Cognitive Shifting in Children with Attention-Deficit Hyperactivity Disorder: A Near Infrared Spectroscopy Study

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Introduction

Attention-deficit hyperactivity disorder (ADHD) is a developmental disease characterized by inattention, hyperactivity, and impulsivity [1]. Several studies have shown that core ADHD symptoms are related to deficits in executive function [2,3]. Executive function is an assemblage of high-level cognitive domains, including inhibition, working memory, planning, fluency, and shifting, that facilitate the inhibition of incorrect behaviors and the selection of appropriate behaviors according to context and goals [4]. Cognitive shifting is the mental ability to switch between thinking about one concept and begin thinking of another; it is often measured using the Wisconsin Card Sorting Test [5-8]. Kado et al. [6] previously demonstrated that children with ADHD make many more Nelson-type preservative errors (PENs) than do typically developing children (TDC). PENs are persistently repeated responses in accordance with a patient’s initial preference or card sorting pattern that would have been correct during the immediately preceding stage or during the first stage of the task. During the WCST, participants need to change their responses according to shape, color, or number following feedback from the experimenter. However, using the WCST as a specific measure of cognitive shifting is problematic because this task requires additional cognitive processes, thus making it hard to elicit a specific shifting effect. Conversely, the Dimensional Change Card Sort (DCCS) task is specific to cognitive shifting and entails instructions that are easily understood by individuals from a wide range of ages and intellectual abilities [9]. In a previous study using DCCS, 3-year-old children tended to respond to a new stimulus in the same manner as an old stimulus, despite changes in rules or goals. This lack of behavioral flexibility has also been observed among older children and adults with damage to the prefrontal cortex. In contrast, typically developing 5-year-old children alter their responses according to rule changes [10]. Therefore, the DCCS task can measure developmental changes in executive function. A recent study also demonstrated that children with ADHD experienced difficulty in a modified DCCS task with more frequent rule switches [11].

In addition to deficits in the modified DCCS task, some ADHD patients display weaker prefrontal cortex (PFC) activation during a stop-signal task [12]. Recent neuroanatomical research suggests that 8- to 12-year-old children with ADHD exhibit a marked delay in the...
maturity of prefrontal areas [13]. These studies suggest that patients with ADHD may have functional and anatomical deficits in the prefrontal cortex. However, there are few brain-imaging studies of young children with ADHD. Simultaneous neurobehavioral testing using DCCS and neuroimaging may help determine why older children with ADHD exhibit the same deficits in cognitive shifting observed in very young TDC. In the present study, we evaluated cognitive shifting in children with ADHD during the DCCS task while simultaneously measuring patterns of PFC activity using near-infrared spectroscopy (NIRS)-a non-invasive measure of cerebral blood oxygenation that does not require children to be in a fixed position. Therefore, this method can be used to assess young, hyperactive children [15,16]. We hypothesized that children with ADHD would make more errors during a cognitive shifting task than would TDC due to the impulsivity associated with ADHD. However, we posited that children with ADHD would demonstrate similar reaction times on the task as compared to TDC.

Method

Participants

The ADHD group (Table 1) consisted of 22 participants (mean age: 10 years 3 months, standard deviation: 2 years 0 months, 15 boys and 7 girls, all but three right-handed) who were diagnosed by 2 pediatric neurologists according to DSM-IV-TR criteria [1]. The IQs of the ADHD group were evaluated using the third edition of the Japanese version of the Wechsler Intelligence Scale for Children [17]. No individual had an IQ score lower than 80. A total of 37 TDC (mean age: 10 years 10 months, standard deviation: 1 years 8 months, 19 boys and 18 girls, all but four right-handed) were recruited as controls. All participants had normal or corrected-to-normal vision and no history of neurological disorders. All participants from both groups and their mothers provided written informed consent before the experiment, the protocol for which was approved by the Ethics Committee of the National Center of Neurology and Psychiatry (Japan). We gathered the clinical sample from the National Center of Neurology and Psychiatry (Japan) and the control sample from an after-school care program at Sukage Nursery.

