Frontal corticostriatal functional connectivity reveals task positive and negative network dysregulation in relation to ADHD, sex, and inhibitory control

Aki Nikolaidis\textsuperscript{a,*}, Xiaoning He\textsuperscript{a}, James Pekar\textsuperscript{b,c,d}, Keri Rosch\textsuperscript{b,e,f}, Stewart H. Mostofsky\textsuperscript{b,e,g}

\textsuperscript{a} Center for the Developing Brain, Child Mind Institute, USA
\textsuperscript{b} Center for Neurodevelopmental and Imaging Research, Kennedy Krieger Institute, USA
\textsuperscript{c} F.M. Kirby Center for Functional Brain Imaging, Kennedy Krieger Institute, USA
\textsuperscript{d} Department of Radiology, Johns Hopkins University School of Medicine, USA
\textsuperscript{e} Department of Psychiatry and Behavioral Sciences, Johns Hopkins University School of Medicine, USA
\textsuperscript{f} Department of Neuropsychology, Kennedy Krieger Institute, USA
\textsuperscript{g} Department of Neurology, Johns Hopkins University School of Medicine, USA

**Article Info**

**Keywords:**
- ADHD
- Corticostriatal
- FMRI
- Development
- Sex

**Abstract**

Frontal corticostriatal circuits (FCSC) are involved in self-regulation of cognition, emotion, and motor function. While these circuits are implicated in attention-deficit/hyperactivity disorder (ADHD), the literature establishing FCSC associations with ADHD is inconsistent. This may be due to study variability in considerations of how fMRI motion regression was handled between groups, or study specific differences in age, sex, or the striatal sub-regions under investigation. Given the importance of these domains in ADHD it is crucial to consider the complex interactions of age, sex, striatal subregions and FCSC in ADHD presentation and diagnosis. In this large-scale study of 362 8–12 year-old children with ADHD (n = 165) and typically developing (TD; n = 197) children, we investigate associations between FCSC with ADHD diagnosis and symptoms, sex, and go/no-go (GNG) task performance. Results include: (1) increased striatal connectivity with age across striatal subregions with most of the frontal cortex, (2) increased frontal-limbic striatum connectivity among boys with ADHD only, mostly in default mode network (DMN) regions not associated with age, and (3) increased frontal-motor striatum connectivity to regions of the DMN were associated with greater parent-rated inattention problems, particularly among the ADHD group. Although diagnostic group differences were no longer significant when strictly controlling for head motion, with motion possibly reflecting the phenotypic variance of ADHD itself, the spatial distribution of all symptom, age, sex, and other ADHD group effects were nearly identical to the initial results. These results demonstrate differential associations of FCSC between striatal subregions with the DMN and FPN in relation to age, ADHD, sex, and inhibitory control.

## 1. Introduction

Frontal corticostriatal circuits (FCSC) are key in the development of self-regulation and learning of behavior ranging from motor to cognitive and emotion functions (Postuma and Dagher, 2006; Graybiel, 2008; Arnsten and Rubia, 2012; Nikolaidis et al., 2014; Graybiel and Grafton, 2015). Both learning and regulatory mechanisms of the FCSC are thought to be closely tied to reward through dopaminergic pathways (Volkow, 2009; Alexander, DeLong, and Strick, 1986; Tost, Alam, and Meyer-Lindenberg, 2010; Doyon et al., 2009). Understanding the diverse roles of FCSC in relation to diagnostic, symptomatic, and behavioral variation in healthy and psychiatric populations has become an important area of research in clinical neuroscience. Researchers have recognized that atypical development of FCSC may contribute to the pathophysiology of neurodevelopmental disorders, in particular attention-deficit/hyperactivity disorder (ADHD; Mennes et al., 2012; Castellanos and Proal, 2012). ADHD is associated with deficient self-regulation of attentional/cognitive (Castellanos et al., 2006), emotional (Shaw et al., 2014; Da Fonseca et al., 2009) and motor (Mostofsky, Newschaffer, and Denckla, 2003; Mostofsky et al., 2006;...
Macneil et al., 2011; Gilbert et al., 2011) responses, which contribute to core symptoms of inattention, hyperactivity, and impulsivity.

Given the crucial role that FCSC plays in behavioral regulation, researchers have long hypothesized they play a crucial role in ADHD pathophysiology (Heilman, Voeller, and Nadeau, 1991; Denckla, 1991). However, neuroimaging studies of the intrinsic network connectivity of FCSC have yet to converge. Of the existing studies that have probed anomalous FCSC in ADHD, results have been relatively heterogeneous. Some studies have observed increases in FCSC functional connectivity with diagnosis and symptom effects (Oldehinkel et al., 2016a, 2016b; Sanefuji et al., 2017; Ma et al., 2016; Dias et al., 2013, 2015; Rosch et al., 2018; Damiani et al., 2021; Mennes et al., 2012; Di Martino et al., 2013; Yang et al., 2018), while others have observed decreases (Hong et al., 2015; Cao et al., 2006; Posner et al., 2013), both increases and decreases in FCSC in ADHD (Tomasi and Volkow, 2012; Cao et al., 2009), or no differences compared to typically developing children (Oldehinke et al., 2016a). This inconsistency of results may reflect variability in FCSC and associations with ADHD and symptom effects due to the particular striatal subregions investigated or variability in sample age and sex composition. Rosch 2018 was the first study to simultaneously consider the impact of sex and ADHD of FCSC and found key differences by sex as well as sex by diagnosis interaction in FCSC. Furthermore, regions in the FCSC are known to exhibit significant structural developmental changes tied to puberty timing (Raznahan et al., 2014), and developmental changes in some FCSC regions are associated with ADHD (Barber et al., 2019). Finally, striatal subregions exhibit heterogeneity in their structural connectivity and functional relationships to cortex (Elliott et al., 2021), yet few prior studies have explicitly considered the cortical connectivity of striatal subregions in their analyses. Taken together, these findings suggest that characterization of differences in age, sex, and striatal subregion are essential in the pursuit of robust FCSC based biomarkers of ADHD.

