Research Article

Altered Small-World Functional Network Topology in Patients with Optic Neuritis: A Resting-State fMRI Study

Ke Song,1,2 Juan Li,3 Yuanqiang Zhu,4 Fang Ren,4 Lingcan Cao,1 and Zi-Gang Huang1

1The Key Laboratory of Biomedical Information Engineering of Ministry of Education, Institute of Health and Rehabilitation Science, School of Life Science and Technology, Xi’an Jiaotong University, The Key Laboratory of Neuro-informatics & Rehabilitation Engineering of Ministry of Civil Affairs, Xi’an, Shaanxi 710049, China
2Department of Equipment, Xi’an People’s Hospital (Xi’an Fourth Hospital), China
3Shaanxi Eye Hospital, Xi’an People’s Hospital (Xi’an Fourth Hospital), Affiliated Guangren Hospital, School of Medicine, Xi’an Jiaotong University, Xi’an, 710004, China
4Department of Radiology, Xijing Hospital, Fourth Military Medical University, Xi’an, 710032 Shaanxi, China

Correspondence should be addressed to Zi-Gang Huang; huangzg@xjtu.edu.cn

Aim. This study investigated changes in small-world topology and brain functional connectivity in patients with optic neuritis (ON) by resting-state functional magnetic resonance imaging (rs-fMRI) and based on graph theory.

Methods. A total of 21 patients with ON (8 males and 13 females) and 21 matched healthy control subjects (8 males and 13 females) were enrolled and underwent rs-fMRI. Data were preprocessed and the brain was divided into 116 regions of interest. Small-world network parameters and area under the integral curve (AUC) were calculated from pairwise brain interval correlation coefficients. Differences in brain network parameter AUCs between the 2 groups were evaluated with the independent sample t-test, and changes in brain connection strength between ON patients and control subjects were assessed by network-based statistical analysis.

Results. In the sparsity range from 0.08 to 0.48, both groups exhibited small-world attributes. Compared to the control group, global network efficiency, normalized clustering coefficient, and small-world value were higher whereas the clustering coefficient value was lower in ON patients. There were no differences in characteristic path length, local network efficiency, and normalized characteristic path length between groups. In addition, ON patients had lower brain functional connectivity strength among the Rolandic operculum, medial superior frontal gyrus, insula, median cingulate and paracingulate gyri, amygdala, superior parietal gyrus, inferior parietal gyrus, supramarginal gyrus, angular gyrus, lenticular nucleus, pallidum, superior temporal gyrus, and cerebellum compared to the control group (P < 0.05).

Conclusion. Patients with ON show typical “small world” topology that differed from that detected in HC brain networks. The brain network in ON has a small-world attribute but shows reduced and abnormal connectivity compared to normal subjects and likely causes symptoms of cognitive impairment.

1. Introduction

Optic neuritis (ON) is a condition affecting 115 out of every 100,000 persons [1]; it is characterized by inflammation and demyelination of the optic nerve as a result of infection or systemic autoimmune disease. The main clinical symptoms are sudden loss of visual acuity in one or both eyes within a short period of time, relative afferent pupil disorder (RAPD), papillary edema, pain during eye rotation, and visual field defect. ON is closely related to demyelinating diseases of the central nervous system such as optic neuromyelitis and multiple sclerosis, among others [2]. Thinning of the retinal nerve fiber layer around the optic papilla in ON is observable by optical coherence tomography. ON can cause severe visual impairment but the pathogenesis is not fully understood, although it is thought to involve inflammation or immune factors that lead to optic nerve damage and ganglion cell apoptosis. In addition to demyelination, ON patients have abnormal activity in many brain regions [3]. For example, brain atrophy was observed in patients with chronic...
recurrent solitary ON [4], along with Wallerian degeneration in the optic tract, cerebellum, thalamus, posterior cingulate, and other brain areas [5], indicating that specific brain areas are affected in ON. Given that the visual loss caused by ON can negatively affect the quality of life of patients, it is important to clarify the pathogenesis and associated changes in the brain. Resting-state functional magnetic resonance imaging (rs-fMRI) is a safe and widely used method for evaluating brain activity based on the detection of the balance between local segregation and global integration of signals associated with interconnected neurons. The normal human brain network has a short path length and high transmission efficiency, known as small-world attributes [6]; these enable the brain network to meet local and global demands and balance functional integration and segregation in order to achieve synchronization of neural activity between different brain regions. Thus, small-world attributes allow efficient information transmission at low wiring cost [7].