All participants completed Raven’s Colored Progressive Matrices (RCPM) [18] to determine non-verbal intelligence. In addition, we measured sentence comprehension using the Kaufman Assessment Battery for Children (K-ABC). The Swanson, Nolan, and Pelham Scale version IV (SNAP-IV) [19-21] was completed by the mother of each participant in the TDC and ADHD groups in order to verify both the current severity of ADHD symptoms and the absence of symptoms in the control group. The SNAP-IV test assessed three cognitive/behavioral deficits: the inattention domain, the hyperactivity/impulsivity domain, and the oppositional defiant disorder (ODD) domain. The Pervasive Developmental Disorders Autism Society Japan Rating Scale [22] was also administered by interviewing the mother of each participant in the TDC and ADHD groups to verify ASD severity since ADHD and ASD are often combined. The PARS test included two question categories: questions about clinical conditions during infancy (PARS-Infant) and questions about current clinical conditions (PARS-Present). There were no significant differences between groups in mean age, K-ABC sentence comprehension score, or RCPM nonverbal intelligence score (age: t(57)=1.13, n.s.; reading comprehension: t(57)=1.61, n.s.; RCPM: t(57)=0.96, n.s.; see Table 1). However, significant differences were observed between groups on the three SNAP-IV subscores (inattention: t(57)=7.01, p<0.001; hyperactivity/impulsivity: t(57)=5.76, p<0.001; ODD: t(57)=3.69, p<0.01; Table 1). Moreover, significant differences were observed between groups on the two PARS subscores (Infant: t(57)=6.72, p<0.001; Present: t(57)=7.20, p<0.001; Table 1).

<table>
<thead>
<tr>
<th></th>
<th>TDC (n = 37)</th>
<th>ADHD (n = 22)</th>
<th>t value (df = 57)</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>10.10 (1.8)</td>
<td>10.3 (2.0)</td>
<td>1.13</td>
<td>.263</td>
</tr>
<tr>
<td>Reading comprehension (K-ABC) a</td>
<td>19.84 (3.81)</td>
<td>17.91 (5.38)</td>
<td>1.61</td>
<td>.113</td>
</tr>
<tr>
<td>Non-verbal intelligence (RCPM) b</td>
<td>29.54 (3.86)</td>
<td>30.50 (3.50)</td>
<td>.96</td>
<td>.343</td>
</tr>
<tr>
<td>Inattention (SNAP) c</td>
<td>5.81 (4.36)</td>
<td>15.73*** (6.51)</td>
<td>7.01</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Hyperactivity/ impulsivity (SNAP) d</td>
<td>2.05 (2.38)</td>
<td>8.64*** (6.26)</td>
<td>5.76</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>ODD (SNAP) e</td>
<td>3.16 (3.74)</td>
<td>8.36** (7.11)</td>
<td>3.69</td>
<td>&lt;.01</td>
</tr>
<tr>
<td>PARS (infant) f</td>
<td>1.32 (1.68)</td>
<td>7.18** (4.86)</td>
<td>6.72</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>PARS (present) g</td>
<td>1.51 (2.22)</td>
<td>8.36*** (5.05)</td>
<td>7.20</td>
<td>&lt;.001</td>
</tr>
</tbody>
</table>

Table 1: Characteristics of typically developing children (TDC) and children with attention-deficit hyperactivity disorder (ADHD).

Differences were assessed using Student’s t-tests (two-tailed)

4raw score on reading comprehension as measured by the Kaufman Assessment Battery for Children (K-ABC)

5raw score on Raven’s Colored Progressive Matrices test (RCPM) for non-verbal intelligence

6raw score on the Swanson, Nolan, and Pelham Scale (SNAP-IV) questionnaire items on inattention

7raw score on SNAP-IV questionnaire items on hyperactivity/impulsivity

8raw score on SNAP-IV questionnaire items on oppositional defiant disorder

9raw score on infant items of the Pervasive Developmental Disorders Autism Society Japan Rating Scale (PARS)

10raw score on present items of the PARS
Behavioral task

Each participant performed the DCCS task on a computer. Participants sat 50 cm away from a 15-inch liquid crystal display (LCD) screen with a gray background and performed a series of baseline trials and DCCS task trials.

Baseline task

A white square was shown on the lower right or lower left side of the 15-inch LCD screen, and participants indicated which side the square appeared by pressing the corresponding arrow button (Figure 1A). The next square appeared randomly 1 s after the participants pressed the button. A 15-s rest period was allowed before beginning the task; during this time, a small white circle was shown in the center of the screen, and participants were instructed to watch it closely. Task duration was 30 s.

