Age-related connectivity differences between attention deficit and hyperactivity disorder patients and typically developing subjects: a resting-state functional MRI study

Jisu Hong1,2, Bo-yong Park1,2, Hwan-ho Cho1,2, Hyunjin Park3,4,*
1 Department of Electronic, Electrical and Computer Engineering, Sungkyunkwan University, Suwon, Korea
2 Center for Neuroscience Imaging Research (CNIR), Institute for Basic Science, Suwon, Korea
3 School of Electronic and Electrical Engineering, Sungkyunkwan University, Suwon, Korea

How to cite this article: Hong J, Park BY, Cho HH, Park H (2017) Age-related connectivity differences between attention deficit and hyperactivity disorder patients and typically developing subjects: a resting-state functional MRI study. Neural Regen Res 12(10):1640-1647.

Funding: This work was supported by the Institute for Basic Science [grant No. IBS-R015-D1] and the National Research Foundation of Korea [grant No. NRF-2016R1A2B4008545].

Abstract
Attention deficit and hyperactivity disorder (ADHD) is a disorder characterized by behavioral symptoms including inattention/impulsivity among children, adolescents, and adults. These ADHD related symptoms are influenced by the complex interaction of brain networks which were under explored. We explored age-related brain network differences between ADHD patients and typically developing (TD) subjects using resting state fMRI (rs-fMRI) for three age groups of children, adolescents, and adults. We collected rs-fMRI data from 184 individuals (27 ADHD children and 31 TD children; 32 ADHD adolescents and 32 TD adolescents; and 31 ADHD adults and 31 TD adults). The Brainnetome Atlas was used to define nodes in the network analysis. We compared three age groups of ADHD and TD subjects to identify the distinct regions that could explain age-related brain network differences based on degree centrality (DC), a well-known measure of nodal centrality. The left middle temporal gyrus showed significant interaction effects between disease status (i.e., ADHD or TD) and age (i.e., child, adolescent, or adult) (P < 0.001). Additional regions were identified at a relaxed threshold (P < 0.05). Many of the identified regions (the left inferior frontal gyrus, the left middle temporal gyrus, and the left insular gyrus) were related to cognitive function. The results of our study suggest that aberrant development in cognitive brain regions might be associated with age-related brain network changes in ADHD patients. These findings contribute to better understand how brain function influences the symptoms of ADHD.

Key Words: nerve regeneration; attention deficit and hyperactivity disorder; cognitive function; connectivity; resting-state fMRI; Brainnetome Atlas; whole brain analysis; disease-aging interaction effect; neuroscience; neural regeneration

Introduction
Attention deficit and hyperactivity disorder (ADHD) is a brain disorder that is characterized by the symptoms of inattention and hyperactivity/impulsivity (Schneider et al., 2006; Subcommittee on Attention-Deficit/Hyperactivity Disorder, 2011; Castellanos and Proal, 2012). In addition to inattentive or hyperactive behaviors, ADHD is also known to be highly associated with cognitive dysfunction (Wilens et al., 1999; Sagen, 2006; Rostain and Ramsay, 2006; Solanto et al., 2008; Knouse and Safren, 2010; Castellanos and Proal, 2012). Castellanos et al. (2006) suggested that ADHD-related studies should consider cognitive deficits in ADHD patients to better quantify their neurobehavioral symptoms. Previous studies have adopted cognitive behavioral treatment (CBT) approaches to treat ADHD patients (Wilens et al., 1999; Rostain and Ramsay, 2006; Solanto et al., 2008; Knouse and Safren, 2010). Solanto et al. (2008) found enhanced executive skills in ADHD patients who received CBT and others found a significant reduction in ADHD-related symptoms after receiving combined medication and CBT (Rostain and Ramsay, 2006). These studies suggested that ADHD is highly related to dysfunctions in cognitive processes.

