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Yoshifumi Mizuno, M.D., Ph.D, Weidong Cai, Ph.D, Kaustubh Supekar, Ph.D, Kai Makita, Ph.D, Shinichiro Takiguchi, M.D., Ph.D, Timothy J. Silk, Ph.D, Akemi Tomoda, M.D., Ph.D, Vinod Menon, Ph.D

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Yoshifumi Mizuno M.D., Ph.D.\textsuperscript{a,b,c,d,*}, Weidong Cai Ph.D.\textsuperscript{a,f,g}, Kaustubh Supekar Ph.D.\textsuperscript{a,f,g}, Kai Makita Ph.D.\textsuperscript{b,c}, Shinichiro Takiguchi M.D., Ph.D.\textsuperscript{c,d}, Timothy J. Silk Ph.D.\textsuperscript{h,i}, Akemi Tomoda M.D., Ph.D.\textsuperscript{b,c,d}, Vinod Menon Ph.D.\textsuperscript{a,c,d,f,g,*}

\textsuperscript{a}Department of Psychiatry & Behavioral Sciences, Stanford University School of Medicine, Stanford, CA 94304, USA.
\textsuperscript{b}Research Center for Child Mental Development, University of Fukui, Fukui, 910-1193, Japan.
\textsuperscript{c}Division of Developmental Higher Brain Functions, United Graduate School of Child Development, University of Fukui, Fukui, 910-1193, Japan.
\textsuperscript{d}Department of Child and Adolescent Psychological Medicine, University of Fukui Hospital, Fukui, 910-1193, Japan.
\textsuperscript{e}Department of Neurology and Neurological Sciences, Stanford University, Stanford, CA 94304, USA.
\textsuperscript{f}Wu Tsai Neurosciences Institute, Stanford University, Stanford, CA 94304, USA.
\textsuperscript{g}Maternal & Child Health Research Institute, Stanford University, Stanford, CA 94304, USA.
\textsuperscript{h}Centre for Social and Early Emotional Development and School of Psychology, Deakin University, Geelong, VIC, 3125, Australia.
\textsuperscript{i}Murdoch Children's Research Institute, Parkville, VIC, 3052, Australia.

Yoshifumi Mizuno and Weidong Cai contributed equally to this work
Vinod Menon and Akemi Tomoda contributed equally to this work

*Corresponding authors:

Yoshifumi Mizuno, M.D., Ph.D.
Department of Psychiatry & Behavioral Sciences
Stanford University School of Medicine
1070 Arastradero Rd., Palo Alto, CA 94304, USA
Tel: +1-650-736-3699
E-mail: mizunoy@stanford.edu

Vinod Menon, Ph.D.
Department of Psychiatry & Behavioral Sciences
Stanford University School of Medicine
1070 Arastradero Rd., Palo Alto, CA 94304, USA
Tel: +1-650-736-3699
E-mail: menon@stanford.edu
Short title

Methylphenidate effects on neural activity in ADHD

Key words

methylphenidate, attention-deficit/hyperactivity disorder, nucleus accumbens, salience network, default mode network, spontaneous neural activity
Abstract

Background
Methylphenidate, a first-line treatment for attention-deficit/hyperactivity disorder (ADHD), is thought to influence dopaminergic neurotransmission in the nucleus accumbens (NAc), and its associated brain circuitry, but this hypothesis has yet to be systematically tested.

Methods
We conducted a randomized, placebo-controlled double-blind crossover trial with 27 children with ADHD. Children with ADHD were scanned twice with resting-state functional MRI under methylphenidate and placebo conditions, along with assessment of sustained attention. We examined spontaneous neural activity in the NAc and the salience, frontoparietal, and default mode networks, and their links to behavioral changes. Replicability of methylphenidate effects on spontaneous neural activity was examined in a second independent cohort.

Results
Methylphenidate increased spontaneous neural activity in the NAc, and the salience and default mode networks. Methylphenidate-induced changes in spontaneous activity patterns in the default mode network were associated with improvements in intra-individual response variability during a sustained attention task. Critically, despite differences in clinical trial protocols and data acquisition parameters, the NAc, and the salience and default mode networks showed replicable patterns of methylphenidate-induced changes in spontaneous activity across two independent cohorts.

