Brain Relatively Inert Network: Taking Adult Attention Deficit Hyperactivity Disorder as an Example

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Resting-state functional MRI (rs-fMRI) has been increasingly applied in the research of brain cognitive science and psychiatric diseases. However, previous studies only focused on specific activation areas of the brain, and there are few studies on the inactivation areas. This may overlook much information that explains the brain’s cognitive function. In this paper, we propose a relatively inert network (RIN) and try to explore its important role in understanding the cognitive mechanism of the brain and the study of mental diseases, using adult attention deficit hyperactivity disorder (ADHD) as an example. Here, we utilize methods based on group independent component analysis (GICA) and t-test to identify RIN and calculate its corresponding time series. Through experiments, alterations in the RIN and the corresponding activation network (AN) in adult ADHD patients are observed. And compared with those in the left brain, the activation changes in the right brain are greater. Further, when the RIN functional connectivity is introduced as a feature to classify adult ADHD patients from healthy controls (HCs), the classification accuracy rate is 12% higher than that of the original functional connectivity feature. This was also verified by testing on an independent public dataset. These findings confirm that the RIN of the brain contains much information that will probably be neglected. Moreover, this research provides an effective new means of exploring the information integration between brain regions and the diagnosis of mental illness.

Keywords: activation network, relatively inert network, functional MRI, group ICA, functional connectivity, adult ADHD

INTRODUCTION

The identification of resting-state networks (RSNs) in the brain from resting-state functional MRI (rs-fMRI) time series (TC) and the analysis of functional connectivity have revealed much about the macroscopic spatio-temporal organization of the brain. The brain’s acquisition of information and decision-making is a manifestation of a range of neural activities. These processes involve a wide range of functional brain networks. Activation responses of voxels are often associated with the multifunctional collaboration of the brain. So even in the same network, different regions may show different levels of activation, and different activation levels can reflect different responses of neurons to different tasks. However, in previous studies, more have been conducted on activated areas of the brain (activation values above a certain threshold). Few have researched relatively inert
regions of the brain, ignoring important information about the presence of relatively inert areas. Therefore, this prompted us to study it and try to discover its important role in the discovery of biomarkers of mental illness through this study.

In this paper, we define relatively inactive regions in the RSN as relatively inert networks (RINs), and we explore the important role of RINs in brain cognition in terms of network and functional connectivity. The method of brain activation detection includes correlation analysis (Bandettini, 2010), general linear model (Friston et al., 2010), cluster analysis (Smolders et al., 2007; Slavakis et al., 2018), principal component analysis (PCA) (Andersen et al., 1999; Smith et al., 2014), and independent component (IC) analysis (ICA) (Mckeown et al., 1998; Lin et al., 2010; Du and Fan, 2013; De Blasi et al., 2020). Among them, correlation analysis and general linear models require a priori knowledge, whereas their more suitable for data analysis. For resting-state data, the detection of activation is equivalent to the identification of RSN. ICA is one of the most widely used methods for estimating brain functional connectivity. For resting-state data, the detection of RIN also plays an important role in the discovery of biomarkers related to their neural mechanisms.

This paper is the first to study the RIN of the brain and to use adult ADHD data as an example to explore its application in the research and diagnosis of mental diseases. Figure 1 shows the entire research process. In the first step (Figure 1A), the GICA method is used to obtain RSN, and then it is selected. In the second step (Figure 1B), the RIN and AN were identified, and then the corresponding TC were estimated using our proposed method. Finally, (Figure 1C) the extracted features were analyzed and used for the training and testing of the support vector machine (SVM) classifier. We observed that RIN and AN were significantly different between adult ADHD patients and healthy controls (HCs). Also, we compared the differences in functional network connectivity (FNC) between the two groups of subjects.

MATERIALS AND METHODS

Independent Component Analysis
The ICA method assumes that the observed random signal $X$ obeys the model as follows (Mckeown et al., 1998):

$$X = AS$$

Since the number of voxels in spatio-temporal fMRI data is much more than the number of time points, the spatial ICA method is usually more suitable for modeling given fMRI data. $X = (x_1, x_2, \ldots, x_L)$ represents a random variable in the spatial ICA, $K$ is the number of time points, $x_i$ is a vector of length $L$, and $X$ represents a given 3D fMRI image with $L$ voxels at time $K$. $S = (S_1, S_2, \ldots, S_N)$ is $N$ unknown and mutually independent source signals (corresponding to the function network), $A$ is an unknown mixing matrix, and the purpose of ICA is to estimate the mixing matrix $A$ and the source signal $S$ only through $X$. The ICA model has two assumptions about the source signal: one is that the observed signal is a linear combination of several statistical independent source signals and the other is that the distribution of the source signal is non-Gaussian (at most one can be Gaussian).

