Alterations of dynamic redundancy of functional brain subnetworks in Alzheimer’s disease and major depression disorders

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A B S T R A C T

The human brain is not only efficiently but also “redundantly” connected. The redundancy design could help the brain maintain resilience to disease attacks. This paper explores subnetwork-level redundancy dynamics and the potential of such metrics in disease studies. As such, we looked into specific functional subnetworks, including those associated with high-level functions. We investigated how the subnetwork redundancy dynamics change along with Alzheimer’s disease (AD) progression and with major depressive disorder (MDD), two major disorders that could share similar subnetwork alterations. We found an increased dynamic redundancy of the subcortical-cerebellum subnetwork and its connections to other high-order subnetworks in the mild cognitive impairment (MCI) and AD compared to the normal control (NC). With gained spatial specificity, we found such a redundancy index was sensitive to disease symptoms and could act as a protective mechanism to prevent the collapse of the brain network and functions. The dynamic redundancy of the medial frontal subnetwork and its connections to the frontoparietal subnetwork was also found decreased in MDD compared to NC. The spatial specificity of the redundancy dynamics changes may provide essential knowledge for a better understanding of shared neural substrates in AD and MDD.

1. Introduction

Resting-state fMRI (rs-fMRI)-derived functional connectivity (FC) reflects information exchanges and coordination among different brain regions, resulting in large-scale brain functional networks (Sporns, 2013). It has been suggested that brain FC is time-varying and non-stationary, possibly reflecting adaptive control and attention fluctuations during resting state (Chang & Glover, 2010). Such a dynamic connectome (a.k.a., “chronnectome”) could be altered by many brain diseases and mental disorders (Marusak, et al., 2017). Many studies have used graph theory as a powerful tool to investigate the brain network’s efficiency and how brain diseases alter it (Xuan, et al., 2017). Among many efficiency metrics, the characteristic path length (Liu, et al., 2017) (i.e., the shortest paths) and its derivatives (e.g., assortativity and resilience), including the small-worldness (Newman, 2006; Ravasz & Barabási, 2003; Achard, et al., 2006) have been extensively investigated. These metrics, including their dynamics, have performed well on analyzing sparse brain networks that often include a few strong connections. On the contrary, in denser networks where weaker connections are also included, redundancy could become a dominant property.

However, characterizing brain network’s redundancy and its changes over a short time (i.e., redundancy dynamics) is largely neglected in the previous brain research. For instance, one may calculate the number of independent paths in the brain network as a redundancy measurement to quantify the robustness of the network under possible attacks that could occur brain wide. While such a redundancy analysis on many natural networks has been extensively conducted (Corson, 2010; Härkägård & Glad, 2005), brain network studies are still very scarce. A magnetoencephalography (MEG)-based FC study (Di Lanzo, et al., 2012) used various redundancy metrics at each frequency band and revealed that the functional brain network was more redundant than a random network. In the study, the average number of alternative

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paths was used as the redundancy metric. In another electroencephalogr
(EEG) study on spinal cord injured patients, a redundancy measuremen
t of the entire FC network was used but showed no signifi
changes compared to healthy subjects (Fallani, et al., 2011). It was also
found that education level could strengthen the redundancy of dif
MRI-derived brain structural connectivity network of Alz
zheimer’s disease (AD) patients compared to normal controls (NC) (Yoo,
et al., 2015). A major depressive disorder (MDD) study showed a red
ction in the brain network’s redundancy in MDD patients compared
to NC during stimulus processing (Leistritz, et al., 2013).