In contrast to the variable findings from studies of FCSC connectivity in ADHD, a fairly consistent finding that has emerged from studies of cortico-cortical network connectivity is that adults and children with ADHD demonstrate hyperconnectivity of the default mode network (DMN), which is preferentially activated when an individual is not actively engaged in a task (Konrad and Eickhoff, 2010; Henry and Rosch, 2016). This inconsistency of results may reflect variability in FCSC and associations with ADHD and symptom effects due to the particular striatal subregions investigated or variability in sample age and sex composition. Rosch 2018 was the first study to simultaneously consider the impact of sex and ADHD of FCSC and found key differences by sex as well as sex by diagnosis interaction in FCSC. Furthermore, regions in the FCSC are known to exhibit significant structural developmental changes tied to puberty timing (Raznahan et al., 2014), and developmental changes in some FCSC regions are associated with ADHD (Barber et al., 2019). Finally, striatal subregions exhibit heterogeneity in their structural connectivity and functional relationships to cortex (Elliott et al., 2021), yet few prior studies have explicitly considered the cortical connectivity of striatal subregions in their analyses. Taken together, these findings suggest that characterization of differences in age, sex, and striatal subregion are essential in the pursuit of robust FCSC based biomarkers of ADHD.

The current sample included 362 children and adolescents with either a diagnosis of ADHD (n = 165; 47 girls) or typically developing (TD) children (n = 197; 61 girls). All participants were between ages 8–13 (mean 10.27, SD 1.27). Summary demographics for included participants are provided in Table 1. Participants were recruited from local schools, pediatricians (electronically via MyChart), community centers using flyers and word-of-mouth. Participants with ADHD were also recruited from local outpatient clinics. Study protocols were reviewed and approved by Johns Hopkins Medicine Institutional Review Board.

2. Methods

2.1. Participants

The current sample included 362 children and adolescents with either a diagnosis of ADHD (n = 165; 47 girls) or typically developing (TD) controls (n = 197; 61 girls). All participants were between ages 8–13 (mean 10.27, SD 1.27). Summary demographics for included participants are provided in Table 1. Participants were recruited from local schools, pediatricians (electronically via MyChart), community centers using flyers and word-of-mouth. Participants with ADHD were also recruited from local outpatient clinics. Study protocols were reviewed and approved by Johns Hopkins Medicine Institutional Review Board.
All parents completed an initial telephone screening to determine eligibility. Children with a history of intellectual disability, seizures, traumatic brain injury, neurological illnesses, prenatal exposure to teratogens, genetic disorders, or other neurodevelopmental disorders (e.g., Autism Spectrum Disorders) were excluded from participation. Eligible participants completed two laboratory sessions for each visit. Sessions occurred within a period of six months (with nearly all occurring within 4 weeks) to maintain validity of data collected between Session 1 and Session 2. Additional inclusion criteria applied across groups following study participation include the following: 1) successful fMRI scan; (2) Full Scale Intelligence Quotient (FSIQ) above 80 using either the Wechsler’s Intelligence Scales for Children current at the time of testing (WISC-IV or WISC-V) or the Wechsler Abbreviated Scale for Intelligence, 2nd Edition (WASI-II) (Wechsler, 1992), and 3) mean framewise displacement (FD) (Power et al., 2012) across all fMRI scans are within 3 standard deviations from the group mean FD.

At each visit, a diagnosis of ADHD was determined using a structured semi-structured parent interview, either the Diagnostic Interview for Children and Adolescents (DICA-IV) (Reich, 2000) or the Kiddie Schedule for Affective Disorders and Schizophrenia (K-SADS) (Kaufman et al., 2016); the ADHD Rating Scale (ADHD-RS) (DuPaul et al., 1998) and the Conners Parent Rating Scale-Revised or Version 3 (Conners, 2000) were used to confirm diagnosis and to provide dimensional measures of ADHD symptom severity. Parents of all participants provided written consent, and all participants provided assent. All children taking stimulant medication (n = 97; 26 girls; see Table 1) were asked to withhold medication on the day prior to and day of testing. Children taking psychotropic medications other than stimulant medication (n = 15; 3 girls) did not discontinue their medication for study visits. Additionally, parents were instructed on both the diagnostic interview and report forms to make ratings based on their children’s symptoms off of their regularly prescribed medication.

Participants were included in the ADHD group if they: (1) met criteria for an ADHD diagnosis either on the DICA-IV or K-SADS and (2) received a T-score of 60 or higher on the DSM Inattentive or DSM Hyperactive-Impulsive scales on the Conners Parent or Teacher (when available) rating scales (revised or 3rd edition), or a score of 2 or 3 (i.e., symptoms rated as occurring ‘often’ or ‘very often’) on at least 6/9 items on the Inattentive or Hyperactivity/Impulsivity scales of the ADHD-RS Home or School (when available) Version. Children with ADHD were allowed to meet criteria for comorbid psychiatric diagnoses on the DICA-IV or K-SADS including oppositional defiant disorder (ODD; n = 53; 17 girls), anxiety disorders (n = 38; 12 girls) and depressive disorders (n = 5; all boys) (Supplementary Table S1). Girls and boys with ADHD did not differ in comorbid diagnoses of ODD (p > .05), anxiety (p > .05) or depression (p > .05). Master’s level clinicians conducted all diagnostic interviews and integrated information from rating scales to inform diagnoses under the supervision of licensed doctoral level clinical psychologists.

Participants were included in the control group if they: (1) did not meet criteria for any psychiatric disorders on the DICA-IV or K-SADS, (2) had below clinically significant scores (T < 60) on the Conners Parent and Teacher (when available) rating scales, and ADHD-RS Home and School (when available) Versions, and (3) did not have immediate family with ADHD.

