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Effects of d-Methylphenidate, Guanfacine, and their Combination on EEG Resting State EEG Spectral Power in ADHD

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ABSTRACT

Objective: Psychostimulant medications are the gold standard of treatment for attention-deficit/hyperactivity disorder (ADHD), however, a significant minority (~30%) of individuals with ADHD fail to respond favorably. Noradrenergic agents are increasingly used as ADHD monotherapies or adjuncts for sub-optimal stimulant response, yet knowledge of their cortical effects is limited. This study is the first to examine comparative effects of guanfacine (an alpha adrenergic 2A agonist), psychostimulant, and their combination on resting state cortical activity in ADHD.

Method: The sample comprised 179 participants aged 7- to 14- years old with ADHD (113 boys, 55 girls). Participants were randomized to one of three blinded conditions: guanfacine (GUAN), d-methylphenidate (DMPH) or the combination (COMB). Electroencephalography (EEG) was performed pre-, mid-, and post- medication titration, with concomitant assessment of behavioral and cognitive functioning.

Results: Analyses of spectral power measures during resting EEG suggested that each medication condition displayed a distinct profile of effects on cortical activity. Significant time effects suggested that GUAN decreased global alpha-band (8-12 hertz [Hz]) power, DMPH and COMB increased centro-parietal beta-band (13-21 Hz) power, and COMB resulted in decreased theta-band (4-7 Hz) power. Relative to other medication groups, COMB was associated with significantly lower theta power and DMPH with higher beta-band power compared to the GUAN group. Medication-related changes in theta power were correlated with improvements in behavioral and cognitive functioning.

Conclusions: These data revealed distinct underlying medication-related effects on neural mechanisms. The COMB condition uniquely exhibited an EEG profile that was associated with improved behavioral and cognitive functioning.

Clinical Trial Registration Information:

URL: http://clinicaltrials.gov/show/NCT00429273
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Keywords: electroencephalography, children, medication, treatment, stimulants
BACKGROUND

Attention-deficit/ hyperactivity disorder (ADHD) is one of the most common psychiatric disorders in children, affecting approximately 5-10% of youths. The core symptoms of this neurodevelopmental disorder include inattention, hyperactivity, and impulsivity and result in significant academic, psychological, and social impairment. While considerable progress has been made in elucidating genetic, neurobiological, and cognitive correlates of ADHD, a complete mechanistic model of ADHD has been elusive. The same can be said regarding our understanding of the mechanisms of successful treatment. A host of brain imaging methodologies has been applied to the study of ADHD and its treatment, revealing a number of abnormalities at the level of brain structure, function, connectivity, and neurochemistry. Among brain imaging modalities, electroencephalography (EEG) has long been used to assess the neural correlates of ADHD. EEG is especially well-suited for these studies as it is more tolerant of ADHD-related movement artifacts, is much less expensive and more easily administered than other brain imaging modalities such as fMRI, and has been used extensively to identify resting state differences in cortical activity among children with ADHD.

One of the most consistent neurophysiologic correlates of ADHD relative to typically developing children is higher frontocentral slow-wave power in the theta frequency band (4-7 Hertz [hz]). In meta-analytic studies of EEG that include over 1400 ADHD-affected subjects, resting theta band power is significantly increased by 32% on average versus controls, with an effect size of 1.31 (95% confidence interval, 1.14 - 1.48). In addition to higher frontocentral theta band power, lower global alpha and beta band activity have also been associated with ADHD. However, findings in these frequency bands are more variable and may differ according to age, ADHD subtype, and psychiatric co-morbidities. Although EEG measures such as theta power as well as the ratio of theta power to beta power, i.e. theta/beta ratio (TBR) have been suggested as having clinical utility in
ADHD, recent studies suggest that neither have the sensitivity/specificity needed to function as stand-alone diagnostic markers\(^5\).