Graph theory, which is the study of the topologic structure of networks, provides a means of quantifying network changes in the brain [8] by considering these as a set of general elements sharing a specific relationship beyond anatomic connections [9]. Graph theory can be applied to rs-fMRI to characterize the functional connectivity and obtain a structural map of the brain from functional data, which can provide insight into the anatomic basis of brain dysfunction [10] and thus serves as an important reference for the diagnosis and treatment of diseases. Graph theory analysis has been widely used in studies on the mechanisms of posttraumatic stress disorder [11], Alzheimer’s disease [12], schizophrenia [13], stroke [14], epilepsy [15], and concussion [16].

Most previous studies on brain abnormalities in ON have focused on abnormal activity in specific brain regions. However, changes in small-world attributes and brain functional connectivity caused by ON have not been assessed. This was investigated in the present study by comparing small-world attributes and brain network connectivity in patients with ON and normal subjects by rs-MRI and the application of graph theory.

2. Materials and Methods

2.1. Subjects. A total of 21 ON patients (8 males and 13 females) were recruited at the Department of Ophthalmology, the First Affiliated Hospital of Nanchang University Hospital. Inclusion criteria were as follows: (1) sudden loss of visual acuity in one or both eyes within a short period of time; (2) positive RAPD or abnormal visual evoked potentials; (3) no visual field abnormalities related to nerve fiber injury; (4) none of the subjects had a history of psychiatric or neurological disorders; (5) no acute visual loss caused by other ophthalmologic or nervous system diseases; (6) no history of mental disorders, diabetes, hypertension, and taking psychotropic drugs; (7) no history of drug, smoking, or alcohol addiction; and (8) average somatotype and weight (Figure 1). We also recruited 21 age-, sex-, and weight-matched healthy control (HC) subjects (8 males and 13 females) who met the following criteria: (1) no pathway or brain parenchyma abnormalities observed by head MRI, (2) no ophthalmic disease and maximum corrected visual acuity > 1.0, (3) no neuropsychiatric abnormalities or headache, and (4) no contraindications for MRI. After being informed of the nature of the study, all patients (or their guardians for participants under 18 years old) provided written and informed consent before participating. The study was approved by the research ethics committee of the First Affiliated Hospital of Nanchang University, and the protocol was in accordance with the Helsinki Declaration (CDYFY-LL-2015-29).

2.2. MRI Data Acquisition. A 3.0 T TrioTim (Siemens, Munich, Germany) MR scanner and 8-channel head coil were used to collect rs-fMRI data and 3-dimensional high-resolution T1-weighted anatomic images. Participants were instructed to avoid drinking alcohol or coffee before the scan and those with intracranial lesions were excluded. During the scan, the subjects lay quietly with their eyes closed, breathe evenly and remain still, and avoid any mental activities insofar as possible. The subject was used a sponge pad to fix the head to reduce head movement, and the subject wore earplugs to block noise. For rs-fMRI, the parameters were as follows: repetition time (TR) = 2000 ms, echo time (TE) = 40 ms, flip angle = 90°, field of view (FOV) = 230 mm × 230 mm, matrix = 64 × 64, slice thickness = 4 mm, and slice number = 240 mm. The scanning parameters for T1-weighted structural images were as follows: TR = 1900 ms, TE = 2.26 ms, flip angle = 9°, FOV = 250 × 250 mm, matrix = 256 × 256, layer thickness = 1 mm, and layer number = 176.

