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# Neuroimaging-Aided Prediction of the Effect of Methylphenidate in Children with Attention-Deficit Hyperactivity Disorder: A Randomized Controlled Trial

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Although methylphenidate hydrochloride (MPH) is a first-line treatment for children with attention-deficit hyperactivity disorder (ADHD), the non-response rate is 30%. Our aim was to develop a supplementary neuroimaging biomarker for predicting the clinical effect of continuous MPH administration by using near-infrared spectroscopy (NIRS). After baseline assessment, we performed a double-blind, placebo-controlled, crossover trial with a single dose of MPH, followed by a prospective 4-to-8-week open trial with continuous MPH administration, and an ancillary I-year follow-up. Twenty-two drug-naïve and eight previously treated children with ADHD (NAÏVE and NON-NAÏVE) were compared with 20 healthy controls (HCs) who underwent multiple NIRS measurements without intervention. We tested whether NIRS signals at the baseline assessment or  $\Delta$ NIRS (single dose of MPH minus baseline assessment) predict the Clinical Global Impressions-Severity (CGI-S) score after 4-to-8-week or I-year MPH administration. The secondary outcomes were the effect of MPH on NIRS signals after single-dose, 4-to-8-week, and I-year administration. ΔNIRS significantly predicted CGI-S after 4-to-8-week MPH administration. The leave-one-out classification algorithm had 81% accuracy using the NIRS signal.  $\Delta$ NIRS also significantly predicted CGI-S scores after I year of MPH administration. For secondary analyses, NAÏVE exhibited significantly lower prefrontal activation than HCs at the baseline assessment, whereas NON-NAÏVE and HCs showed similar activation. A single dose of MPH significantly increased activation compared with the placebo in NAÏVE. After 4-to-8-week administration, and even after MPH washout following I-year administration, NAÏVE demonstrated normalized prefrontal activation. Supplementary NIRS measurements may serve as an objective biomarker for clinical decisions and monitoring concerning continuous MPH treatment in children with ADHD. Neuropsychopharmacology (2015) 40, 2676-2685; doi:10.1038/npp.2015.128; published online 27 May 2015

#### INTRODUCTION

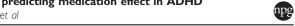
Individualized medicine (Erder *et al*, 2012) using biomarkers such as neuroimaging ('third-generation' or 'next-generation' neuroimaging) is increasingly important in psychiatry (Borgwardt and Fusar-Poli, 2012). The psychostimulant methylphenidate hydrochloride (MPH) is an effective first-line treatment for children with attention-deficit hyperactivity

disorder (ADHD) (Buitelaar and Medori, 2010). However, ~30% of children with ADHD do not respond to MPH (Wilens *et al*, 2008) and experience no benefit or only adverse side effects (Cho *et al*, 2007; Rapport *et al*, 1994). Moreover, although positron emission tomography studies of MPH exposure suggest that catecholamine dysfunction at least partially mediates the behavioral and cognitive features of ADHD (Hannestad *et al*, 2010; Krause, 2008; Volkow *et al*, 2002), how MPH affects the nervous system remains unclear, especially over long periods.

Several studies have attempted to predict MPH efficacy using clinical characteristics including age (Buitelaar *et al*, 1995; Zeiner *et al*, 1999), IQ (Buitelaar *et al*, 1995), symptoms, disorder severity (Buitelaar *et al*, 1995; Zeiner *et al*, 1999), and

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**Table I** Demographics of the NAÏVE, NON-NAÏVE, and Healthy Control Subjects at Baseline Assessment

	NAÏVE (N = 22)	NON-NAÏVE (N=8)	HCs (N = 20)	P-value	Post hoc P-value		
					HCs: NAÏVE	HCs: NON- NAÏVE	NAÏVE: NON-NAÏVE
Age, mean (SD), year	8.6 (1.4)	9.3(1.8)	8.1 (1.6)	0.1871			
Sex, n (%)	M 18 (81.8) F 4 (18.2)	M 8 (100) F 0 (0)	M 14 (70) F 6 (30)	0.1924			
IQ, mean (SD)	100.1 (16.0)	105.0 (15.8)	107.0 (15.7)	0.3753			
SST performance, mean (SD), %	61.8 (15.7)	72.5 (8.9)	68.0 (10.6)	0.1025			
MRT (SST), mean (SD), ms	545.0 (110.4)	571.4 (88.1)	568.8 (67.7)	0.6464			
ADHD-RS-IV all, mean (SD)	31.5 (10.9)	30.5 (7.8)	6.9 (5.9)	< 0.0001	< 0.0001	< 0.0001	0.9624
ADHD-RS-IV inattention, mean (SD)	18.8 (6.2)	18.8 (4.4)	5.2 (5.2)	< 0.0001	< 0.0001	< 0.0001	0.9999
ADHD-RS-IV hyperactivity, mean (SD)	12.7 (6.1)	11.8 (4.7)	1.7 (1.8)	< 0.0001	< 0.0001	< 0.0001	0.8785
Handedness, n (%)	Right 22 (100)/Left 0 (0)	Right 6 (75)/Left 2 (25)	Right 18 (90)/ Left 2 (10)	0.0757			
Subtype, n (%)	inattention, 4 (18.2)/ combined, 18 (81.8)	inattention, I (12.5)/ combined, 7 (87.5)					