Conclusions
We provide reproducible evidence demonstrating that methylphenidate enhances spontaneous neural activity in NAc and cognitive control networks in children with ADHD, resulting in more
stable sustained attention. Findings identify a novel neural mechanism underlying methylphenidate treatment in ADHD and inform the development of clinically useful biomarkers for evaluating treatment outcomes.

**Trial Registration**

umin.ac.jp/ctr Identifier: UMIN000027533

https://center6.umin.ac.jp/cgi-open-bin/ctr_e/ctr_view.cgi?recptno=R000031550
Introduction

Methylphenidate is a widely used first-line medication for alleviating clinical symptoms of inattention, hyperactivity, and impulsivity in children with attention deficit hyperactivity disorder (ADHD) (1–3). Altered dopamine signaling has been hypothesized to be a key mechanism underlying the therapeutic effects of methylphenidate in ADHD (4). Individuals with ADHD display low dopamine receptor availability in the corticolimbic pathway (5,6) and methylphenidate has been hypothesized to ameliorate ADHD symptoms by increasing extracellular dopamine in the nucleus accumbens (NAc) (7). In neurotypical individuals, dopamine acts as a reinforcer to facilitate motivated behaviors and goal-driven adaptive control (8) via its action on the NAc and cognitive control systems that it regulates (9–12). However, despite decades of its effective use in clinical practice, the precise brain mechanisms underlying the therapeutic effects of methylphenidate are poorly understood as no consistent findings have emerged to date (13). Specifically, the parallel effects of methylphenidate-induced changes on NAc and its interconnected cognitive control networks, and their relation to attentional deficits in childhood ADHD remain unknown.

Dopaminergic pharmacology has been most consistently mapped in the NAc where dopamine receptors and transporters are particularly dense (7,13). Low dopamine receptor density in the NAc has been linked to the severity of inattention symptoms in adults with ADHD (5). At the brain network level, integrative PET-MRI analyses in neurotypical adults have further revealed that mesolimbic dopamine function influences connectivity of the salience network (SN) and the default mode network (DMN) (14). The SN is important for identifying biologically and
cognitively salient events, and guiding attention and goal-directed behaviors (15–18). The SN, together with the frontoparietal network (FPN), and the default mode network (DMN) constitute a triple-network system (15) which plays a crucial role in a wide range of cognitive tasks that require moment-by-moment changes in adaptive cognitive control (15,19–21). Task-based fMRI studies of inhibitory control in children with ADHD have suggested that psychostimulants increase activation in the right insula/inferior frontal cortex (22), a key SN node implicated in inhibitory control (16,17). The SN as a locus of deficits in childhood ADHD has been further bolstered by network connectivity analysis of a Go/NoGo task which identified SN-FPN connectivity as a common locus of deficits in cognitive control and clinical measures of inattention symptoms (23). DMN impairments have also emerged as a prominent feature of ADHD, consistent with theoretical models which have proposed that aberrant engagement of the SN leads to a lack of active suppression and disengagement of the DMN and inattention (24–26). Together, these observations suggest that aberrancies in the NAc together with the SN, FPN and DMN cognitive control networks may underlie the clinical symptoms of ADHD and constitute specific brain targets for remediation using methylphenidate.

Here we use a randomized placebo-controlled double-blind crossover design (Figure S1) to investigate the effect of methylphenidate on spontaneous neural activity in the NAc, as well as the SN, FPN and DMN, and their links to the behavioral effects of medication in children with ADHD. We used amplitude of low-frequency fluctuation (ALFF) to capture the regional intensity of spontaneous fluctuations in fMRI signals (27). Multimodal PET-MRI studies have suggested that spontaneous fluctuations in fMRI signals arise from metabolic demands associated with ongoing fluctuations in synaptic currents and action potential propagation
(28,29). ALFF has been widely used to probe the integrity of brain region-level functioning in psychiatric and neurological disorders (28, 31–33). We used ALFF to test the hypothesis that methylphenidate increases spontaneous neural activity in the NAc, a key node in the dopaminergic reward system, and associated cognitive control circuitry.