### Table 1

Demographic and clinical characteristics of ADHD and TD groups overall and within sex. Cohen’s d is used to measure effect size.

|               | TD Girls | Boys | TD Girls vs. TD Boys | ADHD Girls | Boys | ADHD Girls vs. ADHD Boys | TD Girls vs. ADHD Girls | TD Boys vs. ADHD Boys | TD All vs. ADHD All |
|---------------|----------|------|----------------------|------------|------|--------------------------|------------------------|----------------------|---------------------|
| **Age**       |          |      |                      |            |      |                          |                        |                      |                     |
| Mean (SD)     | 10.09 (1.01) | 10.43 (1.21) | -0.30   | 10.19 (1.32) | 10.21 (1.43) | -0.01          | -0.08 (0.17)          | 0.17 (0.10)          |                     |
| **IQ**        | 113.13 (10.31) | 116.60 (12.28) | -0.30   | 107.68 (14.25) | 105.90 (12.54) | 0.14           | 0.45* (0.87*** 0.73*** |                     |                     |
| **SES Family**| 54.54 (8.99)  | 53.97 (9.93)  | 0.06    | 51.96 (8.95)  | 51.14 (9.94)  | 0.08           | 0.29 (0.29* 0.29**)   |                     |                     |
| **Inattention T-Score** | 46.46 (6.37)  | 45.27 (5.71)  | 0.20    | 82.35 (8.56)  | 72.77 (9.45)  | 1.04***        | -4.86*** -3.58*** -3.67*** |                     |                     |
| **Hyperactive-Impulsive T-Score** | 46.18 (5.92)  | 46.83 (5.72)  | -0.12   | 76.98 (13.48) | 72.92 (12.58) | 0.32           | -3.2*** -2.73*** -2.85*** |                     |                     |
| **GNG RT variability** | 105.87 (48.94)  | 92.22 (48.82)  | 0.28    | 128.68 (56.66) | 148.83 (99.66) | -0.23          | -0.43* -0.69*** -0.61*** |                     |                     |
| **GNG ComRate** | 0.35 (0.18)  | 0.41 (0.18)  | -0.30   | 0.38 (0.18)  | 0.51 (0.19)  | -0.70***       | -0.13 -0.56*** -0.45*** |                     |                     |
| **GNG Rate**  | 0.04 (0.03)  | 0.04 (0.02)  | 0.01    | 0.05 (0.03)  | 0.05 (0.03)  | 0.18           | -0.43* -0.29* -0.33**  |                     |                     |
| **Displacement** |          |      |                      |            |      |                          |                        |                      |                     |
| % Stim Med    | 0         | 0    | n/a                  | 55.32%     | 60.17% | -0.10         | n/a                   | n/a                  |                     |
| % Non Stim Med | 0        | 0    | n/a                  | 6.38%      | 10.17% | -0.15        | n/a                   | n/a                  |                     |
| % ODD         | 0         | 0    | n/a                  | 36.73%     | 30.89% | -0.12        | n/a                   | n/a                  |                     |
| % Anxiety     | 1.64%     | 2.21%| -0.04                | 23.40%     | 19.49% | 0.09          | -0.73*** -0.59*** -0.63*** |                     |                     |
| % Depression  | 0         | 0    | n/a                  | 4.24%      | n/a    | n/a          | n/a                   | n/a                  |                     |
registered to the MNI152 template (3 mm isotropic). Following recommendations of Ciric et al. (2017), time series were bandpass filtered at 0.01–0.08 Hz and nuisance signal removal was performed for 24 regressors derived from the parameters estimated during motion realignment, and physiological noise was modeled using the 5 principal components with the highest variance from a CompCor decomposition of white matter and CSF time series (Behzadi et al., 2007). Mean, squared, delayed, and squared delayed regressors were also used for the global signal, and WM/CSF signal. Extracted time series were then normalized before further analysis. Ciric et al. (2017) shows that median functional connectivity associations with motion are reduced more by despiking than scrubbing. Our analyses focused on moderate distance connections between the striatum and frontal cortex, therefore we applied AFNI 3D-despike, rather than scrubbing as our motion correction approach. Furthermore, given that head motion is an known phenotypic indicator for ADHD, scrubbing would have resulted in fewer frames for our ADHD sample than the TD sample, resulting in potential biases through systematically different levels of reliability in functional connectivity estimates between groups.

Extensive prior work has shown that the length of fMRI acquisition has a significant impact on its reliability (Birn et al., 2013; Cho et al., 2021; Laumann et al., 2015; Gratton et al., 2018; Nikolaidis et al., 2020) that task-specific activity accounts for a low percentage of variance in the fMRI signal (Gratton et al., 2018), and that concatenating rest and task data together can improve the scan reliability (Cho et al., 2021). Given these findings, we used FSL to concatenate multiple resting state and task-based fMRI scans after preprocessing to aggregate the longest possible scan for each participant (Smith et al., 2014). The average total aggregate scan length per subject was 9.89 min (ranging from 5 to 22.83 min). 351 subjects had available rs-fMRI data with an average length of 6.31 min per subject (ranging from 5 to 6.5 min); 84 subjects had 4 available task-fMRI data (GNG1, GNG2, GNGr1, GNGr2), each had an average length of 4.07 min per subject (ranging from 3.79 to 4.08 min). ADHD participants and TD participants did not differ in rs-fMRI (p > .05), task-fMRI (p > .05) or aggregated (p > .05) scan time. Girls and Boys did not differ in rs-fMRI (p > .05) scan time, but differed in task-fMRI (p < .05) scan time and aggregated (p < .001) scan time, with girls having longer scan time than boys. The aggregated scans are then down sampled to 2 mm, and spatially smoothed at 6 mm using FSL (Smith et al., 2014).

2.3. Functional connectivity and randomise analyses

Frontal cortical and striatal regions were defined using Harvard-Oxford cortical and subcortical atlases (Kennedy et al., 2016). The frontal cortical region consisted of the following bilateral cortical regions: precentral gyrus, superior frontal gyrus, middle frontal gyrus, paracingulate gyrus, cingulate gyrus anterior division, inferior frontal gyrus, and pars triangularis, frontal pole, subcallosal cortex, juxtapositional lobule cortex (formerly supplementary motor cortex), and frontal orbital cortex. To assess spatial specificity of corticostriatal interactions in relation to self-regulation of emotion, cognition, and motor functions implicated in ADHD (Armstein and Rubia, 2012), we created three striatal maps (limbic, executive (cognitive), and motor) based on the Harvard-Oxford probabilistic subcortical atlas. These regions were defined using the following bilateral subcortical regions: limbic striatum: ventral putamen, nucleus accumbens; executive striatum: anterior caudate, dorsal putamen; motor striatum: posterior and dorsal putamen (Kennedy et al., 2016).