DCCS task

We used the “border version” of the DCCS task to evaluate cognitive shifting among participants. The task consists of participants having to frequently alternate their responses according to the color or shape of the stimulus in line with rule changes [9]. Three cards were displayed in the form of a pyramid on the same screen (Figure 1 B). On each white card, a diamond or a star shape was overlaid in red or blue. Participants were required to select which of the two cards at the bottom of the pyramid (selection cards) matched the card at the top of the pyramid (reference card) according to the given rule (shape or color). The color and shape of the two selection cards differed from those of the reference card. If the reference card was hemmed in black line, the correct response was to select the selection card that matched the reference card in shape. When no black hemming was shown, then the correct response was to select the selection card that matched the reference card in color. The reference and selection cards kept appearing on the screen until the participant responded, and the next set of cards appeared at random, 0.5 s after participants made their selection.

Participants were allowed a 15-s rest period before each successive trial. During the rest period, participants stared at a small white circle on the monitor screen, as in the baseline task. Task duration was 30 s.

We performed the DCCS and baseline tasks twice, in alternation (i.e., baseline task i. DCCS task ii. baseline task as DCCS task). In addition, before the actual task, participants underwent a practice trial (5-s rest period followed by a 20-s trial). We provided feedback to participants during the practice trial and confirmed that participants completely understood the instructions. During the actual task, feedback was not provided, and participants’ responses and reaction times were recorded automatically.

Behavioral Data Analysis

The number of correct answers and errors as well as the reaction times for correct responses and percentage of correct answers were assessed for each participant.

A significance level of p<0.05 (two-sided) was adopted for all analyses.

NIRS Recording and Analysis

While participants performed the DCCS task, we recorded activity in the lateral PFC by noting changes in oxygenated hemoglobin (ΔHbO2) using a multi-channel NIRS system (OEG-16; Spectratech Inc., Tokyo, Japan). Near-infrared laser diodes emitted near-infrared light (approximately 770 and 840 nm); re-emitted light was detected by avalanche photodiodes located 30 mm from the emitters with a temporal resolution of 655 ms at a measurement depth of approximately 3 cm below the scalp. In our system, six emitters and six detectors were placed at alternate points on a 2 × 6 grid (Figure. 2). This configuration enabled us to detect signals from 16 differential channels spread over the bilateral frontal region. The center of the probe matrix was placed on Fpz (International 10-10 system), and the bottom left and bottom right corners were located around F7 and F8, respectively, in line with a previous report [23]. Detected NIRS signals were sent to a separate data collection computer. Presentation time of baseline symbols or DCCS cards was transmitted to the data collection computer from the task-control computer through a local area network with a UDP communication. A band-pass filter was set at 0.01-0.1 Hz during the task, using fast Fourier transformation to reject NIRS data distorted by minor movement artifacts. Data from each channel were converted into z-scores because raw data of highly variable amplitudes cannot be compared across participants and channels [15]. Z-scores were calculated using the mean and standard deviation of ΔHbO2 during the last 6 s of the orienting period (when participants stared at a white dot at the center of the computer screen). The mean and standard deviation were adjusted to a z-score of 0 and 1 for every channel. Trials with scores more than two standard deviations above or below the mean were excluded because of the possibility of motion artifacts.
Figure 2: Near-infrared spectroscopy (NIRS) probes were attached to the scalp over the prefrontal cortex in the configuration shown here. The center of the probe matrix was placed on Fpz (International 10-10 system), and the bottom left and bottom right corners were located around F7 and F8, respectively.

Figure 3: Behavioral results of the Dimensional Change Card Sort task. The percentage of correct answers for each group during the task is plotted. Error bars indicate the standard error. *p<0.05, **p<0.01

Figure 4: Behavioral results of the Dimensional Change Card Sort task. The number of errors for each group during the task is plotted. Error bars indicate the standard error. *p<0.05, **p<0.01

Figure 5: Behavioral results of the Dimensional Change Card Sort task. Reaction times for each group during the task are plotted. Error bars indicate the standard error. *p<0.05, **p<0.01

Results

Behavioral results

The percentage of correct answers ($t(57) = 2.93, p<0.01$, Figure 3) and the number of errors ($t(57) = 2.80, p<0.01$, Figure 4) on the DCCS differed significantly between the ADHD and TDC groups; although, there were no differences in mean reaction times for correct responses ($t(57) = 1.97, p> .05$, Figure 5).
NIRS results

During the baseline task, differences in the HbO2 signal were not observed between the ADHD and TDC groups. During the DCCS task, the HbO2 signal at channels 1, 2, and 13 were significantly lower in the ADHD group than in the TDC group (#1: t(57) = 3.53, p = 0.001, #2: t(57) = 2.46, p = 0.017, #13: t(57) = 2.22, p =0.03, Figure 6).

