Resting-state functional connectivity predicts individual language impairment of patients with left hemispheric gliomas involving language network

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A B S T R A C T

Language deficits following brain tumors should consider the dynamic interactions between different tumor growth kinetics and functional network reorganization. We measured the resting-state functional connectivity of 126 patients with left cerebral gliomas involving language network areas, including 77 patients with low-grade gliomas (LGG) and 49 patients with high-grade gliomas (HGG). Functional network mapping for language was performed by construction of a multivariate machine learning-based prediction model of individual aphasia quotient (AQ), a summary score that indicates overall severity of language impairment. We found that the AQ scores for HGG patients were significantly lower than those of LGG patients. The prediction accuracy of HGG patients ($R^2 = 0.27$, permutation $P = 0.007$) was much higher than that of LGG patients ($R^2 = 0.09$, permutation $P = 0.032$). The rsFC regions predictive of LGG's AQ involved the bilateral frontal, temporal, and parietal lobes, subcortical regions, and bilateral cerebro-cerebellar connections, mainly in regions belonging to the canonical language network. The functional network of language processing for HGG patients showed strong dependence on connections of the left cerebro-cerebellar connections, limbic system, and the temporal, occipital, and prefrontal lobes. Together, our findings suggested that individual language processing of glioma patients links large-scale, bilateral, cortico-subcortical, and cerebro-cerebellar functional networks with different network reorganizational mechanisms underlying the different levels of language impairments in LGG and HGG patients.

1. Introduction

A brain tumor is a mass or growth of abnormal cells in the brain. Gliomas are the most common primary brain tumors. Diffusive and progressive glioma infiltration of healthy tissues leads to significant functional reshaping and behavioral or cognitive deficits (e.g., language impairment or aphasia). Most previous structural or functional MRI (fMRI) studies (Huang et al., 2018; Lu et al., 2017; Weng et al., 2018; Zhang et al., 2016), as well as intraoperative electrocortical stimulation studies (Duffau et al., 2014; Picart et al., 2018; Sanai et al., 2008; Wu et al., 2015), attempted to localize the surrounding eloquent functional areas to maximize the extent of tumor resection while reducing the risk of postoperative cognitive deficits. The patterns of functional and structural alterations following glioma growth have been explored, including the recruitment of perilesional regions as local reorganization, homotopic reorganization of gray matter volume (Almairac et al., 2018), activation changes in the contralateral homologous region (Chivukula et al., 2018; Desmurget et al., 2007), and the remote recruitment of other ipsihemispheric and contrahemispheric regions, such as the cerebellum (Zhang et al., 2016). These local

Abbreviations: LGG, low-grade glioma; HGG, high-grade glioma; ABC, Aphasia Battery of Chinese; AQ, aphasia quotient; SS, spontaneous speech

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and remote adaptive mechanisms have been thought to be physiological basis for the high level of functional reorganization and are used to explain why tumor infiltration within the so-called “eloquent” areas often does not result in detectable neurological deficits. However, relatively few studies have focused on language network reorganization of the brain following glioma growth.

In contrast to the long-lasting localizationist view of language neurobiology as residing solely in Broca’s and Wernicke’s areas, numerous pieces of convincing evidences indicated that language function relies on interconnected cortico-subcortical regions and bilaterally-distributed connections, forming the language network (Duffau et al., 2014; Fedorenko and Thompson-Schill, 2014; Luo et al., 2019; Torres-Prioris et al., 2019; Tremblay and Dick, 2016). In this delocalized network-based framework, language regions in the frontal, temporal and parietal lobes, and the right cerebellum are structurally and/or functionally connected, forming large-scale language (sub) networks for linguistic processing.

Due to the infiltrative, diffusive, and migratory characteristics of gliomas, pathological perturbations by tumor are rarely confined to a single locus but instead influence other regions via axonal pathways (Fornito et al., 2015; Hillary and Grafman, 2017; Stam, 2014). Numerous reports have shown that brain tumors can lead to various cognitive deficits-related alterations in local and interhemispheric resting-state functional connectivity (rsFC) across different functional networks (Fox and King, 2018; Ghinda et al., 2018). We have shown that left hemispheric gliomas induce language-related structural and functional alterations in cerebellum via cerebro-cerebellar circuits (Zhang et al., 2018). Outside the language network, diffused functional reorganization may occur in the contralesional hemisphere and other functional systems (Fox and King, 2018; Ghinda et al., 2018), allowing both long-distance and global lesion effects. Brain tumors also cause significant alterations in global network topology, such as global network efficiency (Aerts et al., 2016).