Abbreviations: ADHD-RS-IV, attention-deficit hyperactivity disorder rating scale IV; HCs, healthy controls; IQ, intelligence quotient; MRT, mean reaction time; NAÏVE, ADHD-drug naïve; NON-NAÏVE, ADHD patients who had taken methylphenidate for several months; SST, stop signal task.

neuropsychological test scores (Lee *et al*, 2009). Recently, more objective biomedical predictors (Hermens *et al*, 2006) of brain function (An *et al*, 2013; Cho *et al*, 2007; Ilgin *et al*, 2001; la Fougere *et al*, 2006; Sangal and Sangal, 2004; Schweitzer *et al*, 2003) have been used (see also Supplementary Materials). Most neuroimaging MPH trials have included previously medicated subjects. However, recent studies have revealed differences in brain structure between drug-naïve and previously treated ADHD patients (Frodl and Skokauskas, 2012; Shaw *et al*, 2009).

Here, we primarily aimed to develop an objective marker using near-infrared spectroscopy (NIRS) that predicts the efficacy of continuous MPH administration for children with ADHD. NIRS imaging is safe, portable, and allows easy examination of subjects in a natural sitting position (Ishii-Takahashi et al, 2013; Monden et al, 2012; Takizawa et al, 2008). To elicit prefrontal activation, we used the stop signal task (SST), an inhibitory task that reflects the pathophysiology of ADHD (Hart et al, 2013; Rubia et al, 2014). The single-dose MPH administration phase used a randomized, double-blind, placebo-controlled, crossover design, and the subsequent outcome was assessed in a 4-to-8-week open trial and 1-year follow-up. To our knowledge, no previous study has followed patients for 1 year to predict the effect of MPH using neuroimaging. We hypothesized that prefrontal activation at baseline assessment, or the difference in NIRS signal between a single administration of MPH and the baseline assessment, would predict clinical improvement after mid-term (4-to-8-week) and long-term (1-year) MPH treatment in drug-naïve patients with ADHD. Notably, we divided children with ADHD into those who had not received any medication (drug naïve) and those who had received chronic treatment with MPH (non-naïve) because the NIRS signals at baseline may be altered by previous MPH

Our secondary purpose was to test whether multiple NIRS measurements are useful in monitoring prefrontal activation

during continuous MPH treatment in children with ADHD. For this purpose, we recruited healthy controls (HCs) and monitored their NIRS signals at multiple times without any intervention. Then, we tested the effect of MPH treatment on the NIRS signals after single-dose, mid-term, and long-term administration in ADHD children.

### MATERIALS AND METHODS

### **Participants**

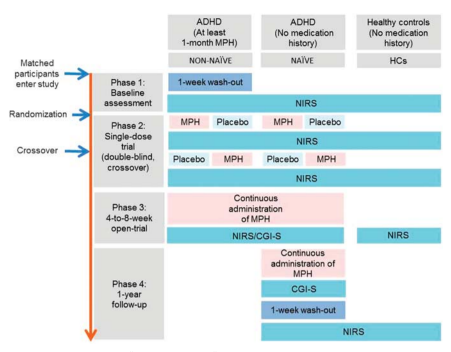
We examined 30 children with ADHD (mean age:  $8.6\pm1.4$  years) and 20 age-, sex-, and IQ-matched HCs (Table 1), all recruited from an outpatient unit at the University of Tokyo Hospital, who consisted of two groups: drug-naïve ADHD (NAÏVE) and ADHD patients taking MPH for at least 1 month (NON-NAÏVE; Figures 1 and 2). HCs underwent multiple neuroimaging measurements without any clinical intervention.

Child psychiatrists diagnosed ADHD through comprehensive clinical assessments and in accordance with the DSM-IV. Current and lifetime DSM-IV diagnoses were determined by a consensus decision based on the results of independent clinical interviews, clinical data, and the Mini-International Neuropsychiatric Interview for Children and Adolescents (Otsubo *et al*, 2005). Additional inclusion criteria were age (6–12 years) and IQ (above 70) determined with the Wechsler Children Intelligence Scale, 3rd edition. The exclusion criteria are described in Supplementary Materials.