ADHD is a lifetime mental disorder and it has been found that patients show distinct behavioral symptoms across different age groups (Bresnahan and Barry, 2002; Schneider et al., 2006; Hurtig et al., 2007; Subcommittee on Attention-Deficit/Hyperactivity Disorder, 2011; Castellanos and Proal, 2012; Park et al., 2016). These ADHD related symptoms are influenced by the complex interaction of brain networks which are typically explored using neuroimaging approaches (Zang et al., 2007; Tian et al., 2008; Cortese et al., 2012). Most ADHD studies have focused on exploring the differences in brain function in limited age groups (i.e., only in children or adolescents) and studies investigating brain networks among a wide spectrum of age groups (i.e., from children to adults) have been largely lacking (Wilens et al., 1999; Castellanos et al., 2006; Knouse and Safren, 2010; Konrad and Eickhoff, 2010; Uekermann et al., 2010). ADHD...
patients show age dependent alterations in brain networks which have not been fully explored. Here, we aimed to explore the age-related functional changes in brain networks in ADHD patients.

We explored the age-related brain network differences between ADHD patients and typically developing (TD) subjects using resting state functional magnetic resonance imaging (rs-fMRI). Rs-fMRI is an effective tool for analyzing neuro-behavioral disorders such as ADHD (dos Santos Siqueira et al., 2014). One study reported that rs-fMRI demonstrated enhanced brain activation in the sensory-related cortices of adolescent ADHD patients (Tian et al., 2008). Another study found that a feature derived from rs-fMRI known as amplitude of low-frequency revealed significant differences between children with ADHD and TD children (Zang et al., 2007).

We assessed functional brain network differences using a network centrality measure which has been widely used to assess regional importance (Bullmore et al., 2009; Rubinov and Sporns, 2010; Ferreira and Busatto, 2013). We hypothesized that there would be age-related functional network differences between ADHD patients and TD subjects. In this study, we aimed to explore functional brain network changes related to ADHD among a wide spectrum of age groups.

Subjects and Methods

Subjects and imaging data

The Institutional Review Board (IRB) of Sungkyunkwan University approved our retrospective study (#2015-09-007). Our study was performed in full accordance with the principles of the Declaration of Helsinki, and informed consent was obtained from all subjects. We collected raw T1-weighted structural MRI and rs-fMRI data from the ADHD-200 database (ADHD-200 Consortium, 2012; Belloc et al., 2017). We also obtained structural and functional MRI data from the Human Connectome Project (HCP) database (Van Essen et al., 2013). The ADHD-200 database provided the child and adolescent data and the HCP database provided the adult data. The subjects were recruited via advertisement and further details were available (ADHD-200 Consortium, 2012; Van Essen et al., 2013). Scores related to ADHD symptoms were measured using Conner’s Parent Rating Scale Revised, Long Version (CPRS-LV) for the ADHD-200 dataset and Diagnostic and Statistical Manual of Mental Disorders, Fourth Edition (DSM-IV) for the HCP data (American Psychiatric Association, 1994; Conners et al., 1998). With both the ADHD-200 and HCP datasets, subjects with T-scores greater than or equal to 65 on at least one measure of the ADHD-related index were selected as ADHD patients. Subjects with a secondary diagnosis were excluded along with subjects who did not have ADHD-related scores. Based on these criteria, we classified subjects into the ADHD (n = 90) and TD groups (n = 94). Each group was further divided into child, adolescent, and adult groups based on age. Subjects under 10 years of age were considered children, and subjects between 10 and 19 years of age were classified as adolescents. Finally, 27 ADHD children, 32 ADHD adolescents, 31 ADHD adults, 31 TD children, 32 TD adolescents, and 31 TD adults were included in the study. Comparison of the sex ratio did not yield significant differences among the groups. Detailed participant information is given in Table 1. Although the ADHD-200 database consists of eight data collection sites, data from only two sites were retained after adopting the criteria mentioned above: the Kennedy Krieger Institute (KKI) and New York University Child Study Center (NYU). The T1-weighted structural data from the KKI were acquired with the following imaging parameters: repetition time (TR) = 8.0 ms; echo time (TE) = 3.7 ms; field of view (FOV) = 256 × 256 mm²; and voxel resolution = 1.0 × 1.0 × 1.0 mm³. The rs-fMRI functional data from the KKI were acquired with the following imaging parameters: TR = 2,500 ms; TE = 30 ms; FOV = 256 × 256 mm²; number of slices = 72; and voxel resolution = 2.67 × 2.67 × 3.0 mm³. The T1-weighted structural data from NYU were acquired with the following imaging parameters: TR = 2,530 ms; TE = 3.25 ms; FOV = 256 × 256 mm²; and voxel resolution = 1.3 × 1.0 × 1.3 mm³. The rs-fMRI functional data from NYU were acquired with the following imaging parameters: TR = 2,000 ms; TE = 15 ms; FOV = 240 × 192 mm²; number of slices = 33; and voxel resolution = 3.0 × 3.0 × 4.0 mm³. The T1-weighted structural data from the HCP were acquired with the following imaging parameters: TR = 2,400 ms; TE = 2.14 ms; FOV = 224 × 224 mm²; and voxel resolution = 0.7 × 0.7 × 0.7 mm³. Finally, the rs-fMRI functional data from the HCP were acquired with the following imaging parameters: TR = 720 ms; TE = 3.31 ms; FOV = 208 × 180 mm²; number of slices = 72; and voxel resolution = 2.0 × 2.0 × 2.0 mm³. The TD-child group included 16 subjects from the KKI site and 15 subjects from the NYU site. The TD-adolescent group included 16 subjects from the KKI site and 16 subjects from the NYU site. The TD-adult group included 31 subjects from the HCP site. The ADHD-child group included 6 subjects from the KKI site and 21 subjects from the NYU site. The ADHD-adolescent group included 5 subjects from the KKI site and 27 subjects from the NYU site. The ADHD-adult group included 31 subjects from the HCP site.