Notably, the severity of cognitive deficits, network disruptions, and the degree of network reorganization are related to the speed of infiltration and the extent to which the tumor infiltrates healthy functional systems (Aerts et al., 2016; Fox and King, 2018; Ghinda et al., 2018; Zhang et al., 2018). Tumor grade is an important predictor of patients’ clinical and cognitive deficits (Taphoorn and Klein, 2004). In the World Health Organization (WHO) grading system, grade I tumors are the least malignant and grade II or IV tumors are the most malignant. Grade I and II brain tumors are often referred to as low-grade gliomas (LGG), whereas grade III and IV are described as high-grade gliomas (HGG). Previous studies have shown that LGG patients with slow infiltration (over a time period of years) typically have extensive functional reshaping and only slightly impaired or undetectable language deficits (Duffau, 2005; Zhang et al., 2018). In contrast, HGG patients (infiltration over a time period of months) exhibit more severe language and other cognitive impairments than LGG patients (Desmurget et al., 2007; Noll et al., 2015; van Kessel et al., 2017; Zhang et al., 2018). Additionally, several studies reported that LGG and HGG patients had different plastic mechanisms of brain networks (van Dellen et al., 2012; Zhang et al., 2016). However, no study has attempted to investigate the language network basis at the whole brain level to determine the language network reorganization differences between LGG and HGG patients.

In this work, a large sample of 126 patients, including 77 patients with LGGs and 49 patients with HGGs, were subjected to machine learning-based network mapping for language. All patients included in this study have left cerebral gliomas involving language network areas. For each patient, we measured rsFC using a whole-brain parcel (Shen et al., 2015) after excluding the patient’s tumor extent. For each group, we then built a machine learning-based rsFC prediction model of individual language scores. The language-related network for each model was identified based on predictive contribution (i.e., the highest predictive weights). We hypothesized that glioma would induce widespread network disruptions, including the language network. The perilesional intact language regions, the ipsilesional and contralesional (homologous) language regions, and the cerebellum and cerebro-cerebellar circuits may participate in language processing. The reliability of the findings was evaluated by further considering different pre-processing/analysis strategies (i.e., the parcellation scheme, covariates, global signal removal, feature selection threshold and generalizability of the prediction model).

2. Materials and methods

2.1. Participants

Patients were prospectively screened and collected from two centers: 1) Huashan Hospital (from 2011 to 2016), and 2) the First Affiliated Hospital of Zhengzhou University (from 2014 to 2018). Because we used the same system of MRI scanner (Siemens Magnetom Verio 3.0 T MRI), as well as identical imaging protocols and inclusion criteria, the data from the two centers were analyzed together and a potential site effect was controlled by adding the site factor as a single regressor in the statistical analysis.

Patient inclusion criteria: (1) Pathologically confirmed glioma (based on 2007 WHO classification system) in left cerebral hemisphere (Louis et al., 2007); (2) No chemo and/or radiation treatment history; (3) Location of the glioma overlapped or was within the language network areas proposed by Fedorenko et al. (2010); (4) Patients’ age was 18–75 years old; (5) Right-handedness confirmed by the Edinburgh Handedness Inventory; (6) No symptoms of motor impairment, as indicated by a score of grade V on the Medical Research Council (MRC) Scale for Muscle Strength (Paterson-Stirling et al., 2008); (7) Chinese Han nationality; (8) No history of brain surgery; (9) More than nine years of education; (10) No midline shift in structural images, as confirmed by the in situ location of midline structures (corpus callosum, septa pellucida, third ventricle, hypothalamus, and pineal region) of the brain; (11) Both structural and functional images covered the whole brain, especially the whole cerebellum; (12) Good cooperation during the linguistic/cognitive evaluation; (13) No history of other major neurological or psychiatric disorders; and (14) No history of alcohol or drug abuse.

In total, 126 patients met all inclusion criteria and completed the entire preoperative clinical, language, and imaging data acquisition. The patients were further grouped based on the malignancy of brain tumors: 77 patients with grade I (n = 2) and II (n = 75) brain tumors were designated as low-grade gliomas (LGG) and 49 patients with grade III (n = 26) and IV (n = 23) brain tumors were designated as high-grade gliomas (HGG). For the two cases of grade I in LGG group, one patient was diagnosed as ganglioglia and the other patient was diagnosed as neuroepithelial glioma.

All patients underwent comprehensive assessment of Karnofsky performance status, language, motor, and general cognitive function (Zhang et al., 2018). The language function was assessed using the Aphasia Battery for Chinese speakers (ABC). ABC is the Chinese standardized adaptation of the Western Aphasia Battery (Lu et al., 2013; Wu et al., 2015; Zhang et al., 2018), and includes spontaneous speech (range, 0–20), comprehension (range, 0–230), repetition (range, 0–100), and naming (range, 0–100) scores. The Aphasia Quotient (AQ) score (range, 0–100) can be calculated from these items to reflect the global severity. The demographics and clinical information are summarized in Table 1. The language scores from the ABC tests for the two groups are presented in Supplementary Fig. 1.

All processes strictly followed the requirements of the Declaration of Helsinki. This study was approved and supervised by the Ethics Committee of Huashan Hospital and the Ethics Committee of the First Affiliated Hospital of Zhengzhou University, respectively. Written informed consents were obtained from all subjects and/or their legal guardians.
2.2. Image acquisition

For LGG, high resolution T1-weighted and T2-weighted fluid-attenuated inversion recovery (T2-FLAIR) images were acquired with the following parameters. T1-weighted images: axial magnetization-prepared rapid gradient echo (MPRAGE) sequence, repetition time (TR) = 1900 ms; echo time (TE) = 2.93 ms; flip angle (FA) = 9°; inversion time (TI) = 900 ms; field of view (FOV) = 250 × 219 mm; matrix size = 256 × 215; slice thickness = 1 mm; voxel size = 1 × 1 × 1 mm³; slice number = 176; and scanning time = 7 min 47 s. T2-FLAIR: TR = 9000 ms; TE = 99 ms; FA = 150°; TI = 2500 ms; FOV = 240 × 214 mm; matrix size = 256 × 160; slice thickness = 2 mm; voxel size = 0.9 × 1.3 × 2.0 mm³; slice number = 66; and scanning time = 7 min 30 s. For HGG, the T1-weighted sequence images with contrast (gadopentetate dimeglumine) were acquired with the same parameters.