An average time series was extracted from each striatal mask, and FCSC connectivity was calculated as the voxel-wise Pearson correlation between the striatal seeds and the frontal regions using the Nilearn package (Abraham et al., 2014). Fisher Z transform (Fisher, 1921) was then applied to the correlation values. Using FSL’s Randomise (version 2.9) we applied non-parametric permutation testing to assess the relationships between frontostraital connectivity, age, ADHD diagnosis and symptom severity, GNG performance, and sex (Nichols and Holmes, 2002; Winkler et al., 2014). We used 5000 permutations in Randomise, and statistical thresholding was performed with FSL’s threshold-free cluster enhancement (TFCE) with a family-wise error rate (FWE) of p < .05. Temporally demeaned data before model fitting and variance smoothing for t-stats were selected.

We used Randomise to test for two-way interactions of diagnosis, age, and sex (Table 2). We also tested the main effects of diagnosis, age, and sex, as well as main effects of Inattention T-score, Hyperactivity T-score, and GNG performance. We also tested interaction effects of GNG with age, sex, and diagnosis to probe the neural correlates of ADHD-related sex differences in GNG performance (Table 1).

To investigate the spatial distribution of FCSC overlap with canonical correlation networks (Yeoo et al., 2011), we compared overlap of these results with each of the networks and report percentage of each statistical map covering each cortical network (Supplemental Table 2). The frontal cortical region of interest is comprised of differing amounts of each of the 7 Yeo network, (e.g. 5% for the Dorsal Attention Network to 26% for the DMN). Therefore, we also show the amount of overlap with these networks as well as the difference in percentage of significant voxels for each analysis compared to that which would be expected due to chance (Supplemental Table 3). We assess the similarity of spatial results across analyses and report the spatial correlations of these results (Supplemental Figure 8).

Table 2
Randomise results summary across three striatal subregions. Figure numbers associated with each result are included in the final column.

| Regions                  | Figure | Figure# |
|--------------------------|--------|---------|
| **Limbic striatum**      | 1      |         |
| **Executive striatum**   | 2      |         |
| **Motor striatum**       | 3      |         |

| Main effects             | Regions                  | Figure | Figure# |
|--------------------------|--------------------------|--------|---------|
| Main Age                 | Positive Age effect      | 1      |         |
| Male Dx                  | Positive Age effect      | 2      |         |
| Female Dx                | Positive Age effect      | 2      |         |
| ADHD symptom severity effects | Inattention T-Score | –      | Positive effect | 2 |
|                          | Hyperactive-Impulsive T-Score | –     | Positive effect | 2 |
|                          | ADHD Hyperactive-Impulsive T-Score | –     | Positive effect | 2 |
| Go/no-go main effects    |                          |        |         |
| GNG RT variability       | Negative effect          | 3      |         |
| GNG ComRate              | Negative effect          | 3      |         |
| ADHD GNG ComRate         | Positive effect          | 3      |         |
| Go/no-go interaction effects |                       | –      |         |
| Dx*GNG RT variability    | F > M positive slope     | 3      |         |
| Sex*GNG ComRate Interaction |                       | –      |         |
| TD Sex*GNG ComRate       | F > M positive slope     | 3      |         |
| ADHD Sex*GNG ComRate     | F > M positive slope     | 3      |         |
2.4. Go/No-Go task

A subset of participants completed a standard GNG task \((n=298)\) assessing response inhibition and variability (DeRonda et al., 2021; Wodka et al., 2007). The task stimuli consisted of a green spaceship for "Go" trials (80% of trials) and a red spaceship for "No-Go" trials (20% of trials) presented on-screen for 300 ms with an interstimulus interval of 2000 ms. Participants were instructed to press the spacebar with their index finger as quickly as possible in response to green spaceships. There were 11 practice trials followed by 217 experimental trials lasting 8 min and 19 s. Responses and reaction times (RT) were recorded to calculate the commission error rate (ComRate), defined as incorrectly pressing for a red spaceship, and RT variability, using an ex-Gaussian parameter quantifying the skewed tail of the RT distribution (tau) as an index of response variability separate from response speed (Epstein et al., 2011; Kolier et al., 2013; Tamm et al., 2012).

3. Results

3.1. Demographics, ADHD symptoms, and GNG performance by sex and diagnosis

Descriptive statistics and results of t-tests comparing demographic characteristics and GNG task performance between diagnostic groups across and within sex are provided in Table 1. Cohen’s \(d\) is reported as an estimate of effect size (Cohen, 2013).

Diagnostic groups did not differ in age across or within sex, whereas intellectual reasoning ability (full scale IQ) was significantly lower in the ADHD group for both boys and girls \((d=0.73, p<.001)\). Socioeconomic status (SES) was also significantly lower in boys with ADHD compared to TD boys \((d=0.29, p<.010)\), whereas it did not significantly differ among girls with and without ADHD, although a similar effect size was observed. As expected, parent-rated symptoms of inattention and hyperactivity/impulsivity T-scores were much higher among children with ADHD compared to TD children (inattention \(d=3.67, p<.001\); hyperactivity/impulsivity \(d=2.85, p<.001\)) with similar diagnostic effects among girls and boys. In addition, inattention T-scores were greater among girls compared to boys with ADHD \((d=1.04, p<.001)\), suggesting higher severity of inattentive symptoms among ADHD females.

For the GNG task, RT variability was higher for ADHD compared to TD \((d=0.61, p<.001)\) with significant effects among boys \((d=0.69, p<.001)\) and girls \((d=0.43, p<.05)\). GNG ComRate was also greater among children with ADHD compared to TD children \((d=0.43, p<.001)\), but this significant effect was specific to boys \((d=0.56, p<.001)\) and not observed among girls \((d=0.13, p>.05)\); furthermore, boys with ADHD showed significantly greater GNG ComRate compared to ADHD girls \((d=0.70, p<.001)\).

