Altered functional connectivity in patients with post-stroke memory impairment: A resting fMRI study

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Abstract. Post-stroke memory dysfunction (PMD) is one of the most common forms of cognitive impairment among stroke survivors. However, only a limited number of studies have directly investigated the neural mechanisms associated with memory decline. The aim of the present study was to identify dynamic changes in the functional organization of the default mode network (DMN) and the dorsal attention network of patients with PMD. A total of 27 patients with PMD who experienced a stroke in the right hemisphere were enrolled in the current study, along with 27 healthy control subjects matched by age, sex, and educational level. A behavioral examination and functional magnetic resonance imaging scan were performed. The data were analyzed using an independent component analysis method. The results revealed a significantly increased functional connectivity between the DMN and prefrontal cortex (left middle/inferior frontal and left precentral gyri), temporal regions (left superior temporal gyrus), and bilateral and posterior cingulate gyrus/precuneus (P<0.001). There was also a significantly decreased functional connectivity between the DMN and right middle temporal gyrus, left uvula, and right inferior parietal lobule, and between the dorsal attention network and prefrontal cortex (left precentral/inferior and right inferior/middle frontal gyri), right inferior parietal gyrus, and right insula (P<0.001). These results suggest that the stroke affected both the lesioned and contralesional hemispheres. The prefrontal cortex, temporal regions, insula, and posterior cingulate gyrus/precuneus serve a crucial role in memory processing.

Introduction

Each year >15 million people suffer from stroke, and almost two-thirds of stroke patients exhibit cognitive decline in different domains, including attention, memory and executive function (1,2). Post-stroke memory dysfunction (PMD) is one of the most prevalent cognitive impairments among survivors of stroke, as reported previously (3). PMD greatly influences the restoration of patient motor function and activities of daily living as patients with PMD are unable to fully comprehend complex rehabilitation training instructions and perform the instructions later (4). Rasquin et al (5) and Hochstenbach et al (6) have reported that cognitive impairment after stroke (including subtle deficits) are one of the main causes of disability in daily activities. Tatemichi et al (7) reported that cognitive impairments are a significant independent risk factor for dependent living after stroke. Furthermore, PMD may have a potentially devastating economic and social impact on patients, ultimately leading to a decreased quality of life and increased likelihood of mortality (8).

Although mechanisms leading to general cognitive impairment have previously been reported (9-11), a limited number of studies have directly investigated the neural mechanisms associated with memory decline. Tuladhar et al (12) found that following a stroke, patients presented with delayed memory dysfunction, decreased functional connectivity in the left medial temporal lobe, posterior cingulate and medial prefrontal cortical areas within the DMN compared with controls. However, this study only used one scale to test delayed memory and the mechanism of PMD remains unclear. Considering the high frequency of memory dysfunction, patient complaints and the potentially negative impact of cognitive impairment on the patient and society, the development of viable and effective therapeutic methods to treat PMD is of great importance. To facilitate accurate and timely diagnoses, prognoses and more effective treatments, identifying the underlying mechanisms associated with the development of PMD is of paramount significance.
In addition to the location of PMD-related brain damage, post-stroke cognitive impairment may be attributable to brain regions far removed from the lesion (9). A possible reason for these remote effects may be the disruption of brain networks that occurs following stroke (13,14). Recently, the importance of the segregation and integration functions of the brain has been identified (15). Dacosta-Aguayo et al (9) found that, compared with healthy control, patients following a stroke have greater DMN activity in the left precuneus and the left anterior cingulate gyrus and also present with a marked impairment in the functional connectivity between the DMN nodes, including the left superior frontal gyrus and posterior cingulate cortex. Following a stroke, individuals also demonstrate positive correlations with the mini mental state examination (MMSE) scores, which is a scale to test the general cognitive function. Lawrence et al (10) also indicated that whiter matter lesions are associated with the cognitive function, including executive function and processing speed in cerebral small vessel disease. Ding et al (11) demonstrated that the functional connectivity of posterior cingulate cortex/precuneus is associated with cognitive function in stroke patients. In order to understand the mechanism(s) associated with PMD, both structural damage to the brain and changes in its functional connectivity must be considered. Neuroimaging studies have revealed a set of dynamically interrelated brain networks that are considered to serve vital roles in cognition such as dorsal attention network (DAN) (16) and memory such as default mode network (DMN) (12,17). These two brain networks are two diametrically opposed; the DMN consists of regions that typically exhibit higher levels of activity during the resting state, whereas activity in the DAN is higher during task-related activity (18). However, there is no unified conclusion about how PMD affects these two brain networks.