For all patients, rs-fMRI images were acquired using the following parameters: TR = 2000 ms; TE = 35 ms; FA = 90°; FOV = 240 × 240 mm²; matrix size = 64 × 64; thickness/gap = 4 mm/1 mm; voxel size = 3.3 × 3.3 × 5.0 mm³; slice number = 33; scanning time = 8 min; and number of time points = 240. All subjects were required to remain still with their eyes closed and not fall asleep during the time of scanning.

2.3. Structural data processing

Lesion mapping. For each patient, manual tumor drawing on the 3D-T1 images was performed based on the contrast-enhancing tumor areas or the FLAIR hyperintense areas (necrotic areas were included but not peritumoral edema). For 3D-T1 images without glioma enhancement, T2-FLAIR images were first coregistered to the 3D-T1 images to serve as visual reference. After manually tracing the tumor, we created a 3D-T1 volume lacking the tumor area (set to 0) for each patient. This procedure was performed manually using RANO criteria as reference (Wen et al., 2010). The accuracy of the manual tracing was then confirmed by a senior neurosurgeon. Each 3D-T1 volume without tumor area was segmented into gray matter (GM), white matter (WM), cerebrospinal fluid (CSF), bone, soft tissue, and air/background using SPM12 (https://www.fil.ion.ucl.ac.uk/spm). The cost function masking method was used to improve the precision of spatial normalization in glioma patients (Andersen et al., 2010). We then used DARTEL (Diffeomorphic Anatomical Registration using Exponentiated Lie algebra) for registration, normalization, and modulation (Ashburner, 2007). A customized template was generated using the average tissue probability maps across all subjects, and then each subject’s segmented maps were

| Table 1 Demographic information and language scores of LGG and HGG patients |
|----------------------|----------------------|----------------------|
|                      | LGG (n = 77)         | HGG (n = 49)         |
| Age                  | 38.84 ± 9.65         | 43.27 ± 14.71        |
| Gender (M/F)         | 38:43               | 29:20                |
| Education            | 10.92 ± 4.07         | 11.35 ± 4.28         |
| Tumor volume (cm³)   | 82.34 ± 49.99        | 77.22 ± 49.58        |
| FD                   | 0.07 ± 0.03          | 0.08 ± 0.04          |
| WHO grade            | 1/2:75               | 11/16:26/23          |
| AQ                   | 95.23 ± 4.84         | 91.75 ± 8.11         |
| Spontaneous speech   | 18.86 ± 1.48         | 18.12 ± 2.39         |
| Comprehension        | 222.7 ± 13.09        | 213.92 ± 22.55       |
| Repetition           | 96.82 ± 6.64         | 94.51 ± 10.11        |
| Naming               | 94.06 ± 5.64         | 88.05 ± 10.83        |

WHO, World Health Organization; AQ, Aphasia Quotient; FD, framewise displacement.

* P-value obtained using Pearson’s chi-square test.

Fig. 1. Lesion topography of the 77 LGG patients and 49 HGG patients. The color bar represents the number of patients with a lesion on a specific voxel.
warping into the template. This procedure was repeated until the best study-specific template was generated. The images were then modulated according to the Jacobian determinants to ensure conservation of regional differences in the absolute amounts of GM. Finally, the registered images were transformed to Montreal Neurological Institute (MNI) space. The native tumor mask was spatially normalized to standard MNI space by applying the deformation field estimated by segmentation. Tumor masks of each group (LGG or HGG) in MNI space were then stacked and binarized to construct a tumor-overlapping image (Fig. 1), in which each voxel was identified as part of the tumor region from at least one patient.

2.4. Functional data processing

Only fMRI signals in intact voxels were considered in the following analyses. The first 10 volumes were discarded, and then slice timing and motion correction were performed. No patients exhibited a head motion > 3 mm maximum translation or 3° rotation. The motion-corrected functional images were first co-registered to the 3D-T1 images and were then spatially normalized into the MNI space by applying the deformation field estimated in segmentation. The normalized images were spatially smoothed using a Gaussian kernel (Full width at half maximum of 6 mm). The linear trend and the nuisance signals (36 parameters, including the $x$, $y$, $z$ translations and rotations) + WM/CSF/global time courses (9 parameters), plus their temporal derivatives (9 parameters) and the quadratic terms of 18 parameters (Circi et al., 2017; Satterthwaite et al., 2013) were removed by linear regression from each voxel’s time course. Then, temporal band-pass filtering (0.01–0.1 Hz) was performed on the residuals. A ‘scrubbing’ procedure was additionally adopted to remove any head motion artifact (Power et al., 2012; Yan et al., 2013). Specifically, rs-fMRI volumes that showed sudden head motion, based on the criterion of framewise displacement (FD) above 0.5 mm (Power et al., 2012, 2015, 2014) were discarded, together with one volume before and two volumes after the bad volume. No patient had volumes less than 140. Voxel-wise temporal signal-to-noise ratio (tSNR) was estimated after spatial normalization (Molloy et al., 2014; Triantafyllou et al., 2005), and was averaged over voxels in a whole brain mask. The whole brain mask was the binarized image of a brain parcellation atlas (Shen et al., 2013) used for network construction and feature extraction (described in the next section). Patients were excluded if the whole brain mean tSNR was less than 40.