Since both $A$ and $S$ are unknown, their exact values cannot be calculated in the actual solution. The separation matrix $W$ and
the estimate $Y$ can be estimated by optimizing the independence of $S$. For example, the classic ICA method uses minimizing mutual information (Amari et al., 1996), and Hyvarinen used maximizing non-Gaussian (measured by negentropy or kurtosis) to estimate these components (Hyvarinen, 1999). Specifically, it finds a separation matrix $W$ by constantly updating and iterations:

$$Y = (WA)S = \tilde{S}$$

When $W$ is closer to $A^{-1}$, $Y$ is closer to $S$.

**Group Independent Component Analysis**

An effective method of multi-subject analysis is GICA (Allen et al., 2012), which can identify ICs at the group level and reconstruct the independent specific subject components at the subject level. Compared with ICA for each subject separately, it is easier for GICA to establish a direct correspondence of different subject ICs. First, the group IC (GIC) is calculated by the GICA method, and then specific subject ICs are reconstructed by using the GIC. Common reconstruction methods include GICA
(Chen et al., 2008), constrained ICA (cICA) (Wei and Rajapakse, 2005), spatio-temporal regression (Nickerson et al., 2017), and GIG-ICA (Du and Fan, 2013). In this paper, the GIG-ICA is used to extract brain functional networks in fMRI data, and specifically, the method transformed this process into the multi-objective optimization problem described in (3).

\[
\max \left\{ Q(w_i) = [E[G(Y_i)] - E[G(z)]]^2 
\right\} \\
\text{s.t. } \parallel w_i \parallel = 1
\]

(3)

\(Q(w_i)\) is the negative entropy of \(Y_i\), which is the \(i\)th estimated IC, \(Y_i = w_i^T X; \hat{X}\) is the random vector after PCA whitening, and \(R_i\) represents the spatial reference that is the GIC of the zero mean and unit variance in the previous step after normalization. \(E[.]\) represents a mathematical expectation, \(G[.]\) is an arbitrary non-quadratic function, and \(z\) is a Gaussian variable with zero mean unit variance. \(C(w_i)\), which is specifically defined as \(\phi[Y_i, R_i] = E[Y_i R_i]\), is used to measure the closeness between the estimated component \(Y_i\) and the reference \(R_i\). By solving the multi-objective optimization problem, the optimal separation column vector \(w_i\) is generated to maximize the independence of \(Y_i\) and maximize the proximity of the reference \(R_i\) and the independence of \(Y_i\), at the same time. To solve the multi-objective optimization problem, it is usually necessary to find the Pareto optimal set or its subset and to strictly evaluate which specific compromise solution is more suitable for the researched problem. To prevent the optimization from being dominated by a larger cost function, the \(\text{arctan}\) function is used to normalize \(Q(w_i)\), and the objective function is transformed into the following formula:

\[
w_i^* = \arg \max_{w_i} F (w_i) \\
= \arg \max_{w_i} \left[ a \cdot K (w_i) + (1 - a) \cdot C (w_i) \right] \\
\text{s.t. } \parallel w_i \parallel = 1 \\
0 < a < 1
\]

(4)

Here, \(a\) is the weighting parameter, which can be determined empirically. Use the gradient descent method to continuously iterate for solving \(w_i\), and calculate \(Y_i\). The corresponding TC \(T_i\) is equivalent to the mean of blood oxygenation level-dependent (BOLD) series of all voxels weighted by its associated IC z-score.