2.3. MRI Data Preprocessing. In order to eliminate the impact of magnetic field uniformity on network computing and intrasubject variability, poor-quality data were removed along with the first 10 time points. This study is based on the MATLAB 2014a (Mathworks, Natick, MA, USA) platform and uses the DPARSF V2.3 software package to preprocess the data. The preprocessing steps involved realigning, slice timing, time layer correction, spatial standardization, and smoothing with a 6 × 6 × 6 mm full-width (half-height) Gaussian kernel [17]. Subjects with maximum x, y, or z displacement > 1.5 mm or angular motion > 1.5° were excluded from the analysis. In addition, the mean frame-wise displacement (FD) was calculated as a measure of the microhead motion of each subject. Linear regression was used to remove nuisance variables containing signals from the region of interest (ROI) in the ventricles and areas centered on white matter. After correcting for head motion, the standard echo plane imaging template was used to normalize the fMRI image space to the Montreal Neurological Institute (MNI) space with resampling at a resolution of 3 × 3 × 3 mm. Finally, the data were then detrended to remove linear drift and temporally filtered by hand-pass (0.01–0.08 Hz) to reduce the effects of low-frequency drift and high-frequency noise.

2.4. Construction of Brain Network Structure and Analysis of Topologic Attributes. We divided the whole brain into 116 network nodes. To define the ROI for the node region, the average processing time was calculated as the average of the fMRI time series of all voxels in the region, and the average time series of each region was obtained. We used a weighted
to construct a full-connection weighted network of the brain with sparsity as the threshold. The network analysis was carried out under a sparsity of 0.08–0.48 with an interval of 0.01. For the graph of each subject, we evaluated the whole-brain connectivity between the groups, using an independent two-sample t-test with P < 0.05 and permutations of 5,000.

3. Results

3.1. Demographic and Clinical Characteristics. There were no statistically significant differences between the ON and HC groups in terms of weight ($P = 0.652$), age ($P = 0.821$), height ($P = 0.634$), and BMI ($P = 0.963$), while significant differences were found in VA-Right ($P < 0.001$) and best-corrected VA-Left ($P < 0.001$) between the two groups.

3.2. Analysis of Small-World Properties. Seven topologic small-world parameters were determined under a sparsity of 0.08 to 0.48 with an interval of 0.01. $C_p$, $L_p$, $E_g$, $E_l$, $\sigma$, $\alpha_y$, and $\alpha_\lambda$ were positively correlated whereas $L_p$, $\gamma$, $\lambda$, and $\sigma$ were negatively correlated with sparsity. Both groups had small-world attributes ($\lambda > 1$, $\gamma > 1$, $\sigma > 1$). For the small-world indices, $\alpha_y$ (Figure 2(b)) and $\alpha_\sigma$ (Figure 2(c)) were significantly higher for ON patients than for HC subjects ($P < 0.05$). There were no statistically significant differences in $\alpha_\lambda$ (Figure 2(a)) between groups (Table 1). For the other indicators of brain network topology, $aC_p$ (Figure 3(a)) was lower, whereas $aE_g$ (Figure 3(b)) was higher in ON patients compared to HC subjects (both $P < 0.05$). There were no significant

Figure 1: Eye examination data of ON patients. Notes: (a) The results of fundus photography in ON patients. (b) The results of fundus fluorescein angiography in ON patients. Abbreviations: ON: optic neuritis; FFA: fundus fluorescein angiography.
These results demonstrate that optic nerve inflammation has a far-reaching effect on the functional brain network. Compared to the HC group, ON patients showed decreased brain functional connectivity. These mainly occurred among rolandic operculum (ROL), medial superior frontal gyrus (SFG), insula (INS), median cingulate and paracingulate gyri (DCG), amygdala (AMYG), superior parietal gyrus (SPG), inferior parietal gyrus (IPL), supramarginal gyrus (SMG), angular gyrus (ANG), lenticular nucleus, pallidum (PAL), superior temporal gyrus (STG), and cerebellum (Figure 4). The difference is statistically significant ($P < 0.05$) (Table 2). There were no instances where functional connectivity was higher in the ON group than in the HC group. These results demonstrate that optic nerve inflammation has a far-reaching effect on the functional brain network.