A critical unaddressed question is whether methylphenidate-induced changes in spontaneous neural activity are related to remediation of attention and cognitive control deficits. Intra-individual response variability (IIRV), a quantitative measure of trial-wise performance for behavioral instability, is the most consistent robust behavioral phenotype associated with ADHD (33,34), and that psychostimulant treatment reduces this increased variability (33). We recently reported that IIRV in ADHD is associated with poor sustained attention and problems in cognitive control (35). Here, we used a novel similarity metric to measure the extent to which ALFF in the cognitive control network system are similar between children with ADHD and typically developing (TD) children (34). We specifically focused on the triple-network system encompassing the SN, FPN and DMN in relation to behavioral instability (35) based on extensive evidence for their role in attention and cognitive control (17,21,23,26,36). We hypothesized that children with ADHD, whose post-medication spontaneous activity patterns are more similar to TD controls, would exhibit greater improvements in IIRV with medication.

Finally, to address the replication crisis in ADHD (13), we leveraged resting-state fMRI data from a second independent cohort of children with ADHD who participated in a similar randomized controlled trial involving single-dose methylphenidate treatment (37). We test the hypothesis that multivariate pattern analyses (38) would provide convergent evidence for
reproducible findings of methylphenidate-induced changes in spontaneous activity in the NAc and associated cognitive control circuitry, in the primary and secondary cohorts.

Methods and Materials

Participants and Study Design

This study protocol was approved by the Ethics Committee of the University of Fukui, Japan (Assurance no. 20170005). All participants and their parent(s) provided written informed consent for participation in this study. This study is registered with the University Hospital Medical Information Network (UMIN000027533).

Thirty-four children with ADHD and 65 TD children were recruited at the University of Fukui Hospital, Japan. Figure S1 shows the study design (see Supplemental Methods for details). Children with ADHD were scanned twice, in a randomized placebo-controlled double-blind crossover design. The administration order was counterbalanced across participants to address potential test-retest issues. During the first visit, they were administered osmotic release oral system methylphenidate (OROS-MPH) (1.0 ± 0.1mg/kg) or placebo (lactose) under double-blind conditions as previous studies (39–41). Five to eight hours after administration, when the methylphenidate concentration in the blood is maximal (42), they underwent a resting-state functional MRI (fMRI) scan and performed a standardized continuous performance task (CPT) (43,44) outside the MRI scanner.
During the second visit, within 1 to 6 weeks after the first visit, children with ADHD underwent a resting-state fMRI scan, and performed the CPT after they took the second medicine: children who took OROS-MPH at the first visit took the placebo at the second visit under double-blind conditions, and vice versa. The OROS-MPH and the placebo condition are referred to as ADHD-MPH and ADHD-Placebo, respectively, in this study.

TD children completed the same resting-state fMRI scan once without either OROS-MPH or placebo. The following inclusion criteria were used for both groups: no contraindications for magnetic resonance imaging (MRI), full-scale intelligence quotient (FSIQ) > 70 (to exclude participants with intellectual disability), no history of severe head trauma or neurological abnormalities (e.g. epilepsy, arachnoid cysts). To minimize the potential impact of sex differences, we included only male participants, consistent with previous ADHD imaging studies (37,45–48). Participants with excessive head motion (over 3.0 mm, 3.0 degrees, and mean framewise displacement (FD) 0.3 mm) during the scanning were excluded (45). Seven children with ADHD were excluded because of refusal to participate, arachnoid cysts, and motion during the MRI, while 16 TD controls were excluded because of psychiatric disorders, and neurological abnormalities, leading to a final sample of 27 children with ADHD (age: 10.6±1.8 years, range 7.3-15.5 years) and 49 TD controls (age: 11.1±2.3 years, range 6.1-15.6 years) (Table S1). Nine patients with ADHD had autism spectrum disorder, 6 ADHD patients had oppositional defiant disorder, 2 had specific learning disorder, and 1 had developmental coordination disorder as comorbid disorders. While one of the patients with ADHD was medication-naïve, 25 were medicated with OROS-MPH (medication period was 22.2±15.3 months, range 1-58 months), three with atomoxetine, and two with aripiprazole. Children with ADHD took their regularly
prescribed medications between the two visits, but all participants were medication-free prior to MRI for at least 5 times half-life, including methylphenidate and atomoxetine, consistent with protocols from previous studies (45,49).