### Identification of Relatively Inert Network and Activation Network

During a resting-state scan, the internal network patterns of the brain may change substantially (Chen et al., 2015). If we focus only on the activated areas of the brain, it is easy to overlook the potential information in relatively inert areas. We used the GIG-ICA method to separate the signals from the rs-fMRI data of HCs, and we processed the individual components with a one-sample \(t\)-test. We defined the area higher than the threshold \(\mu\) in each IC (corresponding to a certain RSN) as AN and the other area as RIN. Thus, for each RSN, the corresponding AN and RIN can be further obtained. On this foundation, we propose the following method to estimate the TC corresponding to each AN and RIN, and the sum of their TC is the same as the TC of that original IC.

\[T_{act} = \frac{1}{V_{an}} \sum_{v=1}^{V} (y_{iv} \cdot d_v \cdot f_v)\]

\[T_{inc} = \frac{1}{V_{an}} \sum_{v=1}^{V} (y_{iv} \cdot |1 - d_v| \cdot f_v)\]

\(T_{act}\) represents the TC corresponding to the AN, and \(T_{inc}\) represents the TC corresponding to the RIN. \(V_{an}\) and \(V_{an}\) represent the number of voxels in the RIN and AN of the \(i\)th component, respectively. \(V\) is the total number of voxels, \(f_v\) is the \(v\)th column of \(X\), and \(y_{iv}\) is the \(v\)th value of the \(i\)th component. \(d_v\) is the \(v\)th value of \(d(V \times 1)\), which indicates whether the \(v\)th voxel is activated, 1 indicates activated, and 0 indicates inactive. So the estimation is equivalent to the mean of BOLD series of the corresponding areas voxels weighted by the z-score of their associated IC.

### EXPERIMENTS AND RESULTS

#### Materials and Data Preprocessing

The neuroimaging dataset shared by the University of California, Los Angeles (UCLA) Neuropsychiatric Research Association (Poldrack et al., 2016), which includes 138 healthy subjects, 58 subjects with schizophrenia, 49 subjects with bipolar disorder, and 45 subjects with ADHD, was downloaded. The dataset is shared through the OpenfMRI project and formatted according to the brain imaging data structure (BIDS) standard. The participants were asked to stay relaxed and to keep their eyes open, without giving them any stimuli or asking them to respond, and then were scanned to obtain resting-state data using the 3T Siemens Trio scanner (Siemens Healthineers, Erlangen, Germany). A total of 152 time points were scanned. T2*-weighted echo-planar imaging (EPI) sequence was used to collect functional MRI data with the following parameters: repetition time (TR) = 2 s, echo time (TE) = 30 ms, flip angle (FA) = 90°, field of view (FOV) = 192 mm × 192 mm (64 × 64 matrix), number of slices 34, and slice thickness = 4 mm. For more information about the dataset, please refer to the link: https://openneuro.org/datasets/ds00030. The resting-state data of healthy and ADHD subjects aged 21–50 years were used in this paper. The SPM12 software1 was used to preprocess the data including removal of the first 10 time point images, slice timing correction, motion correction, spatial standardization using the Montreal Neurological Institute (MNI) EPI template, and use of 4-mm full width at half maximum (FWHM) Gaussian kernel to smooth in space. Subjects with excessive motion were excluded, and 42 healthy and 41 ADHD subjects with matching gender and age were selected.

To assess the robustness of our results, our method was replicated on an independent publicly ADHD dataset, publicly available for download at http://icon_1000.projects.nitrc.org/ftpclassic/FcpTable.html. This dataset has 25 ADHD patients and 84 healthy subjects. These subjects were recruited by the

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1https://www.fil.ion.ucl.ac.uk/spm/software/spm12/
adult ADHD program group at the New York University School of Medicine, United States. The subjects laid flat in the MRI machine with their eyes open and in a relaxed awake state. The scan TR = 2 s, TE = 25 ms, scan resolution was 64 × 64, intra-slice resolution was 3 mm × 3 mm, slice thickness = 3 mm, and number of slices was 39, covering the entire brain area; and a total of 197 time points were scanned. Subjects who did not undergo psychiatric evaluation or were younger than 20 were excluded. The same preprocessing method used for the UCLA data was used, and subjects with excessive head movements were excluded. To avoid a confounding effect due to an imbalance in the sample, 23 healthy subjects and 24 patients with ADHD were finally selected. The two groups of subjects were matched in terms of gender and age. All subjects were between 20 and 50 years of age. The distribution of the HC-ADHD population for these two datasets is shown in Figure 2. The distribution of the control–ADHD population. ADHD, attention deficit hyperactivity disorder.

As shown in Table 1, there was no significant difference in age and gender between the two groups for each dataset.