Imaging preprocessing

The neuroimaging data from the ADHD-200 and HCP were preprocessed using AFNI and FSL software (Cox, 1996; Jenkinson et al., 2012). These preprocessing steps consisted of structural and functional preprocessing. The structural preprocessing included the following steps: performing a de-oblique procedure; reorienting into a right posterior inferior (RPI) orientation; skull-stripping; registering the skull-stripped anatomical image onto the template space at 3 × 3 × 3 mm³ resolution; segmentation into cerebral-spinal fluid (CSF), white matter (WM), and gray matter (GM); and constructing WM and CSF masks by binarizing probability masks using a 0.99 threshold. The functional preprocessing included the following steps: performing a de-oblique procedure; reorienting into a RPI orientation; removal of the first 6 echo planar imaging (EPI) volumes; motion correcting EPI volumes; frame scrubbing based on frame-wise displacement (FD) to exclude frames with FD greater
than 0.5 mm (Power et al., 2012); slice timing correction; registration of the mean EPI image onto the corresponding anatomic image; masking the dataset to exclude non-brain tissue; mapping of the fMRI data and mean image onto the anatomic image; filtering of time courses; and blurring the data using a 6 mm full width half maximum (FWHM) Gaussian filter.

**Network construction**

To construct the functional network from the images, connectivity analysis was performed with regions of interest