2.5. Functional connectivity

Whole brain resting-state functional connectivity (rsFC). We adopted a functional brain parcellation atlas including 268 nodes (Shen et al., 2013) for network construction because this parcellation scheme included subcortical regions and cerebellum and was developed based on rsFC homogeneity. Individual functional networks were constructed by calculating the Pearson’s correlation coefficients between pairwise nodes (268 × 268). The $r$ values in each matrix were transformed to $z$ values using Fisher’s $r$-to-$z$ transformation. If a node fully injured, its connections with other nodes were set to 0.

Structural and functional data preprocessing was performed using SPM (http://www.fil.ion.ucl.ac.uk/spm) and Data Processing Assistant for Resting-State fMRI software (DPARSF; Chao-Gan and Yu-Feng, 2010; Yan et al., 2016; available at http://rfmri.org/DPARSF). The network construction was performed using the Graph-theoretical Network Analysis Toolkit (GRETNA; Wang et al., 2016; available at http://www.nitrc.org/projects/gretna/).

2.6. Machine learning-based brain-language prediction model

A schematic overview of our prediction framework is presented in Fig. 2. A rsFC feature selection procedure (Dosenbach et al., 2010) was implemented by ranking features according to their degrees of correlation coefficients with the AQ values, retaining only features with the highest correlation coefficients corresponding to a $P$ value < .005. The initial univariate feature-filtering step has been shown to improve overall model performance (Martino et al., 2008; Pereira et al., 2009). Note that we chose AQ as dependent variable because it reflected the global severity of language impairment.

Linear relevance vector regression (RVR). We selected to use RVR due to its demonstrated high prediction performance in brain-behavior/cognition mapping (Cui and Gong, 2018). RVR is a Bayesian framework for learning sparse regression models. It does not include an algorithm-specific parameter, so does not require extra computational resources for parameter estimation (Tipping, 2000). In RVR, only some samples (fewer than the training sample size), termed the ‘relevance vectors’, are used to fit the model:

$$y(x) = \sum_{i=1}^{m} w_i \tau_i + \epsilon$$

where $\tau$ are basis functions, and $\epsilon$ is normally distributed with mean 0 and variance $\beta$. RVR uses the training data to build a regression model:

$$y = \beta_0 + \epsilon$$

where $y = [y_1, y_2, ... y_n]^T$, $\beta = [\beta_0, \beta_1, ... \beta_n]$, $\beta_i = [\tau_i(x_1), \tau_i(x_2), ... \tau_i(x_n)]^T$. Each vector $\beta_i$ consists of the values of basis function $\tau_i$ for the input vectors and is a relevance vector. The model parameters $\beta$ were determined by using the maximum likelihood estimates from the conditional distribution:

$$p(y \mid x, \beta) = N(y \mid C \beta_0 + \Phi A^{-1} \Phi^T, \tau_0)$$

where $\tau_0$ is a vector of priors on the model parameters and $C = \beta_0 + \Phi A^{-1} \Phi^T$. To generate the RVM favor sparse regression models, prior distributions were assumed for both $\phi$ and $\tau_0$, i.e., $p(\beta_0 \mid \tau_0) = N(0, \tau_0^{-1})$ and resulted in the
same relevance vector machine construction. After finding the optimal point estimates for parameters $\alpha_i$ and $\beta_i$, the point estimate $w$ for weight vector $w$ was obtained from the equation $w = \beta_i (\Phi_i \Phi^T + A)^{-1} \Phi_i y_i$.

**Prediction accuracy and significance.** Leave-one-out-cross-validation (LOOCV) was used to calculate the prediction accuracy (the Pearson correlation coefficient between the predicted and actual labels). For each round of LOOCV, one patient was designated as the test sample and the remaining patients were used to train the model. The predicted score was then obtained from the feature matrix of the tested sample.

The significance level was computed based on 1000 permutation tests. For each permutation test, the prediction labels (i.e., the patients’ AQ) were randomized, and the same RVR prediction process used for the actual data was carried out. After 1000 permutations, a random distribution of accuracies was obtained and the $P$ value was correspondingly calculated as: $P = \frac{\text{number of permutation tests with accuracy less than actual accuracy}}{\text{number of permutation tests} + 1}$.

**Exploration of the weight-based language network.** The feature weight indicates the importance of each feature in the regression model. For the significant prediction models (permutation $P < 0.05$), the most predictive connections with the highest absolute weights were extracted. For each rsFC-based prediction model of AQ, the top 100 rsFC connections were extracted. In the linear RVR model, connections with positive weights indicate that increased rsFC predicts better performance, and connections with negative weights indicate that increased rsFC predicts worse performance.