**Assessment of attention and cognitive control**

A standardized CPT (43,44) was administered to children with ADHD outside the MRI scanner under both methylphenidate and placebo conditions. The task consisted of a Go/NoGo paradigm in which children were presented with either a target or non-target stimulus on the screen for 100 msec, once every 2 seconds for 15 minutes across three 5-minute blocks. The target stimulus was a triangle, while the non-target stimulus was either a circle or a square. Children were required to press a button when a target stimulus was presented, and withhold response to non-targets. The test has been normed to age-adjusted T-scores on four distinct performance measures: omission errors, commission errors, mean response time (RT), and IIRV, which were quantified using RT standard deviation (43,44). We examined medication-induced performance differences using paired t-tests.

**fMRI data acquisition**

Functional images were acquired with a T2*-weighted gradient-echo echo-planar imaging (EPI) sequence via a 3-T scanner (Discovery MR 750; General Electric Medical Systems, Milwaukee, WI) and a 32-channel head coil. In total, 201 volumes were acquired for a scanning time of 7 minutes 42 seconds. Each volume consisted of 40 slices, with a thickness of 3.5 mm and a 0.5-mm gap. The time interval between each successive acquisition of the same slice (repetition
time, TR) was 2300 ms, with an echo time (TE) of 30 ms, and a flip angle (FA) of 81°. The field of view (FOV) was 192 × 192 mm, and the matrix size was 64 × 64, yielding volume dimensions of 3 × 3 mm. The participants were instructed to stay awake with their eyes closed.

fMRI data pre-processing

Resting state fMRI data were analyzed using SPM12 and DPARSF (50). First, the initial 10 volumes were discarded, and slice-timing correction was performed. The signal from each slice was realigned temporally to that obtained from the middle slice using sinc interpolation. The re-sliced volumes were normalized to the Montreal Neurological Institute space with a voxel size of 2 × 2 × 2 mm using the EPI template provided by SPM12. The normalized images were spatially smoothed with a 6-mm Gaussian kernel. Next, the non-neural noise in the time series was controlled, and several sources of spurious variance (e.g., the Friston 24-parameter model) were removed from the data through linear regression.

fMRI data analysis

Our overall analysis is illustrated in Figure 1A and summarized below (see Supplemental Methods for details).

Brain regions and networks of interest. We focused on the NAc, a key node in the reward pathway, and the SN, DMN, and FPN, three core brain systems involved in cognitive control. Probabilistic masks of the bilateral NAc were obtained from an independent high-resolution structural study, and the masks were thresholded at 0.9 (51). The SN, DMN, left FPN, and right FPN maps were obtained from a previous study (24). To test the robustness of our findings, we
applied independent component analysis to generate another set of network masks for SN, DMN, left FPN, and right FPN, using the analytic approach used in our previous study (25).

ALFF analysis. We assessed spontaneous neural activity by computing ALFF in bilateral NAc, SN, DMN, left FPN, and right FPN. Paired t-tests were used to examine the medication effects (ADHD-MPH versus ADHD-Placebo) and two-sample t-tests were used to examine the difference between ADHD and TD controls.

ALFF pattern similarity analysis. We evaluated the extent to which ALFF values are similar between children with ADHD and TD children in the SN, DMN, and FPN. We then determined how ALFF similarity is modulated by medication, and determined its relation with medication-induced changes in behavior. We computed an ALFF similarity metric (52) (Figure 1B) using z-transformed Pearson’s correlation between ALFF values within each brain network (SN, DMN, or FPN) from each child with ADHD and those from the averaged ALFF map in the TD group. This metric captures the similarity of ALFF patterns in each child with ADHD with respect to the expected patterns in the TD group, in each brain region or network of interest. A higher ALFF similarity value indicates that the child with ADHD has a more TD-like ALFF spatial pattern of ALFF. Medication effect was calculated by subtracting z-transformed correlation coefficients in ADHD-Placebo from ADHD-MPH conditions. A positive value indicates that medication leads to a more TD-like ALFF spatial pattern. We tested whether medication effects on the ALFF patterns are associated with a behavioral measure of attention and cognitive control, the IIRV, using Pearson’s correlation.
Replication of methylphenidate effects on spontaneous neural activity patterns using multivariate analysis

Finally, we evaluated the replicability of methylphenidate effects on spontaneous neural activity patterns using a second independent cohort of children with ADHD who participated in a similar randomized controlled study involving single-dose methylphenidate treatment. Details of participants and study design are reported elsewhere (37) and summarized in the Supplemental Methods and Table S3. To overcome the limitations of small sample size in the secondary cohort (N=15), we used a multivariate pattern analysis strategy which facilitates greater reproducibility in comparison to univariate voxel-wise measures (38). Specifically, we sought to determine whether methylphenidate would modulate multivariate patterns of ALFF activity in the NAc, SN, DMN, and FPN.