<table>
<thead>
<tr>
<th>Brain region (index number of the ROI)</th>
<th>DOF</th>
<th>F value</th>
<th>P value (age × disease)</th>
<th>P value (age)</th>
<th>P value (disease)</th>
</tr>
</thead>
<tbody>
<tr>
<td>L. Superior frontal gyrus (9)</td>
<td>2, 178</td>
<td>3.0891</td>
<td>0.047991</td>
<td>0.787615</td>
<td>0.198468</td>
</tr>
<tr>
<td>L. Inferior frontal gyrus (29)</td>
<td>2, 178</td>
<td>4.0198</td>
<td>0.019611</td>
<td>0.811140</td>
<td>0.246761</td>
</tr>
<tr>
<td>L. Inferior frontal gyrus (33)</td>
<td>2, 178</td>
<td>4.1594</td>
<td>0.017161</td>
<td>0.675609</td>
<td>0.253190</td>
</tr>
<tr>
<td>R. Inferior frontal gyrus (36)</td>
<td>2, 178</td>
<td>5.5417</td>
<td>0.006426</td>
<td>0.828528</td>
<td>0.134809</td>
</tr>
<tr>
<td>L. Inferior frontal gyrus (39)</td>
<td>2, 178</td>
<td>3.1641</td>
<td>0.046438</td>
<td>0.872547</td>
<td>0.187522</td>
</tr>
<tr>
<td>R. Precentral gyrus (64)</td>
<td>2, 178</td>
<td>3.6681</td>
<td>0.027475</td>
<td>0.728483</td>
<td>0.734674</td>
</tr>
<tr>
<td>L. Superior temporal gyrus (77)</td>
<td>2, 178</td>
<td>3.4586</td>
<td>0.033605</td>
<td>0.834183</td>
<td>0.360218</td>
</tr>
<tr>
<td>L. Middle temporal gyrus (83)</td>
<td>2, 178</td>
<td>7.2908</td>
<td>0.000905</td>
<td>0.645098</td>
<td>0.061416</td>
</tr>
<tr>
<td>L. Middle temporal gyrus (85)</td>
<td>2, 178</td>
<td>3.5759</td>
<td>0.030019</td>
<td>0.887161</td>
<td>0.542660</td>
</tr>
<tr>
<td>L. Middle temporal gyrus (87)</td>
<td>2, 178</td>
<td>3.271</td>
<td>0.040261</td>
<td>0.977927</td>
<td>0.617160</td>
</tr>
<tr>
<td>R. Postcentral gyrus (156)</td>
<td>2, 178</td>
<td>3.457</td>
<td>0.033657</td>
<td>0.712092</td>
<td>0.820640</td>
</tr>
<tr>
<td>L. Insular gyrus (165)</td>
<td>2, 178</td>
<td>3.6151</td>
<td>0.028909</td>
<td>0.362170</td>
<td>0.551337</td>
</tr>
<tr>
<td>L. Insular gyrus (171)</td>
<td>2, 178</td>
<td>3.1477</td>
<td>0.045352</td>
<td>0.941490</td>
<td>0.891245</td>
</tr>
<tr>
<td>L. Medioventral occipital cortex (191)</td>
<td>2, 178</td>
<td>4.089</td>
<td>0.018355</td>
<td>0.727416</td>
<td>0.968738</td>
</tr>
<tr>
<td>R. Medioventral occipital cortex (192)</td>
<td>2, 178</td>
<td>4.1922</td>
<td>0.016632</td>
<td>0.497655</td>
<td>0.908918</td>
</tr>
<tr>
<td>L. Amygdala (211)</td>
<td>2, 178</td>
<td>5.5692</td>
<td>0.004508</td>
<td>0.903618</td>
<td>0.251414</td>
</tr>
<tr>
<td>L. Amygdala (213)</td>
<td>2, 178</td>
<td>6.6238</td>
<td>0.001680</td>
<td>0.063394</td>
<td>0.494163</td>
</tr>
<tr>
<td>L. Basal ganglia (229)</td>
<td>2, 178</td>
<td>3.8229</td>
<td>0.023681</td>
<td>0.065501</td>
<td>0.620435</td>
</tr>
</tbody>
</table>

**Table 2** Identified regions with significant interaction effects of age-by-status

**Table 1** Demographic data of children, adolescents, and adults in the ADHD and TD groups

<table>
<thead>
<tr>
<th>Site</th>
<th>ADHD score</th>
<th>Age (year)</th>
<th>Sex (n, M/F)</th>
<th>ADHS score</th>
<th>Age (year)</th>
<th>Sex (n, M/F)</th>
</tr>
</thead>
<tbody>
<tr>
<td>KKI (n)</td>
<td>23/8</td>
<td>72.97±10.04</td>
<td>72.97±10.04</td>
<td>19/12</td>
<td>24.94±1.41</td>
<td>21/10</td>
</tr>
<tr>
<td>NYU (n)</td>
<td>21</td>
<td>19/12</td>
<td>24.94±1.41</td>
<td>21/10</td>
<td>24.94±1.41</td>
<td>21/10</td>
</tr>
<tr>
<td>HCP (n)</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