For illustration purpose, we summed the absolute weights for each node of all its top connections, so the size of each node is relative to its total contribution to the model. rsFCs with top weights were then classified into six different types to determine the network lateralization: interhemispheric positive, interhemispheric negative, left or right intrahemispheric positive, and left or right intrahemispheric negative (Siegel et al., 2016). Finally, the numbers of rsFCs belonging to each of the six types were counted for each model.

### 2.7. Validation analysis

We validated our main results (model significance and top predictors) by further considering the following variables: 1) **Parcellation scheme.** To determine if our major results were affected by the specific brain parcellation scheme used, we repeated the prediction analyses using another brain parcellation—the Brainnetome atlas (Fan et al., 2016). The Brainnetome Atlas ($n=246$) was defined using both anatomical landmarks and connectivity-driven information. The anatomical regions defined by Desikan et al. (2006) were parcellated into subregions based on functional and structural connectivity data from 40 adults. Because the Brainnetome atlas only includes cortical and subcortical parcels, we created a whole brain parcellation ($n=271$) by adding the well-established cerebral subregions ($n=25$) created by Buckner et al. (2011) to the Brainnetome Atlas. 2) **Covariates.** We found that age was statistically significantly correlated with language scores for both LGG and HGG patients (Supplementary Table 1). Education showed statistically significant correlations with the language scores of HGG patients. We also found weak correlations of individual mean FD (Power et al., 2012; Yan et al., 2013) and tumor volume with language scores for both LGG and HGG patients. To minimize potential confounding effects of these covariates in model prediction, we calculated the partial correlations between the predicted AQ scores and the observed AQ scores including sex, age, education, FD, tumor volume, and site as covariates. 3) **Global signal removal.** Global signal removal is a controversial preprocessing step and recent work suggested that different preprocessing strategies may provide complementary insights into functional brain organization (Murphy and Fox, 2017). Global signal removal was performed in the main analysis to suppress the motion effect (Power et al., 2015), and we also reanalyzed our data without regressing out the global signal. 4) **Feature selection threshold.** We examined the effects of different feature selection significance thresholds on our main results by comparing the results with different feature selection thresholds (i.e., $P$ value $< .05$ and $P$ value $< .01$).

**Generalizability.** To assess the generalizability of the prediction model, we applied 10-fold cross-validation (Cui and Gong, 2018). For each group (i.e., LGG and HGG), all patients were divided into 10 subsets. To prevent random bias between subsets, we sorted the subjects according to their AQ scores and then assigned individuals with a rank of (1st, 11th, 21th, ...) to the first subset, (2nd, 12th, 22th, ... ) to the second subset, ..., and (10th, 20th, 30th, ...) to the tenth subset. Of the 10 subsets, 9 were combined as the training set, and the remaining subset was used as the testing set. A prediction model was constructed using all the training samples and then used to predict the scores of the testing samples. The Pearson correlation coefficients between the actual scores and the predicted scores were computed to quantify the accuracy of the prediction. The training and testing procedures were repeated 10 times so that each of the 10 subsets was used once as the testing set. To yield the final accuracies, we averaged the correlations across the 10 iteration. The significance level was computed based on 1000 permutation tests.

### 3. Results

#### 3.1. Demographics and language deficits in LGG and HGG patients

There was no significant difference in gender, education, FD, and tumor volumes between LGG and HGG patients (Table 1). Patients in the HGG group were of significantly higher age than those in the LGG group (two sample $t$-test, $t(124) = 2.144, P = 0.037$). More than 20% (17/77) of LGG patients had AQ scores less than 93.8, a clinical criterion of an aphasic subject. More than 45% (22/49) patients had AQ scores less than 93.8 (Supplementary Fig. 1). HGG patients had significantly lower AQ (two sample $t$-test, $t(124) = 3.02, P = 0.003$), spontaneous speech (two sample $t$-test, $t(124) = 2.13, P = 0.035$), comprehension (two sample $t$-test, $t(124) = 2.63, P = 0.009$), and naming scores (two sample $t$-test, $t(124) = 4.09, P = 0.001$) compared to the LGG patients. The replication scores of HGG patients were lower than those of LGG patients, but did not have statistical significance (two sample $t$-test, $t(124) = 1.56, P = 0.12$).

To see whether the language differences between LGG and HGG groups were due to the age difference or site effect, we then performed two-sample $t$-test for each kind of language test by adding age, sex, education, tumor volume, and site as covariates. After controlling for these potential confounding factors, we found that AQ ($t(119) = 2.81, P = 0.006$), comprehension ($t(119) = 2.75, P = 0.007$), and naming ($t(119) = 3.87, P = 1.7e-4$) scores of HGG patients were significantly lower than those of LGG patients. The spontaneous speech ($t(119) = 1.61, P = 0.01$) and repetition scores ($t(119) = 1.27, P = 0.208$) of HGG patients were lower than LGG patients but did not reach statistical significance.

#### 3.2. Whole-brain rsFCs predict AQ scores for both LGG and HGG patients

As shown in Fig. 3, the generated rsFC-based RVR models significantly predicted AQ scores for both LGG ($r = 0.299, permutation P = 0.032$) and HGG patients ($r = 0.521, permutation P = 0.007$). For each model, the top 100 rsFC features that exhibited the highest predictive power were extracted and projected back onto the brain (Fig. 4).