The ADHD Rating Scale IV (ADHD-RS-IV) was used to assess ADHD symptoms (DuPaul *et al*, 1998). Problematic behaviors were measured from parent reports using the Child Behavior Checklist (CBCL; Achenbach *et al*, 1991).

#### Trial Design

After baseline assessment before MPH administration (Phase 1), we conducted a double-blind, placebo-controlled,



**Figure I** Schematic diagram of the trial protocol. NAÏVE and NON-NAÏVE underwent baseline assessment (Phase I), a single-dose trial (Phase 2), and a 4-to-8-week open trial (Phase 3). NAÏVE participants also underwent a I-year follow-up (Phase 4). NAÏVE was evaluated by NIRS in five sessions (Phase I: I session, Phase 2: 2 sessions, Phase 3: I session, and Phase 4: I session). NON-NAÏVE was evaluated by NIRS signal in four sessions (Phase I: I session, Phase 2: 2 sessions, and Phase 3: I session). We evaluated the NIRS signal of HCs, who did not take MPH, at the same intervals and times as NAÏVE to detect the effects of the repeated measurement. ADHD, attention-deficit hyperactivity disorder; HCs, healthy controls; MPH, methylphenidate hydrochloride; NAÏVE, ADHD-drug NAÏVE; NIRS, near-infrared spectroscopy; NON-NAÏVE, ADHD patients who had taken methylphenidate for several months.

crossover, prospective trial of single-dose MPH administration (Phase 2), with a subsequent 4-to-8-week open trial (Phase 3; Figure 1). We then conducted a 1-year follow-up (Phase 4) as an ancillary study. Prefrontal hemodynamics were measured in NAÏVE and NON-NAÏVE using NIRS at Phase 1, Phase 2 (two times after a single dose of MPH or placebo), and Phase 3. NAÏVE was also assessed at Phase 4. We evaluated the NIRS signals of HCs at the same time points used for NAÏVE (Figure 1). For details about the repeated measurements in HCs, see Supplementary Materials.

The target sample size for this study was ~20 children (NAÏVE) for the primary outcome, after accounting for dropout cases. We estimated the sample size based on an optimal design for functional brain imaging (Carter *et al*, 2008).

All children and parents provided written informed consent after receiving a complete explanation of the study, according to the Declaration of Helsinki. The institutional review board (IRB) and ethical committee of the University of Tokyo Hospital approved this study (approval number: P2007009 and 630-8). This trial is registered as number UMIN000001270.

### **Clinical Interventions**

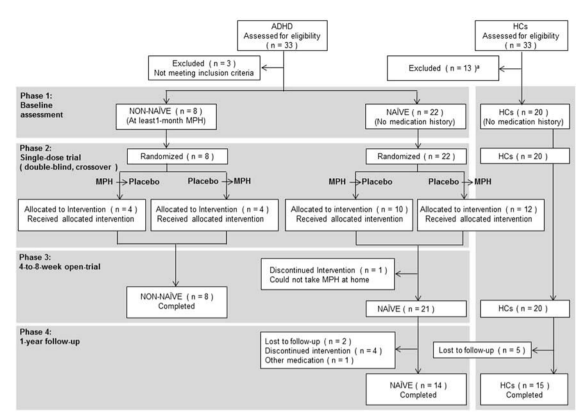
A schematic diagram of the trial protocol is shown in Figure 1. In Phase 1, NON-NAÏVE was instructed to stop taking MPH for 1 week before the single-dose trial began.

In Phase 2, placebo or MPH (18 mg; MPH osmotic-release oral system (OROS), Concerta) was administered at 1-week

intervals. Subjects received a blinded capsule (placebo or MPH) in the front of the investigator (AI-T) 5 h before the NIRS session.

In Phase 3, the optimal dose was administered for 4 weeks. After a 1-week washout period, the NON-NAÏVE group was administered the same amount of MPH that they were taking before participating in the trial. Because the NON-NAÏVE group did not experience a titration period, the length of their open-label trial was 4 weeks. In the NAÏVE group, the dose was determined from the results of the controlled medication trial described below. The ADHD-RS-IV and side-effect rating scale were assessed weekly. A psychiatrist reviewed the data each week to determine the best MPH dose. MPH was titrated to an optimal response from an initial dose of 18 mg. During titration, the dosage was increased by 9 mg at weekly visits. When side effects occurred, the dose was reduced to a level where no side effects occurred, which then was considered the optimal dose. After the titration period, the NAÏVE group was administered the optimal dose for 4 weeks. After titration, the optimal dose for each patient was determined (mean dose: 25.4 mg, 0.91 mg/kg; SD: 5.9; median: 27). Participants continued receiving the optimal dose of MPH for 4 weeks (mean duration including titration: 5.45 weeks; SD: 1.29; median: 5).