The major drawback in the previous brain network redundan
studies is that none of them investigated redundancy in the dynamic
brain functional network that may reflect the higher functional signifi
ance of the brain network redundancy than the static one. Whether the
FC dynamics also lead to dynamics in the network redundancy, and
whether such alterations can be sufficiently sensitive to brain diseases so
that it could be utilized for brain disease studies, remains unknown. As
such, in our recent works, we proposed to measure the dynamic
redundancy of the whole brain network (Ghanbari, et al., 2020; Ghan
bargi, et al., 2021). We defined a new brain state that every pair of brain
regions were connected through at least two independent paths (with no
node shared) as a redundant state, and calculated its dynamics as one of
the major redundancy metrics. We found an increased chance of the
whole-brain redundancy along time in the mild cognitive impairmen
t subjects (MCI), an early stage of AD, compared to the NC (Ghanbari,
et al., 2020, Ghanbari, et al., 2021). We further designed an individu
ized early AD detection method based on a decision tree with the
dynamic redundancy metrics as features, which reached an accuracy of
90% in the classification between MCI and NC (Ghanbari, et al., 2020).

While this method seems promising, it lacks spatial specification, as all
the redundancy metrics were defined on the entire network without know
ing which specific subnetwork has redundancy changes. Disease studi
may be more interested in where such redundancy changes take
place. Meanwhile, by focusing on more dynamic redundancy features,
one could increase the sensitivity in group difference detection, whereas
the whole-brain metric could have reduced sensitivity due to the
average effect. It could be possible that different brain subnetworks, due
to their specific cognitive roles (Smith, et al., 2009), may be affected
differently and show different dynamic redundancy changes. Without
the spatial resolution, whole-brain averaged redundancy may not reveal
such subnetwork-level changes.

Cognitive dysfunction is prevalent in neurological and psychiatric
disorders. Two typical disorders, AD and MDD, could develop cognitive
dysfunction at certain stage of the diseases (Sierksma, et al., 2010) and
may share broad similarities in brain structural changes such as hippo
campal volume loss (Sampath, et al., 2017) as well as brain subnetwork
disconnection, especially within and among a set of cognitive function-
related subnetworks (Menon 2011). In this study, we performed subnetwork-level dynamic redundancy analysis on two disease cohorts (AD and MDD) and investigated the redundancy dynamics of each cognitive function-related brain functional subnetwork and the redu
dancy dynamics of the inter-subnetwork connections. We hypothesized
that the two disease cohorts (AD and MDD) may share similar cognitive
dysfunction symptoms and thus could have similar overlapping underr
lying neural features as revealed by the subnetwork-level redundancy dynamics.

2. Materials and methods

2.1. Datasets

In Study 1, we used the Alzheimer’s Disease Neuroimaging Initiative
(ADNI) dataset. A participant with or without subjective memory com
plaints, verified by a study partner, with Mini-Mental State Exam score
between 24 and 30 inclusive, and also Clinical Dementia Rating equals
0 and Memory Box score 0, belongs to NCs. While participants were
diagnosed as MCI if they expressed a subjective memory concern as
reported by the participant or recalled by study partner or clinician with
Mini-Mental State Exam (MMSE) score between 24 and 30 inclusive, and
also Clinical Dementia Rating equals 0.5 and Memory Box score at least
0.5. However, participants were diagnosed as AD if they expressed a
subjective memory concern as reported by the participant or recalled by
study partner or clinician and Mini-Mental State Exam score between 20
and 24 inclusive with Clinical Dementia Rating equals 0.5 or 1.0 (http:
/adni.loni.usc.edu/). The 7-min rs-fMRI data (140 volumes, repetition
time = 3 s, slice thickness = 3.5 mm preferred, 4.5 mm maximum, flip angle
= 8°, matrix size = 64 × 64) was preprocessed using AFNI (Cox, 1996)
according to a standard pipeline (Yan & Zang, 2010). Specifically, the
first ten volumes are discarded, followed by a rigid-body head motion
correction and a nonlinear spatial registration to the Montreal Neuro-
logical Institutes (MNI) space. The subjects with large head motion (i.e.,
larger than 2 mm or 2°) and those who had more than 2.5 min rs-fMRI
data with large (greater than 0.5 mm) frame-wise displacement were
excluded (Chen, et al., 2017). A total of 6 ADs, 3 MCIs and 17 NCs were
excluded due to excessive head motion. Mean rs-fMRI time series of each
brain region defined by Shen’s 268-region atlas (Shen, et al., 2013), was
band-pass filtered (0.015–0.15 Hz), before it was further processed to
reduce artifacts by regression analysis (nuisance regressors included
head motion parameters according to the “Friston-24” model, the mean
rs-fMRI signal in the white matter, and that in the cerebrospinal fluid).
A total of 49 NC (26 males and 23 females, age 73.1 ± 8.0, MMSE 29.1 ±
1.0), 49 MCI (26 males and 23 females, age 74.3 ± 9.8, MMSE 27.9 ±
1.6), and 49 AD (26 males and 23 females, age 73.3 ± 8.5, MMSE 23.1 ±
2.5) subjects were selected from the ADNI0 and the ADNI2. Specifi-
cally, we found that 49 AD subjects had baseline fMRI data, while the
NC and MCI groups had more subjects than the AD group. Therefore, we
selected the baseline data (or the follow-up data that was closest to the
baseline data, if the baseline data had excessive head motion) NC and
MCI subjects at the similar age (+2 years) and of the same gender as
each of the AD subjects to make sure a good match of subjects among the
three groups. Due to the limited sample size (N = 49) of ADs, we selected
the same amount (N = 49) of NC and MCI subjects, respectively, to make
sure as many matched data as possible were used. The three groups were
age-matched (p = 0.752, one-way analysis of variance (ANOVA)) and
gender-matched.