Table 1: Age and gender indices of two independent datasets.

|         | UCLA | New York |
|---------|------|----------|
| Age (years) | 32.45 ± 9.05, 32.85 ± 10.89 | 34.89 ± 8.387, 34.75 ± 9.712 |
| Gender | 23/21, 20/21 | 14/9, 19/5 |
| Statistical test | 0.85/0.91, 0.96 (t-test)/0.19 (chi-square test) |

Group Independent Component Analysis and Component Identification

The PCA is used to reduce the dimensionality of each subject’s data for simplifying the ICA algorithm. Then the time cascade method is used to combine all the subjects’ reduced data. Another PCA (group level) was performed to further reduce the time dimension of the group data to the number of IC, and then the Infomax algorithm (Bell and Sejnowski, 1995) was applied to extract the GIC. GIFT software (version 3.0c)² was used to perform GICA in the two groups of subjects. Meanwhile, the minimum description length (MDL) standard (Li et al., 2007) was utilized to estimate the dimensions to determine the number of components. HCs were estimated with 30 components, and ADHD with 29. In order to obtain a reliable GIC, the ICASSO method (Himberg and Hyvärinen, 2003) was used to repeat the ICA 20 times. And the GIG-ICA method was used to reconstruct the corresponding TC and the unique IC of each subject. The IC that we needed was identified and picked out. In the first step, the correlation between the spatial map of each component and the prior mask map of gray matter, white matter, and cerebrospinal fluid (CSF) provided in the DPABI software package (Yan et al., 2016) was calculated in the MNI standardized brain space. Since components with high correlation with prior CSF or white matter or low correlation with the gray matter may be affected by human factors (Assaf et al., 2010), they were excluded to ensure that the selected components are mainly concentrated in the gray matter area. In the second step, due to the low-frequency characteristics of the BOLD signal, the power spectrum of the network should show higher low-frequency spectrum power. Therefore, the ratio of the low-frequency power to the high-frequency power of each component was calculated, and the component with a higher ratio was screened out (most ratios of low-frequency to high-frequency power of some networks are greater than 10). The third step is to calculate the spatial correlation between the remaining components and the eight core intrinsic connectivity networks (ICNs) reported by Beckmann et al. (2005) and to select the eight best-matched components with the visual inspection and correlation coefficients. These eight networks are (a) medial visual network (MVN), (b) lateral occipital visual cortex (LO), (c) auditory area network (auditory), (d) sensorimotor area (sensorimotor), (e) DMN, (f) executive control network (ECN), (g) and (h) right-lateralized fronto-parietal network (RFPN) and left-lateralized fronto-parietal network (LFPN). Figure 3 shows the sagittal, coronal, and cross-sectional planes of the eight best-matched ICNs. Figure 3A (left) shows the activation peak coordinates of eight ICNs from a to h in the HC group, and Figure 3B (right) shows the activation peak coordinates of 8 ICNs from a to h in the ADHD group. It is worth noting that after comparing the activation peak coordinates of the eight large-scale networks of the two groups of subjects, it is found that the network with the closest peak coordinates in the sensorimotor area is followed by LO, LFPN, and DMN, while the network with the largest difference in activation peak coordinates in the auditory area is followed by RFPN and ECN.

Comparing Group Differences in the Relatively Inert Networks and the Activation Networks

A one-sample t-test was used to determine the RINS and ANS of HCs and ADHD patients. In this way, each RSN could be divided

2 http://icatb.sourceforge.net/, version 3.0c.
into AN and RIN. The areas with different activation levels in the two groups of subjects were counted, and a two-sample t-test was further performed. Figure 4 shows the areas where ANs and RINs differed significantly between the two groups of subjects when the threshold $\mu = 3$ ($p < 0.05$). There are two scenarios for this difference. One is that the areas (indicated by cold areas) are the intersections of the RINs of HCs and the ANs of adult ADHD; i.e., HCs show no activation, whereas adult ADHD patients show significant activation. These areas show abnormally enhanced activation in adult ADHD patients; the second (indicated by hot area) is that the areas are the intersections of ANs of HCs and RINs of adult ADHD; i.e., HCs show activation, but ADHD patients show no activation. In other words, the activation in these areas is significantly weakened in adults with ADHD. Some responses in cognitive function are usually reflected in the brain activation. Statistical analyses were conducted in these abnormal areas. Compared with those in HCs, the areas of enhanced activation in adult ADHD patients were concentrated in the precentral gyrus (Brodmann areas 4 and 6), insula (Brodmann area 13), postcentral gyrus (Brodmann areas 3, 43, and 40), cuneus (Brodmann areas 7, 18, and 19), and superior temporal gyrus (Brodmann area 41). The areas of weakened activation were mainly concentrated in the insula (Brodmann area 13), superior temporal gyrus (Brodmann area 22), medial frontal gyrus (Brodmann areas 8, 9, and 10), and inferior parietal lobule (Brodmann areas 7, 39, and 40). Furthermore, when analyzing RSN activation in both groups of subjects, it was found that the activation difference of adult ADHD patients is distributed in both the left and right hemispheres, compared with that of the HCs. However, Figure 4 shows that there are more areas of discrepancy in the right hemisphere than in the left hemisphere.