Phase 4 was a 1-year follow-up as an ancillary study. We followed NAÏVE participants who continued MPH administration for at least 1-year (mean duration: 14.7 months; SD: 2.3; median: 14.5) after the 4-to-8-week open trial. The same child psychiatrist who evaluated the children in the baseline assessment evaluated the severity and clinical response after 1-year-MPH administration using the Clinical Global



**Figure 2** Trial profile. For the primary outcome, only NAÏVE (N=21) was assessed. For the secondary outcome, NAÏVE (N=22) and NON-NAÏVE (N=8) groups were evaluated for the effect of the single-dose trial. NAÏVE (N=21), NON-NAÏVE (N=8), and HCs (N=20) were evaluated in the analysis for the effect of the 4-to-8-week open trial. For the ancillary analysis, naïve was analyzed to determine the predictor of CGI-S after I year of MPH administration (N=14). NAÏVE (N=14) and HCs (N=15) were analyzed for the effect of I year of MPH administration after washout. (a) After participants were matched for age, sex, IQ, and SST performance among the NAÏVE, NON-NAÏVE, and HCs, 20 members of HCs were selected. ADHD, attention-deficit hyperactivity disorder; HCs, healthy controls; MPH, methylphenidate hydrochloride; NAÏVE, ADHD-drug naïve; NIRS, near-infrared spectroscopy; NON-NAÏVE, ADHD patients who had taken methylphenidate for several months; SST, stop signal task.

Impressions-Severity (CGI-S). The NAÏVE stopped taking MPH for 1 week before undergoing the last NIRS measurement.

The washout period, timing of MPH administration, titration procedure, and mean dose of MPH are described in the Supplementary Materials.

#### **Outcomes**

We used NIRS to measure changes in the mean oxygenated hemoglobin concentration ([oxy-Hb]) in the bilateral inferior frontal cortex (IFC: Supplementary Figure S1) during the SST, which served as an explanatory variable. The procedures for the SST and NIRS measurements are described in detail in Supplementary Materials and Supplementary Figure S2, and in our previous report (Ishii-Takahashi *et al*, 2013).

We also used the CGI-S (Guy, 1976) scores after 4-to-8-week administration (Phase 3) and 1-year administration (Phase 4) of the optimal dose of MPH as outcome measures (Figure 1). We did not use the ADHD-RS-IV as a primary outcome measure in the clinical trial registry because ADHD-RS-IV was translated to Japanese in 2007 and has not been validated in Japanese patients. A detailed explanation about the adoption of CGI-S as an assessment tool of primary outcome was reported in Supplementary Materials. The child psychiatrist (AI-T) reviewed the severity and clinical response as assessed by CGI-S (Buitelaar *et al*, 1995; Cho *et al*, 2007).

For the primary analysis, regression analysis was conducted to examine whether [oxy-Hb] measured at the baseline assessment or  $\Delta$ [oxy-Hb] (single dose of MPH minus baseline assessment) predicted CGI-S scores after the 4-to-8-week open trial and after 1 year of MPH administration.

For secondary analysis, we examined at Phase 1, whether brain function differed among NAÏVE, NON-NAÏVE, and HCs; at Phase 2, whether a single dose of MPH or placebo-affected [oxy-Hb]; at Phase 3, whether brain function differed among NAÏVE, NON-NAÏVE, and HCs during the 4-to-8-week administration of MPH; and at Phase 4, whether administration of MPH for 1-year affected [oxy-Hb] after washout.

#### Randomization and Blinding of Phase 2

After baseline assessments, a study investigator (AI-T) assigned the participants to an intervention. An IRB member (Y.A.) generated a random allocation sequence to blind participants assigned an intervention and stratified the participants into two groups according to their medications. The patients were randomly assigned in a 1:1 ratio to one of the two groups. In the first group, the patients first received a single dose of MPH, and 1 week later, they received a single dose of placebo. In the second group, the order was reversed. Participants, care providers, and investigators were double-blinded to the intervention. Placebos were sucrose capsules



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similar to the MPH capsules. The investigators were informed of patient assignments after Phase 3 was completed.

# Statistical Analyses

Statistical analyses were conducted after completing the trials and opening the allocation; the analyses followed the intention-to-treat principle. All reported P-values are two-tailed. Clinical and behavioral results were considered significant at P < 0.05. The 95% confidence interval (95% CI) and effect size (d) are shown when applicable. Analyses were conducted with IBM SPSS software (version 20.0). At the baseline assessment, between-group differences in clinical scores and SST performance were tested using a one-way analysis of variance (ANOVA) with Tukey's Honestly Significant Difference (HSD) post hoc test. The  $\chi^2$ -test was used to determine differences in sex and handedness. Mean reaction time during the SST, SST performance, and ADHD-RS-IV were analyzed using a two-way repeated-measures ANOVA with post hoc Tukey's HSD test.