In Study 2, we used a dataset collected from our collaborative hos-
pital, the First Affiliated Hospital of Guangzhou University of Chinese
Medicine, Guangdong, China from September 2015 to June 2018. After
an initial screening utilizing the 17-item Hamilton Rating Scale for
Depression (HDRS-17) with a total score larger than 18 (Hamilton,
1967), two expert psychologists (with at least ten years experience
separately diagnosed the MDD patients according to the Diagnostic and
Statistical Manual (DSM-5, American Psychiatric Association, 2013).
Only if both psychologists diagnosed a patient as MDD, then that subject
was recruited. The inclusion criteria of MDD are as follows: (a) aged
between 18 and 55 years old, (b) right-handed native Chinese speaker,
(c) firstly diagnosed with MDD and had no history of any neurological
illness or any other forms of psychiatric disorders, and (d) head motion
smaller than 2 mm of translation or 2° of rotation in any direction during
the rs-fMRI scan. Exclusion criteria included (a) a history of significant
medical illness (15 participants were excluded), (b) alcohol abuse (a
total score > 8 in Alcohol Use Disorders Identification Test (Saunders,
et al., 1992)), (0 participants were excluded), and (c) contraindications
to fMRI scan (6 participants were excluded). The rs-fMRI and 3D T1-
weighted images (3D-T1WI) in the MDD dataset had the following pa-
rameters: repetition time / echo time = 2,000/30 ms, flip angle = 90°,
matrix size = 64 × 64, field of view (FOV) = 240 mm × 240 mm, slices
number = 33, slice thickness = 4.0 mm, scanning time = 8′20″ (250
volumes) for the rs-fMRI, and TR/TE = 10.4±4.3 ms, flip angle = 15°,
slice thickness = 1.0 mm, slice gap = 0 mm, matrix size = 256 × 256,
FOV = 256 mm × 256 mm, and slices number = 156 for the 3D-T1WI.
The image preprocessing was performed using SPM12 (www.fil.ion.ucl.)
several nuisance signals, including the Friston-24 head motion parameter, half-maximum Gaussian kernel, and further denoised by regressing out signals from cerebrospinal fluid and white matter, before linear detrending and temporal band-pass filtering (0.01–0.08 Hz). The same data was previously used in another study (Li, et al., 2020), which included 66 NC (35 males and 31 females, age 29.3 ± 10.1) and 66 MDD (41 males and 25 females, age 29.5 ± 9.9). The two groups were age (p = 0.931, two-sample t-test) and gender-matched (p = 0.379, chi-square test). The MDD group had a Hamilton Depression Rating Scale (HAM-D, a measurement of disease severity) of 22.3 ± 3.6 and a disease duration of 9.1 ± 1.9 months. This study was conducted in accordance with the Declaration of Helsinki. All participants provided written informed consent and the study was approved by the local ethics committee.