**Group Differences in Functional Network Connectively**

The FNC reflects, to some extent, the information interaction between different networks (Power et al., 2011). In this study, the method in section “Identification of Relatively Inert Network and Activation Network” was used to estimate the TC corresponding to the RINs and ANs for each participant. Applying Pearson’s correlation coefficient to calculate the FNC, a $16 \times 16$ FNC matrix was obtained for each subject. Figure 5 shows the mean FNC for each group of subjects. The bottom left corner of the matrix shows that there is some connectivity between RINs and ANs. Also, comparing Figures 5A,B, it was observed that AN and RIN of the same functional network are highly connected for both ADHD patients and HCs. And the connectivity between RINs is higher than the connectivity between ANs. In addition, the differences in functional connectivity between different networks were more pronounced in HCs than in ADHD patients.

Accurate identification of disease-specific induced changes in functional connectivity is considered an important task that can highlight the underlying mechanisms of the disease. Therefore, a two-sample t-test was performed for each element of the FNC matrix to compare the different functional connectivity between the two groups. Figure 6A shows the FNCs with significant differences between HCs and adult ADHD ($\mu = 3$, $p < 0.05$,
false discovery rate (FDR) corrected]. Figure 6B shows the number of altered connections for each network. It can be found that the auditory area and RFPN changed the most. Also, the enhancement and weakening of the FCN were further counted. Figure 6C shows that the connectivity of ECN\textsubscript{i} was significantly weakened in the ADHD group compared with the HC group. Meanwhile, compared with HCs, adult ADHD patients show a very strong connection in most changed FNCs, especially those involving the auditory area.

**Classification Based on Relatively Inert Networks and Activation Networks**

**Classification Within University of California, Los Angeles, Dataset**

In order to verify the importance of RINs in assisting the diagnosis of mental disorders, four different strategies were used to construct features, namely, the connectivity of the original functional network (OriginalFNC), the functional connectivity of ANs (ActFNC), the functional connectivity of RINs (InertFNC), and the functional connectivity of both ANs and RINs (AllFNC). SVM models were utilized to classify HC and ADHD. Thirty percent of the UCLA dataset was used as the test set, and the remaining part as the training set. Feature selection using XGBoost (Chen and Guestrin, 2016) and 10-fold cross-validation was used to train the models. The predicted values of each strategy were obtained for McNemar’s test. As a result, the AllFNC classification performance was significantly higher than the OriginalFNC at \( \mu = 1, 2, \) and 3 (\( p < 0.05 \)). Further, the data label was randomly shuffled, and the classification was repeated 100 times to test the significance along the massive permutation. Figure 7 shows the box plots of accuracy, sensitivity, specificity, and F1-score for 100 classifications of the four features with different values of the threshold \( \mu \).

When \( \mu = 2 \), AllFNC offers the highest classification (the average accuracy = 94.2%, sensitivity = 92.1%, specificity = 96.3%, and F1-score = 93.7%). The next highest is InertFNC (accuracy = 87.1%, sensitivity = 86.8%, specificity = 87.4%, F1-score = 86.6%) and then OriginalFNC classification (the
FIGURE 6 | Group differences in functional network connectivity. (A) The FNC with significant differences between HCs and ADHD patients under the two-sample t-test. Only the case of $p < 0.05$ (FDR corrected) is shown here, where red indicates that the FNC of ADHD patients is stronger than that of HCs, and blue indicates that the FNC of ADHD patients is lower than that of HCs. (B) The statistical result of a significant alteration in FNC. The horizontal axis is the functional network, and the vertical axis is the number of alterations FNCs. RIN, relatively inert network; AN, activation network. (C) The statistics of enhanced and weakened network connectivity of FNC. Both with $\mu = 3$. FNC, functional network connectivity; HC, healthy control; ADHD, attention deficit hyperactivity disorder; FDR, false discovery rate.

