Hypofrontality in Attention Deficit Hyperactivity Disorder During Higher-Order Motor Control: A Study With Functional MRI

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Objective: Functional magnetic resonance imaging (MRI) was used to investigate the hypothesis that attention deficit hyperactivity disorder (ADHD) is associated with a dysfunction of prefrontal brain regions during motor response inhibition and motor timing. Method: Generic brain activation of seven adolescent boys with ADHD was compared to that of nine comparison subjects equivalent in sex, age, and IQ while they were performing a stop task, requiring inhibition of a planned motor response, and a motor timing task, requiring timing of a motor response to a sensory cue. Results: The hyperactive adolescents showed lower power of response in the right mesial prefrontal cortex during both tasks and in the right inferior prefrontal cortex and left caudate during the stop task. Conclusions: ADHD is associated with subnormal activation of the prefrontal systems responsible for higher-order motor control. Functional MRI is a feasible technique for investigation of neural correlates of ADHD.

pects of higher-order motor control: a “stop” task (21), requiring inhibition of a planned response, and a “delay” task (22), requiring synchronization of a motor response to an intermittently appearing visual stimulus. In a group of normal adults we demonstrated activation by the stop task of a predominantly right hemisphere network, comprising the mesial and inferior frontal cortex and caudate nucleus (21). Performance of the delay task by the same group of subjects elicited dorsolateral prefrontal, mesial frontal, and parietal activation (22).

In the current study we applied these two tests to a controlled investigation of higher-order motor control in adolescent patients with a diagnosis of hyperactivity disorder. In light of our prior findings, we expected to find less activity in the frontal brain regions of the seven hyperactive subjects studied, during both motor response inhibition and motor timing, than in nine healthy comparison subjects. Adolescents, rather than children, with current evidence of hyperactive behavior were chosen in order to reduce the possibility of movement artifacts. The neuropsychological deficits underlying ADHD, including performance on inhibition tasks, have been shown to persist into adolescence (23); we therefore considered that any findings were likely to generalize to the problems of younger children. A subsidiary, but necessary, hypothesis was that functional MRI could reliably be applied to the rather difficult subject group of young people with markedly hyperactive behavior.

METHOD

Subjects

Seven clinically referred right-handed (24) male adolescents aged 12 to 18 years (mean=15.71 years) who met the DSM-IV criteria for ADHD participated in the study. The hyperactive adolescents scored above the conventional thresholds for hyperactivity on the activity scale of a standardized parent interview, the Parental Account of Child Symptoms (25) (mean score=1.79), and on the Connors teacher rating scale (26). The exclusion criteria were comorbidity with any other psychiatric disorder, such as learning or speech disorder, with the exception of conduct disorder, which can be seen as a complication of ADHD (27). Patients with neurological disease were also excluded. The patients were either unmedicated or medicated free for 1 week before scanning. The comparison subjects were nine healthy comparison subjects. Adolescents, rather than children, with current evidence of hyperactive behavior were chosen in order to reduce the possibility of movement artifacts. The neuropsychological deficits underlying ADHD, including performance on inhibition tasks, have been shown to persist into adolescence (23); we therefore considered that any findings were likely to generalize to the problems of younger children. A subsidiary, but necessary, hypothesis was that functional MRI could reliably be applied to the rather difficult subject group of young people with markedly hyperactive behavior.

Experimental Design

Each experimental task consisted of two main conditions (activation and control condition). Each condition lasted 30 seconds and was preceded by a short visual cue (lasting 3 seconds), warning the subject that one of the conditions was about to begin. The total epoch length (cue plus task) was therefore 33 seconds. The control and activation conditions were periodically alternated five times in the course of a single experiment lasting 5.5 min. The control condition was presented first.

Stop task. In the control condition, an airplane appeared on the screen; the interstimulus interval was 1650 msec—the airplane appeared for 1000 msec and was then followed by a 650-msec blank screen—and there were 18 stimuli per epoch. For 50% of the trials the airplane was followed by a zeppelin, which appeared 250 msec after the onset of the airplane, replacing it for 300 msec, and was then followed by a blank screen for 1100 msec. The subject was required to press a button whenever an airplane appeared, whether or not it was followed by a zeppelin. The activation (stop) condition was identical except that a bomb appeared in 50% of the trials, instead of the zeppelin, 250 msec after the airplane. The subject was instructed to press the button if the airplane appeared alone and not to press the button if the airplane was followed by the bomb.

