (corrected p-value = 0.17). Confusion matrices, sensitivity, specificity, balanced accuracy, and Brier score largely agreed with AUC results (see Supplementary Table 3 and Fig. 3). Classification models were also tested using EDSS of 3 as the threshold for defining disability groups (88 pwMS who had no disability vs 12 pwMS who had evidence of disability); the results are largely similar but the models, except the regional eFC model, had overall slightly larger AUCs (see Supplementary Fig. 5).

The drop in the AUC results of the regional eFC model might be due to the major decrease in the number of the subjects of the disability group (33 with EDSS 2 and 12 with EDSS 3) and decreased separation of regional eFC metrics when EDSS 3 was used. We also re-ran the classification models using only CIS and RRMS patients (i.e. five pwMS with progressive disease category were excluded as their disease mechanisms or progression may be different from other subtypes. The results were similar to what we observed with the original dataset: the pairwise and regional eFC outperformed other models (see Supplementary Fig. 6).

#### 3.4. Feature weights

Age and sex had the highest weights in the pairwise and regional eSC and eFC models; being older or male was associated with having evidence of disability. Fig. 5 shows the feature weights for the pairwise and regional eSC and eFC classification models. Connections from the cerebellum to the somatomotor regions and from cerebellum to limbic regions had the highest weights in both pairwise eSC and eFC models but in opposite directions; greater eFC and weaker eSC were associated with no disability. Additionally, weaker eSC between ventral attention and frontoparietal and subcortical networks was associated with evidence of disability while weaker eFC between cerebellum and somatomotor/
visual/dorsal attention networks was associated with evidence of disability.

Weaker eSC node strength in the cerebellum, somatomotor, and limbic networks and stronger eSC node strength in the ventral/dorsal attention and default mode networks were associated with no disability. Stronger eFC node strength in the visual, dorsal attention and fronto-parietal was associated with evidence of disability, while weaker eFC node strength in the cerebellum and somatomotor networks was associated with evidence of disability.

Fig. 6 shows the feature weights for the observed SC and FC models, which both had lower classification accuracy and only provided them here for comparison to the eFC/eSC models. Weaker SC between the dorsal attention and visual networks and weaker FC between cerebellum and subcortical, ventral attention and default mode networks were associated with greater disability. Weaker SC between dorsal attention and cerebellum/somatmotor networks and weaker FC between default mode and dorsal attention networks was associated with no disability. Weaker SC and FC to/from the right cerebellum to the rest of the brain was also found to be associated with having evidence of disability (Tozlu et al., 2020a). There was a moderate correlation between the feature weights from the regional FC and eFC models ($r = 0.52$), while the correlations between pairwise FC vs eFC, as well as pairwise/regional SC vs eSC were non-significant and weak ($r < 0.1$) (See Supplementary Table 1). When the features were analyzed at network level, weaker SC and FC in the cerebellum was commonly found as associated with disability.

4. Discussion

In this study, we compared the performance of models using observed SC and FC (derived from advanced MRI directly in pwMS) against models using estimated SC and FC (derived from clinically...
acquired lesion masks and the NeMo Tool) in classifying pwMS into no disability vs evidence of disability groups. We also identified the regions and/or connections that were most different between and important in classifying pwMS into disability groups and compared feature weights over the estimated and observed models. Our main findings were (1) models based on regional eSC and pairwise/regional eFC outperformed models based on their observed counterparts, (2) models based on pairwise eSC performed just as well as models based on pairwise observed SC, (3) models based on regional eSC and eFC performed the best out of all models considered (4) stronger eFC node strength of regions in the visual network and weaker eSC node strength of regions in the default mode and ventral attention networks was associated with disability, and (5) feature weights computed from the regional eFC and FC models were moderately, significantly correlated.