**Table 2** Identified regions with significant interaction effects of age-by-status

**Table 1** Demographic data of children, adolescents, and adults in the ADHD and TD groups

<table>
<thead>
<tr>
<th>Brain region (index number of the ROI)</th>
<th>DOF</th>
<th>F value</th>
<th>P value (age × disease)</th>
<th>P value (age)</th>
<th>P value (disease)</th>
</tr>
</thead>
<tbody>
<tr>
<td>L. Superior frontal gyrus (9)</td>
<td>2, 178</td>
<td>3.0891</td>
<td>0.047991</td>
<td>0.787615</td>
<td>0.198468</td>
</tr>
<tr>
<td>L. Inferior frontal gyrus (29)</td>
<td>2, 178</td>
<td>4.0198</td>
<td>0.019611</td>
<td>0.811140</td>
<td>0.246761</td>
</tr>
<tr>
<td>L. Inferior frontal gyrus (33)</td>
<td>2, 178</td>
<td>4.1594</td>
<td>0.017161</td>
<td>0.675609</td>
<td>0.253190</td>
</tr>
<tr>
<td>R. Inferior frontal gyrus (36)</td>
<td>2, 178</td>
<td>5.5417</td>
<td>0.006426</td>
<td>0.828528</td>
<td>0.134809</td>
</tr>
<tr>
<td>L. Inferior frontal gyrus (39)</td>
<td>2, 178</td>
<td>3.1641</td>
<td>0.046438</td>
<td>0.872547</td>
<td>0.187522</td>
</tr>
<tr>
<td>R. Precentral gyrus (64)</td>
<td>2, 178</td>
<td>3.6681</td>
<td>0.027475</td>
<td>0.728483</td>
<td>0.734674</td>
</tr>
<tr>
<td>L. Superior temporal gyrus (77)</td>
<td>2, 178</td>
<td>3.4586</td>
<td>0.033605</td>
<td>0.834183</td>
<td>0.360218</td>
</tr>
<tr>
<td>L. Middle temporal gyrus (83)</td>
<td>2, 178</td>
<td>7.2908</td>
<td>0.000905</td>
<td>0.645098</td>
<td>0.061416</td>
</tr>
<tr>
<td>L. Middle temporal gyrus (85)</td>
<td>2, 178</td>
<td>3.5759</td>
<td>0.030019</td>
<td>0.887161</td>
<td>0.542660</td>
</tr>
<tr>
<td>L. Middle temporal gyrus (87)</td>
<td>2, 178</td>
<td>3.271</td>
<td>0.040261</td>
<td>0.977927</td>
<td>0.617160</td>
</tr>
<tr>
<td>R. Postcentral gyrus (156)</td>
<td>2, 178</td>
<td>3.457</td>
<td>0.033657</td>
<td>0.712092</td>
<td>0.820640</td>
</tr>
<tr>
<td>L. Insular gyrus (165)</td>
<td>2, 178</td>
<td>3.6151</td>
<td>0.028909</td>
<td>0.362170</td>
<td>0.551337</td>
</tr>
<tr>
<td>L. Insular gyrus (171)</td>
<td>2, 178</td>
<td>3.1477</td>
<td>0.045352</td>
<td>0.941490</td>
<td>0.891245</td>
</tr>
<tr>
<td>L. Medioventral occipital cortex (191)</td>
<td>2, 178</td>
<td>4.089</td>
<td>0.018355</td>
<td>0.727416</td>
<td>0.968738</td>
</tr>
<tr>
<td>R. Medioventral occipital cortex (192)</td>
<td>2, 178</td>
<td>4.1922</td>
<td>0.016632</td>
<td>0.497655</td>
<td>0.908918</td>
</tr>
<tr>
<td>L. Amygdala (211)</td>
<td>2, 178</td>
<td>5.5692</td>
<td>0.004508</td>
<td>0.903618</td>
<td>0.251414</td>
</tr>
<tr>
<td>L. Amygdala (213)</td>
<td>2, 178</td>
<td>6.6238</td>
<td>0.001680</td>
<td>0.063394</td>
<td>0.494163</td>
</tr>
<tr>
<td>L. Basal ganglia (229)</td>
<td>2, 178</td>
<td>3.8229</td>
<td>0.023681</td>
<td>0.065501</td>
<td>0.620435</td>
</tr>
</tbody>
</table>
(ROIs) specified by the Brainnetome Atlas. The Brainnetome Atlas is a structural atlas that consists of 246 regions (Fan et al., 2016). Connectivity information was assessed with a graph structure using nodes and edges. The nodes were 246 ROIs derived from the Brainnetome Atlas. Pearson correlation values of the time series between two nodes were used as edges. The edge values were filled into the matrix as elements and the matrix was referred to as the correlation matrix. We adopted the weighted and un-directional network model. Soft thresholding was used to prevent binarization of the correlation matrix using the following equation (1).

\[ w_{ij} = \left( \frac{r_{ij} + 1}{2} \right)^{\beta} \]  

The \( r_{ij} \) term denotes the edge value between the node \( i \) and \( j \) (Mumford et al., 2010; Schwarz and McGonigle, 2011). The \( \beta \) value was set to 6 to ensure scale-free topology (Mumford et al., 2010).