The primary analyses were conducted for NAÏVE. A stepwise multiple regression analysis was conducted with CGI-S scores obtained after the 4-to-8-week open trial as a dependent variable with the [oxy-Hb] at the baseline assessment in the right IFC (RIFC) and left IFC (LIFC),  $\Delta$ [oxy-Hb] (single dose of MPH minus baseline assessment) in the RIFC and LIFC, and 11 demographic and clinical variables as independent variables: MPH dosage, ADHD-RS-IV scores ('Inattention' and 'Impulsivity and Hyperactivity' sub-scores and total score), CBCL scores ('Internalizing' and 'Externalizing' sub-scores and total score), SST task performance, IQ, age, and sex. We used t-tests to identify differences in  $\Delta$ [oxy-Hb] (single dose of MPH minus baseline assessment) between subjects with a CGI-S score  $\leq 3$  and  $\geq 4$ . We then submitted the resulting values to a parametric Fisher linear discriminant analysis classification algorithm (Ponseti et al, 2012) to discriminate CGI-S  $\leq$  3 from CGI-S ≥ 4. Additional information about the definition of responder and the cross-validation methods are presented in the Supplementary Materials. We also performed a stepwise multiple regression analysis after administration of MPH for 1 year with CGI-S scores as the dependent variable, and mean [oxy-Hb] change at the baseline assessment,  $\Delta$ [oxy-Hb] (single dose of MPH minus baseline assessment), and the 11 demographic and clinical variables described above as independent variables.

For the secondary analyses, to detect differences among NAÏVE, NON-NAÏVE, and HCs at the baseline assessment (Phase 1) and after the 4-to-8-week open-trial (Phase 3), the [oxy-Hb] was subjected to a repeated-measures ANOVA with group (NAÏVE, NON-NAÏVE, and HCs) as a betweensubject factor and NIRS session (Phase 1 and 3) and laterality (RIFC and LIFC) as within-subject factors. At Phase 2, we identified differences in NIRS signals between MPH and placebo administration in the single-dose trial; the [oxy-Hb] was subjected to a repeated-measures ANOVA with group (NAÏVE and NON-NAÏVE) as a between-subject factor and NIRS session (single dose of MPH and placebo) and laterality (RIFC and LIFC) as within-subject factors. To identify differences in NIRS between NAÏVE and HCs at the baseline assessment and after washout following MPH administration for 1 year, [oxy-Hb] was subjected to a repeated-measures ANOVA with group (NAÏVE and HCs) as a between-subject factor and NIRS session (Phase 1 and 4) and laterality (RIFC and LIFC) as within-subject factors. Subsequently, we used the Bonferroni correction for specific *post hoc* contrasts (significance at Bonferroni-corrected P < 0.05).

#### **RESULTS**

#### **Participants**

The study began on 1 August 2008 and ended with the 1-year follow-up termination on 29 June 2011. Table 1 presents the baseline demographic characteristics for the study participants. Of the 33 potential participants with ADHD, 3 were excluded for not meeting the inclusion criteria (Figure 2). Thus, we examined 30 patients in Phase 2 (22 NAÏVE and 8 NON-NAÏVE). One NAÏVE participant was excluded because of an inability to continue taking MPH during Phase 3. Among 21 NAÏVE patients who participated in Phase 3, 14 completed Phase 4, 4 stopped taking MPH, 2 stopped attending the hospital, and 1 changed medications. In addition, 20 HCs were included in the analysis after assessment of 33 children for eligibility; 15 completed the follow-up, and 5 stopped attending the study. NAÏVE, NON-NAÏVE, and HCs did not differ in SST performance (Table 1). The ADHD-RS-IV scores of both NAÏVE and NON-NAÏVE were significantly higher compared with those of HCs. Changes in performance and severity (baseline assessment and 4-to-8-week open trial), the comparison of demographic characteristics at the baseline assessment between NAÏVE and HCs participated in 1-year follow-up, and changes in performance and severity (baseline assessment and 1-year follow-up) in NAÏVE are described in Supplementary Materials, and Supplementary Table S1-S3.

#### **Outcomes**

*Primary results* 

*Predictor of CGI-S after 4-to-8-week open trial.* In the 4-to-8-week open trial, there were 4 participants in CGI-S 2, 12 in CGI-S 3, and 5 in CGI-S 4. Multiple regression analysis for CGI-S scores after the 4-to-8-week open trial revealed a significant regression only for  $\Delta$ [oxy-Hb] in LIFC (R = 0.519, P = 0.0160, β = −0.519, 95% CI = −4.680 to −5.440; Figure 3a). The  $\Delta$ [oxy-Hb] in LIFC was significantly lower in participants with CGI-S  $\geqslant$  4 than in those with CGI-S  $\leqslant$  3 (P = 0.0084, d = 1.200, 95% CI = 0.351 to 2.541). The leave-one-out classification algorithm had an 81% (95% CI = 0.782 to 0.813) accuracy (81.3% sensitivity (95% CI = 0.782) using the  $\Delta$ [oxy-Hb] in LIFC.