Results

Methylphenidate improves attention and cognitive control function

Methylphenidate significantly reduced omission errors, mean RT, and IIRV in the CPT in children with ADHD (all ps < 0.001) (see Supplemental Results and Figure S2 for details).

Methylphenidate effects on spontaneous neural activity in NAc
ALFF in the right NAc in ADHD-MPH was significantly higher than ADHD-Placebo condition ($p < 0.05$, Bonferroni corrected Cohen's $d = 0.55$) (Figure 2A) (see Supplemental Results and Figure S3A for comparisons with TD controls). These results suggest that methylphenidate enhances spontaneous neural activity in the right NAc.

**Methylphenidate effects on spontaneous neural activity in SN, FPN, and DMN**

ALFF in the SN and DMN in ADHD-MPH were significantly higher than ADHD-Placebo condition (SN, $p < 0.05$, Bonferroni corrected, Cohen's $d = 0.57$; DMN, $p < 0.01$, Bonferroni corrected, Cohen's $d = 0.66$). There was no significant difference in the left and right FPN ($p > 0.05$) (Figure 2B) (see Supplemental Results and Figure S3B for comparisons with TD controls). Results were replicated using alternate SN, DMN, left FPN, and right FPN masks (see Supplemental Results and Figure S4 for details). These results suggest that methylphenidate enhances spontaneous neural activity in the SN and DMN.

**Relationship between methylphenidate-induced changes in spontaneous neural activity and changes in response variability**

We focused on the SN and DMN as these two networks showed significant effects of medication on the mean ALFF. We found that medication-induced changes in IIRV were significantly correlated with medication-induced changes in spontaneous activity patterns in the DMN ($r = -0.46$, $p < 0.05$, Bonferroni corrected, Figure 3), but not in the SN ($r = -0.34$, $p = 0.080$).
Additional analysis confirmed that the relationship between changes in IIRV and changes in DMN ALFF was robust against several potential confounds (Table 1). Results were replicated using an alternate DMN mask (see Supplemental Results, Figure S5, and Table S2 for details). These results suggest that greater similarity with TD-like ALFF patterns in the DMN post-medication is associated with more stable behavioral performance in children with ADHD.

**Replication of methylphenidate effects on spontaneous neural activity patterns**

Multivariate classification analysis revealed that ALFF differentiated ADHD-MPH and ADHD-Placebo conditions in the primary cohort in right NAc (accuracy = 70%, \( p = 0.02 \)), SN (accuracy = 74%, \( p = 0.002 \)), and DMN (accuracy = 82%, \( p = 0.002 \)). A similar differentiation was observed in the replication cohort: right NAc (accuracy = 87%, \( p = 0.002 \)), SN (accuracy = 73%, \( p = 0.002 \)), and DMN (accuracy = 73%, \( p = 0.002 \)) (Figure 4, Supplemental Tables S4). These analyses demonstrate the robustness of our key findings related to methylphenidate-induced changes in spontaneous neural activity patterns in the NAc, SN, and DMN across two independent cohorts.

**Discussion**

We examined whether methylphenidate alters spontaneous neural activity in the mesolimbic dopaminergic system and cognitive control networks, and how these alterations impact cognitive flexibility in children with ADHD. Using a randomized placebo-controlled double-blind crossover design, with sample sizes larger than extant randomized controlled studies (13), we
show that methylphenidate alters spontaneous activity in the NAc as well as the SN and DMN, two large-scale cognitive control networks implicated in attention and cognitive control deficits in ADHD. Importantly, methylphenidate-induced changes in spontaneous activity patterns in the DMN were associated with improvements in intra-individual response variability during a sustained attention task. Finally, in advance over previous studies, we discovered that methylphenidate alters spontaneous neural activity patterns in the NAc, SN and DMN and demonstrated replication across two independent cohorts of children with ADHD. Together, these findings identify a novel neural mechanism underlying methylphenidate treatment in ADHD.