For LGG patients, nodes with top weights were distributed in the bilateral inferior frontal gyrus (IFG), middle frontal gyrus (MFG), superior frontal gyrus (SFG), anterior temporal lobe (ATL), middle temporal gyrus (MTG), inferior temporal gyrus (ITG), the inferior parietal lobe (IPL), the dorsal anterior cingulate cortex (dACC), fusiform, insula, subcortical regions, the posterior cerebellum, the brainstem, and the right middle occipital gyrus (MOG).

For HGG patients, the nodes with top weights were distributed in the bilateral IFG, MFG, SFG, ATL, MTG, STG, ITG, insula, subcortical...
regions, MOG, calcarine, precentral and postcentral gyrus, middle cingulate gyrus, precuneus, fusiform, the posterior cerebellum, the right IPL, angular gyrus, cuneus, and the brainstem. The right medulla oblongata had the highest weight, with connections to the precentral gyrus, ATL, MFG, and subcortical regions.

The connections of each rsFC-AQ model were further divided into subgroups. As shown in Fig. 5A, there were fewer ipsilesional intrahemispheric connections (15 vs 25) and more interhemispheric connections (55 vs 46) in LGG model than HGG model, and a comparable number of contralesional intrahemispheric connections (30 vs 29) for the two models. When considering the weight sign (positive or negative, Fig. 5B): LGG vs HGG: interhemispheric positive, 0.29 vs 0.27; interhemispheric negative, 0.26 vs 0.19; L intrahemispheric positive: 0.06 vs 0.15; L intrahemispheric negative: 0.09 vs 0.1; R intrahemispheric positive: 0.12 vs 0.15; R intrahemispheric negative: 0.18 vs 0.14.

We then assessed the predictive contribution of each anatomical network by averaging the weights of all nodes belonging to each network. As shown in Fig. 5D, the temporal, prefrontal, cerebellum, limbic, and parietal networks showed the most predictive contributions for the LGG model and the cerebellum, limbic, temporal, occipital, and prefrontal networks showed the most predictive contributions for the HGG model.

For both LGG and HGG models, nodes with top weights in the cerebellum included the bilateral cerebellar lobule VI, Crus I, Crus II, and VIIa. The regions with the highest weights in the right cerebellar lobule were Crus I and Crus II for LGG models and the regions with the highest weights in the left cerebellar lobule were IV-V, VI, VIII, X and the right pons for HGG patients (Figs. 4-6). Given the importance of the cerebro-cerebellar circuits in language processing, we calculated the number of connections from nodes in the cerebellum or brainstem to nodes in cortical and subcortical regions. For the 100 connections with top weights in each model, there were more cerebro-cerebellar connections present in the HGG model than in the LGG model, 37 vs 49 (Fig. 5C). The subtype numbers of cerebro-cerebellar connections in the LGG and HGG models were: ipsilesional intrahemispheric connections: 3 vs 19, interhemispheric connections: 19 vs 20, and contralateral intrahemispheric connections: 15 vs 11 (Fig. 6).

For LGG patients, the rsFC models of spontaneous speech (r = 0.359, permutation P = 0.033) and comprehension (r = 0.299, permutation P = 0.034) scores were significant.

3.3. Validation results

The validation results are summarized in Supplementary Tables 2 and 3.

The effect of Parcellation scheme. When using the Brainnetome atlas, rsFCs significantly predicted HGG patient’s AQ (r = 0.473, permutation P = 0.015) and repetition (r = 0.398, permutation P = 0.016) scores. All models for LGG patients did not perform well.

The effect of Covariates. We found that controlling for potential confounding variables did not change the model significance and correlations between real scores and predicted scores were only slightly changed.

The effect of global signals. Without removal of global signals, there were significant predictive accuracies in the rsFC models for HGG patient’s AQ (r = 0.539, permutation P = 0.003) and LGG patient’s naming (r = 0.344, permutation P = 0.02) scores.

The effect of rsFC feature selection threshold. When using a P value of 0.05 for rsFC feature selection, the predictive accuracies were significant in the rsFC models of HGG patient’s AQ (r = 0.454, P = 0.004), repetition (r = 0.251, P = 0.049) and naming (r = 0.278, P = 0.04) scores. A positive trend of significance was observed for LGG patient’s AQ (r = 0.21, P = 0.06) scores.

When using a P value of 0.01 for rsFC feature selection, the predictive accuracies were significant in the rsFC models of LGG patient’s AQ (r = 0.372, P = 0.01), comprehension (r = 0.237, P = 0.049) scores, and for HGG patient’s AQ (r = 0.478, P = 0.007) scores.
10-folder cross-validation. We found that rsFC-based RVR models significantly predicted AQ scores for HGG patients \( (r = 0.496, \text{permutation } P = 0.013) \) and comprehension scores \( (r = 0.493, \text{permutation } P = 0.002) \) for LGG patients. Positive trends of significance were observed for LGG patient's AQ \( (r = 0.288, P = 0.056) \) and repetition \( (r = 0.283, P = 0.058) \) scores.