In the present study, the DMN and DAN in patients with PMD were examined using independent component analysis (ICA). In particular, the association between network functional connectivity and different brain regions in patients with PMD was examined to clarify the neuropathological mechanism(s) associated with PMD. It was also investigated whether changes in brain network functional connectivity are correlated with the severity of PMD.

Materials and methods

Participants. The present study was approved by the Ethics Committee of the Fujian University of Traditional Chinese Medicine Subsidiary Rehabilitation Hospital (Fuzhou, China; approval no. 2013KY-005-01) and each participant gave written informed consent, in accordance with the provisions of the Declaration of Helsinki. A total of 29 participants with PMD were enrolled in Fujian University of Traditional Chinese Medicine Subsidiary Rehabilitation Hospital (Fuzhou, China) from July 2014 to July 2015; All patients were admitted due to vessel disease. Ding et al (11) demonstrated that the functional connectivity of posterior cingulate cortex/precuneus is associated with cognitive function in stroke patients. In order to understand the mechanism(s) associated with PMD, both structural damage to the brain and changes in its functional connectivity must be considered. Neuroimaging studies have revealed a set of dynamically interrelated brain networks that are considered to serve vital roles in cognition such as dorsal attention network (DAN) (16) and memory such as default mode network (DMN) (12,17). These two brain networks are two diametrically opposed; the DMN consists of regions that typically exhibit higher levels of activity during the resting state, whereas activity in the DAN is higher during task-related activity (18). However, there is no unified conclusion about how PMD affects these two brain networks.

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Functional MRI (fMRI) data were examined to avoid excessive movement or inclusion of MRI artifacts prior to analysis. A total of 2 patients with PMD and 1 healthy subject were excluded due to excessive motion during resting-state fMRI scanning, and 1 healthy subject was excluded due to an absence of resting-state data. Thus, a total of 27 patients with PMD and 27 control subjects were included in the present study. Demographic and clinical characteristics for the included healthy control and PMD groups are presented in Table I. Both groups were well matched in age, education level and sex, with no significant differences between groups.

Neuropsychological data. All subjects underwent MMSE (22) and Wechsler Memory Scale-Chinese revision (WMS-CR) testing (23). The MMSE was performed to assess general cognitive function and the WMS-CR was used to measure memory function. The WMS-CR consisted of three sub functions and 10 subtests as follows: Immediate memory (digit span), short-term memory (including addition, picture, recognition, visual reproduction, associative, learning, touch and comprehension) and long-term memory (forwards and backwards recital of digits 1-100 and addition). Clinical severity of patients with PMD was determined according to the National Institutes of Health Stroke Scale (24). The modified Barthel index was used to measure activity of daily living (25).

SPSS version 18.0 software (SPSS, Inc., Chicago, IL, USA) was used for neuropsychological data analysis. The Shapiro-Wilk test was performed to examine the normality of demographic variables. Continuous variables analysis between PMD and control groups was performed using a t-test for independent and nonparametric independent samples. P<0.05 was considered to indicate a statistically significant difference.

fMRI data acquisition. Images were obtained using a 3.0 Tesla General Electric scanner (GE Healthcare Life Sciences, Chalfont, UK) with an eight-channel phased array head.
The resting-state scan was performed using an echo planar imaging sequence (time resolution=2,100 msec, echo time=30 msec, flip angle=90˚, slice thickness=3 mm with a 0.6-mm gap, 42 slices, 64x64 matrix, field of view=200 mm). Subjects were instructed to lie still with their eyes closed without falling asleep, while staying relaxed and not