**Methylphenidate modulates spontaneous activity in the nucleus accumbens**

Prominent theories of ADHD have emphasized deficits in the reward and motivation system (53–56). This hypothesis is supported by behavioral findings of aberrant delay discounting, i.e. preference of a small immediate reward over a large delayed reward in children with ADHD, and abnormal activation in regions of dopamine reward circuitry during anticipation or processing of rewards in children with ADHD (57,58). As a key node of the dopaminergic reward pathway, the NAc plays an important role in these processes (59,60).

In the present study, we first examined whether methylphenidate alters the spontaneous neural activity of the NAc. We found that, compared to placebo, methylphenidate increased spontaneous activity in the NAc in children with ADHD. Our results converge with findings from PET studies which have reported methylphenidate-induced dopamine increases in the
ventral striatum in adults with ADHD (61). Due to the use of radioactive ligands, PET imaging studies cannot be conducted in children. This is an impediment to investigations of methylphenidate-induced dopamine changes in children with ADHD at ages closer to clinical diagnosis, but ALFF measures may offer a useful alternative. In contrast to PET, results with fMRI provide greater anatomical precision and localize methylphenidate-induced effects specifically to the NAc within the ventral striatum. In line with our results, methylphenidate has been reported to increase spontaneous activity in rodent NAc (62) and a recent study in non-human primates found that the therapeutic effect of methylphenidate on impulsive decision is associated with the pharmacological action on the dopamine transporter in the NAc (63). Similarly, in both children and adults with ADHD, methylphenidate has been reported to modify abnormal striatal activity during reward processing (57,64–66). Together, these findings demonstrate that methylphenidate has a strong effect on spontaneous neural activity in the NAc, a key node in the mesolimbic reward pathway and that the ALFF might be a useful proxy measure to probe methylphenidate effects in children with ADHD.

**Methylphenidate modulates spontaneous activity in the salience and default mode networks**

Next, we examined the parallel effects of methylphenidate-induced changes on the SN, FPN, and DMN, three large-scale cognitive control networks implicated in ADHD and in attention and cognitive control more broadly (13,16,26,35,52). We found that methylphenidate also increases spontaneous activity in the SN and DMN. Key nodes of the SN, including the anterior insula and anterior cingulate cortex, are among the most highly activated regions in a variety of attention
and cognitive control tasks (16,67). Weak activation in the anterior insula and anterior cingulate cortex during cognitive control, especially on error trials, has been reported in children with ADHD (23). Increased attention and cognitive control demand is also accompanied with deactivation in the DMN (21,68,69), and abnormal DMN activity during cognitively demanding tasks is a reproducible feature of ADHD (70). In adults with ADHD, methylphenidate has been shown to increase intrinsic functional connectivity within DMN regions (71), and enhance deactivation of the DMN regions during attentional tasks (72). Our results extend these findings and suggest that one mechanism by which methylphenidate alters cognitive control function is by enhancing spontaneous activity in both the SN and DMN in children with ADHD.

**Methylphenidate improves behavioral performance by modulating spontaneous activity in the DMN**

Cognitive control dysfunction is a prominent feature of ADHD and we recently showed that inattention is correlated with IIRV (35), a key intermediate phenotype of childhood ADHD (73). Several studies have shown that, compared to controls, children with ADHD display increased IIRV during cognitive task performance (33,34). We used a novel multivariate pattern similarity measure (52) to determine whether children with ADHD whose spontaneous activity patterns are more similar to TD controls after methylphenidate treatment would exhibit greater improvements in IIRV with medication. Our analysis revealed that higher similarity of ALFF patterns in DMN between children with ADHD and TD controls was associated with a greater reduction in IIRV in children with ADHD. Previous task-based fMRI studies have reported that activity in DMN during cognitive performance was related to IIRV and psychostimulants alter the DMN activity
in youth with ADHD (74,75). Our results suggest that the alteration in spontaneous activity in the DMN is a plausible mechanism by which methylphenidate alleviates cognitive inflexibility in children with ADHD. Our results also highlight the specificity of the DMN in terms of its unique association with the effects of medication on IIRV, and provide novel evidence that methylphenidate actions on the DMN contribute to remediation of core attention and cognitive control deficits in ADHD (75,76).