4.1. Previous studies comparing estimated and observed connectivity

The original NeMo Tool was used previously to map disruptions of SC to concurrent and future impairments in stroke and MS (A. Kuceyeski et al., 2015a; Kuceyeski et al., 2018, 2014; A. F. Kuceyeski et al., 2015b). As far as we know, no study to date has compared indirectly estimated SC or FC from the NeMo tool or any other techniques to directly observed SC or FC in their respective abilities to predict impairment or disability in pwMS. However, we are aware of several recent studies in stroke that have shown contrasting results. For example, one study showed that lesion masks themselves, in addition to estimated structural disconnectivity and to a lesser extend directly observed FC, predicted multiple post-stroke impairments while estimated functional network disruptions did not (Salvalaggio et al., 2020). In a follow-up replication study with the same dataset, Cohen et al. (Cohen et al., 2021) found that
modifying the way in which estimated functional disruption was calculated resulted in models with explained variance similar to the estimated structural disconnectivity. Although our estimated FC was calculated in a very different manner than these studies, our results agree with the latter one. In fact, our results in pwMS actually show that models based on pairwise eFC outperform models based on pairwise eSC. Our novel approach to estimating FC from lesion masks could be a complimentary or alternative technique to lesion-network dysfunction mapping that may allow even greater explained variance when performing brain-behavior mapping in individuals with neurological disease.

4.2. Estimated vs observed connectivity: Advantages and disadvantages

Collecting and processing dMRI and fMRI in patient populations is difficult, as individuals often cannot tolerate long scan times, scans are expensive, and processing of the images requires a high level of expertise. In addition, anatomical or physiological changes like edema and/or inflammation can add noise to already noisy dMRI and fMRI modalities and further corrupt their accurate measurement. It has been shown that various tractography approaches in pwMS will better allow tracing of streamlines through MS lesions than others (Läpp et al., 2020). This is an especially important consideration when calculating SCs based on diffusion summary statistics like FA since the damaged voxels would be a complimentary or alternative technique to lesion-network dysfunction mapping that may allow even greater explained variance when performing brain-behavior mapping in individuals with neurological disease.

Fig. 4. Classification results. AUC results obtained using pairwise connectivity (left panel) and regional node strength (right panel) to classify pwMS into no disability vs evidence of disability groups. The plots show the median (white dot) and inter quantile range (black bar) of the AUCs over the 500 hold-out test sets. The asterisks show the models that were significantly different after BH correction. SC: structural connectivity, FC: functional connectivity, eSC: estimated structural connectivity, and eFC: estimated functional connectivity.

Previous studies have used SC and FC metrics to distinguish pwMS from HC and/or to classify pwMS by disability level defined by thresholding EDSS scores (Richiardi et al., 2012; Zhong et al., 2017; Zurita et al., 2018). A previous study compared the classification accuracy of SC and FC to discriminate pwMS into disability groups using a threshold EDSS of 2 (Zurita et al., 2018); our models based on observed connectivity metrics had higher AUC (0.47 vs 0.59 for models based on SC and 0.53 vs 0.59 for models based on FC). The difference in results might be due to different clinical stage used in the studies (they used relapsing remitting and our study that included all phenotypes), different classification methods (they used support vector machines and we used a ridge classifier) and different MRI acquisition and processing protocols. In a recent preprint, we used the NeMo Tool’s estimated regional dis-
categories also based on a threshold EDSS of 2 (Tozlu et al., 2020b). An EDSS threshold of 3 was identified as the cut-off for functional reorganization and adaptation in MS (Tommasin et al., 2018). This study found no significant difference in FC between HC and pwMS who had an EDSS $\leq 3$, while FC decreased in the pwMS who had an EDSS $> 3$. Our replication study using a threshold EDSS of 3 and found largely similar results (see Supplementary Fig. 5).

The focus of this work was on comparing the accuracy of observed vs estimated connectivity networks in classifying people with MS according to disability level as well as investigating the regions or pairwise connections which were informative of disability. We did not set out to identify the most accurate classification model; with the limited number
of individuals in the study and large dimensionality of the imaging variables, a model with fewer variables likely will have higher AUC. In fact, a simple logistic regression model using only age, sex, race, disease duration, and total brain lesion volume (on T2 FLAIR) had a median AUC of 0.71, which was similar to the best model (regional eFC with median AUC of 0.68). Future work focusing only on prediction accuracy (and not model inference, as we do here) using this data set may investigate dimensionality reduction techniques applied to high dimensional imaging measures to improve classification performance.