### Table I. Demographic and clinical characteristics for PMD and healthy control groups.

| Characteristics                          | HC (n=27)         | PMD (n=27)        |
|-----------------------------------------|-------------------|-------------------|
| Age, medians (percentile 25-75), years  | 59 (54-62)        | 57 (47-57)        |
| Educational level, mean ± SD, years     | 8.11 (4.74)       | 9.04 (3.65)       |
| Sex, No.                                |                   |                   |
| Male (%)                                | 20 (74)           | 20 (74)           |
| NIHSS, mean ± SD                        | NA                | 7.96±3.66         |
| MBI, mean ± SD                          | NA                | 51.22±25.52       |
| Lesion volume, medians (percentile 25-75), cm³ | NA              | 17.17 (8.15-25.99)|
| Stroke type                             |                   |                   |
| ICH                                      | NA                | 12                |
| Infarction                               | NA                | 15                |
| Anatomical regions affected              |                   |                   |
| Basal Ganglia                           | NA                | 19                |
| Corona Radiata                          | NA                | 9                 |
| Occipital lobes                          | NA                | 4                 |
| Temporal lobes                           | NA                | 9                 |
| Parietal lobes                           | NA                | 9                 |
| Frontal lobes                            | NA                | 9                 |
| Thalamus                                 | NA                | 5                 |

a=t=-0.451 and P=0.652; b=z=0.804 and P=0.425; NIHSS, national institute of health stroke scale; SD, standard deviation; PMD, post-stroke memory dysfunction; HC, healthy control; NA, not applicable; MBI, the Modified Barthel Index; ICH, intracerebral hemorrhage.

### Table II. Neuropsychological tests scores for PMD and HC groups.

| Variable                  | HC (n=27)         | PMD (n=27)        | P-values    |
|---------------------------|-------------------|-------------------|-------------|
| General cognitive function|                   |                   |             |
| MMSE                      | 27 (25-29)        | 21 (18-22)        | <0.01*      |
| Total WMS score           | 88 (25-29)        | 47 (32-60)        | <0.01*      |
| Immediate memory          |                   |                   |             |
| Digit span                | 7.93 (2.21)       | 4 (2.27)          | <0.01*      |
| Short-term memory         |                   |                   |             |
| Picture                   | 9 (7-10)          | 6 (5-8)           | <0.01*      |
| Recognition               | 9 (7-12)          | 2 (0-5)           | <0.01*      |
| Visual reproduction       | 9 (7-12)          | 2 (0-4)           | <0.01*      |
| Associative learning      | 7 (6-9)           | 5 (1-7)           | 0.011*      |
| Touch memory              | 9 (7-10)          | 6 (6-6)           | <0.01*      |
| Comprehension memory      | 8.70 (2.05)       | 4.70 (2.25)       | <0.01*      |
| Long-term memory          |                   |                   |             |
| Forward recite digit (1-100)| 9 (6-10)        | 3 (0-6)           | <0.01*      |
| Backward recite digit (100-1)| 10.07 (3.54)   | 4.46 (3.65)       | <0.01*      |
| Calculation               | 11 (10-12)        | 7 (2-9)           | <0.01*      |

All values are the means (standard deviation) or medians (percentile 25-75). n=27 for each group. *P<0.05. MD, post-stroke memory dysfunction; HC, healthy control; MMSE, mini mental state examination.
thinking of anything in particular. T1 three-dimensional magnetization-prepared rapid gradient-echo imaging was also performed in the same session (echo time=1.764 msec, flip angle=15°, inversion time=450 msec, slice thickness=1 mm, field of view=240 mm and 164 slices per acquisition). Both behavioral examinations and fMRI scanning were completed within 1 week of enrollment.

fMRI preprocessing. fMRI data preprocessing and analysis were performed using the Oxford Centre for Functional MRI of the Brain's (FMRIB) Software Library (http://www.fmrib.ox.ac.uk/fsl). The current study

performed regression analysis (28) using the network connectivity change maps between patients and healthy control subjects and corresponding changes in clinical outcomes. A threshold of voxel-wise Z>2.3 and a cluster correction significance threshold of P<0.05 was used. Furthermore, for both analyses, age, sex, and education level were not considered covariates of interest in this GLM.

**Results**

**Neuropsychological results.** Patients with PMD had a statistically significant impairment in general cognitive function as measured by the MMSE (P<0.01), as well as in memory function measured by the WMS-CR (P<0.01) compared with healthy controls (Table II).

**Between-group analysis of DMN and DAN functional connectivity.** Demographic and neuropsychological data were analyzed using SPSS (Tables I and II). The DMN was obtained from ICA data in conformity with earlier studies (27) and included the medial prefrontal, anterior cingulate, bilateral parietal, and precuneus (Pcu)/posterior cingulate cortices (PCC; Fig. 2A). The DAN in the present study was consistent with that of a previous study (29), consisting bilaterally of the intraparietal, precentral, and superior frontal sulci and ventral precentral gyri and middle frontal gyrus (Fig. 2B).