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2.2. Subnetwork definitions

We adopted a widely used functional parcellation atlas (Shen, et al., 2013). The 268 brain regions had been clustered using a group-wise spectral clustering algorithm, generating eight functional subnetworks. They were further identified and named as the medial frontal, frontoparietal, default mode, subcortical-cerebellum, motor, visual I, visual II, and visual association networks. In this study, we evaluated the dynamic redundancy of four subnetworks in the AD study, including the medial frontal, frontoparietal, default mode, and subcortical-cerebellum networks, since many studies have shown alterations in these four subnetworks due to AD (Cui, et al., 2018; Xue et al., 2019). In the MDD study, since the previous studies indicated that the first three subnetworks could be involved (Bludau, et al., 2016; Yan, et al., 2019); we only focused on the first three subnetworks. The four subnetworks were visualized in Fig. 1.

2.3. Network redundancy

Similar to our previous studies (Ghanbari, et al., 2020; Ghanbari, et al., 2021), we used a set of well-adopted connectedness metrics in the field of graph theory. Specifically, a network G with a node set V (G) and an edge set E(G) is connected if there is a path between every pair of its nodes. A connected network G is called 2-connected if, for every two nodes x, y ∈ V (G), there are at least two “independent” paths between x and y that do not share any node(s) except x and y themselves. Fig. 2A depicts a connected graph but it is not 2-connected because of the existence of a shared node. Fig. 2B shows a 2-connected graph because there are at least two independent paths for every pair of nodes. Therefore, this is a redundant network that is resilient to any attacks (the graph is still connected after removing any node or link). Please note that such a redundancy definition is quite strict as it defines the redundancy of the entire network. While previous graph theory analysis also measured brain network’s resiliency by calculating the reduction of network efficiency after targeted removal of high centrality nodes or after random removal certain number of nodes (Joyce, et al., 2013); these methods were still efficiency-based and not as strict as our method.

2.4. Dynamic network redundancy

We calculated dynamic binary brain functional networks and quantified dynamic network redundancy on each selected subnetwork and inter-subnetwork connections between each pair of the subnetworks. First, for each subnetwork (Fig. 3A) of each subject in each group (we did the same for the graph consisting of inter-subnetwork connections between each pair of the subnetworks), we generated T sliding windows to calculate the dynamic FC based on pairwise Pearson’s correlation (Fig. 3B). The window length was set to 60 s (20 volumes for the AD dataset and 30 volumes for the MDD dataset), and the step size was set to one volume (3 s for the AD dataset and 2 s for the MDD dataset). Second, for every window t, 1 ≤ t ≤ T, we applied D density levels to generate D binary networks (Fig. 3C). Third, we defined redundancy states (either state #1 or state #2) for every time window (Fig. 3D). Let r = {r_t} (1 ≤ t ≤ T) be the redundancy state vector where r_t = 1 if the minimally connected network in t is not 2-connected and r_t = 2, otherwise. Of note, the density level to have the minimal 2-connected network is higher than or equal to the density level that the network is minimally connected. This is the reason that we defined redundancy states as whether the density level of the minimal 2-connected network is the same as the density level of the minimal connected network. In other words, we set the redundancy index for a network to “2” if the minimal 2-connected and the minimal connected networks have the same density level and we “1” if the minimal connected network is not 2-connected. For example, r = {1, 2, 2, ..., 1} describes redundancy state changes in Fig. 3D. Finally, we quantified the dynamic redundancy by calculating the occurrence frequency of the state #2 during the entire scan time from r, generating a “redundancy index” for a subnetwork or a pair of subnetworks. This index measures the probability (between 0 and 1) that a network is also 2-connected when it is minimally connected (with the lowest density to ensure its connectedness) through time. Thus, the more the redundancy index is, the more frequent back-up paths exist for every pair of the network. In other words, the network with a higher redundancy index is more likely to have an optimized topology along time that makes it robust to any attacks. We obtained ten redundancy indices for each subject in the AD study, four for each subnetwork, and six for all pairs of the subnetworks. In the MDD study, six redundancy indices were obtained for every subject (since three subnetworks were...
Please note, since the redundancy index was determined by focusing on the minimal connected network at every time window, this study was irrelevant to a specific network density level.