FIGURE 7 | Box plots comparing the performance of classification using different features. Red represents the OriginalFNC, green represents the ActFNC, blue represents the InertFNC, and purple represents the AllFNC. (A–D) The horizontal axis is the threshold $\mu$; vertical axis represents the accuracy, sensitivity, specificity, and F1-score. * indicates $p < 0.05$, and ** indicates $p < 0.01$. The average accuracy = 82.0%, sensitivity = 82.3%, specificity = 82.1%, F1-score = 81.5%). A two-sample t-test was used, and the results showed that both AllFNC and InertFNC were significantly better classified than the OriginalFNC ($p < 0.01$). Similar results can be obtained when $\mu$ takes other values.

As with most psychiatric disorders, there are no specific physiological indicators to diagnose adult ADHD (Cortese et al., 2018). To locate mutations more accurately, a single FNC as a feature was used to classify adults with ADHD and HCs, and the experiment was repeated 20 times. Features with F1-scores...
higher than 0.65 were selected under four thresholds (μ). Table 2 shows the mean of the classification performance of the four best-performing features that were finally screened. When they were examined, it was found that all four features were significantly different, and all showed enhanced ADHD functional connectivity. The classification results of the four best-performing features are shown in Table 3. Notably, all four features were associated with the Auditory_a network. The FNC between Auditory_a and Sensorimotor_i was used as a feature for classification with an accuracy of 0.87 and an area under the curve (AUC) of 0.91.

**Generalization to New York Dataset**

Sample size has a profound impact on the variability of accuracy and accuracy estimates (Flint et al., 2021). To avoid exaggeration of statistical power and classification parameters, independent publicly available datasets were used to compare the performance of the original functional connectivity and the functional connectivity with the introduction of RINs and ANs on the classification of adult ADHD. Specifically, with the complete UCLA dataset was trained and tested in New York dataset, which features different acquisition parameters. It is worth noting that in practical application scenarios, the labels of the samples are not available before classification. Therefore, for all test sets, cIcA was utilized to compute a specific set of eight RSNs, which avoids the restriction in GICA that requires grouping the data in advance. The computation of features in the UCLA data is then replicated. Similarly, the XGBoost method is employed to select the features. Meanwhile, the data were randomly shuffle, SVM was repeated 100 times for the classification results shown in Figure 8. The experimental results show that the average classification accuracy, sensitivity, specificity, and F1-score of the OriginalFNC100 are 0.438, 0.472, 0.402, and 0.459, respectively, while the average accuracy, sensitivity, specificity, and F1-score of the AllFNC100 classifications with the introduction of RIN and AN are 0.647, 0.675, 0.620, and 0.663, respectively. A two-sample t-test was further performed on the results of the two methods, and it was found that their differences were significant (p < 0.01). Although it is difficult to avoid the degradation of classification accuracy in the independent dataset because more work related to feature engineering was not performed (using only functional connectivity), our method still has a greater advantage over the OriginalFNC.

**Threshold Analysis**

After extraction of the IC (corresponding to the functional network of the resting brain) using the ICA method, the results are usually Z-transformed. For the determination of brain activation areas, the area above the threshold is AN, while the area below the threshold is RIN. The threshold μ can be determined empirically. To assess the effect of the threshold μ on the study results, the experiments compared four different FNCs under different threshold conditions. As the threshold value increases, the range of AN becomes smaller. Therefore, the threshold value is usually not set too large. In this study, ECN_i is empty when μ is not less than 4. Therefore, the differences in FNCs between the two groups of subjects were compared under four conditions with threshold μ equal to 1, 2, 3, and 4. Figure 9 shows the FNC with significant differences between the adult ADHD and HC groups at different threshold value μ. The results show that when μ is taken at different values, there is only a small change in functional connectivity with significant differences between the two groups of subjects. This also suggests that the changes in FNC differences between the adult ADHD and HC groups are within acceptable limits when the thresholds are within a certain range.

To further exclude the effect of different thresholds on the classification performance of HCs and ADHD patients, the classification accuracy, sensitivity, specificity, and F1-score were calculated when the threshold μ was taken with four different values of 1, 2, 3, and 4 in the experiments in section “Classification Within University of California, Los Angeles, Dataset.” As shown in Figure 7, the classification performance using AllFNC is always the best when μ takes different values. Also, the performance of classification using only the functional connectivity features between RINs in most situations is higher than the performance of the original network functional connectivity. Table 4 shows the average accuracy, sensitivity, specificity, and F1-score values

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**Table 2** | The comprehensive results of the classification performance for a single feature.

| FNC                  | Mean F1-score |   |
|----------------------|---------------|---|
| RFPN_i-Auditory_i    | 0.69          |   |
| RFPN_a-Auditory_i    | 0.72          |   |
| RFPN_i-Sensorimotor_i| 0.72          |   |
| RFPN_i-Auditory_a    | 0.74          |   |

FNC, functional network connectivity.