Predictor of CGI-S after 1-year of MPH administration. At the 1-year follow-up, there were six participants in CGI-S 2, six in CGI-S 3, and two in CGI-S 4. Multiple regression analysis for the CGI-S score at the 1-year follow-up revealed that  $\Delta$ [oxy-Hb] in LIFC also predicted the CGI-S score (R = 0.716, P = 0.0040,  $\beta$  = -0.716, 95% CI = -6.527 to -1.562; Figure 3b).

#### Secondary results

Group comparison at baseline assessment and effect of 4-to-8-week MPH administration (Phase 1 and 3). We

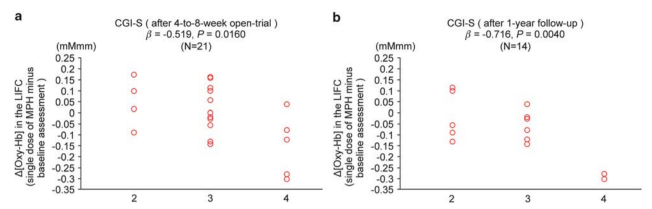


Figure 3 Results for primary outcomes: predictor of CGI-S in NAÏVE. (a) Correlation between CGI-S scores (4-to-8-week open trial: 2, borderline mentally ill (N=4); 3, mildly ill (N=12); 4, moderately ill (N=5)) and  $\Delta$ [oxy-Hb] (single dose of MPH minus baseline assessment) in the LIFC ( $\beta=-0.519$ , P=0.0160). (b) Correlation between CGI-S (after I-year follow-up: 2, borderline mentally ill (N=6); 3, mildly ill (N=6); 4, moderately ill (N=2)) and  $\Delta$ [oxy-Hb] (single dose of MPH minus baseline assessment) in the LIFC ( $\beta=-0.716$ , P=0.0040). ADHD, attention-deficit hyperactivity disorder; CGI-S, Clinical Global Impression-Severity; HCs, healthy controls; LIFC, left inferior frontal cortex; MPH, methylphenidate hydrochloride; NAÏVE, ADHD-drug naïve; NIRS, near-infrared spectroscopy; NON-NAÏVE, ADHD patients who had taken methylphenidate for several months; [oxy-Hb], oxygenated hemoglobin concentration; RIFC, right inferior frontal cortex.

observed a significant interaction for group (NAÏVE vs NON-NAÏVE vs HC) × NIRS session (baseline assessment vs 4-to-8-week open trial) × laterality (RIFC vs LIFC; F = 3.743, P = 0.0312). At the baseline assessment, NAÏVE demonstrated a significantly lower activation in the RIFC (Bonferroni-corrected P = 0.0455, d = -0.778, 95% CI = -1.452 to -0.178) and a tendency toward lower activation in the LIFC (corrected P = 0.0908, d = -0.729, 95% CI = -1.424 to -0.153) compared with HCs (Figure 4a). However, NON-NAÏVE and HCs did not differ significantly (Figure 4a). After the 4-to-8-week open trial, [oxy-Hb] did not differ significantly among NAÏVE, NON-NAÏVE, and HCs (Figure 4c). Additionally, HCs showed significantly lower bilateral IFC activation in the 4-to-8-week open trial than in the baseline assessment (RIFC; Bonferroni-corrected P = 0.0017, d = 0.888, 95% CI = 0.204-1.499 and LIFC; Bonferroni-corrected P = 0.0011, d = 1.170, 95% CI = 0.256 to 1.557). The details about the interaction of laterality and session in each group are described in the Supplementary Materials.

Effect of single-dose MPH administration (Phase 2). In the single-dose trial, a significant interaction was observed for group (NAÏVE vs NON-NAÏVE) × NIRS session (MPH vs placebo) × laterality (RIFC vs LIFC; F = 5.058, P = 0.0326). NAÏVE demonstrated significantly higher activation in only the RIFC during the SST with MPH compared with the placebo (Bonferroni-corrected P = 0.0281, d = 0.579, 95% CI = 0.009 to 1.218; Figure 4b). In NON-NAÏVE, the effects of MPH and placebo did not differ significantly in the bilateral IFC.

Effect of 1-year MPH administration after washout (Phase 4). After the 1-year follow-up, a significant interaction was observed for group (NAÏVE vs HCs) × NIRS session (baseline assessment vs 1-year follow-up; F=8.160, P=0.0081). NAÏVE showed significantly lower activation in the bilateral IFC compared with the HCs group (Bonferronicorrected P=0.0121, d=-0.938, 95% CI=-1.741 to -0.201; Supplementary Figure S3) at the baseline assessment.