Psychostimulant medications (e.g. methylphenidate, amphetamine) have been first line ADHD treatments for over 70 years; however, a substantial number (25-35\%) of children do not respond satisfactorily to initial stimulant trials\(^6\). Noradrenergic alpha-2 adrenoceptor agonists (\(\alpha\) agonists), such as guanfacine and clonidine, have been increasingly used both as alternative monotherapies or adjunctive treatments in cases of inadequate stimulant response\(^7,8\). The \(\alpha_2\text{A}\) mechanism of action of these agonists have been supported as treatments for disorders associated with prefrontal cortical (PFC) dysfunction based on preclinical studies that demonstrated guanfacine increased firing in PFC neurons during task-related activities and strengthened functional network connectivity between the PFC and other cortical regions\(^9\). Furthermore, a recent study using an animal model for ADHD, the spontaneously hypertensive rat (SHR), reported that guanfacine achieves its effects through stimulation of the postsynaptic adrenergic \(\alpha_{2\text{A}}\) receptors in the prefrontal cortex as opposed to \(\alpha_{2\text{B}}\) or \(\alpha_{2\text{C}}\) receptors or through presynaptic \(\alpha_{2\text{A}}\) receptor-regulated noradrenaline release\(^10\). Results of randomized clinical trials indicate that guanfacine, a selective \(\alpha_{2\text{A}}\) noradrenergic agonist, is effective in reducing the core symptoms of ADHD as a monotherapy\(^11,12\) and as an adjunctive treatment for those individuals with sub-optimal responses to psychostimulants alone\(^8\). Further work on the mechanistic actions of \(\alpha_{2\text{A}}\) agonists, however, is needed to better understand, the cognitive and neural effects of these medications as monotherapies and in combination with stimulants in human subjects with ADHD.

The overall effect of psychostimulants has been shown to improve, but not normalize, many features of ADHD-related abnormal EEG activity. Specifically, MPH tends to decrease theta band and increase beta band power\(^13-15\), particularly when associated with medication-related improvements in cognition\(^16,17\). The effects of non-stimulant medications on cortical activity in ADHD have not been well studied. One study examined the effects of atomoxetine, a selective
norepinephrine reuptake inhibitor, on EEG\textsuperscript{18}. Atomoxetine appears to have some similarities to psychostimulants, such as its ability to significantly reduce posterior theta band power and increase absolute beta band power with acute administration, however, these changes were modest, and differences versus reported stimulant effects were also seen in the increase in slow-wave, delta power\textsuperscript{17}. Thus, some shared effects associated with partial normalization of the EEG profile apparent in ADHD may be associated with both psychostimulants and non-stimulant medications used to treat ADHD. To date, the cortical effects of $\alpha_2\text{A}$ agonists compared to and in combination with psychostimulants in children with ADHD have not been reported.

The current study takes a step in that direction by testing the effects of the single enantiomer stimulant d-methylphenidate (DMPH), the $\alpha_2\text{A}$ agonist guanfacine (GUAN), and the combination of DMPH and GUAN (COMB) on resting state EEG measures in youth with ADHD. Goals of this study are to contrast cortical activity effects under each medication condition during resting state and test whether acute changes in cortical activity during rest are predictive of cognitive functioning. Given that the two medication classes, psychostimulants and $\alpha_2\text{A}$ agonists, have different proposed neural mechanisms that underlie their clinical benefits, we hypothesized that these mechanistic differences would be evident in differential EEG effects on resting state spectral power. The COMB condition was predicted to have additive effects of the monotherapies, and hence, more beneficial effects on the spectral power profile. We hypothesized that significant medication-related EEG changes will be associated with improvements in behavioral and cognitive functioning.

**METHODS**

*Sample.* The sample consisted of 179 participants (113 boys, 55 girls), aged 7- to 14-years old who were diagnosed with ADHD. All participants were enrolled in the “Translational Research to Enhance Cognitive Control” ADHD (TRECC; McCracken et al., under review) project. Participants were recruited from clinic referrals, radio and newspaper advertisements, community organizations
(CHADD; www.chadd.org), local schools, and primary care physicians. After receiving verbal and written explanations of study requirements, and prior to any study procedures, all parents/participants provided written informed permission/assent as approved by the UCLA Institutional Review Board. The study was also overseen by a local Data Safety and Monitoring Board (DSMB), which provided a tri-annual review.

Procedure. As part of their participation in the TRECC study, participants underwent phenotypic assessment including diagnostic interviews and EEG recording. For a detailed description of diagnostic and cognitive assessment, please see McCracken et al, submitted; Bilder et al, submitted). Briefly, individuals were evaluated based on a semi-structured diagnostic interview with the primary caretaker (usually mother) and a direct interview if 8 years of age or older using the Schedule for Affective Disorders and Schizophrenia for School-Age Children (KSADS-PL). Teacher reports were solicited and used to supplement clinical interview data. Psychiatric disorders were considered ‘present’ if the participant currently met full DSM-IV diagnostic criteria. All interviews were conducted by clinical psychologists or other highly trained interviewers with extensive experience in psychiatric diagnoses and training in using the KSADS. ‘Best estimate’ diagnoses were determined after individual review of diagnoses, symptoms, and impairment level by senior clinicians (JJM, JP). Participants were included if they had a current diagnosis of ADHD, any subtype, and excluded if positive for any neurological disorder, head injury resulting in concussion, diagnoses of autism, chronic tic disorder, current major depression, panic disorder, lifetime bipolar disorder or psychosis, or estimated Full Scale IQ < 80. Participants were off medication for baseline assessments.