Fig. 6. The feature weights computed with SC and FC models. The relative weights of the (A) pairwise and (B) regional (node strength) SC and FC obtained in classifying pwMS according disability. Feature weight metrics were scaled by the maximum of the absolute value over all models. Positive weights (hotter colors) indicate those connections/regions where stronger connectivity was associated with having evidence of disability class, while negative weights (cooler colors) indicate those connections/regions where stronger connectivity was associated with being in the no disability class. The circle plots summarize the pairwise feature weights by representing the mean of the positive and negative pairwise feature weights between functional network assignments (7 networks from the Yeo atlas, plus cerebellum and subcortex) The radial plots summarize the feature weights of regional models by representing the mean of the positive and negative weights in the same functional networks used for the circle plot. The absolute value of the average negative connections were presented in the radial plots.
4.4. Comparison of feature weights: Estimated vs observed connectivity models

Even though the feature weights from the regional FC and eFC models are moderately correlated (Pearson’s r = 0.52, p-value < 10e-7), the correlation between the raw values of regional FC and eFC was weak (r = 0.09, p-value < 10e-16), therefore the interpretation of the feature weights of the eFC variables which were estimated via a deep learning model trained on healthy controls should be made with caution. Increased FC and eFC in cerebellum, subcortex, somatomotor, and ventral attention networks were commonly found to be associated with no disability while decreased FC and eFC in dorsal attention and default mode networks were commonly found to be associated with no disability. Moreover, higher node strength in many regions (>35) in the FC and eFC models were commonly associated with disability. This finding is consistent with previous studies showing upregulation of FC reflecting possible compensation in pwMS having lower EDSS compared to those with greater EDSS (Tommasin et al., 2019).

4.5. Assessment of feature importance

A recent study (Tian et al., 2021) showed that coefficients of models predicting cognition and sex using observed FC can be unreliable. In their paper, they argue that increasing sample size, applying the Haufe transformation and using non-sparse regularization can all improve the feature weight reliability; furthermore they found that mass univariate results can be more reliable than model coefficients. In our study, we used (non-sparse) ridge regularization, however the Haufe transformation cannot be applied as our model contains binary/categorical demographic variables (such as sex and spinal cord lesion category). Therefore, here we focus on reporting classification model coefficients that also agreed with the mass univariate group comparison results.

4.6. eSC in the default mode and ventral attention and eFC in the visual networks had the highest importance for disability classification in MS

In our study, weaker eSC in the default mode and ventral attention networks was commonly found as associated with disability in both univariate test and the classification model. Structural disruption and functional alterations in the default mode network were previously found as associated with poorer cognitive abilities and cognitive rehabilitation outcomes (Fuchs et al., 2020; Has Silemek et al., 2020). However, this is the first study that showed the relationship between disability and connectivity changes in these networks in MS. In our study, we also showed that greater eFC in the regions of the visual network was associated with disability. Our results were in concordance with a previous study (Rocca et al., 2018) that also showed greater FC of the visual network in pwMS as compared to HCs and interpreted as the presence of cross-modal plasticity mechanisms after visual impairment seen in early MS (Rocca et al., 2005).

4.7. eSC and eFC between the cerebellum and somatomotor networks is central to accurate disability classification

It was not surprising that the eSC and eFC from the somatomotor network and cerebellum had the highest weights in identifying pwMS’s disability level, as these networks are central to motor and movement. In addition to the cerebellum-somatomotor/limbic connection, a decrease in the connectivity metrics in the cerebellum was also associated with having more probability of having disability. Previous studies have shown relationships between cerebellar pathology and impairments in motor control and cognition (D’Ambrosio et al., 2017; Weier et al., 2014). The presence of cerebellum-related symptoms at the onset of MS such as coordination issues or tremor were i) shown to be associated with shorter time to an EDSS of 6 (Weinschenker et al., 1991) and ii) related to earlier onset of progressive disease diagnosis (Novotna et al., 2015). Atrophy in the anterior cerebellum was associated with motor dysfunction (D’Ambrosio et al., 2017) in pwMS. In our previous study, the cerebellum was also shown as the most important region in classifying pwMS by disability level, where eSC was computed using only hyperintense rim lesions from quantitative susceptibility imaging (Tozlu et al., 2020b). Another study that investigated FC in the cerebellum found that the loss of functional cerebellar connections was related to disability. Our results and previous findings may suggest that decreasing structural deterioration and functional upregulation in the cerebellum may be a new target for clinical trials (Tommasin et al., 2018). Among demographics and clinical variables, age and sex appeared to be an important predictor for disability classification; specifically, being older and male was associated with a higher probability of being in the disability group. It has been shown previously that male patients tend to have more severe disease onset with accelerated clinical progression in MS (Bove et al., 2012).