No significant difference was observed between NAÏVE and HCs after washout of 1-year administration (Figure 4d). HCs demonstrated significantly lower bilateral IFC activation at the 1-year follow-up than at the baseline assessment (Bonferroni-corrected P = 0.0102, d = 0.719, 95% CI = -0.017 to 1.460).

#### Harm

Side effects included loss of appetite (N=18), difficulty falling asleep (N=2), and transient tics (N=2). These side effects diminished during titration. No serious side effects occurred.

# **DISCUSSION**

This clinical trial is the first to demonstrate that a difference in prefrontal hemodynamic responses after single administration of MPH relative to the baseline level significantly predicted mid-term (4-to-8-week) and long-term (1-year) clinical efficacy of MPH in children with ADHD (Figure 3); the leave-one-out classification algorithm produced 81% accuracy. Additionally, (1) at the baseline assessment, NAÏVE exhibited significantly lower prefrontal activation than HCs, whereas NON-NAÏVE and HCs showed similar activation; (2) a single dose of MPH significantly increased activation compared with the placebo in NAÏVE; and (3) after the 4-to-8-week administration, and even after MPH washout following 1-year administration, naïve demonstrated normalized prefrontal activation (Figure 4). Collectively, these results indicate that supplementary NIRS measurements may serve as a safe and simple objective biomarker for clinical decisions and monitoring for continuous MPH treatment for children with ADHD. The simplicity of NIRS measurement is also an important advantage for its clinical application in child psychiatry.

The strengths of our study include its randomized, double-blind, placebo-controlled, crossover design; 1-year follow-up prospective study; and comparison of drug-naïve and MPH-treated children. The follow-up rate for the primary outcome was high (96.7%). Therefore, the risk of biased conclusions

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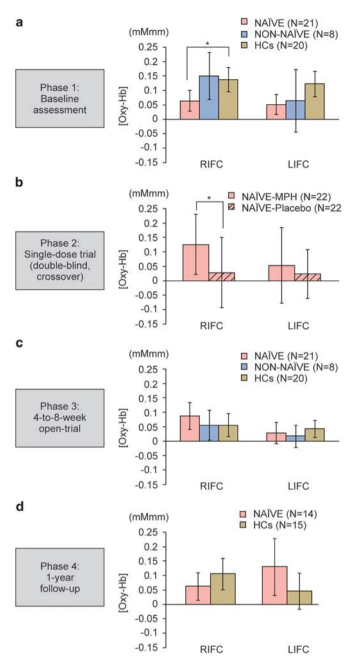


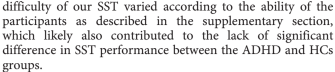
Figure 4 Results for secondary outcomes: effects of the 4-to-8-week open trial of MPH, effects of a single-dose trial, and effects of 1 year of MPH administration observed after washout. (a) At the baseline assessment (RIFC, Bonferroni-corrected P = 0.0455; LIFC, Bonferroni-corrected P = 0.0908). (b) Differences in [oxy-Hb] changes during the SST in the single-dose trial between MPH and placebo treatments in NAÏVE (RIFC: Bonferronicorrected P = 0.0281). (c) Differences in [oxy-Hb] changes during the SST between the HCs and ADHD groups after the 4-to-8-week open trial. Error bars indicate 95% Cl. (d) Differences in [oxy-Hb] changes during SST between the HCs and NAÏVE after 1-year follow-up (baseline assessment: Bonferroni-corrected P = 0.0121; after 1-year follow-up: Bonferronicorrected P=0.6386). Error bars indicate 95% Cl. ADHD, attentiondeficit hyperactivity disorder; CGI-S, Clinical Global Impression-Severity; HCs, healthy controls; LIFC, left inferior frontal cortex; MPH, methylphenidate hydrochloride; NAÏVE, ADHD-drug naïve; NIRS, near-infrared spectroscopy; NON-NAÏVE, ADHD patients who had taken methylphenidate for several months; [oxy-Hb], oxygenated hemoglobin concentration; RIFC, right inferior frontal cortex; SST, stop signal task.

was low. The timing of NIRS (5 h after taking MPH) was selected to achieve the peak serum level of MPH OROS.