Correlations

Negative correlations were observed between the HbO2 signal at channel #1 and both PARS scales (Infant: r=-0.30, p<0.05; Present: r=0-.29, p<0.05; Figures 7 and 8, respectively). Moreover, negative correlations were observed between the HbO2 signal at channel #1 and two of the SNAP subscales (inattention: r=-0.39, p<0.01; hyperactivity/impulsivity: r=0-.34, p<0.01; Figures 9 and 10, respectively). Positive correlations were observed between one of the SNAP subscales and number of errors (inattention: r=0.27, p<0.05; Figure 11).

Discussion

Group differences in cognitive shifting

During the DCCS task, children with ADHD made significantly more incorrect responses than did age- and IQ-matched normally developing children. Reaction times for all responses did not differ significantly between groups. These results indicate a deficit in cognitive shifting among children diagnosed with ADHD, consistent with previous studies [6]. The DCCS task requires participants to alter responses according to rule changes (color matching or shape matching). It is believed that higher numbers of incorrect responses reflect the inattention and hyperactivity characteristic of ADHD. In contrast, children with ADHD understood the rules of the task because their accuracy rate exceeded 75%. In other words, the ADHD children understood the rules of the task but exhibited impairments in cognitive shifting. Compared with the TDC group, children with ADHD required more time to shift to the new rule, resulting in a greater number of incorrect responses. We believe that the DCCS task is more suitable to the evaluation of cognitive shifting than numerous other measures because its simplicity (i.e., only two classifications) minimizes the load on working memory. This contrasts with other tests of executive function, such as the WCST, which uses three classifications (color, shape, and number) [24]. In addition, the language impairment observed in some instances of ADHD should not affect the DCCS task because of its simple instructions and rules [25].
Neural findings

Using NIRS, we found that cortical activity over channels 1, 2, and 13 was significantly lower in the ADHD group than in the TDC group during the DCCS task but not during the baseline task. The area surrounding regions #1 and #2 is the right inferior frontal gyrus (rIFG). This region is thought to play a significant role in inhibition [26-28]. Previous studies using an inhibition task (e.g., Stroop, Go/No-go) have indicated that lack of inhibition is a core symptom of ADHD. In the present study, the larger number of errors made by children with ADHD may reflect problems with both inhibition and cognitive shifting. In addition, previous studies have suggested that children with ASD exhibit lower brain activation during cognitive shifting tasks [14, 29]. Therefore, both ADHD and ASD seem to be associated with lower brain activation during this task. In order to clarify differences in the deficit, there is a need for further study of brain connectivity. For instance, the rIFG shows lower activation among three-year-old children who were unable to complete the DCCS task; this result was not found among five-year-old children [10]. Thus, because improvements in rIFG activity may not occur during natural development among individuals with ADHD, new methods for alleviating ADHD symptoms that target other brain areas are required.
Correlations

We observed correlations between ASD symptoms and brain activity among children with ADHD. These correlations are perhaps explained by the fact that developmental disorders make up a spectrum, whereby ASD and ADHD symptoms fall at different points along the same continuum. These results may also be associated with the persistence of the DCCS task [14]. Furthermore, the correlations among ASD symptoms, ADHD, and brain activity may be associated with inhibition problems present among children with ADHD [28].

Advantages and Limitations

To measure activity of the prefrontal cortex during cognitive shifting, we measured ΔHbO2 using NIRS, a neuroimaging technique employed in several previous studies to assess frontal lobe function among children and adults during the WCST. The NIRS technique has several advantages over functional magnetic resonance imaging (fMRI), most notably the lower cost and sensitivity to motion artifacts [8,30-32]. We note, however, that the NIRS detects hemodynamic changes only at the surface of the brain (approximately 3 cm beneath the skull). Thus, some potential differences in subcortical responses cannot be assessed using NIRS. Moreover, NIRS has a relatively low spatial resolution compared with fMRI. Despite these shortcomings, the NIRS is becoming a key imaging modality in developmental neuroscience studies involving newborns [33], preschool children [32], and school-aged children [15] because of its excellent safety and relative insensitivity to body movements. We believe that we can adequately measure neural activity patterns associated with executive function tasks using NIRS in children who have yet to develop full language capacity.

Conclusions

Deficits in cognitive shifting were observed among children with ADHD during the DCCS task. Children with ADHD exhibited more errors, as well as lower prefrontal cortex activity as compared to TDC.

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Contributions

All results were obtained through discussion among the authors. Every author reviewed the manuscript.

Conflict of Interest

The authors declare no conflicts of interest.

References