Delay task. This test alternated between two synchronization tasks that differed exclusively in their interstimulus interval (short- and long-event-rate conditions). In the short-event-rate condition a visual stimulus appeared on a computer screen with an interstimulus interval of 600 msec. The subject had to produce high-frequency movements (tapping) in order to synchronize his motor response to the visual stimulus. In the long-event-rate (delay) condition a visual stimulus appeared with an interstimulus interval of 5 sec and the subject had to synchronize his motor response to the visual stimulus, generating intermittent movements. In both conditions the subject was instructed to synchronize his motor response with the appearance of the stimulus by pressing a response button with his right hand at the same time as or shortly after seeing the visual stimulus. In order to be able to synchronize, especially in the long-event-rate condition, the subject was instructed to monitor the time elapsed since presentation of the previous stimulus.

The computerized activation images were visually presented to the subject in the scanner by means of a mirror from a liquid crystal diode projector. Throughout acquisition of the MR images, the subject responded to the stimuli by means of a right-handed button press, which was recorded by means of an MR-compatible interface to a personal computer.

Functional MRI Data Acquisition and Analysis

Image acquisition. Gradient-echo echoplanar MR images were acquired by using a 1.5-T GE Signa system (General Electric, Milwaukee) fitted with Advanced NMR hardware and software (Advanced NMR Systems, Woburn, Mass.) at the Maudsley Hospital, London. Daily quality assurance was carried out to ensure a high signal-to-noise ratio and excellent temporal stability by using an automated quality control procedure (29). A quadrature birdcage head coil was used for radio frequency transmission and reception. In each of 15 noncontiguous planes parallel to the anterior-posterior commissure, 100 T2*-weighted MR images depicting blood-oxygenation-level-dependent (BOLD) contrast (30) were acquired with TE=40 msec, TR=3000 msec, flip angle=90°, in-plane resolution=3.1 mm, slice thickness=5 mm, slice-skip=0.5 mm. Head movement was limited by foam padding within the head coil and a restraining band across the forehead. At the same session, a 43-slice, high-resolution imaging recovery echoplanar image of the whole brain was acquired in the intercommissural plane with TE=40 msec, TI=180 msec, TR=16,000 msec, in-plane resolution=1.5 mm, slice thickness=3 mm, slice-skip=0.3 mm.

Movement estimation and correction. Before image analysis a two-stage motion correction procedure was used in order to reduce the impact of slight subject motion (31). Since head movement can adversely affect functional MRI analysis and differences in the degree of motion between groups can account for apparent differences in physiological response, the amounts of stimulus-correlated motion in the two groups were compared by using the mean of each of six rigid body movement modes (three translations, three rotations), individually estimated for each subject.

Generic brain activation mapping. The methods used for functional MRI time series analysis have been described elsewhere in de-
The power of periodic signal change at the frequency of alternation between the control and activation conditions (1/60 Hz) was modeled by the sum of a sine wave and cosine wave at that frequency. The amplitudes of the sine and cosine waves, \(\gamma\) and \(\delta\) respectively, were estimated by fitting a sinusoidal regression model to the movement-corrected functional MRI time series at each voxel. The model was fit by using an iterated least-squares procedure: the residuals of an ordinary least-squares fit were modeled as a first-order autoregressive process, the terms of the sinusoidal regression model were transformed by the estimated first-order autoregressive coefficient, and the transformed model was fit again by ordinary least squares. This procedure can be regarded as a technique for prewhitening model residuals or noise. The sum of squared amplitudes of an ordinary least-squares fit were modeled as a first-order autoregressive process, while controlling for the possibly confounding effects of variabilities in the mean power of response to the activation condition. Voxels activated in phase with the activation condition were estimated by fitting a sinusoidal regression model to the delay task performance during scanning. Maps of median power of response in the stop condition were observed in the right prefrontal cortex (Brodmann area 8/32), right medial/inferior prefrontal cortex (Brodmann areas 9/45 and 45), right supplementary motor area (Brodmann area 6), and right and left caudate nuclei (figure 1). Generic activation in the comparison subjects during the stop condition (\(p<0.003\)) was observed in the right mesial frontal cortex (approximate Brodmann area 8/32), right medial/inferior prefrontal lobe (Brodmann areas 9/45 and 45), right supplementary motor area (Brodmann area 6), and right and left caudate nuclei (figure 1). Generic activation in the subjects with ADHD during the stop condition was observed in the right pre- and postcentral gyrus (Brodmann area 4/3/2/1), right inferior parietal lobe (Brodmann area 40), and right caudate nucleus (figure 1). We used analysis of variance, as described earlier more formally, to test the null hypothesis of zero between-group difference in the mean fundamental power quotient at each voxel generically activated in one group or both. There was no overlap between the activation patterns of the two groups. The search volume for this comparison was 196 voxels, and the voxel-wise probability of a false positive test was \(p<0.05\). At this size of test and with an assumption of in-
Dependence, we expect 10 false positive tests. In fact, we observed significant differences at 46 voxels.