4.8. Limitations and future work

One of the limitations of the present study was in the quality of the advanced MRI collected directly in pwMS. The fMRI acquisition time was relatively short and had a longer TR (6 min, TR = 2.3 s) and the dMRI acquisition had only a single b-value of 800 and only 55 directions. These limitations may mean less accurate observed FC or SC, which could have negatively impacted their classification accuracy. A future study will be needed to compare eSC and eFC to SC and FC from high-quality fMRI and dMRI acquisitions, however if high-quality SC and FC are superior than eSC and eFC, it may still be preferable to use estimates given feasibility issues. The estimates of SC and FC from the NeMo tool are based on a database of controls and thus cannot capture an individual’s specific SC anatomy or the particulars of the relationship between SC and FC under the influence of lesion pathology. Additionally, there may be pathologies in the white matter of pwMS that are not contained in the lesion masks and are thus not considered in the NeMo Tool’s estimates. However, MS lesions disrupt diffusion MRI signals and add noise to tractography results, and the current findings indicate that the NeMo Tool is a good alternative to performing tractography directly in pwMS. This study investigates cross-sectional relationships between disability and connectivity; a future study mapping baseline connectivity to future disability is needed to better understand the prognostic ability of estimated connectomes. The secondary goal of this study was to compare the various models’ feature importance patterns, and dimensionality reduction techniques would not permit easy comparison. However, future work may explore using dimension reduction to improve model performance.

4.9. Conclusions

This is the first study to show that models using estimated SC and FC based only on lesion masks and the NeMo Tool outperform models based on observed SC and FC extracted from advanced MRI in classifying pwMS into disability categories. Models based on eFC’s regional node strength had the highest overall classification performance. Stronger eFC node strength of regions in the visual network and weaker eSC node strength of regions in the default mode and ventral attention networks was associated with disability. A deeper understanding of the role of the connectome in MS is needed if we are to gain a comprehensive view of the disease, develop more accurate prognostic or therapeutic tools and ultimately improve clinical outcomes in pwMS. This work provides a viable alternative to performing high-cost, advanced MRI in patient populations, brings the connectome one step closer to the clinic.

5. Citation gender diversity statement

We used classification of gender based on the first names of the first and last authors, with possible combinations including male/male,
male/female, female/male, and female/female. The gender balance of papers cited within this work was quantified using gender-api.com. The authors with a gender estimation accuracy lower than 90% were checked using manual gender determination from authors’ publicly available pronouns. Among the 58 cited works, 2 articles had only one author. Among the 56 cited works with more than one author, 46% (n = 26) were MM, 34% (n = 19) were WM, 11% (n = 6) were MW, and 9% (n = 5) were WW.

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**CRediT authorship contribution statement**

**Ceren Tozlu:** Formal analysis, Investigation, Visualization, Writing - original draft. **Keith Jamison:** Data curation, Software, Investigation, Writing - review & editing. **Zijn Gu:** Formal analysis, Writing - review & editing. **Susan A. Gauthier:** Conceptualization, Methodology, Writing - review & editing. **Amy Kuceyeski:** Conceptualization, Methodology, Software, Supervision, Writing - review & editing.

**Declaration of Competing Interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

**Appendix A. Supplementary data**

Supplementary data to this article can be found online at https://doi.org/10.1016/j.nicl.2021.102827.

**References**

Anderson, J.L.R., Graham, M.S., Zolod, E., Sotopouloou, S.N., 2016. Incorporating outlier detection and replacement into a non-parametric framework for movement and distortion correction of diffusion MR images. Neuroimage 141, 556–572. https://doi.org/10.1016/j.neuroimage.2016.06.058.

Anderson, J.L.R., Sotopouloou, S.N., 2016. An integrated approach to correction for off-resonance effects and subject movement in diffusion MR imaging. Neuroimage 125, 1063–1078. https://doi.org/10.1016/j.neuroimage.2015.10.019.

Barkhof, F., 2002. The clinic-neurological paradox in multiple sclerosis revisited. Curr. Opin. Neurol. 15 (3), 239–245.