### Table 1: The AUC of the small-world parameters in patients with ONs and HC.

| Network properties | ON         | HCs        | t      | P values |
|--------------------|------------|------------|--------|----------|
| aCP                | $0.245 \pm 0.100^*$ | $0.253 \pm 0.009$ | -2.714 | 0.01     |
| aLP                | $0.712 \pm 0.027$ | $0.728 \pm 0.269$ | -1.865 | 0.069    |
| aEI                | $0.310 \pm 0.006$ | $0.313 \pm 0.008$ | -1.239 | 0.223    |
| aEg                | $0.234 \pm 0.005^*$ | $0.229 \pm 0.006$ | 2.812  | 0.008    |
| Small-world attribute | —       | —         | —      | —        |
| aλ                 | $0.425 \pm 0.008$ | $0.429 \pm 0.007$ | -1.551 | 1.129    |
| aγ                 | $0.655 \pm 0.087^*$ | $0.587 \pm 0.071$ | 2.768  | 0.008    |
| aσ                 | $0.610 \pm 0.084^*$ | $0.550 \pm 0.057$ | 2.664  | 0.01     |

Notes: significant at $^*P < 0.05$, independent t-test. P, P value between ON and HC. Abbreviation: ON: optic neuritis; HCs: healthy controls; AUC: area under curve; aCP: the AUC of clustering coefficient; aLP: the AUC of characteristic path length; aγ: the AUC of normalized clustering coefficient; aλ: the AUC of normalized characteristic path length; aσ: the AUC of small-worldness; aEg: the AUC of global network efficiency; aEi: the AUC of local network efficiency.

### 4. Discussion

This is the first study to use graph theory and NBS to analyze the small world and brain functional connectivity strength in ON. Inflammation and immune activation can cause demyelination of the optic nerve, leading to reduced signal transmission and visual impairment [21]. The demyelination and axon damage associated with ON was shown to result in the destruction of functional brain networks in a small-world study of craniocerebral trauma.

In the present work, we found that ON patients retained small-world characteristics ($\lambda > 1, \gamma > 1, \sigma > 1$) although some network parameters were altered. Compared to HCs, Eg, γ, and σ were increased whereas Cp was decreased in ON patients; moreover, Cp, El, and Eg were positively correlated and Lp, γ, λ, and σ were negatively correlated with sparsity. The lower Cp in ON patients may reflect a reduced...
functional connectivity in some brain regions caused by extreme demyelination, which has also been observed in cases of axonal injury. Patients with long-term disturbance of consciousness also show alterations in small-world parameters. Lp measures the capacity for global information transmission and is related to cognitive function [22], while Eg represents global network efficiency. In our study, Eg was higher in ON patients than in HC subjects, suggesting greater efficiency in network information transmission. Patients with ON often have monocular disease. Insufficient stimulation of the visual cortex from the decreased visual acuity in one eye can lead to compensatory activation of the contralateral brain region [23]. We observed an increase in the amplitude of low-frequency fluctuation (ALFF) value of the left superior temporal gyrus in ON patients, consistent with findings from other MRI studies of ON. This implies that the increase in Eg in ON patients is a mechanism to offset brain network dysfunction. The decreased λ value was accompanied by a compensatory reduction in Lp. The injury caused by ON alters small-world attributes of brain connectivity networks. The parameter σ is a clustering coefficient that represents approximate shortest path length (Y. [24]). A lower σ represents a greater tendency toward a random brain network; in concussion, these were shown to be more susceptible to pathologic insults than small-world networks.

In this study, σ was >1 in ON patients, indicating that they have small-world characteristics; however, the value was higher than that in HC subjects. The optic nerve is rich in macrophages and T cells [25]; thus, the increase in σ may be attributable to inflammation or damage. γ is the standardized clustering coefficient and is used to measure dispersion within the network, with a higher value indicating a higher degree of grouping. In patients with ON, damage to the optic nerve may lead to damage to the related cerebral hemispheres, which may lead to changes in the value of brain function areas and cognitive dysfunction. The significant difference in the static state connection network between ON patients and HC subjects emphasizes the role of NBS in multivariate comparisons.