A comparison between PMD and control subjects (stroke-control) revealed significant differences in functional connectivity between the DMN and the following regions (P<0.001): The bilateral PCC/Pcu and bilateral cingulate, left precentral, left inferior/middle frontal and left superior temporal gyri (Fig. 3; Table III). A comparison of HCs with PMD subjects (control>stroke) found significant functional connectivity differences between the DMN and right middle temporal gyrus, left uvula, and right inferior parietal lobule. Furthermore, no significant functional connectivity differences were observed between the DAN and cortical areas between the PMD and control groups, whereas functional connectivity between the DAN and left precentral, left inferior frontal, right inferior/middle frontal, and right inferior parietal gyri and right insula was significantly higher in the control group as compared with the PMD group (Fig. 4; Table III).

**Correlations between functional connectivity and WMS-CR scores.** Regression analysis of the DMN and WMS-CR scores revealed a positive association between the DMN and right medial frontal gyrus, and left anterior cingulate and a negative association between the DMN and left claustrum and the bilateral cingulate, left inferior frontal and left precentral gyrus (Fig. 3; Table IV). Fig. 3 also depicts the correlation between brain region (cingulate gyrus) FC alterations and WMS score changes. That is, Z-Score (functional connectivity alterations between DMN and cingulate gyrus) was negatively associated with the corresponding WMS score changes; the higher the Z-Score, the lower WMS score. It demonstrated that the function of the connection change has a linear correlation with the behavior changes. Regression analysis between the DAN and WMS-CR identified a positive association between the DAN and right precentral, right middle frontal, and right inferior frontal gyri and right insula (Fig. 4; Table IV). Fig. 4 depicts the
correlation between brain region (right inferior frontal gyrus) FC alterations and WMS score changes. Demonstrating that the Z-Score (functional connectivity alterations between DAN and prefrontal) was associated with the corresponding WMS score changes; the higher the Z-Score, the higher WMS score. It demonstrated that the functional connectivity change has a linear correlation with the behavior changes to some extent.

No negative significant associations were observed between the DAN and WMS-CR.

Discussion
In the present study, ICA methods were used to improve understanding regarding the underlying pathophysiology
of resting-state brain networks and brain regions of patients who experienced their first-ever stroke. Increased functional connectivity was observed between the DMN and frontal cortex (left middle/inferior frontal and left precentral gyri), temporal regions (left superior temporal gyrus), bilateral cingulate gyrus, and PCC/Pcu, and decreased functional connectivity was detected between the DMN and right middle temporal gyrus, left uvula and right inferior parietal lobule in patients who had experienced their first ever stroke. Furthermore, decreased functional connectivity was identified between the DAN and prefrontal cortex (left precentral/inferior frontal and right inferior/middle frontal gyri), right inferior parietal gyrus and right insula.

Analysis of functional connectivity via resting-state fMRI is attracting more researchers to the study of brain networks as a result of its less complex task design. As a data-driven

### Table III. Brain regions demonstrated significant FC differences between PMD and HC groups.

| Brain region Cluster size | X | Y | Z | Peak Z-Score | P-value |
|--------------------------|---|---|---|--------------|---------|
| Stroke > control         | R. cingulate gyrus  3,355  | 14 | -36 | 34 | 4.3 | <0.001 |
| Stroke > control         | R. precentral gyrus 1,616 | -50 | -6  | 30 | 4.12 | <0.001 |
| Stroke > control         | L. mid frontal gyrus 1,202 | -18 | 20  | 66 | 4.04 | <0.001 |
| Stroke > control         | R. mid temporal gyrus 1,878 | 60  | -4  | -16 | 5.01 | <0.001 |
| Stroke > control         | L. uvula 1,168 | -10 | -90 | -38 | 4.57 | <0.001 |
| Stroke > control         | R. inferior parietal lobule 1,159 | 46 | -66 | 50 | 4.32 | <0.001 |
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**Table IV.** Brain regions demonstrated significant associations between WMS and FC.