Connectivity measures
We used degree centrality (DC) to assess the regional connectivity of brain networks (Lohmann et al., 2010; Fransson et al., 2011). The DC value for a node \( i \) is defined as the number of links connected directly to the node (Rubinov and Sporns, 2010). We used MATLAB (version 2016; Mathworks Inc., Natic, MA, USA) to compute the DC values (The Mathworks Inc., 2016).

Multi-site effect
Since our neuroimaging data was acquired from different sites, we adopted a dummy coding regression model to remove multi-site effects from the DC values using the following equation (2).

\[ Degree \text{ centrality} = \sum_{i=1}^{3} \beta_{site_i} x_{site_i} + \varepsilon \]  

The \( X_{site_i} \) term is the dummy vector denoting different data collection sites, \( \beta_{site} \) is the regression coefficient of the \( i^{th} \) dummy vector, and \( \varepsilon \) is the residual DC value (Hardy, 1993). The regression model removed the multi-site effects and the residuals were used for further analyses.

Statistical analysis
We used MATLAB for statistical analysis (version 2016; Mathworks Inc.). The two-way analysis of variance (ANOVA) was used to explore differences in age-related DC patterns between the ADHD and TD groups (Fujikoshi, 1993). The DC values were set as the dependent variables, and disease status (ADHD or TD) and age group (child, adolescent, or adult) were set as the independent variables. The significance of the interaction effects of disease and age group was quantified using \( P \) values (\( P < 0.001 \)). We adopted an uncorrected \( P \) value of 0.001 due to the exploratory nature of our study. We applied a stringent \( P \) value threshold of 0.001 compared to the conventional 0.05 since our study was an exploratory study investigating 246 regions covering the whole brain. We also reported results using a relaxed \( P \) value of 0.05. Chi-square tests were applied to assess differences in sex among comparison groups.

Results
Motion scrubbing
We calculated the FD for each volume from the rs-fMRI data. Two children from the TD group had part of the frames scrubbed. Figure 1 shows the FD of these two subjects. We removed 13 frames from one child in the TD group and 5 frames from the other child in the TD group.

Connectivity differences
We performed a two-way ANOVA to determine the brain regions that showed significant interaction effects of disease status and age. The left superior frontal gyrus, left inferior frontal gyrus, right inferior frontal gyrus, right precentral gyrus, left superior temporal gyrus, left middle temporal gyrus, right postcentral gyrus, left insular gyrus, left medio-ventral occipital cortex, right medioventral occipital cortex, left amygdala, and left basal ganglia showed significant interaction effects (\( P < 0.05 \) (Table 2). Figure 2a shows the locations and their \( P \) values of the identified regions. Among the identified regions, the left middle temporal gyrus showed the most significant interaction effects (\( P < 0.001 \); Figure 2b). Further post-hoc tests were not conducted because there were no significant main effects of disease status and age.

Age-related patterns
We show regions that have significant age-by-status interaction for each age group in Figure 3.

Discussion
The main purpose of this study was to determine if there were age-related network differences between ADHD patients and TD subjects. We divided the subjects into six groups based on disease status (i.e., ADHD or TD) and age (i.e., children, adolescents, and adults) to form comparison groups. From the two-way ANOVA results, we found significant interaction effects of disease status and age. Since this was an exploratory study investigating hundreds of brain regions, we relaxed the constraint of the \( P \) value and found significant interaction effects of disease status and age based on functional connectivity.

Among the identified regions, the left inferior frontal gyrus, the left middle temporal gyrus, which showed the most significant interaction effects, and the left insular gyrus were known to be related to cognitive function (Vandenberghe et al., 1996; Goel and Dolan, 2001; Swick et al., 2008; Fan et al., 2016). Swick et al. (2008) reported that subjects with damage in the left inferior frontal gyrus and left insula region had higher error rates than controls in a response inhibition task. The response inhibition task is known as a major task that can discriminate between ADHD and TD subjects (Nigg, 1999; Epstein et al., 2001; Tamm et al., 2004). Tamm et al. (2004) also found significant differences in brain activation in the middle temporal gyrus between subjects with ADHD and TD subjects in a behavioral response inhibition task. Figure 4B shows the locations of these regions. There is a noticeable overlap between the region previously reported.
in the literature and the region we found as shown in Figure 4A. The cognitive system plays an important role in typical development from childhood through adolescence to adulthood (Blakemore and Choudhury, 2006). Thus, aberrant development in the cognitive brain regions that we identified between ADHD and TD groups implies that impairment in cognitive function might be associated with age-related brain network changes in ADHD patients.