Bassi, M.S., Gli, L., Buttari, F., Maffei, P., Marfia, G.A., Restivo, D.A., Centone, D., Iezzi, E., 2017. Remodeling functional connectivity in multiple sclerosis: A challenging therapeutic approach. Front. Neurosci. https://doi.org/10.3389/fnins.2017.00710.

Benjamini, Y., Hochberg, Y., 1995. Controlling the False Discovery Rate: A Practical and Powerful Approach to Multiple Testing. J. R. Stat. Soc. Ser. B (57), 289–300. https://doi.org/10.1111/j.2517-6161.1995.tb02031.x.

Boes, A.D., Prasad, S., Liu, H., Liu, Q.i., Pascual-Leone, A., Caviness, V.S., Fox, M.D., 2013. Predicting future brain tissue loss from white matter connectivity disruption in ischemic stroke. Stroke 45 (3), 717–722. https://doi.org/10.1161/STROKEAHA.113.003645.

Kuceyeski, A., Maruta, J., Niogi, S.N., Gharaj, J., Raj, A., 2011. The generation and validation of white matter connectivity importance maps. Neuroimage 58, 159–191. https://doi.org/10.1016/j.neuroimage.2011.05.087.

Kuceyeski, A., Maruta, J., Relkin, N., Raj, A. 2013. The Network Modification (NeMo) Tool: Elucidating the Effect of White Matter Integrity Changes on Cortical and Subcortical Structural Connectivity. Brain Connect. 3 (5), 451–463. https://doi.org/10.1089/brain.2013.0147.

Kuceyeski, A., Monohan, E., Emori, F., Fujimoto, K., Vargas, G., Sautha, I., 2018. Baseline biomarkers of connectome disruption and atrophy predict future progression in early multiple sclerosis. Neuroimage Clin. 19, 417–424. https://doi.org/10.1016/j.nicl.2018.05.003.

Kuceyeski, A., Navi, B.B., Kamel, H., Relkin, N., Villanueva, M., Raj, A., Toiglia, J., O’Dell, J., Iadecola, C., 2015a. Exploring the brain’s structural connectome: a quantitative stroke lesion-dysfunction mapping study. Hum Brain Mapp 36, 2147–2160. https://doi.org/10.1016/j.humbrainm.2015.02.024.

Kuceyeski, A., Shah, S., Dyke, J.P., Bickel, S., Abdellour, F., Schiff, N.D., Voss, H.U., Raj, A., 2016. The application of a mathematical model linking structural and functional connectonles in severe brain injury. Neuroimage Clin. 11, 635–647. https://doi.org/10.1016/j.nicl.2016.04.006.

Kuceyeski, A.F., Vargas, W., Dayan, M., Monohan, E., Blackwell, C., Raj, A., Fujimoto, K., Guthe, A., 2018. Baseline relationship among gray matter atrophy, abnormalities in connecting white matter, and cognitive performance in multiple sclerosis. Am. J. Neuroradiol. 39 (6), 709–729. https://doi.org/10.3174/jnr.2017.A4165.

Li, Y., Zhan, J., Kim, Y., Chen, Y., Moon, A., Aramo, D., Troiani, L., Markovic-Plese, S., Lin, W., Shen, D., 2013. Diffusion tensor imaging based network analysis detects alterations of neuroconnectivity in patients with clinically early relapsing-remitting multiple sclerosis. Hum. Brain MAPP. 34 (12), 3376–3391. https://doi.org/10.1002/hbm.23218.

Lipp, I., Parker, G.D., Tallantyre, E.C., Goodall, A., Gama, S., Patitucci, E., Heveron, P., Tomassini, V., Jones, D.K., 2020. Tractography in the presence of multiple sclerosis lesions. Neuroimage 209, 116471. https://doi.org/10.1016/j.neuroimage.2019.116471.

Muthuraman, M., Fleischer, V., Kolber, P., Luesi, F., Zipp, F., Groppa, S., 2016. Structural Brain Network Characteristics Can Differentiate CIS from Early RRMS. Front. Neuosci. 10, 14. https://doi.org/10.3389/fnins.2016.00014.