We observed a decrease in the connection strength of multiple brain regions in ON patients. The INS is located in the deep part of the lateral sulcus at the boundary between the annular sulcus and frontal, temporal, and parietal lobes. Abnormal activation of the insular-interstitial area has been reported in ON patients [26]; additionally, the latency of visual evoked potentials decreased with REHO signal in the

---

**Figure 3: Comparison of structural properties of brain network between ON patients and HCs.** Notes: compared with HCs, the value of Cp in patients with ON was significantly lower (a) and the value of Eg was significantly higher (b), and the difference was statistically significant (P < 0.05). There was no significant difference in the values of Lp (c) and EI (d) between the two groups (P > 0.05). Abbreviation: ON: optic neuritis; HCs: healthy controls; AUC: area under curve; aCp: the AUC of node clustering coefficient; aLp: the AUC of characteristic path length; aEg: the AUC of global network efficiency; aEl: the AUC of local network efficiency.
INS, which could decrease the connection strength between this and other brain regions [3]. The cerebellum is located in the inferior part of the brain posterior to the medulla oblongata and pons. Cerebellar demyelination has been reported in ON [27], which could explain the reduced connectivity between the cerebellum and other brain regions in patients.

The STG is located in the temporal lobe between the lateral and superior temporal sulci and plays a key role in sound processing. An fMRI study revealed that the ALFF signal was decreased in the superior temporal gyrus of ON patients, which may be related to the severity of ON. The STG is also involved in visual searching and decreased magnetic resonance-related signals in this region have been observed in patients with retinal detachment [28]. An impaired STG in patients with ON could result in decreased connectivity with surrounding brain regions. The AMYG is located in the dorsomedial part of the anterior temporal lobe, slightly anterior to the top of the hippocampus and inferior horn of the lateral ventricle. As part of the limbic system, the AMYG plays an important role in generating, identifying, and regulating emotion, and it is among the key brain areas responsible for normal and pathologic stress responses. A decreased connection strength between the AMYG and other brain areas in ON patients may indicate a reduced ability to respond to pathologic events, leading to the destruction of the brain network structure. The SMG contributes to the maintenance of short-term auditory-language, motor, and visual-spatial memory sequence [29]; the reduced connectivity between the SMG and surrounding areas in ON patients suggests that the normal perception of visual space is disrupted.

The ANG, which is the visual language (reading) center, is arched around the end of the supratemporal sulcus in the temporal lobe. The ANG integrates incoming sensory and cognitive information, responds to stimuli in memory and learning, and functions in memory retrieval [30]. The activity of ANG-related neural circuits is increased during eye-to-eye communication, and both the ANG and STG have been implicated in Wernicke’s (sensory) aphasia. Therefore, the decreased connectivity between the ANG and STG in ON may be associated with reading dysfunction. The SFG, located in the upper part of the prefrontal lobe, is involved in motor coordination, working memory, and resting-state and cognitive control. The fractional ALFF signal of the SFG was shown to be positively correlated with perceived stress, and the gray matter structure of the SFG has been