In this study, we used multi-center neuroimaging data to obtain a sufficient number of samples. Although the differences in imaging parameters were relatively small, this could lead to different amounts of noise and distortions in the data, making a fair comparison difficult. The high-resolution data from the HCP were resampled and pre-processed to low-resolution ADHD-200 data so that data can be fairly compared. We applied the common image processing steps performed on the low quality (i.e., low resolution ADHD-200) spatial reference space so that high quality (i.e., high resolution HCP) data was effectively rendered to low quality data. Such approaches have been successfully applied in other studies (Fennema-Notestine et al., 2007; Di Martino et al., 2014; Bellec et al., 2017). We visually confirmed similar image qualities by computing the average of T1-weighted structural data for each subgroup as shown in Figure 5 and they all appeared similar in the low resolution common space. Furthermore, we used the correlation of rs-fMRI time series between two different brain regions as the main feature in this study. Each region contained over hundreds of voxels, hence the regional average time series might reduce the potential differences in image quality. Finally, we also performed a multi-site regression using the dummy-coding to remove the potential multi-site effects from the centrality measurement. The dummy-coding regression approach has been applied in other studies comparing data from different sites (Hardy, 1993; Sanfilipo et al., 2004).

We used Brainnetome atlas to specify the ROIs for child, adolescent, and adult groups. The Brainnetome atlas was derived from adults and thus application to adults is natural. We investigated if a single atlas could specify the ROIs for various age groups. We registered T1 anatomical images onto a common space for each age group and then compared averaged images with one another. The average images for each age group appeared quite similar and those

Figure 1 The plot of the FD values for two children from TD group whose frames were partly censored. (A) The FD of one subject (13 frames removed). (B) The FD of another subject (5 frames removed). The lower sub-figures are the results after removing the frames. FD: Frame-wise displacement; TD: typically developing.

Figure 2 Brain regions that showed a significant age-by-status interaction by the two-way analysis of variance (ANOVA) test using degree centrality (DC). The $P$ values from analysis of variance (ANOVA) are visualized using the color bar (from red to yellow) in the middle of the plot. (A) The left superior frontal gyrus, left inferior frontal gyrus, right inferior frontal gyrus, right precentral gyrus, left superior temporal gyrus, left middle temporal gyrus, right postcentral gyrus, left insular gyrus, left medioventral occipital cortex, right medioventral occipital cortex, left amygdala, and left basal ganglia showed a significant effect of interaction ($P < 0.05$). (B) The left middle temporal gyrus region showed a significant effect of interaction ($P < 0.001$). The ADHD and TD groups showed different patterns based on age as inferred by the two-way ANOVA. ADHD: Attention deficit and hyperactivity disorder; TD: typically developing; L: left; R: right.
Figure 3 Age-related degree centrality patterns of the identified regions.
The mean and standard error of degree centrality for brain regions that showed significant age-by-status interactions for each age group from Table 2 ($P < 0.05$). The green plot denotes the mean degree centrality values of the TD groups and the blue plot denotes the mean degree centrality values of the ADHD groups. L.: Left; R.: right; TD: typically developing; ADHD: attention deficit and hyperactivity disorder.

Figure 4 Comparison between identified regions and known regions related to cognitive functions.
(A) Regions (orange mask) that showed a significant age-by-status interaction by the two-way analysis of variance (ANOVA). (B) Regions (blue mask) related to cognitive function from previous studies (Van den Bergh et al., 1996; Goel and Dolan, 2001; Swick et al., 2008; Fan et al., 2016). L: Left; R: right.

Figure 5 The average T1 structural images of each subgroup after we applied the common anatomical preprocessing steps.
Results of two axial slices are shown. The first and third rows show average T1 images and the second and fourth rows show overlaid region of interest (ROI) information from the Brainnetome atlas. All six subgroups showed visually similar average T1 images and the overlaid ROIs matched well with known structural boundaries. ADHD: attention deficit and hyperactivity disorder; TD: typically developing.

for each group were compared with overlaid ROIs from the atlas and they seemed reasonable as well.