3.2. FCSC connectivity – associations with age

We found a significant positive main effect of age that was highly consistent across analyses using the three striatal subregions (Table 2; Pearson’s spatial correlation = 0.66–0.85; Supplemental Figure 8). By counting voxel-wise overlap between these significance maps and the Yeo 7 canonical intrinsic connectivity networks, we found that areas with significant age-associated increases in FCSC connectivity were primarily localized to the DMN (24.6%–28.4%), FPN (18.7%–23.2%), and somatomotor networks (15.0%–20.7%; Supplemental Table 2). These frontal cortex regions included the supplementary motor area (SMA), paracingulate gyrus, superior frontal gyrus, middle frontal gyrus, pre-central gyrus, cingulate gyrus, frontal pole, and frontal orbital cortex (Fig. 1). Although there was no Diagnosis × Age interaction for FCSC connectivity across striatal subregions, we conducted an exploratory analysis examining age-related change in FCSC connectivity within our ADHD and TD groups given prior findings of maturational lag in functional connectivity of cortical networks in ADHD (Sripada et al., 2014). Interestingly, while we found significant age-associated increases with FCSC connectivity for ADHD in all three striatal subregions, for TD children significant age-associated increases with FSCS were localized to the limbic striatum (Fig. 1). Furthermore, we found notable differences in the spatial distribution of age effects between the limbic-frontal connectivity effects for TD and ADHD. ADHD children showed relatively more age-associated effects in the somatomotor, dorsal attention, and ventral attention networks while showing less in the default and limbic networks.

![Fig. 1. ADHD Age Effects](image-url)

The figure shows the spatial distribution of age effects in ADHD across striatal subregions. Dark blue, teal, and yellow correspond to the limbic, executive, and motor striatum respectively. Main age effects include significant voxels in the superior frontal gyrus, middle frontal gyrus, supplementary motor cortex, precentral gyrus, frontal orbital cortex, frontal pole, frontal medial cortex, and subcallosal cortex. For the ADHD group effects include significant voxels in the superior frontal gyrus, supplementary motor cortex, middle frontal gyrus, precentral gyrus, frontal orbital cortex, frontal pole. TD age effects were relatively restricted, including significant voxels in the subcallosal cortex, frontal medial cortex, and frontal pole. At least 50% of the voxels in each region had significant voxels in order to be mentioned here.
networks compared to TD children.

### 3.3. FCSC connectivity – associations with ADHD diagnosis and symptom severity

The effect of diagnosis on FCSC connectivity was specific to boys, such that boys with ADHD showed greater FCSC connectivity selectively with the limbic striatum and regions of the DMN (70%), with nearly no significant regions in the FPN (3.5%; Fig. 2). To assess the robustness of all Randomise results, we also repeated all analyses while controlling for both total scan length (which differed between boys and girls) and mean framewise displacement, which differed among ADHD and TD groups (Table 1). Although the ADHD>TD results in males were no longer significant when including scan length and FD as a covariate, we found high concordance in the spatial maps of our other analyses and those corrected for motion (Pearson’s spatial correlation; average = 0.97; range 0.88–0.99; See Supplemental Table S1 and Supplemental Fig. S5). Correcting for both scan length and motion also showed high spatial similarity with our original analyses (Pearson’s spatial correlation; average = 0.92; range 0.82–0.98). We chose to not match the groups on mean FD given evidence that head motion is correlated with ADHD symptomatology and that head motion and ADHD may have similar genetic loadings (Couvy-Duchesne et al., 2016), suggesting that co-varying for FD accounts for variance attributable to ADHD. We include spatial maps of significant results controlling for only motion (Supplemental Figs. 1–3) as well as both motion and scan length (Supplemental Figures 4–6).

Analysis of ADHD symptom associations across the sample revealed a significant positive relationship between parent-rated inattention (T-score) and FCSC connectivity between the motor striatum and regions of the DMN and FPN (54.9%–22.9%) including the frontal pole, superior frontal gyrus, middle frontal gyrus, and paracingulate cortex (Fig. 2). These effects were spatially segregated from the effects of diagnosis as evidenced by the relatively low spatial overlap of the inattention T-score effects and the diagnosis effects (Pearson’s r = 0.23). The significant inattention effects seemed to be driven mainly by the ADHD group, with a broader range of areas demonstrating a significant positive relationships between inattention T-score and FCSC connectivity in the motor and executive striatum and regions of the DMN (26.8%–34.6%) and FPN (21.9%–23.9%) including the superior frontal gyrus, middle frontal gyrus, paracingulate gyrus, cingulate gyrus, precentral gyrus, and frontal pole (Fig. 2). The frontal regions showing significant associations were highly consistent across striatal subregions (Pearson’s spatial correlation = 0.61). We found no relationship between FCSC connectivity and the hyperactivity/impulsivity T-score.

### 3.4. FCSC connectivity – associations with GNG task performance

Analysis of GNG task associations with FCSC connectivity showed a negative relationship with GNG RT variability (tau) that reproduced across all striatal regions and regions of the frontal cortex in the DMN (42.7%–43.0%) and FPN (21.3%–22.9%) including the paracingulate gyrus, superior frontal gyrus, and frontal pole (Pearson’s correlation = 0.50–0.82). In other words, greater RT variability was associated with less FCSC connectivity. The executive striatum and motor striatum showed a tight similarity of findings which also included the middle frontal gyrus, cingulate gyrus, frontal orbital cortex and frontal medial cortex (Pearson’s correlation = 0.82 Fig. 3). The limbic striatum showed a moderately different relationship to FCSC connectivity, with a reduced spatial overlap with the executive and motor striatum effects (Pearson’s correlation 0.50 and 0.58 respectively), and more notably nearly all regions significantly associated with RT variability were located in the DMN (72.6%). This significant negative relationship between FCSC connectivity and RT variability seemed to be mostly driven by the ADHD group, who showed largely a similar spatial significance pattern, while the TD group did not show a significant relationship between FCSC connectivity and RT variability. Spatial correlation between the full sample maps and ADHD sample alone calculated pairwise for each striatal seed shows these maps to be highly consistent (Pearson’s correlation = 0.69–0.92; Fig. 3).