**Table 3** | Best classification performance of a single feature.

| FNC                  | Mean F1-score |   |
|----------------------|---------------|---|
| RFPN_i-Auditory_i    | 0.69          |   |
| RFPN_a-Auditory_i    | 0.72          |   |
| RFPN_i-Sensorimotor_i| 0.72          |   |
| RFPN_i-Auditory_a    | 0.74          |   |

FNC, functional network connectivity.

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**Figure 8** | Box plot of predicted performance using independent datasets. Two-sample t-test showing significant differences in all indicators.
for 100 classifications at different threshold values ($\mu$). It can be demonstrated above that there is no significant effect on the classification results when the threshold value is 1–4.

**DISCUSSION**

In this paper, we propose a method for the identification and estimation of RINs and apply it to the study and diagnosis of the disease using fMRI data of adult ADHD as an example. Compared with the traditional methods, our method focuses not only on the activated areas of the brain but also on the relatively inert areas of the brain. The activated areas of the brain are usually delineated by setting a threshold value. However, brain areas encode information in multivariate responses, and neural computations are nonlinear (Cannon et al., 2011). Strictly speaking, the portion below the threshold (the relatively inert area) is not completely inactive, and weaker or nonlinear reactions may be present. It is likely that these responses are not decomposed by ICA (or other methods of matrix decomposition), since these methods usually assume that the obtained BOLD signal is a linear accumulation of the source signal. As a result, we believe that the relatively inert areas also contain important information that can be used for disease diagnosis.

Specifically, by comparing ANs and RINs, we identified abnormal activation areas in the brains of adults with ADHD compared with healthy subjects. Mainly, the activation was significantly enhanced in the precentral gyrus (Brodmann areas 4 and 6), postcentral gyrus (Brodmann areas 3, 40, and 43), and superior temporal gyrus (Brodmann area 41), while the activation was significantly weakened in the superior temporal gyrus (Brodmann area 22), medial frontal gyrus (Brodmann areas 8, 9, and 10), insula (Brodmann area 13), and cuneus and inferior parietal lobule (Brodmann areas 7, 39, and 40). This suggests that the increased activation in Brodmann areas 4, 6, and 40 in children with ADHD (Suskauer et al., 2008) did not disappear in adult patients either. Abnormalities in Brodmann area 4, which controls behavioral movements, and Brodmann area 8, which together with Brodmann area 6 constitutes the premotor cortex, are likely to be associated with overactive symptoms in ADHD patients. In addition to this, we found changes in both right- and left-brain activation in adult ADHD patients and more extensive changes in right brain activation. This is similar to the changes in the right hemisphere in childhood ADHD patients (Zou and Yang, 2019).

The results in Figure 5 show that there is a correlation between RINs and ANs as well as different areas of different networks. The exchange of information between these areas can be further explained by the fact that brain processing of complex cognitive tasks is not only related to ANs, but RINs are also involved. We also found some weakened connectivity of the auditory area with other networks in adult ADHD patients. For example, the functional connection between Auditory_a and Sensorimotor_i, RFPN_i, and Auditory_i is weakened. This coincides with studies related to ADHD patients showing deficits in the auditory area (Serrallach et al., 2016; Bijlenga et al., 2017). And these

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**TABLE 4** Classification performance of different features under different thresholds.

| Feature | Mean accuracy | Mean sensitivity | Mean specificity | Mean F1-score |
|---------|---------------|-----------------|-----------------|---------------|
| OriginalFNC | 0.820 | 0.823 | 0.821 | 0.815 |
| ActFNC | 0.820 | 0.855 | 0.796 | 0.815 |
| InertFNC | 0.910 | 0.916 | 0.909 | 0.908 |
| AllFNC | 0.979 | 0.981 | 0.979 | 0.979 |
| ActFNC | 0.849 | 0.874 | 0.826 | 0.848 |
| InertFNC | 0.871 | 0.868 | 0.874 | 0.866 |
| AllFNC | 0.942 | 0.921 | 0.963 | 0.937 |
| ActFNC | 0.886 | 0.891 | 0.882 | 0.886 |
| InertFNC | 0.833 | 0.802 | 0.873 | 0.820 |
| AllFNC | 0.930 | 0.925 | 0.935 | 0.928 |
| ActFNC | 0.853 | 0.860 | 0.850 | 0.849 |
| InertFNC | 0.838 | 0.820 | 0.856 | 0.832 |
| AllFNC | 0.957 | 0.960 | 0.951 | 0.966 |