4. Discussion

In this study, we performed functional network mapping of language by predicting aphasia scores based on the connectome-based functional connectivity for patients with left cerebral glioma involving language network. The main findings were as follows: 1) HGG patients showed more severe language impairments than LGG patients; 2) The rsFC model of HGG patients accounted for >27% of language variance, a percentage that was much higher than that of LGG patients (model only accounted for 0.09 of language variance); 3) For LGG patients, regions predictive of language impairments involved canonical language regions distributed in the left frontal, temporal, and parietal lobes, the subcortical regions, the right cerebellum and homotopic regions, whereas the language-related network of HGG patients showed heavy dependence on the left cerebellum, the right brainstem, the limbic system, and the temporal, prefrontal, and occipital lobes.

To our knowledge, this is the first study using a machine-learning prediction model to investigate the large-scale whole brain language network basis of patients with left cerebral glioma involving language network.

4.1. Individualized language prediction of glioma patients

We found that the language impairments of glioma patients could be significantly predicted by whole-brain functional connectivity, especially for HGG patients with severe language deficits. Our results are consistent with those of previous studies which found that language deficits after focal brain damage, especially tumors, traumatic brain injury, and stroke, could be predicted based on disruptions of network connectivity (Fang et al., 2015; Siegel et al., 2016; Warren et al., 2014; Yourganov et al., 2016). We also found much higher model accuracy of HGG patients than that of LGG patients. Numerous results have shown that slow infiltrative LGGs (generally over years) induce great plasticity within the central nervous system, often resulting in undetectable or only slight neurological deficits (Duffau, 2005; Zhang et al., 2018).
means that although there may be marked structural or functional changes in LGG patients (Almairac et al., 2018; Zhang et al., 2018), there was weak or no imaging-behavioral correlation. The strong brain-behavior relationship in HGG patients may be because the rapid (generally several months) and diffused growth of tumor cells infiltrating the normal brain tissue overwhelms the network plasticity and induces more severe and diverse language deficits as well as network disruptions.

4.2. Different network bases for language processing in LGG and HGG

In this work, we showed the functional connections of delocalized regions to support language processing for glioma patients. For both LGG and HGG, rsFC regions predictive of AQ involved canonical cerebral language areas, including the left IFG, STG, MTG, TP, the superior and posterior parietal cortex, insula, subcortical regions, and their homotopic regions. The recruitment of the right homotopic language regions has been well studied. In healthy (right-handed) subjects, right hemisphere language homologues, mainly the right IFG and the temporal lobe, participate in semantic, phonological, and syntactic processing (Tan et al., 2005; Vigneau et al., 2006; Kwok et al., 2019). For patients with left hemisphere damage, language-related structural and functional alterations have been frequently observed in the right hemisphere language homologues (Saur et al., 2008; Turkeltaub et al.,

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**Fig. 5.** The distributions of the 100 most predictive connections. A: Connections are divided into three groups: ipsilesional intrahemispheric (L), interhemispheric (Inter), and contralesional intrahemispheric (R). (B): Connections are further divided into six groups based on the sign of the connection weight (+: positive; -: negative). (C): The percentages of the cerebrocerebellar connections in each model. (D): The relatively predictive contributions of each predefined network based on anatomy. For each network, the weights of all nodes belonging to this network were summed.

**Fig. 6.** The most predictive cerebro-cerebellar connections and regions for the two rsFC-AQ models.
Third, di
tlanguage network after left hemisphere glioma than HGG patients.
showed a higher degree of integration of the right hemisphere into the
tumor-grade related shift from the left to the right, i.e., LGG patients
4.3. The cerebro-cerebellar circuits implicated in language processing for
processing for LGG and HGG patients. First, there are di

temperal and prefrontal regions showed the most predictions in LGG models, but
connections in the cerebellum, limbic system, the temporal, occip-
tal, and prefrontal lobes were leading predictors in the HGG model.
Second, based on the distribution and predictive contribution of rsFC, the language network of LGG is right-lateralized, but no obvious la-
teralization was observed in HGG patients. These observations are
consistent with the findings by Thiel et al. (2005) who showed that a
tumor-grade related shift from the left to the right, i.e., LGG patients
showed a higher degree of integration of the right hemisphere into the
language network after left hemisphere glioma than HGG patients. Third, different cerebro-cerebellar circuits are implicated in language
processing for LGG and HGG patients.

4.3. The cerebro-cerebellar circuits implicated in language processing for
both LGG and HGG patients

The roles of the cerebellum and cerebro-cerebellar circuits in lan-
guage processing have been verified in neuroanatomical, neuroima-
ging, and clinical studies (Buckner, 2013; De Smet et al., 2013; 
Marién et al., 2017). Anatomically, the cerebro-cerebellar circuit cons-
ists of two loops in an approximately homotopic manner. The feed-
backward (output) loop originates from the cerebellum and projects to the
motor (motor and premotor cortex) and non-motor cerebral associa-
tion areas (e.g., prefrontal cortex, the supplementary motor area, the
superior temporal and posterior parietal regions, cingulate gyrus, the
parahypocampal region and the limbic cortices) via the motor and non-
motor nuclei of the thalamus. The feed-forward (input) loop originate-
from the cerebral cortex and sends projections or inputs to the cen-
tralateral cerebellar cortex via the ipsilateral pons (corticopontine and
pontocerebellar mossy fiber pathways). Disruptions in specific cerebro-
cerebellar circuits could contribute to different behavioral symptoms.
The right cerebellum (i.e., the right cerebellar lobule VI, Crus I, Crus II)
is mainly involved in language, with a topographic organization of the
lateralized linguistic cerebellum (Marién et al., 2001). Cerebellar da-
mage induces functional suppression to distal regions that are anato-
mically and functionally connected supratentorial regions due to the
decrease or loss of transmission of excitatory impulses from the deep
nuclei of the cerebellum via the cerebello-ponto-thalamo-cerebral

pathways to the supratentorial brain regions, a phenomenon named
cerebello-cerebral diachisis.