In this study, \( D = 19 \) (from 5% to 95% with a step size of 5%), \( T = 111 \) for AD study, and \( T = 151 \) for MDD study. All the analyses were implemented in MATLAB 2018b (the Mathworks, Inc.), SAGE 8.6 (based on Python 2.7), and SPSS v23 (IBM, Inc.).

### 2.5. Statistical analysis

Statistical analysis was carried out to compare the redundancy index among different groups (Fig. 3E). We used the Kruskal-Wallis test, a non-parametric version of ANOVA for group comparisons among NC, MCI, and AD groups on each subnetwork and each subnetwork pair (\( p < 0.05 \), false discovery rate (FDR) corrected). If a significant group difference was detected, then pairwise post-hoc analysis will be conducted with Mann-Whitney U-tests, a non-parametric version of two-sample \( t \)-test (\( p < 0.05 \) after Bonferroni correction). As a different analysis, we also conducted Spearman’s correlation analysis between each subject’s MMSE score and the redundancy index across all the subjects (NC and MDD) to characterize the possible association between redundancy and cognitive ability in the AD continuum. In the MDD study, we conducted the Mann-Whitney U-tests to detect significant differences in the redundancy indices between NC and MDD groups (\( p < 0.05 \), FDR corrected). We also conducted Spearman’s correlation analysis between the duration of MDD disease and the redundancy index across all the subjects (NC and MDD) to find if there would be any associations between these two metrics.

### 3. Results

#### 3.1. Group differences in subnetwork’s redundancy (the AD study)

We found significant group differences in dynamic redundancy for the default mode and subcortical-cerebellum subnetworks among NC, MCI and AD groups (Table 1, Fig. 4). For subnetwork pairs, we also found group differences in four subnetwork pairs, mostly involving the connections with the subcortical-cerebellum subnetwork and those with the medial frontal subnetwork (Fig. 4C-F). Post-hoc analysis revealed that, in most cases, MCI subjects had elevated redundancy compared to NC, while such an elevation tends to be kept in AD subjects. The only exception was the default mode subnetwork, whose redundancy only decreased in the AD stage compared to NC.

#### 3.2. Association between subnetwork’s redundancy and cognitive performance (the AD study)

We found significant correlations between the subnetwork’s redundancy index and disease symptoms (MMSE score) across all 147 participants in the AD continuum. They were summarized in Table 2.
All the results indicate negative correlations between MMSE and the redundancy index. In other words, the redundancy was increased when the symptom was getting worse. By increasing the MMSE, the tendency of redundancy decreases in some of the subnetworks. More importantly, all the significant associations involved either the subcortical-cerebellum subnetwork or the connections with this subnetwork (especially, to the default mode, frontoparietal, and medial frontal subnetworks).

### 3.3. Group differences in subnetwork’s redundancy (the MDD study)

We found significantly decreased subnetwork redundancy (see Fig. 6A) of the medial frontal (\( p = 0.016 \)) and that of the connections between the medial frontal and frontoparietal subnetworks (\( p = 0.013 \)) in the MDD compared to NC after FDR corrections, while other subnetworks and subnetwork pairs did not (Zhi, et al., 2018) show significant changes.

### 3.4. Correlation between disease duration and subnetwork’s redundancy (the MDD study)

Of all the studied subnetworks, we found a significant negative correlation (\( r = -0.249 \) and \( p = 0.044 \), uncorrected) between the default mode subnetwork’s redundancy index and the disease duration of the MDD subjects (Fig. 6B). As disease duration increased, the redundancy index decreased.