| Brain region Cluster size | X | Y | Z | Peak Z-Score | P-value |
|--------------------------|---|---|---|--------------|---------|
| The FC between the DMN and WMS |
| Positive WMS R. med frontal gyrus 1,475 | 6 | 52 | -24 | 4.14 | <0.001 |
| Negative WMS R. cingulate gyrus 3,705 | 14 | -36 | 36 | 4.7 | <0.001 |
| L. claustrum 3,552 | -30 | 6  | 10 | 4.31 | <0.001 |
| L. Inferior frontal gyrus 3,552 | -48 | 4  | 14 | 3.15 | <0.001 |
| The FC between the DAN and WMS |
| Positive WMS R. precentral gyrus 1,330 | 52 | 6  | 26 | 4.4 | <0.001 |

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**Table IV.** Brain regions demonstrated significant associations between WMS and FC.
ICA is able to determine distinct components by capturing spatial independence and time-courses of resting-state data and thereby reliably defines different resting-state networks (30). In the present study, ICA was used on PMD and control subjects to better define DMN and DAN pathways common to both study groups (31). DMN and the DAN were chosen as they are associated with cognitive and memory function (12,17). The DMN and DAN are competitive in function and distinct in spatial and sub-serving cognitive processing, respectively (32). The DMN is one of the most widely researched networks using resting-state fMRI analysis (33). Previous studies have suggested that the DMN is associated with general cognition (34) and PMD (12); however, the sample sizes of those studies were small and their results remain controversial. Anomalous alterations in the DAN have been reported in patients with Alzheimer's disease (35), Parkinson's disease (36), epilepsy (37), and patients who have suffered a stroke. Perry et al (38) speculated that episodic memory is associated with inattention to related information. Despite considerable interest in the DAN, at present there is limited information regarding its possible changes in patients with PMD.

Figure 3. (A) Red color indicates brain cortices (left precentral gyrus, left middle/inferior frontal gyrus, bilateral cingulate gyrus/posterior cingulate cortices/precuneus) that exhibited increased FC with the DMN in patients with PMD compared with the HC group. (B) Green color indicates brain regions where DMN functional connectivity alterations were negatively associated with the corresponding WMS score changes in all participants. (C) Scatter plots depict the correlation between brain region (cingulate gyrus) FC alterations and WMS score changes. FC, functional connectivity; DMN, default mode network; PMD, post-stroke memory dysfunction; HC, healthy control; WMS, Wechsler Memory Scale; L, left; R, right.

Figure 4. (A) Blue color indicates the brain cortices left precentral/inferior frontal gyrus, right inferior/middle frontal gyrus, cingulate gyrus, right inferior parietal gyrus and right insula that exhibited decreased in FC with the DAN in the PMD group compared with the HC group. (B) Yellow color indicates brain regions (right inferior/middle frontal gyrus) where DAN FC alterations were positively associated with the corresponding WMS score changes across all participants. (C) Scatter plots depict the correlation between brain region (right inferior frontal gyrus) FC alterations and WMS score changes. FC, functional connectivity; DAN, dorsal attention network; PMD, post-stroke memory dysfunction; WMS, Wechsler Memory Scale; HC, healthy control; L, left; R, right.
In the present study, connectivity increments were identified between the DMN and bilateral PCC/Pcu. This finding is in agreement with a previous functional neuroimaging study, which reported increased functional connectivity strength in the PCC/Pcu of patients with post-stroke cognitive impairment compared with controls (11). Notably, patients with Alzheimer's disease have also demonstrated increased functional connectivity in the PCC/Pcu (11,39). In the present study, it was determined that patients with PMD had increased functional connectivity in the bilateral cingulate gyrus, left claustrum, and left frontal cortex, which was correlated with poor performance in memory function tests. Similar results have been identified in studies regarding cognitive impairment and the DMN (9,40). The functional connectivity analysis and the regression analysis together prove that cingulate gyrus serves an important role in memory function. These findings suggest that functional changes in the cingulate gyrus and left frontal cortex are involved in this type of deficit in patients who have suffered a stroke.