---

**Figure 4:** Graph theory analysis of alterations in brain functional connectivity. Notes: The figure shows the subnetwork with decreased connectivity in ON patients compared to controls, identified by the NBS. Decreased brain functional connectivity in individuals with optic neuritis (ONs) compared to healthy controls. (NBS: $T > 3.92, P < 0.05$ and 5,000 permutations). Abbreviation: ON: optic neuritis; HCs: healthy controls; ROL: rolandic operculum; SFGmed: superior frontal gyrus, medial; INS: insula; DCG: median cingulate and paracingulate gyri; AMYG: amygdala; SPG: superior parietal gyrus; IPL: inferior parietal gyrus; SMG: supramarginal gyrus; ANG: angular gyrus; PAL: lenticular nucleus, pallidum; STG: superior temporal gyrus; CRBL: cerebellum Crus1; CRBL 6: cerebellum Crus6; L: left hemisphere; R: right hemisphere.
implicated in the processing of early and recent life stress events. The reduced SFG connection strength in ON patients may be due to stress caused by optic nerve inflammation. The SFG is located in the dorsomedial parietal lobe anterior to the parietal-occipital sulcus and above the parietal sulcus. In the posterior part of the retrocentral sulcus, the SFG participates in stereoscopic visual processing and plays a key role in defining visual space in language and motor areas. Additionally, the SFG controls eye movement. Eye rotation pain is common in ON and may also be associated with changes in brain connectivity; however, in our study, the strength of the connection between the SFG and surrounding brain regions was decreased, suggesting that there was damage to the area corresponding to eye movement pain. The PAL is located in the lentiform nucleus of the striatum. Lesions involving the extrapyramidal system and pyramidal tract may cause movement disorder and nystagmus in the eyes. The DCG, which is crescent-shaped and surrounds the corpus callosum, is a major component of the limbic system and is related to memory and spatial orientation. The reduced connection strength between the DCG and other brain areas in patients with ON could affect their capacity for spatial localization. ROL is the cortex adjacent to the insular, which is one of the major regions involved in the language processing system, and it also involves in motor, sensory, autonomic, and cognitive processing. Relevant research data also confirms the role of rolandic operculum and neighboring areas (such as insular) in processing sensory signals related to other conscious operations (such as visual awareness). The decrease of the connection strength between ROL and the surrounding brain area may reflect the impairment of visual function. IPL is a part of the parietal lobe and is related to visual recognition and selective scanning targets. Finally, the observed changes in the IPL imply the impairment of stereoscopic visual function. Taken together, these results demonstrate that optic nerve inflammation has far-reaching effects on the functional brain network.

4.1. Limitations and Strengths. There were some limitations to this study. Because of the small sample size, we did not examine the correlation between brain topologic characteristics and clinical manifestations of ON or between changes in brain structure and function. In the future, brain network changes in ON will be analyzed in a larger cohort by multimodal analysis.

5. Conclusion

The results of this study show that the functional brain network of ON patients has small-world properties, but that these are significantly impaired relative to HC subjects. The changes in small-world properties observed in ON may be caused by demyelination resulting from inflammation and could reflect functional impairment in the brain. Our findings have found that optic neuritis may have a certain impact on the functional areas of the brain. Based on this, we can diagnose optic neuritis through brain image analysis and prevent possible complications related to brain dysfunction.

Data Availability

The datasets generated during and/or analyzed during the current study are available from the corresponding author on reasonable request.

Consent

All authors agree to publish.

Conflicts of Interest

This was not an industry-supported study. The authors report no conflicts of interest in this work.

Authors’ Contributions

KS, JL, and YQZ designed the current study. FR collected the data. ZGH analyzed the data. KS wrote the manuscript. All the authors read and approved the final manuscript.

Acknowledgments

This manuscript has been released as a preprint at bioRxiv 2020.06.09.141432, Altered small-world functional network topology in patients with optic neuritis: a resting-state fMRI study. Ke Song, Juan Li, Yuanqiang Zhu, Fang Ren, Lingcan Cao, Yi Shao, Zi-Gang Huang. bioRxiv 2020.06.09.141432; doi:10.1101/2020.06.09.141432. This work was supported
by NSFC (No. 11975178), the Project Supported by Natural Science Basic Research Plan in Shaanxi Province of China (No. 2020JM-058), Natural Science Basic Research Program of Shaanxi (No.2020JQ-096), and the Scientific Research Foundation of Xi’an People’s Hospital (Xi’an Fourth Hospital) (No. FZ-45). ZGH acknowledges support of K. C. Wong Education Foundation.

References

[1] H. Wilhelm and M. Schabet, “The diagnosis and treatment of optic neuritis,” Deutsches rzteblatt International, vol. 112, no. 37, p. 616, 2015.