Novotny, J., Vaz Sobr, M., M.M., Abou Zeid, N., Kale, N., Tunuceu, M., Crusan, D.J., Atkinson, E.J., Siva, A., Keegan, B.M., Pirko, I., Pittock, S.J., Lucchini, C.F., Noseworthy, J.H., Weinsenker, B.G., Rodriguez, M., Kancitari, O.H., 2015. Poor early relapse recovery affects onset of progressive disease course in multiple sclerosis. Neurology 85 (8), 722–729. https://doi.org/10.1212/WNL.0000000000001858.

Pagani, E., Bammer, R., Horsfield, M.A., Rovaris, M., Gass, A., Ciccarelli, O., Filippi, M., 2007. Diffusion MR imaging in multiple sclerosis: Technical aspects and challenges. J. Neuroradiol. 24, 211–212. https://doi.org/10.1055/s-2006-950432.

Pagani M.A. Roca E. De Meo M.A. Horsfield B. Colombo M. Rodegher G. Comi M. Filippi Structural connectivity in multiple sclerosis and modeling of disconnection 10 2020 220 222 303 https://doi.org/10.1017/S1200082820000077.

Perera, J.M.S., Xiong, L., Acosta-Cabronero, J., Pengas, G., Williams, G.B., Nestor, P.J., 2010. Registration accuracy for VBM studies varies according to region and degenerative disease group. Neuroimage 49 (3), 2205–2215. https://doi.org/10.1016/j.neuroimage.2009.12.068.
multiple sclerosis patients from resting state functional connectivity. Neuroimage 62 (3), 2021–2033. https://doi.org/10.1016/j.neuroimage.2012.05.078.
Rocca, M.A., Colombo, B., Falini, A., Ghezzi, A., Martinelli, V., Scotti, G., Comi, G., Filippi, M., 2005. Cortical adaptation in patients with MS: a cross-sectional functional MRI study of disease phenotypes 4. https://doi.org/10.1016/S1474-4422(05)00717-X.
Rocca, M.A., Valasina, P., Leavitt, V.M., Rodegher, M., Radaelli, M., Riccitelli, G.C., Martinelli, V., Martinelli-Boneschi, F., Falini, A., Comi, G., Filippi, M., 2018. Functional network connectivity abnormalities in multiple sclerosis: Correlations with disability and cognitive impairment. Mult. Scler. J. 24 (4), 457–471. https://doi.org/10.1177/1352458517749975.
Rossi, S., Studer, V., Motta, C., De Chiara, V., Barbieri, F., Bernardi, G., Centonze, D., 2012. Inflammation inhibits GABA transmission in multiple sclerosis. Mult. Scler. J. 18 (11), 1653–1655. https://doi.org/10.1177/1352458512440207.
A. Salvaglio M. De Filippo De Grazia M. Zorzi M. Thiebaut de Schotten M. Corbetta Post-stroke deficit prediction from lesion and indirect structural and functional disconnection 143 7 2020 2021 2173 2188 10.1093/brain/awaa156.
Santos, M.S., Soares, J.P., Abreu, P.H., Araujo, H., Santos, J., 2018. Cross-Validation for Imbalanced Datasets: Avoiding Overoptimistic and Overfitting Approaches [Research Frontier]. IEEE Comput. Intell. Mag. 13 (4), 59–76. https://doi.org/10.1109/MCI.2018.2866730.
Sarwar, T., Tian, Y., Yeo, B.T.T., Ramamohanaroo, K., Zalesky, A., 2021. Structure-function coupling in the human connectome: A machine learning approach. Neuroimage 226, 117609. https://doi.org/10.1016/j.neuroimage.2020.117609.
Shu, N., Liu, Y., Li, K., Duan, Y., Wang, J., Yu, C., Dong, H., Ye, J., He, Y., 2011. Diffusion Tensor Tractography Reveals Disrupted Topological Efficiency in White Matter Structural Networks in Multiple Sclerosis. Cereb. Cortex 21, 2565–2577. https://doi.org/10.1093/cercor/bhr039.
Smith, R.E., Tournier, J.-D., Calamante, F., Connelly, A., 2015. The effects of SIFT on the feature weights reliable? bioRxiv 2021.05.27.446059. https://doi.org/10.1101/2021.05.27.446059.