### 4. Discussions

Graph theory has become a powerful tool in investigating brain network topological changes in diseased populations, including AD (Farahani, et al., 2019) and also MDD (Zhi, et al., 2018). This paper investigated connectedness, a graph theory metric that measures reachability from any node to any other node in the network. If pairs of nodes are reachable through more than one independent path, then the increased reachability makes the network 2-connected, and making the network redundant. The redundancy of the functional brain network
we did not investigate the previous brain network studies. Previous studies may use connectedness as a prerequisite to quantify other network attributes (Rubinov & Sporns, 2010). They did not directly study connectedness and redundancy and its dynamics in diseased conditions. Importantly, we focused on characterizing truly redundant paths (those sharing no nodes) and redundancy for all node pairs in the network. The result showed the effectiveness and sensitivity of our method in disease studies.

Despite the novelty of the method, in this paper, we mainly focused on detecting which brain subnetwork(s) had altered dynamic redundancy due to two different diseases (AD and MDD), because it had been shown that complex psychiatric and neurological disorders were characterized by structural and functional abnormalities in a few brain subnetworks (Menon, 2011). With this consideration, in the current study, we did not investigate the global redundancy of the entire network but the redundancy of the specific brain subnetworks.

We first applied the method to characterize group differences among groups at different AD progression stages. The gradual reduction in the redundancy of the default mode network found along the AD continuum (Fig. 4A) was consistent with the previous findings with other methods (Xue et al., 2019), (Grieder, et al., 2018). This finding indicates that the AD pathology may target the default mode network. However, we did not find significant reductions in the MCI stage, possibly due to a good resilience of the default mode network, which makes its robustness last longer and finally collapse in the AD stage. This is also different from our previous study characterizing the redundancy of the entire brain network, where the MCI shows elevated redundancy compared to NC and AD. One possible explanation for such a difference is that the default mode subnetwork is losing its redundancy, but other subnetworks are gaining redundancy to compensate for it. This could make the entire network of MCI have increased redundancy. This study, however, further revealed the subnetwork-level changes under the AD pathological attacks.

The generally increased subnetwork redundancy in MCI, as one of the major findings in the current study, is consistent with our previous finding of the elevated whole-brain network redundancy in MCI, arguably caused by the compensatory effect to the AD pathological attack in the early stage. We note that the most involved subnetwork was the subcortical-cerebellum subnetwork. The redundancy of this subnetwork (Fig. 4B) and that of the inter-subnetwork connections with this subnetwork (Fig. 4D-F) were increased in MCI compared to NC. The function of this subnetwork is complex, including memory. Many AD pathology-related regions, such as the hippocampus, are included in this subnetwork. Due to its fundamental role in AD pathology, the connections within this subnetwork could be less efficient; to maintain cognitive ability in the presence of AD-related neurodegeneration, the elevated redundancy in this subnetwork could be the key in the MCI stage. Besides, we found that all three major “high-order” subnetworks (medial frontal, frontoparietal and default mode subnetworks) tended to have more redundant connections to the subcortical-cerebellum regions (Fig. 4D-F). This may further support the “compensatory hypothesis”, as the additional paths to these three high-order subnetworks could prevent a significant drop in the cognitive performance. In addition, we found that such elevations were still maintained in the AD stage, whereas it was not the case in our previous whole-brain analysis. This could be due to the smaller scale of the brain network we were currently focused on. Of note, the elevated redundancy was also observed between the medial frontal and the frontoparietal subnetworks, showing the backup paths could also help to maintain the interactions between the high-order subnetworks for high-level cognitive functions in MCI.

Another main finding of the redundancy changes in the AD study lies in the association between disease symptom (general cognitive ability, as measured by MMSE) and the redundancy index in the AD continuum, related to the correlation of the redundancy index and MMSE of the subjects in the AD progression study. By Fig. 5, this negative correlation was consistent with our group comparison results, indicating increased redundancy when the symptom was getting worsened. This result indicates that our proposed subnetwork redundancy index could be sensitive to an early sign of AD, which is better than the whole-brain redundancy. Such a protective mechanism through enhanced redundancy could be used to prevent or delay AD progression in future studies.