Although there was no evidence of FCSC connectivity in relation to GNG ComRate across the ADHD and TD groups (Table 2), given the behavioral finding of ADHD-related sex differences for GNG ComRate, such that only boys with ADHD showed increased ComRate compared to

---

**Fig. 2.** ADHD Diagnosis and Symptom Severity Effects. Dark blue, teal, and yellow correspond to the limbic, executive, and motor striatum respectively. Diagnosis effects in males were found in the subcallosal cortex, and frontal medial cortex. Inattention T-Score effect was found in the inferior frontal gyrus, pars triangularis, inferior frontal gyrus, pars opercularis, paracingulate gyrus, and superior frontal gyrus. Inattention T-score effects for the ADHD sample were found in the paracingulate gyrus, superior frontal gyrus, anterior cingulate gyrus, and middle frontal gyrus. At least 50% of the voxels in each region had significant voxels in order to be mentioned here.
TD boys and girls with ADHD, we tested whether sex moderated the relationship between GNG ComRate and FCSC connectivity. Results indicated a significant sex*GNG ComRate interaction in the motor subregion of the striatum, with girls having a significantly stronger positive relationship between FCSC connectivity and GNG ComRate than boys in both the full sample and within the ADHD sample (Fig. 3). Significant regions for the full sample were highly consistent with the ADHD sample (Fig. 3) indicating FCSC connectivity of the limbic and motor striatum with subregions of the FPN (31.3–45.9%) and DMN (13.9–34.8%) including the frontal pole, middle frontal gyrus, paracingulate gyrus, superior frontal gyrus (Pearson’s correlation = 0.68).

3.5. Intrinsic network specificity in frontal striatal correlates of ADHD and development

The cortical networks with the greatest overlap with the regions with significant cortico-striatal interactions with Inattention, ADHD, and the GNG task were the DMN and FPN. Interestingly, the loadings on each of these two networks are strongly negatively related (Pearson’s correlation = −0.72), when frontal-striatal connectivity shows significant loadings in the FPN it tends not to do so in the DMN and vice versa. For example, the GNG tau and Inattention T-score results show a significant relationship between striatal connectivity and the DMN but very little connectivity to the FPN, while the sex by GNG interaction effects load significantly onto the FPN but not the DMN.

4. Discussion

4.1. Overview

ADHD is a heterogeneous disorder in terms of symptom presentation and associated deficits in cognitive, emotional, and motor control. Understanding the full range brain-behavioral pattern can provide a crucial roadmap for unraveling ADHD heterogeneity. Therefore, the current study investigated associations between fMRI-based measurement of FCSC connectivity and ADHD in a large sample of 362 school-age children and considered effects of sex and age. Addressing our first hypothesis, we found that, across the entire population (both ADHD and TD children), FCSC connectivity increased with age across much of the frontal cortex, including regions in the somatomotor network (SMN), FPN and DMN. Importantly, these age-related patterns of FCSC connectivity were largely distinct from those FCSC connectivity associated with diagnosis, symptom severity, and GNG performance, showing uniquely significant FCSC connectivity patterns in the subcallosal cortex, supplementary motor cortex, frontal orbital cortex, and precentral gyrus. Addressing our second hypotheses, we found that ADHD diagnosis was associated with greater FCSC connectivity but this was specific to boys with ADHD and connectivity between the limbic striatum with the DMN, extending prior work demonstrating cortico-cortico hyperconnectivity of the DMN in ADHD (Konrad and Eickhoff, 2010; Henry and Cohen, 2019; Posner et al., 2014; Sripada et al., 2014; Duffy et al., 2021; McCarthy et al., 2013; Barber et al., 2015; Hoekzema et al., 2014; Elton et al., 2014; Zhao et al., 2021). Interestingly, we found that parent-rated inattentive symptom ratings were associated with increased connectivity of the executive and motor striatum with regions of the DMN and FPN cortical networks, suggesting spatially distinct
effects from the observed diagnostic group difference in boys. Finally, RT variability during a GNG task (which was elevated among children with ADHD, regardless of sex) was associated with decreased FCSC connectivity with DMN and FPN across striatal subregions. In contrast, sex differences were observed for associations between FCSC connectivity and GNG inhibition errors (which were elevated among boys with ADHD only), such that the positive association was stronger among girls. Overall, our results provide evidence that heterogeneity in ADHD related to sex differences, symptom presentation, and cognitive task performance may be linked to dysregulation of task positive and task negative network connectivity with the striatum.

4.2. FCSC and age

Regarding our first hypothesis, FCSC connectivity increased with age across striatal subregions when examined in the full sample of ADHD and TD children. These age-related increases were observed across a large set of regions covering the SMN, DMN, FPN, and Dorsal Attention networks. Although the Diagnosis × Age interaction was not significant, there were notable differences in the spatial distribution of age-associations for ADHD vs. TD children. ADHD children showed an expanded set of FCSC regions with significant positive associations with age compared to TD children, with localized effects involving the limbic striatum, suggesting that changes in FCSC connectivity may be more widespread among children ADHD in this developmental period. Consistent with these findings, one previous investigation of FCSC connectivity age-effects in children with (and without) ADHD found similar evidence for a much more extensive set of FCSC patterns associated with age in children with ADHD and found limited overlap of FCSC patterns associated with age versus those associated with ADHD severity (Barber et al., 2019). The findings thereby suggest that while age associations with FCSC connectivity are quite extensive, they are also sensitive to ADHD and should be considered whenever investigating FCSC connectivity links to ADHD diagnosis and symptom severity.

4.3. FCSC connectivity and ADHD

Our second hypothesis, that ADHD diagnosis and symptom severity would be associated with increased striatal-DMN connectivity and decreased striatal-FPN connectivity, and that these associations may be stronger for girls with ADHD, was only partially supported. Consistent with this hypothesis, we found an effect of ADHD diagnosis for connectivity between the limbic striatum and DMN cortical regions, but only for males. These findings are inconsistent with our previous study (Rosch et al., 2018), reporting greater FCSC in girls with ADHD. However, we did not examine striatal subregions with different functional roles in our previous analyses and focused instead on connectivity with select cortical regions, which may have contributed to the discrepant results. It is also important to note that there were no Diagnosis × Sex interactions in our current analyses, suggesting that the diagnosis effects may be similar, but weaker in girls and not significant due to the smaller sample. These inconsistent findings may otherwise relate to variability in the methods (e.g., the use of group ICA and analysis of specific cortical regions in our prior study) and sample (e.g., elimination of comorbid mood and anxiety disorders in our prior study), highlighting the need for further research to understand ADHD-related sex differences in brain structure, function, and behavior. In addition to diagnostic group differences in FCSC connectivity, executive and motor striatal subregions both demonstrated a significant positive association between functional connectivity to DMN and FPN regions inattention t-score in ADHD children. Thus, although girls with ADHD did not significantly differ from TD girls in FCSC connectivity, they do display greater levels of inattention symptom severity, that relates to greater FCSC connectivity between the motor and executive striatum with regions of the DMN and FPN.