In the present study, patients with PMD exhibited reduced functional connectivity between the DMN and right middle temporal gyrus, left uvula and right inferior parietal lobule. However, there were no significant correlations between the WMS-CR and Z-scores in voxels exhibiting decreased functional connectivity in these brain regions. As well as differences in memory between the PMD and healthy groups, other confounding factors exist. For example, 25-50% of stroke patients suffer from dysphagia (41) and a recent study (42) reported that patients with dysphagia alone had decreased functional connectivity in the DMN. As such, it cannot be assumed that the decrease in functional connectivity between the DMN and these brain regions in the present study was due solely to the effects of stroke on memory function, as the current study included 8 patients with mild to moderate dysphagia, which may have affected functional connectivity changes.

The authors of the present study were only able to identify three recently published studies that investigated DMN and post-stroke cognitive performance via resting-state fMRI. All three reported that patients who had suffered a stroke exhibited increased functional connectivity in the left frontal cortex compared with healthy controls, which is in accordance with the results of the present study (9,11,40). However, using a region of interest-based analysis and ICA, Tuladhar et al (12) reported that patients who experienced their first ever right or left hemisphere stroke had decreased activity in the left frontal lobe and posterior cingulate cortex compared with controls. Their study assessed the neuropathophysiological mechanism(s) of memory impairment following stroke. Multiple factors may have contributed to the heterogeneity between the results of the present study and those of Tuladhar et al (12). For example, Tuladhar et al (12) included patients with both right and left hemisphere injury. However, the results of the present study are unable to confirm whether brain injury from the left side leads to the same alterations in functional connectivity as injury to the right (43). Furthermore, other clinical variables, including the effects of concomitant pharmacological therapies may have contributed to these contradictory results.

Previous studies have suggested that the prefrontal cortex serves a role in spatial and working memory (44), which is in accordance with the results of the present study that exhibits connectivity changes between the DAN and left precentral, inferior frontal, and right inferior/middle frontal gyri. In the present study, it was also observed that there was decreased functional connectivity between the DAN and right insula that was related to memory performance. The functional connectivity analysis and the regression analysis together indicate that the prefrontal cortex serves an important role in memory function. Christopher et al (45) reported that the fronto-insular cortex is associated with cognitive function. Notably, right frontal areas and adjacent insula associated with negative WMS-CR scores are regions within the DAN and right frontoparietal networks, which suggests that these two networks serve a role in memory processes (27,29,46).

The frontal lobe is one of the most complicated brain regions and is involved in a variety of cognitive functions, including language, attention, emotion and memory (47,48). Different types of memory, including working, spatial and episodic memory have different characteristics (49). Parietal fiber projections accepted by the dorsal prefrontal cortex and subsequently integrated into spatial information transmitted to the dorsal prefrontal region, ultimately forming spatial memory (50). One fMRI study reported that speech memory activates Broca's area, whereas spatial memory activates the prefrontal cortex of the right hemisphere (51). Results from studies involving spatial orientation tasks have suggested that episodic memory-encoding processes primarily depend on the prefrontal cortex, particularly in time-related information recall (47). Furthermore, Jodo et al (52) demonstrated that the brain network between the frontal lobe and hippocampus is associated with memory, particularly working memory. Despite variable findings, the present study determined that DMN activity in patients with PMD is increased in brain regions including the inferior frontal and precentral gyri, whereas DAN activity is reduced. Together, these findings confirm and complement those of previous studies (17,18).

There were various limitations of the present study. The aim of the present study was to investigate the common underlying mechanism(s) of memory impairment for all of stroke patients who suffered from varying degrees of memory decline. Therefore, in the present study, 27 patients with PMD and 27 healthy control subjects were enrolled within 1 year following the onset of stroke to investigate the pathophysiology of PMD. However, different stroke courses may have some heterogeneity (53). Desmond et al (54) reported that typically, cognitive function may exhibit a long-term improvement after stroke when compared with assessments at 3 months and then annually. Another study (55) also suggested that 30% of individuals who had mild cognitive impairments between 0 and 6 months after stroke may improve, and be cognitively intact at 12-18 months. Therefore, future studies should have stricter inclusion and exclusion criteria, in particular taking into account consistent stroke course, to clarify the aforementioned issues. In conclusion, the results of the present study suggest that resting-state fMRI may help researchers to better understand the neuropathophysiological mechanisms of PMD. In particular, it has been determined that the prefrontal cortex, temporal regions, insular cortex, and PCC/Pcu serve crucial roles in memory processing. The alterations described herein suggest that stroke events affect not only the lesioned hemisphere, but also the contralesional hemisphere.
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