Compared to the hyperactive group, the comparison subjects showed significantly greater power of functional response in the right mesial frontal cortex (Brodmann area 8/32), right inferior and medioinferior frontal lobe (Brodmann areas 45 and 9/45), and predominantly left caudate nucleus (table 1, figure 2).

**Delay task.** Generic activation in the comparison subjects during the delay condition was observed predominantly in the right mesial frontal lobe (Brodmann area 32) and in the posterior cingulate gyrus (Brodmann area 31), right supplementary motor area (Brodmann area 6), and right and left extrastriate cortex (Brodmann area 18/19) (figure 1).

Generic activation in the subjects with ADHD during the delay condition was observed mainly in both the left and right putamen, the right supplementary motor area (Brodmann area 6), and the right and left extrastriate cortex (Brodmann area 18) (figure 1).

The search volume for ANCOVA was 148 voxels, and the voxel-wise probability of a false positive test was p<0.05. At this size of test and with an assumption of independence, we expect seven false positive tests. We observed significant differences at 67 voxels.

The comparison subjects showed greater power of response in the anterior (Brodmann area 32) and posterior (Brodmann area 31) cingulate gyrus. A small focus of greater power of response in the hyperactive group was observed in the right supplementary motor area (Brodmann area 6) (table 1, figure 2).

**DISCUSSION**

The neuroactivation pattern observed in the hyperactive adolescents differed quantitatively and qualitatively from that of the comparison subjects during performance of two tasks testing high-level executive control. Overall, the hyperactive adolescents showed less brain activity, predominantly in the right hemisphere mesial frontal cortex during both tasks and in the right inferior prefrontal cortex and left caudate nucleus during the stop task.

Hypofrontality in the hyperactive adolescents is in line with the hypothesis of a maturational lag as the cause of ADHD, in particular, late development of the frontal lobes (38). The brain region that was activated in the comparison but not in the hyperactive subjects during performance of both tasks was the right mesial frontal gyrus—at the border with the anterior cingulate in the stop task and in the anterior part of the anterior cingulate in the delay task. The activation of this area in both tasks suggests that it subserves higher-order motor control functions, such as motor attention and response selection, common to both tasks. This is
in line with reports from neuroimaging studies of adults that the anterior cingulate is involved in a wide range of motor attentional and motor control functions, including output-related attention and response selection (39, 40). In subjects with ADHD, the structural development of this area has been related to performance on selective attention (12) and subnormal cerebral glucose metabolism has been observed during performance on sustained attention (15). The underfunctioning of a structure responsible for motor attention may underlie the deficits in different executive functions in ADHD. Less activation in the posterior cingulate during the delay task suggests, for the first time to our knowledge, that not only prefrontal but also posterior parts of the midline attentional system are affected in ADHD.

The comparison subjects’ activation in the right inferior frontal lobe and caudate nucleus during the stop task seems to be more specifically related to response inhibition. The right inferior frontal lobe—and its projections to the caudate—has been related to response inhibition in recent functional neuroimaging studies (12, 41). It thus seems that the brake system of the brain is localized to the right prefrontal lobe, and its underactivation in ADHD seems to be the neural correlate of a less efficient inhibitory motor control. The right hemisphere pattern of hypofrontality in the hyperactive subjects in both experiments and the caudate underactivation in the stop task support the existing structural and functional neuroimaging evidence for right hemisphere frontal and striatal dysfunction and/or dysplasia in ADHD (10–12, 17, 18).

The “qualitatively” different patterns of activation in the two groups was not specifically part of our hypothesis and should therefore be interpreted with caution. It could be argued that the lack of prefrontal activation has been compensated for in the hyperactive subjects by activation in posterior frontal brain regions. Further studies are needed to confirm this hypothesis.

In summary, it has been shown that functional MRI is a feasible technique for investigating the underlying neural mechanisms of cognitive functioning in a difficult developmental disorder such as ADHD. Our findings of mesial hypofrontality in adolescents with ADHD during performance of two different executive tasks suggest a task-unspecific deficit in higher-order attentional regulation of the motor output. Lower than normal activation of the right inferior prefrontal cortex and caudate nucleus during the stop task may be responsible for poor inhibitory control in ADHD.

**REFERENCES**

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