Our findings of atypical limbic FCSC in ADHD is consistent with prior studies implicating this circuit in ADHD. Increasing evidence from neuroimaging studies for anomalous limbic circuitry in ADHD, including ventromedial/orbitofrontal cortical regions and subcortical regions including the ventral striatum and amygdala (Dias et al., 2013; Tomasi and Volkow, 2012; Mennes et al., 2012; Cao et al., 2006, 2009; Posner et al., 2013). Our results suggest that limbic FCSC connectivity may be particularly sensitive to group-level diagnostic differences in boys, which are primarily examined in the ADHD literature as most studies include predominantly or exclusively male samples thereby limiting examination of sex differences. Our findings indicate, for the first time, novel dysregulation between DMN regions and the striatum in ADHD (specifically, the limbic striatum), something that has not been well characterized previously. Collectively, these findings suggest that disruptions in FCSC connectivity with functionally distinct striatal subregions may differentially relate to overall diagnostic group differences (involving the limbic striatum) and heterogeneity in symptom presentation (involving the motor and executive striatum).

ADHD has been widely regarded as a neurodevelopmental disorder involving dysregulation of FCSC connectivity as well as task-positive/ task-negative network integration (Barber et al., 2015; Castellanos et al., 2008; Sripada et al., 2014; Hoekzema et al., 2014; Elton et al., 2014). Our findings show that FCSC connectivity associations with ADHD, sex, symptom severity and cognitive control are mostly located in the FPN and DMN. Furthermore, we found across all analyses that the FPN and DMN FCSC connectivity associations were themselves negatively correlated. These findings suggest that FCSC connectivity may contribute to the task-positive/task-negative dysregulation often characterized in ADHD. DMN dysregulation is often purported to contribute to ADHD-associated difficulties with sustaining attention. Our findings suggest that DMN-striatum integration may play a role in DMN dysregulation associated with ADHD. FCSC connectivity are also closely involved in the emergence and integration of motor, cognitive, and affective skills over development. Given the key role of these circuits in child and adolescent cognitive maturation, and the neurodevelopmental nature of ADHD, dysregulation in FCSC connectivity involved in task positive/negative integration may contribute to the development of previously-reported differences in cortico-cortico dysregulation. Furthermore, it may be possible that alterations in cortico-cortical circuits may be an early hallmark of emergent dysfunction in cortico-cortico DMN-related alterations in ADHD. Future longitudinal work may be able to clarify the emergence of task-positive task negative dysregulation in cortico-cortico and corticostriatal networks.

4.4. FCSC connectivity and response control

Our third hypothesis, that GNG response inhibition errors would be associated with differences in striatal connectivity to FPN, whereas GNG RT variability would be associated with differences in striatal connectivity to DMN, and that this may differ among girls and boys, was partially supported. We found that across the sample (ADHD and TD), increased GNG RT variability associated with decreased striatal-DMN and FPN connectivity with similar effects across striatal subregions and this effect was driven by the ADHD group. In addition, a significant Sex × GNG ComRate interaction for FCSC connectivity was observed across the sample and within the ADHD group, suggesting ADHD-related sex differences in this relationship. Specifically, girls with ADHD showed a stronger positive relationship between functional connectivity of the striatum and FPN frontal cortical regions and GNG ComRate. Thus, although as a group, girls with ADHD did not show anomalous FCSC connectivity or response inhibition, within group heterogeneity did reveal associations between GNG ComRate and FCSC connectivity. Recent work by Duffy et al. (2021) found no association between ComRate and a graph-theory based participation coefficient metric of connectivity between the DMN and subcortical seeds taken from the Power atlas (Power et al., 2011). Differences in the seed regions used, sample size, and the analytical approach may have contributed to these
findings contrasting the current work.

4.5. Limitations

While the current study aimed to investigate corticostriatal associations with ADHD in a large sample of school-aged children, some limitations are worth noting. First and foremost, the cross-sectional nature of this sample limits our ability to conduct a thorough mapping of the development of corticostriatal circuits and their association with the emergence of ADHD symptoms. Future work collecting longitudinal data (with three time points) is currently ongoing and will be critical for re-evaluating the consistency of corticostriatal associations with age, sex, diagnosis, and symptom severity. Another important limitation worth mentioning is that the amount of fMRI data collected was also limited to ten minutes on average, whereas reliable sampling of individual differences in resting state connectivity (ICC > 0.8) requires functional data acquisitions of over 25 min (Gordon et al., 2017; Lautmann et al., 2015; O’Connor et al., 2017; Cho et al., 2021). Collecting this much data in a young sample can be challenging, though it may be possible that as this sample gets into middle and late adolescence that we are able to acquire more high-quality functional imaging data.

4.6. Conclusions

In conclusion, we found support for our hypotheses that FCSC connectivity would be sensitive to age, sex, ADHD diagnosis and symptom severity, and cognitive task performance. In a large sample of 362 school-aged children we found experimental evidence to support theoretical accounts that ADHD would be reflected in dysregulated FCSC connectivity patterns of the putamen in medication-naïve children with attention deficit hyperactivity disorder. Brain Res. 1303, 195–206. https://doi.org/10.1016/j.brainres.2009.08.029.

Castellanos, F.X., Aoki, Y., 2016. Intrinsinc functional connectivity in attention-deficit/hyperactivity disorder: a science in development. Biol. Psychiatry Cogn. Neurosci. Neuroimaging 1 (3), 252–261. https://doi.org/10.1016/j.bpsc.2016.03.004.

Choi, Y., Chmelis Santiago, J.R., Bixler, K.A., Aalsma, M., Yu, M., Hulvershorn, L.A., 2021. Sex-specific frontal-striatal connectivity differences among adolescents with externalizing disorders. Neuroimage Clin. 32, 102789. https://doi.org/10.1016/j.nicl.2021.102789.