Santos, M.S., Soares, J.P., Abreu, P.H., Araujo, H., Santos, J., 2018. Functional network connectivity abnormalities in multiple sclerosis: Correlations with disability and cognitive impairment. Mult. Scler. J. 24 (4), 459–470. https://doi.org/10.1177/1352458517749975.
Rocca, M.A., Colombo, B., Falini, A., Ghezzi, A., Martinelli, V., Scotti, G., Comi, G., Filippi, M., 2005. Cortical adaptation in patients with MS: a cross-sectional functional MRI study of disease phenotypes 4. https://doi.org/10.1016/S1474-4422(05)00717-X.
Rocca, M.A., Valasina, P., Leavitt, V.M., Rodegher, M., Radaelli, M., Riccitelli, G.C., Martinelli, V., Martinelli-Boneschi, F., Falini, A., Comi, G., Filippi, M., 2018. Functional network connectivity abnormalities in multiple sclerosis: Correlations with disability and cognitive impairment. Mult. Scler. J. 24 (4), 457–471. https://doi.org/10.1177/1352458517749975.
Rossi, S., Studer, V., Motta, C., De Chiara, V., Barbieri, F., Bernardi, G., Centonze, D., 2012. Inflammation inhibits GABA transmission in multiple sclerosis. Mult. Scler. J. 18 (11), 1653–1655. https://doi.org/10.1177/1352458512440207.
A. Salvaglio M. De Filippo De Grazia M. Zorzi M. Thiebaut de Schotten M. Corbetta Post-stroke deficit prediction from lesion and indirect structural and functional disconnection 143 7 2020 2021 2173 2188 10.1093/brain/awaa156.
Santos, M.S., Soares, J.P., Abreu, P.H., Araujo, H., Santos, J., 2018. Cross-Validation for Imbalanced Datasets: Avoiding Overoptimistic and Overfitting Approaches [Research Frontier]. IEEE Comput. Intell. Mag. 13 (4), 59–76. https://doi.org/10.1109/MCI.2018.2866730.
Sarwar, T., Tian, Y., Yeo, B.T.T., Ramamohanaroo, K., Zalesky, A., 2021. Structure-function coupling in the human connectome: A machine learning approach. Neuroimage 226, 117609. https://doi.org/10.1016/j.neuroimage.2020.117609.
Shu, N., Liu, Y., Li, K., Duan, Y., Wang, J., Yu, C., Dong, H., Ye, J., He, Y., 2011. Diffusion Tensor Tractography Reveals Disrupted Topological Efficiency in White Matter Structural Networks in Multiple Sclerosis. Cereb. Cortex 21, 2565–2577. https://doi.org/10.1093/cercor/bhr039.
Smith, R.E., Tournier, J.-D., Calamante, F., Connelly, A., 2015. The effects of SIFT on the feature weights reliable? bioRxiv 2021.05.27.446059. https://doi.org/10.1101/2021.05.27.446059.
Santos, M.S., Soares, J.P., Abreu, P.H., Araujo, H., Santos, J., 2018. Cross-Validation for Imbalanced Datasets: Avoiding Overoptimistic and Overfitting Approaches [Research Frontier]. IEEE Comput. Intell. Mag. 13 (4), 59–76. https://doi.org/10.1109/MCI.2018.2866730.
Sarwar, T., Tian, Y., Yeo, B.T.T., Ramamohanaroo, K., Zalesky, A., 2021. Structure-function coupling in the human connectome: A machine learning approach. Neuroimage 226, 117609. https://doi.org/10.1016/j.neuroimage.2020.117609.
Shu, N., Liu, Y., Li, K., Duan, Y., Wang, J., Yu, C., Dong, H., Ye, J., He, Y., 2011. Diffusion Tensor Tractography Reveals Disrupted Topological Efficiency in White Matter Structural Networks in Multiple Sclerosis. Cereb. Cortex 21, 2565–2577. https://doi.org/10.1093/cercor/bhr039.
Smith, R.E., Tournier, J.-D., Calamante, F., Connelly, A., 2015. The effects of SIFT on the feature weights reliable? bioRxiv 2021.05.27.446059. https://doi.org/10.1101/2021.05.27.446059.
Santos, M.S., Soares, J.P., Abreu, P.H., Araujo, H., Santos, J., 2018. Cross-Validation for Imbalanced Datasets: Avoiding Overoptimistic and Overfitting Approaches [Research Frontier]. IEEE Comput. Intell. Mag. 13 (4), 59–76. https://doi.org/10.1109/MCI.2018.2866730.