Our study adopted an uncorrected $P$ value of 0.001 for statistical significance. We had limited samples but explored hundreds of regions and thus we chose to adopt an uncorrected $P$ value. Use of an uncorrected $P$ value is rather common in many exploratory studies involving the whole brain (Konishi et al., 1999; Anand et al., 2005).

Our study has some limitations. First, rs-fMRI was the only modality used. Using multi-modal imaging data might provide complementary information that could better describe the differences between the ADHD and TD groups. Second, the sample size might be insufficient due to the limitation of available cases from the online databases. A future study performed on a larger cohort is necessary to confirm our findings with higher statistical power. Third, we could not compute the correlation between DC and ADHD scores, because we used two types of ADHD related scores coming from two research databases. Finally, we did not consider longitudinal data for our study, as we are not aware of any openly accessible research database housing longitudinal ADHD neuroimaging data. Thus, we performed our study
Data were provided by the Neuro Bureau, the ADHD 200 consortium, and Virginia Tech’s ARC. Data were also provided by the Human Connectome Project, WU-Minn Consortium (Principal Investigators: David Van Essen and Kamil Ugurbil; U54MH091657) funded by the 16 NIH Institutes and Centers that support the NIH Blueprint for Neuroscience Research and by the McDonnell Center for Systems Neuroscience at Washington University.

Author contributions: JH and HP designed the study and collected, analyzed, and interpreted the data. BYP reviewed the paper. HHIC contributed to the discussion and edited the paper. HP is the guarantor of this work and, as such, had full access to all the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis. All authors approved the final version of this paper.

Conflicts of interests: None declared.

Research ethics: The Institutional Review Board (IRB) of Sungkyunkwan University approved our study (#2015-09-007). Our study was performed in full accordance with local IRB guidelines and the principles and the Declaration of Helsinki, and informed consent was obtained from all subjects.

Declaration of participant consent: The authors certify that they have obtained all appropriate consent forms of participant or their guardians. In the form, participants or guardians have given their consent for participants’ images and other clinical information to be reported in the journal. Participants or guardians understand that participants’ names and initials will not be published and due efforts will be made to conceal their identity, but anonymity cannot be guaranteed.

Data sharing statement: Datasets analyzed during the current study are available from the corresponding author on reasonable request. Plagiarism check: Checked twice by iThenticate.

Peer review: Externally peer reviewed.

Open access statement: This is an open access article distributed under the terms of the Creative Commons Attribution-NonCommercial-Share-Alike 3.0 License, which allows others to remix, tweak, and build upon the work non-commercially, as long as the author is credited and the new creations are licensed under identical terms.

Open peer reviewer: Hao Chen, Shanghai 6th Peoples Hospital Affiliated to Shanghai Jiao Tong University School of Medicine, China.

Additional file: Open peer review report 1.

References


Copied by Li CH, Song LP, Zhuo M
Open peer review report 1 on “Neurological and neuropsychological consequences of electrical and lightning shock: review and theories of causation”.

Reviewer: Gentian Vyshka, University of Medicine in Tirana, Albania

Comments to the authors: This manuscript used rs-fMRI to explore age-related brain network differences and indicate that aberrant development in cognitive brain regions might be associated with age-related brain network changes in ADHD patients. It indeed has a certain novelty and is a good work in neuroimaging field, but more or less doesn't conform to the scope of the NRR. Anyway, it still needs some revisions as follows:

1. Page 4, line 21, "DSM-IV" is an abbreviation, please provide its full name. And provide the cited reference of "Conner's Parent Rating Scale Revised".
2. The imaging parameters used in KKI and NYU, and HCP database were inconsistent. So would it affect the reliability of the results? If so, how to avoid or reduce this difference?
3. There are two charts in Fig. 1 a or b, which one indicates the result after removal of the frames? Please mark it.
4. Both Fig. 2 description in the results section and its caption missed the direction "Left".
5. The left or right direction should be marked in Fig. 2 and 3.
6. The first paragraph in discussion section is still a part of results, so it should be moved in this section.
7. This paper lacked further discussion. It would be better if the author can propose a possible mechanism to explain their findings.