**Methylphenidate modulates multivariate spontaneous neural activity patterns:**

**Replication across two independent cohorts**

Lack of converging evidence across independent studies is a challenge in clinical neuroscience research, especially in the domain of pharmacological interventions (13). To address this challenge, we sought to replicate key findings in a second cohort of participants from a previously published study (37). Because of the small sample size in the replication cohort (N=15), we used a multivariate pattern analysis approach which has been shown to yield more replicable results than univariate methods (38). Using such an approach, here we report the unprecedented replication of findings in two neuroimaging clinical trial cohorts acquired independently. Our analyses revealed that multivariate patterns of spontaneous activity in the NAc as well as in the SN, DMN are modulated by methylphenidate in children with ADHD in both the primary and secondary cohorts. To the best of our knowledge (13), our replication is the first of its kind and provides confirmatory evidence that methylphenidate alters spontaneous neural activity in key reward and cognitive control systems implicated in childhood ADHD.
Limitations and future work

One limitation of the present study is that fMRI measures cannot establish direct links to changes in dopamine. Future work with hybrid PET-MRI techniques that enable multimodal imaging of different neurotransmitter systems (77) is needed to investigate the impact of methylphenidate on dopamine as well as other neurotransmitters such as norepinephrine and their relation to spontaneous fluctuations in fMRI signals. Because PET-fMRI studies can only be conducted in adults due to the use of radioactive ligands, the characterization of methylphenidate effects on different neurotransmitter systems in children remains a challenge. As with extant ADHD brain imaging studies, children with ADHD in our study were not drug naïve, were male, and spanned a wide range from 5 to 16. Larger multi-cohort studies that include drug naïve males and females with ADHD are needed to determine how medication history, sex, and development stage modulate methylphenidate effects and to further assess the robustness of the effects reported here.

Conclusion

Our randomized, placebo-controlled double-blind crossover study revealed that methylphenidate increases spontaneous brain activity in the reward system at the regional level, and in the salience and default mode networks at the network level. Using a novel ALFF similarity metric, we show that the methylphenidate effect on spontaneous activity patterns in the default mode network is associated with the effect of medication on intra-individual response variability.
Strikingly, multivariate analysis demonstrated replicable patterns of methylphenidate-induced changes in spontaneous activity patterns in two independent cohorts of children with ADHD. Our findings advance the current understanding of the neurobiological mechanisms underlying methylphenidate treatment in children with ADHD and may lead to clinically useful biomarkers for evaluating treatment response. Finally, our study provides a template for investigations of the effects of methylphenidate on task-related neural activity in striatal reward and related cognitive control circuitry in children with ADHD.
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Disclosures

The authors report no biomedical financial interests or potential conflicts of interest.

Author contributions statement

Y.M. and A.T conceived and designed the experiments; W.C. and V.M. designed the data analysis strategy; Y.M., K.M, S.T. conducted the experiments; T.S. provided the replication cohort data; Y.M. and W.C. analyzed the data; Y.M., W.C., K.S., A.T., and V.M. wrote the manuscript; all authors edited the manuscript.
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42. *Concerta® Tablets (Methylphenidate Hydrochloride), Common Technical Document in Japan(October 26 2007, CTD2.7.6.8)* (2007):
Figure legends

**Figure 1. (A) Data analysis pipeline.** We first computed ALFF within the bilateral NAc and three brain networks implicated in ADHD: SN, DMN, and left and right FPN. Paired t-tests were used to examine the medication effects (ADHD in methylphenidate versus placebo conditions) and two-sample t-tests were used to examine the difference between ADHD and TD controls. Second, we conducted ALFF pattern similarity analysis (illustrated in detail in Panel B) to quantify the extent to which ALFF values are similar between children with ADHD and TD children, and examined whether children with ADHD whose post-medication spontaneous activity patterns are more similar to TD children would exhibit greater improvement in IIRV with medication. Third, we used classification analysis to test whether the multivariate pattern of ALFF in the NAc and the three brain networks could distinguish children with ADHD in medication or placebo conditions (primary cohort) and crucially whether this can be replicated in another independent dataset (replication cohort). (B) **Overview of ALFF pattern similarity analysis between children with ADHD and TD controls.** We first computed the correlation between ALFF values within SN or DMN from each child with ADHD and those from the mean ALFF map in the TD group. The correlation coefficient was standardized using Fisher’s r-to-z transformation. Next, we calculated methylphenidate-induced changes in the similarity measures of ALFF in the SN or DMN between ADHD-Placebo and ADHD-MPH conditions. Higher values indicate that medication leads to more TD-like spontaneous neural activity patterns. ADHD-MPH: children with attention-deficit/hyperactivity disorder under methylphenidate administration; ADHD-Placebo: children with attention-deficit/hyperactivity disorder under placebo; ALFF: amplitude of low-frequency fluctuations; DMN: default mode network; IIRV: intra-individual response variability; LFPN: left frontoparietal network; NAc: nucleus
accumbens; RFPN: right frontoparietal network; ROI: region of interest; SN: salience network; TD: typically developing.