Our previous work also showed the vital role of the cerebellum and
cerebro-cerebellar circuits in language network reorganization for
glioma patients following supratentorial lesion (Zhang et al., 2018). In
LGG patients, the cerebellar areas with significantly decreased sponta-
neous activity exhibited significantly increased rsFC with high-order cogni-
tive related supratentorial areas, including the right thalamus, ACC,
and posterior cingulate cortex. We extended our previous findings
and showed that a left supratentorial tumor lesion induced network
reorganization of cerebro-cerebellar circuits for both LGG and HGG
patients. In LGG patients, a right lateralized cerebro-cerebellar con-
nectivity pattern was observed, with the right cerebellar lobule Crus I as
the most predictive region, suggesting a nearly normal cerebro-cere-
bellar circuit for language processing. In HGG patients, more left
hemispheric cerebro-cerebellar connections were identified, with the
most predictive regions as the left cerebellar lobule IV-V, VI, VIII and X,
and the right medulla oblongata. The shifting of the cerebro-cerebellar
circuits to the ipsilesional side as observed in HGG patients suggested
disruptions by the left supratentorial glioma of the functionally later-
alized cerebro-cerebellar circuits for language and a homotopic cer-
bro-cerebellar circuits emerged to support language processing.

4.4. Non-language functional systems that support language processing

In addition to the language network, several non-language func-
tional systems were also found, such as the limbic system for both LGG
and HGG models and the occipital and motor networks for HGG models.
The implication of them may be because the language network showed
high interactions with the functional systems involved in visual pro-
cessing, motor processing, emotion processing, executive control,
working memory, and default model network (Duffau et al., 2014; 
Hernández et al., 2019; Siegel et al., 2016; Xu et al., 2016). Language
deficits can be induced by the disruptions of bilaterally distributed
support processes, such as visual attention required for reading or
motor planning for speech (Connor et al., 2000; Otten et al., 2012), and
language recovery is highly related to the recovery of memory and
attention (Ramsey et al., 2017).

5. Limitations

There are several limitations of this study.
First, prediction accuracy and stability of machine learning models
are both exponentially related to the sample size, where a minimum
sample size of 200 is needed to achieve highest model performance
(Cui and Gong, 2018). Although we adopted data from two centers, the
two study groups were small (LGG: 77 patients; HGG: 49 patients). A
large sample size from other centers will be needed to reach stable and
higher model performance and also ensure the representativeness of the
results (Corbetta et al., 2015).
Second, the overall prediction accuracies of the two groups, espe-
cially the LGG group, are relatively low (r values, 0.2–0.5). Focal tumor
damage can induce both structural and functional changes and joint
prediction with cooperating multi-modal imaging features (e.g., GM
volume and white-matter connectivity) can improve prediction accu-
cy (Dai et al., 2012; Liem et al., 2017). However, considering the
slight language impairments in most LGG patients, multi-modal fea-
tures may not improve overall model accuracy due to the weak brain-
behavior relationship.
Third, most of our patients were enrolled before 2016 and thus the
pathological stratification in this study was based on the 2007 WHO
classification system. Future studies that utilize the new 2016 WHO
classification system are needed to delineate the language network re-
organization for each molecular glioma subtype.
6. Conclusions

The whole-brain rsFCs prediction model can significantly predict the language variance of preoperative patients with left cerebral glioma, especially for HGG patients with more severe language impairments. For both LGG and HGG patients, large-scale language connections were identified which bilaterally distributed in prefrontal, temporal, and parietal lobes, posterior cerebellar lobule, limbic system, brainstem, and subcortical regions. Our results suggest a significant role of right hemisphere language homologues for LGG and a strong dependence of ipsilesional cerebro-cerebellar circuits for HGG. These results advance our understanding of the large-scale language network reorganization of preoperative glioma patients, and provide insight into why areas thought to be critical in language processing can be resected without inducing permanent deficit.

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CRediT authorship contribution statement

Binke Yuan: Data curation, Writing - original draft. Nan Zhang: Data curation, Writing - original draft. Jing Yan: Writing - original draft. Jingliang Cheng: Conceptualization, Supervision, Writing - original draft. Junfeng Lu: Conceptualization, Supervision, Data curation, Writing - original draft. Jinsong Wu: Conceptualization, Supervision, Data curation, Writing - original draft.

Declaration of Competing Interest

The authors declare no competing financial interests.

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Supplementary materials

Supplementary material associated with this article can be found, in the online version, at doi:10.1016/j.jic.2019.102023.

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