[2] C. A. Wicki, P. Manogaran, T. Simic, J. V. M. Hanson, and S. Schippling, “Bilateral retinal pathology following a first-ever clinical episode of autoimmune optic neuritis,” Neurology Neuroimmunology & Neuroinflammation, vol. 7, no. 2, article e671, 2020.

[3] Y. Shao, F. Cai, Y. Zhong et al., “Altered intrinsic regional spontaneous brain activity in patients with optic neuritis: a resting-state functional magnetic resonance imaging study,” Neuropsychiatric Disease & Treatment, vol. 11, pp. 3065–3073, 2015.

[4] L. N. Cantó, S. C. Boscá, C. A. Vicente et al., “Brain atrophy in relapsing optic neuritis is associated with crión phenotype,” Frontiers in Neurology, vol. 10, p. 1157, 2019.

[5] A. I. Colpak, A. T. Kurne, K. K. Oguz, A. C. Has, A. Dolgun, and T. Kansu, “White matter involvement beyond the optic nerves in CRION as assessed by diffusion tensor imaging,” International Journal of Neuroscience, vol. 125, no. 1, pp. 10–17, 2015.

[6] L. T. Chen, X. L. Fan, H. J. Li et al., “Disrupted small-world brain functional network topology in male patients with severe obstructive sleep apnea revealed by resting-state fMRI,” Neuropsychiatric Disease & Treatment, vol. 13, pp. 1471–1482, 2017.

[7] L. X. Wang, F. Guo, Y. Q. Zhu et al., “Effect of second-generation antipsychotics on brain network topology in first-episode schizophrenia: a longitudinal rs-fMRI study,” Schizophrenia Research, vol. 208, pp. 160–166, 2019.

[8] Z. A. Gaal, R. Boha, C. J. Stam, and M. Molnár, “Age-dependent features of EEG-reactivity–Spectral, complexity, and network characteristics,” Neuroscience Letters, vol. 479, no. 1, pp. 79–84, 2010.

[9] D. Papo, J. M. Buldú, S. Boccaletti, and E. T. Bullmore, “Complex network theory and the brain,” Philosophical Transactions of the Royal Society B: Biological Sciences, vol. 369, no. 1653, article 20130520, 2014.

[10] F. Miraglia, F. Vecchio, and P. M. Rossini, “7T, EEG characteristics in ‘eyes open’ vs ‘eyes closed’ conditions: Small world network architecture in healthy aging and age-related brain degeneration,” Clinical Neurophysiology Official Journal of the International Federation of Clinical Neurophysiology, vol. 127, no. 4, pp. e134–e1268, 2016.

[11] J. A. Rowland, J. R. Stapleton-Kotloski, D. L. Dobbins, E. Rogers, D. W. Godwin, and K. H. Taber, “Increased Small-World Network Topology Following Deployment-Acquired Traumatic Brain Injury Associated with the Development of Post-Traumatic Stress Disorder,” Brain Connectivity, vol. 8, no. 4, pp. 205–211, 2018.

[12] C. A. Frantzidis, A. B. Vivas, A. Tsołaki, M. A. Klados, M. Tsołaki, and P. D. Bamidis, “Functional disorganization of small-world brain networks in mild Alzheimer’s disease and amnestic mild cognitive impairment: an EEG study using relative wavelet entropy (RWE),” Frontiers in Aging Neuroscience, vol. 6, 2014.

[13] A. Anderson and M. S. Cohen, “Decreased small-world functional network connectivity and clustering across resting state networks in schizophrenia: an fMRI classification tutorial,” Frontiers in Human Neuroscience, vol. 7, p. 520, 2013.

[14] F. Vecchio, F. Miraglia, A. Romano, P. Bramanti, and P. M. Rossini, “Small world brain network characteristics during EEG Holter recording of a stroke event,” Clinical Neurophysiology, vol. 128, no. 1, pp. 1–3, 2017.

[15] R. D. Bharath, G. Chaitanya, R. Panda et al., “Reduced small world brain connectivity in probands with a family history of epilepsy,” European Journal of Neurology, vol. 23, no. 12, pp. 1729–1737, 2016.