**Figure 2.** Methylphenidate modulates spontaneous neural activity in the nucleus accumbens (NAc) and cognitive control networks. (A) Methylphenidate increases ALFF in the right NAc ($p < 0.05$, Bonferroni corrected, Cohen's $d = 0.43$), but not in the left NAc. (B) Methylphenidate increases ALFF in the default mode network (DMN) ($p < 0.01$, Bonferroni corrected, Cohen's $d = 0.66$) and salience network (SN) ($p < 0.05$, Cohen's $d = 0.57$), but not in the left and right frontoparietal network (FPN). ADHD-MPH: children with attention-deficit/hyperactivity disorder under methylphenidate administration; ADHD-Placebo: children with attention-deficit/hyperactivity disorder under placebo; ALFF: amplitude of low-frequency fluctuation. **$p < 0.01$; *$p < 0.05$; n.s., not significant.

**Figure 3.** Methylphenidate modulation of spontaneous neural activity in the default mode network (DMN) predicts the medication effect on intra-individual response variability (IIRV) ($r = -0.46$, $p = 0.016$). MPH: methylphenidate.

**Figure 4.** Methylphenidate modulates spontaneous neural activity in the nucleus accumbens (NAc), salience network (SN), default mode network (DMN), and frontoparietal network (FPN) in children with ADHD: Replicable evidence from multivariate
classification analyses of primary and replication cohorts. In both the primary and replication cohorts, multivariate patterns of ALFF in the right NAc, SN, DMN, and right FPN (RFPN) distinguish ADHD-MPH from ADHD-Placebo conditions. Statistical significance of classification accuracy was estimated using permutation tests. ADHD-MPH: children with attention-deficit/hyperactivity disorder under methylphenidate treatment; ADHD-Placebo: children with attention-deficit/hyperactivity disorder under placebo; LFPN: left frontoparietal network; **p < 0.01; *p < 0.05.
Table 1. Multiple linear regression analysis revealed that only methylphenidate modulation of ALFF similarity pattern within DMN is significantly associated with medication effects on IIRV.

<table>
<thead>
<tr>
<th>Methylphenidate effects on ALFF similarity pattern within DMN</th>
<th>β</th>
<th>t</th>
<th>p</th>
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<tbody>
<tr>
<td>Methylphenidate induced difference in IIRV</td>
<td></td>
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</tr>
<tr>
<td>DMN</td>
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</tbody>
</table>

ALFF: amplitude of low-frequency fluctuation; DMN: default mode network; FSIQ: full scale intelligence quotient; IIRV: intra-individual response variability. *p < 0.05.
A. Data analysis pipeline

ALFF analysis

Classification analysis
in primary and replication cohorts

Similarity analysis

Brain-behavior relation

B. Similarity analysis

Correlation

Similarity of ADHD-MPH with TD

Correlation

Similarity of ADHD-Placebo with TD

MPH effects on ALFF similarity pattern
A. ALFF in left and right nucleus accumbens (NAc)

![Graph showing ALFF in left and right NAc for ADHD, MPH, and Placebo groups.](image)

B. ALFF in salience network (SN), default mode network (DMN), left frontoparietal network (LFPN), and right FPN (RFPN)

![Graphs showing ALFF in SN, DMN, LFPN, and RFPN for ADHD, MPH, and Placebo groups.](image)
MPH effects on DMN spontaneous activity

MPH effects on IIRV