The main finding of our study is that changes in prefrontal activation after a single dose of MPH predicted the efficacy of long-term MPH administration (Figure 3). Increases in LIFC activity were previously observed in patients with ADHD who performed an error-monitoring task after acute (Rubia et al, 2011) and 6 weeks-MPH treatment during interference inhibition (Bush et al, 2008). A change in activation in the single-dose trial toward the level in HCs indicated a better response to continuous administration of MPH. These results are in line with a magnetic resonance imaging (MRI) study that predicted the response to continuous MPH administration using resting-state signals (An et al, 2013). We observed no correlation between MPH efficacy and baseline characteristics such as ADHD-RS-IV, CBCL internalizing score, age, IQ, or sex, which is consistent with the outcomes reported for a previous event-related potential study (Hermens et al, 2005). Although other neuroimaging modalities have predicted the effects of MPH (An et al, 2013; Cho et al, 2007; Ilgin et al, 2001; la Fougere et al, 2006; Sangal and Sangal, 2004; Schweitzer et al, 2003), we used NIRS because of its simplicity and safety, which are beneficial in a clinical setting. Because only NAÏVE were used to predict the effect of MPH, supplementary NIRS measurements may be applied in clinical situations in which a physician must decide whether a child should begin taking the medication.

Although it was not a primary focus of our study, we confirmed previous findings, obtained using other neuroimaging modalities, of prefrontal abnormalities in ADHD and their normalization by MPH treatment. First, NAÏVE demonstrated lower activation in the RIFC at the baseline assessment (Figure 4a), consistent with previous findings in drug-naïve children with ADHD (Hart et al, 2013). In contrast, NON-NAÏVE demonstrated activation similar to that of HCs at the baseline assessment, thus indirectly indicating the effect of previously administered MPH, which is in line with the findings of a previous functional MRI (fMRI) study in adults (Schlochtermeier et al, 2011; Stoy et al, 2011). Second, a single dose of MPH increased the right frontal NIRS signal (Figure 4b), thus confirming that MPH increased RIFC activation during an inhibitory control task (Rubia et al, 2014). Third, the change toward a level of prefrontal activation similar to that for HCs after the 4-to-8week open trial (Figure 4c) is consistent with the results from a fMRI study of adults with ADHD (Bush et al, 2008) and an ERP study in children with ADHD (Sawada et al, 2010). Finally, our finding (Figure 4d) that long-term (1 year) MPH treatment affected brain development in ADHD children is consistent with the findings of a prospective MRI study (Konrad et al, 2007).

In the current study, no significant differences in SST performance were observed between children with ADHD and HCs. To detect the differences between ADHD and control groups using only SST as in previous studies (de Vries and Geurts, 2014; Gupta and Kar, 2009), the task would need to contain more trials than those included in our task. We designed our cognitive task such that the performance of the different groups did not significantly differ because we intended to detect significant differences in the NIRS signal during a short cognitive task. Additionally, the degree of



Some limitations of our study should also be highlighted. First, methodological limitations include the smaller number of participants for NON-NAÏVE (N=8), lack of blindness in Phase 3, and non-negligible dropout rate (7 out of 21 naïve) at the 1-year follow-up. Second, NIRS does not detect activity in deeper cortical structures, such as the anterior cingulate cortex, which is part of the neural network that subserves response inhibition (Duann et al, 2009). Third, as we expected, the NIRS signal amplitude decreased significantly on repeated measurement. We evaluated the NIRS signal of untreated HCs at the same time points as children with ADHD to control for the effects of repeated measurements. Because the crossover design was also applied in the comparison between MPH and placebo, any potential effects of learning on our results were considered to be resolved. Moreover, the primary aim of our study was the prediction of the effect of continuous administration of MPH based on the NIRS signal, and the use of repeated measurements was thus considered unlikely to affect our findings in this context. Although the clinical use of NIRS may be especially suitable in children because of its simplicity and safety, to date, no study has evaluated the reliability of repeated measurement of NIRS in children. It is necessary to further study the reliability of repeated NIRS measurement when using SST or other tasks in children. Finally, more study is required before this technique can be officially approved and applied in realworld clinical settings. The present findings should be replicated and validated in independent and larger subject groups in future multi-site studies. In particular, the differences among age, sex, and ADHD subtype should be evaluated in a large sample. It is also necessary to compare factors for predicting the effect of MPH combined with other medications such as atomoxetine in order to select the best treatment regimen for clinical application. Furthermore, a longitudinal study that follows the participants to adulthood, including those participants who stop treatment, is necessary to determine predictors for long-term success of

In conclusion, this study illustrates an innovative application of 'next-generation' neuroimaging to child psychiatry. Greater change in prefrontal activation after a single dose of MPH predicts greater effectiveness with continuous administration for 4-8 weeks and 1 year. The present NIRS system may serve as a supplemental neuroimaging modality to aid in the clinical decision-making concerning continuous MPH administration in children with ADHD.

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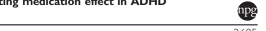
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## Disclaimer

The sponsors had no role in the design and conduct of the study; data collection, analysis, or interpretation; or in preparation, review, or approval of the manuscript. AI-T and KK had full access to all data in the study and take responsibility for the integrity of the data and the accuracy of the data analysis.

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