We also demonstrated the effectiveness of our method in the MDD study and found that the redundancy loss in the medial frontal subnetwork, as well as the connections between the medial frontal and frontoparietal subnetworks, could be associated with MDD. Previously, in MDD studies, disrupted brain network’s efficiency properties have been demonstrated (Zhang, et al., 2011). The most targeted subnetworks in the MDD studies include three high-order networks, i.e., the medial frontal, frontoparietal, and default mode network, as well as their interactions (Kaiser, et al., 2015). Our redundancy index also indicated disrupted medial frontal subnetwork, as well as the medial frontal-frontoparietal interactions (Fig. 6A). Specifically, MDD’s subnetwork seemed to have less tendency to be redundant in these subnetworks. The MDD’s pathology or neuromechanics have mainly been related to aberrant neurotransmitters in the subcortical (e.g., the striatum) regions (Kumar, et al., 2014), but the disrupted cortical (especially the prefrontal cortex (Zuo, et al., 2018)) to subcortical connections have been...
often reported, which involved the medial frontal subnetwork. The functions of this subnetwork also include emotion processes and social behaviors (Waugh, et al., 2014), while the functions of the frontoparietal network mainly include executive control (Veldsman, et al., 2020). The loss of the medial frontal network’s redundancy, as well as the connectivity redundancy between this subnetwork and the frontoparietal network could be the cause of uncontrolled depressive thoughts and negative affections (Ross & Rush, 1981). Besides, from the default mode network where the redundancy level was not significantly affected, we found that increased disease duration could reduce this network’s redundancy. Taken together, this new redundancy index could also help better understand mental disorders.

We think that the number of redundant connections would decrease after the emerging of major depressive disorder due to the disturbance of the disease. This happened to two subnetworks, MF and MF&FP (Fig. 6A), both are believed to be related to the symptom of depression (Kaiser, et al., 2015). On the contrary, the course of AD could be rather long, with many years developing until MCI and many more years to dementia (Petersen, et al., 2018). With AD pathological change in the brain, the symptoms could still be invisible or relatively mild. It is hypothesized that the brains of MCI patients may have adequate time to react to the AD pathological attacks by developing compensatory alterations such as increased FC (Skouras, et al., 2019), which may result in elevated connectivity redundancy, possibly by enhancement of weak FC links or redistribution of FC links, to be able to maintain overall connectedness even with the attacks of AD. The decrease of the redundant connections from MCI to AD (Fig. 4) further favors such a hypothesis. That is, with AD progression, the compensation could not make the entire brain network sufficiently work properly, possibly due to the attack to the redundant connections, which eventually causes collapse and dementia.

Our study has several limitations. First, since our method is based on the frequency of redundancy through the time, this technique can only be applied to dynamic FC in the networks, not the static ones; future works could be carried out based on static FC network or even the structural networks. Second, mediation analysis is considered a very promising method in the study of abnormal aging by treating brain network-derived redundancy (robustness) as a certain protective key metrics in the field of graph theoretic analysis, could be a promising method in the brain network studies.

5. Conclusions

In this study, we carried out a subnetwork-level redundancy analysis that characterized dynamic redundancy within and between specific subnetworks using resting-state fMRI from two disease cohorts (AD and MDD). We revealed the effectiveness and sensitivity of our subnetwork redundancy index in the brain disease studies. Redundancy, as one of the key metrics in the field of graph theoretic analysis, could be a promising method in the brain network studies.

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7. Data availability statement

The data and codes are available from the corresponding authors upon reasonable request.

8. Ethics approval

The experiments and data collection were approved by the local ethics committees, as mentioned in the ADNI data sharing website http://adni.loni.usc.edu. For MDD study, all participants provided written informed consent and the study was approved by the local ethics committee.

9. Consent to participate

Data used from ADNI is publicly available, so this is not applicable. The MDD data are not publicly available due to privacy or ethical restrictions.

Declaration of Competing Interest

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

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