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FIGURE 9 | FNC with significant difference between ADHD and HC groups under different threshold $\mu$. (A–D) Different thresholds and the two-sample t-test $p < 0.05$ (FDR corrected). Red represents that the functional connectivity of ADHD is stronger than that of HC, and blue represents that the functional connectivity of ADHD is weaker than that of HC. FNC, functional network connectivity; ADHD, attention deficit hyperactivity disorder; HC, healthy control; FDR, false discovery rate.
findings could not be obtained using the original functional connectivity analysis.

Across studies, there appear to be significant network disruptions involved in ADHD (Cannon et al., 2011). Therefore, the detection of abnormal connectivity is crucial to further localize areas associated with psychiatric disorders. We further compared the ability to detect abnormal FNCs in ADHD patients using original networks and the introduction of RINs and ANs in Figure 10. In Figure 10A, the FNC changes in ADHD patients are mainly concentrated in the RFPN, while the weakened functional connectivity is reflected in the DMN and ECN. Figure 10B uses pie charts to count the distribution of functional connectivity changes. The two plots on the left depict all changes (including enhancement and weakening). It can be noticed that both results are similar, but more changes in the auditory area can be detected with the introduction of RIN. It is worth noting that the four best-performing features finally filtered in Table 3 are all related to the auditory network, as evidenced by the results in Table 2. Therefore, we have reasons to believe that the auditory changes are real. Moreover, it can also be found in the two plots on the right in subplot (B). Also, for the weakened connections, the original network analysis could not find the weakened LFPN connections. This suggests that the introduction of RINs and ANs may reveal not only the differences analyzed at the original RSN level but also the discovery of additional areas associated with changes in cognitive function in disease. This point is likely to have been overlooked in past studies.

In addition, features constructed based on RINs and ANs (functional connectivity is used in this paper) produce a relatively accurate diagnoses of adult ADHD, with better classification results than those constructed using the original RSN. This effect may be related to the use of the large amount of information present in the RIN. Although it is difficult to avoid the decrease of accuracy in the independent dataset because we did not do more work related to feature engineering (using only functional connectivity), our method still has a greater advantage over the features constructed by the original RSNs. In addition, features based on RINs and ANs, such as dynamic functional connectivity (Sun et al., 2021), dynamic brain fluctuations (Moguilner et al., 2021), or combined with non-image information features (Riaz et al., 2018), are expected to lead to better diagnostic results.

CONCLUSION

The exploration of brain RINs has provided new research ideas to study the integration of information between brain areas and disease diagnosis. In this study, we achieved the identification of brain RINs and ANs. Further experiments showed that brain RINs also contain much information about cognitive functions.
At the same time, we proposed a new method to achieve diagnosis and study of adult ADHD patients using FNC of RINs and ANs. The experimental results showed that our method well improved the accuracy of disease diagnosis (0.98). This is an initial study of the efficacy of RINs. The ICA-based method we used is also a preliminary attempt. In addition, the eight functional networks we used were not customized for adult ADHD but for the most classical cognitive functional networks in the brain. Therefore, our study is also applicable to the study of other psychiatric disorders, such as bipolar disorder, depression, and Alzheimer's disease. Due to time constraints, only functional connectivity was used in this study. In the future, we will implement more features such as effective connectivity and dynamic functional connectivity and will continue to explore more functional brain integration and disease diagnostic outcomes in brain RINs.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/supplementary material, further inquiries can be directed to the corresponding author/s.

ETHICS STATEMENT

The studies involving human participants were reviewed and approved by the Institutional Review Boards at UCLA and the Los Angeles County Department of Mental Health. The patients/participants provided their written informed consent to participate in this study.

AUTHOR CONTRIBUTIONS

HZ and WZ contributed to the conception and design of this research. HZ, JD, and YL derived and designed the theoretical and experimental parts in the article. HZ wrote the first draft of the manuscript. HZ, WZ, YS, and LZ participated in the revision, reading, and approval of the manuscript. All authors contributed to the article and approved the submitted version.

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