[16] Y. A. Yan, J. Song, G. Xu et al., “Correlation between standardized assessment of concussion scores and small- world brain network in mild traumatic brain injury,” Journal of Clinical Neuroscience, vol. 44, pp. 114–121, 2017.

[17] X. Liu, Z. Yan, T. Wang et al., “Connectivity pattern differences bilaterally in the cerebellum posterior lobe in healthy subjects after normal sleep and sleep deprivation: a resting-state functional MRI study,” Neuropsychiatric Disease & Treatment, vol. 11, p. 1279, 2015.

[18] Y. Huang, Y. Liu, D. Zhao et al., “Small-world properties of the whole-brain functional networks in patients with obstructive sleep apnea-hypopnea syndrome,” Sleep Medicine, vol. 62, pp. 53–58, 2019.

[19] F. de Vico Fallani, J. Richiardi, M. Chavez, and S. Achard, “Graph analysis of functional brain networks: practical issues in translational neuroscience,” Philosophical Transactions of the Royal Society B: Biological Sciences, vol. 369, no. 1653, article 20130521, 2014.

[20] A. Zalesky, A. Fornito, and E. T. Bullmore, “Network-based statistic: identifying differences in brain networks,” NeuroImage, vol. 53, no. 4, pp. 1197–1207, 2010.

[21] T. H. Lee, Y. S. Ji, S. W. Park, and H. Heo, “Retinal ganglion cell and axonal loss in optic neuritis: risk factors and visual functions,” Eye, vol. 31, no. 3, pp. 467–474, 2017.

[22] C. Y. Wee, Z. Zhao, P. T. Yap et al., “Disrupted brain functional network in internet addiction disorder: a resting-state functional magnetic resonance imaging study,” Plos One, vol. 9, no. 9, article e107306, 2014.

[23] Y. Shao, X. Huang, F. Cai et al., “Disturbed spontaneous brain-activity pattern in patients with optic neuritis using amplitude of low-frequency fluctuation: a functional magnetic resonance imaging study,” Neuropsychiatric Disease & Treatment, vol. 11, pp. 3075–3083, 2015.

[24] Y. Zhang, J. Liu, L. Li et al., “A study on small-world brain functional networks altered by postherpetic neuralgia,” Magnetic Resonance Imaging, vol. 32, no. 4, pp. 359–365, 2014.

[25] J. D. Glenn, P. Xue, and K. A. Whartenby, “Gemcitabine directly inhibits effector CD4 T cell activation and prevents experimental autoimmune encephalomyelitis,” Journal of Neuroimmunology, vol. 316, pp. 7–16, 2018.

[26] A. T. Toosy, D. J. Werring, E. T. Bullmore et al., “Functional magnetic resonance imaging of the cortical response to photic
stimulation in humans following optic neuritis recovery,” *Neuroscience Letters*, vol. 330, no. 3, pp. 255–259, 2002.

[27] A. Kutzelnigg, J. C. Faber-Rod, J. Bauer et al., “Widespread Demyelination in the Cerebellar Cortex in Multiple Sclerosis,” *Brain Pathology*, vol. 17, no. 1, pp. 38–44, 2007.

[28] Q. Yuan, H. H. Kang, W. Q. Shi et al., “Disturbed interhemispheric functional connectivity in visual pathway in individuals with unilateral retinal detachment: a resting state fMRI study,” *Visual Neuroscience*, vol. 35, p. 35, 2018.

[29] J. T. Zhang, Y. Liu, L. X. Li, K. Li, J. G. Chen, and F. Wang, “Activation of EphB2 in the basolateral amygdala promotes stress vulnerability of mice by increasing NMDA-dependent synaptic function,” *Neuropharmacology*, vol. 167, article 107934, 2020.

[30] A. W. Gilmore, S. M. Nelson, and K. B. Mcdermott, “A parietal memory network revealed by multiple MRI methods,” *Trends in Cognitive Sciences*, vol. 19, no. 9, pp. 534–543, 2015.