Cho, J.W., Korchmaros, A., Vogelstein, J.T., Milham, M.P., Tannock, R., 2015a. Impact of concatingenating fMRI data on reliability of functional connectomics. Neuroimage 226 (117549), 117549. https://doi.org/10.1016/j.neuroimage.2020.117549.

Geric, R., Nomi, J.S., Uddin, L.Q., Satpute, A.B., 2017. Contextual connectivity: a framework for understanding the intrinsic dynamic architecture of large-scale functional brain networks. Sci. Rep. 7 (1), 6537. https://doi.org/10.1038/s41598-017-06866-w.

Cohen, J., 2013. Statistical Power Analysis for the Behavioral Sciences. Routledge. <http://play.google.com/store/books/details?id=cJuH0R33gBI>.

Cole, W.R., Mostofsky, S.H., Larson, J.C.G., Denckla, M.B., Mahone, E.M., 2008. Age-related changes in motor subtile signs among girls and boys with ADHD. Neurology 71 (19), 1514–1520. https://doi.org/10.1212/01.wnl.0000342745.57344.5f.

Conners, C.K., 2008. Conners 3. [http://www.cognitivecentre.com/wp-content/uploads/Conners3_Brochure_2017_isenoexpression.pdf].

Coye-Duchenne, B., Ehejer, J.L., Gillippe, N.A., Duffy, D.L., Hickie, I.B., Thompson, P.M., Lipnick, N.G., de Zubicaray, G.I., McMahon, K.L., Medland, S.E., Wh Farr, M.J., 2016. Head motion and inattention/hyperactivity share common genetic influences: implications for fMRI studies of ADHD. PloS One 11 (1), e0146271. https://doi.org/10.1371/journal.pone.0146271.

Graddock, C., Sikka, S., Cheung, B., Khamu, R., Ghosh, S.S., Yan, C., Li, Q., Lurie, D., Vogelstein, J., Burns, R., Colcombe, S., Mennes, M., Kelly, C., Di Martino, A., Pekar, J.J., 2016. Head motion and inattention/hyperactivity share common genetic influences: implications for fMRI studies of ADHD. PloS One 11 (1), e0146271. https://doi.org/10.1371/journal.pone.0146271.

Harm, B.G., Delain, A., Jablonka, E., Zilles, K., 2017. Toward automated analysis of connectomes: the configurable pipeline for the analysis of connectomes (C-PAC). In: Proceedings of the Front. Neuroinform. Conference. Frontiers in Neuroinformatics. https://doi.org/10.3389/fninf.2014.00014.

Harm, B.G., Delain, A., Jablonka, E., Zilles, K., 2017. Toward automated analysis of connectomes: the configurable pipeline for the analysis of connectomes (C-PAC). In: Proceedings of the Front. Neuroinform. Conference. Frontiers in Neuroinformatics. https://doi.org/10.3389/fninf.2014.00014.

Harm, B.G., Delain, A., Jablonka, E., Zilles, K., 2017. Toward automated analysis of connectomes: the configurable pipeline for the analysis of connectomes (C-PAC). In: Proceedings of the Front. Neuroinform. Conference. Frontiers in Neuroinformatics. https://doi.org/10.3389/fninf.2014.00014.

Harm, B.G., Delain, A., Jablonka, E., Zilles, K., 2017. Toward automated analysis of connectomes: the configurable pipeline for the analysis of connectomes (C-PAC). In: Proceedings of the Front. Neuroinform. Conference. Frontiers in Neuroinformatics. https://doi.org/10.3389/fninf.2014.00014.
data points to neural network dysregulation in adult ADHD. Hum Brain Mapp 35 (4), 1261–1272. https://doi.org/10.1002/hbm.22500.

Herrington, B.B. & Harrison, B.J. Sensation, and the Cortico-striatal circuit. J. Curr. – W., 2015. Functional connectivity of corticostriatal circuitry and differential response to methylphenidate in youth with attention-deficit/hyperactivity disorder. J. Child Neurol. Psychobi. 139 (2013) 02011.

Dias, T.G.C., Wilson, V.B., Bathula, D.R., Iyer, S.P., Mills, K.L., Thurlow, B.L., Stevens, C.A., Musser, E.D., Carpenter, S.D., Gravett, D.S., Mitchell, S.H., J.G.B., Boyce, A., Goldberg, M.C., Pekar, J.J., Macneil, L.K., Xavier, P., Garvey, M.A., Gilbert, D.L., Ranta, M.E., Denckla, M.B., 2012. Neuropsychiatric Differences Between Boys and Girls with ADHD. J. Child Neurol. 6 (Suppl), S76 –359.

Konrad, K. Eckhoff, S.B., 2010. Is the ADHD brain wired differently? A review on structural and functional connectivity in attention deficit hyperactivity disorder. Hum. Brain Mapp 31 (6), 904–916. https://doi.org/10.1002/hbm.20158.

Taylor, E.R., Gordon, K., T.V., Rosen, E.B., Snyder, A.Z., Schlaggar, B.L., Dosenbach, N.U.F., 2015. Precision functional mapping of attention-deficit/hyperactivity disorder in functional networks, gray matter, and white matter. J. Neurosci. Off. J. Soc. Neurosci. 34 (50), 16555–16566. https://doi.org/10.1523/JNEUROSCI.3156-14.2014.

Kofler, B.G. Rapport, M.D., Serer, D.E., Raiker, J.S., Orban, S.A., Friedman, L.M., Kolomey, E.G., 2013. Reaction time variability in ADHD: a meta-analytic review of 319 studies. Clin. Psychol. Rev. 33 (6), 795–811. https://doi.org/10.1016/j.cpr.2013.06.001.

Menssen, M., Potier, N.K., Kelly, C., Di Martino, A., Martin, C., Castellanos, F., 2013. Shared and Distinct Intrinsic Functional Networks in Children With and Without ADHD Based on Reward Processing and executive meso- and nigrostriatal tracts predict impulsivity differences in children with ADHD. Front. Hum. Neurosci. 7, 101101. https://doi.org/10.3389/fnhum.2021.670325.