After confirmation of eligibility, participants were enrolled in a double-blind, comparative clinical trial and randomly assigned to one of three medication conditions: immediate release guanfacine (GUAN) administered twice daily, d-methylphenidate extended release (DMPH) once daily, or the combination of GUAN and DMPH (COMB) (see Figure 1). Treatments were applied sequentially: the first 4 weeks, participants received GUAN or placebo, beginning at 0.5 mg administered twice daily in week 1 and increased as tolerated to 0.5 - 1.5 mg twice daily doses in
subsequent weeks. Participants remained on the optimal GUAN dose (determined by CGI-Improvement ratings, ADHD-RS-IV scores, and side effects for Week 4) for the remainder of the study. Beginning in Week 5, participants initially randomized to GUAN continued taking GUAN with added DMPH or placebo, while participants initially randomized to placebo received added DMPH. Low, medium, and high stimulant doses were assessed weekly for behavioral response and tolerability to determine the “optimal” DMPH dose at Week 7, according to the identical process described above. Participants remained on optimal GUAN and/or DMPH doses for a final study week prior to end-of-study Week 8 assessments. For consort chart summarizing participant completion rates for each medication group as well as adverse side effects reported within each medication group, please see McCracken et al., (under review). EEG was administered at the following time points and medication conditions: at baseline with no medication, at week 4, which compared GUAN and placebo, and at Week 8, which compared one week of optimized treatment with GUAN, DMPH, or COMB.

Insert Figure 1 here

**Symptom assessment:** The ADHD-Rating Scale (ADHD-RS-IV\textsuperscript{20}) was used as a measure of ADHD symptom severity. A clinician blind to medication status completed this measure at baseline, Week 4 and at Week 8 or last visit based on parent, teacher, and other available data.

**Cognitive assessment:** As part of the larger cognitive battery, estimated intelligence (IQ) was assessed using the Wechsler Abbreviated Scale of Intelligence (WASI). Other cognitive variables are factor analytic scores derived from the full cognitive battery that represent four primary cognitive processes of interest: working memory [WM], response inhibition [RI], reaction time [RT], and reaction time variability [RTV]). The individual tests used in the construction of the cognitive factors are described in the accompanying article by Bilder et al (under review) and available as supplemental material.

**Electrophysiologic methods.** EEG recording was carried out using 40 Ag/AgCl surface electrodes that were embedded in an electrode cap arrayed in an extended International 10/20
configuration (ElectroCap, Eaton, Ohio) and was referenced to linked ears. Impedance was below 10 kOhms and EEG signal was recorded using MANSCAN (Sam Technology, San Francisco, CA) hardware and software. EEG data were sampled at a rate of 256 samples per second. Eye movements were monitored by electrodes placed on the outer canthus of each eye for horizontal movements and by electrodes above the eye for vertical eye movements. EEG recording for all subjects consisted of a baseline condition lasting 5-minutes during which participants sat quietly with their eyes closed.

Continuous EEG data were reviewed off-line (i.e., after the recording was complete and the participant had left) by a technician experienced in EEG and all segments containing eye, head movement or muscle artifact were removed from further analysis. In order to be included in the following EEG analyses, at least 30 seconds of artifact free EEG was required. Using a Fast Fourier Transform, average spectral power ($\mu V^2$) was computed for the following frequency bands: theta (4-8 Hz), alpha (8-12 Hz), beta1 or sensorimotor rhythm (12-16 Hz), and beta2 (16-21 Hz). Baseline absolute power for all three medication groups is presented in Supplemental Figure 1. Relative power for each frequency band was calculated (using total power from 1-21 Hz for each electrode as the denominator) and natural log transformed to assume a normal distribution. To reduce the number of comparisons, spectral power was averaged by region as follows: frontal (F3, F4, Fz), central (C3, C4, Cz), and parietal (P3, P4, Pz). EEG technicians were blind to medication group status.

**Data analytic strategy.** Analyses were conducted using IBM SPSS statistics version 21. Because age has significant effects on EEG power, it was used as a covariate in all analyses. In order to control for Type 1 error, two procedures were used. First, we performed analyses on regional spectral power estimates from proximal electrodes to reduce the number of contrasts from 36 individual electrodes to 3 regions. Second, we used the false discovery rate (FDR$^{21}$) to maintain the family-wise experimental error at $p<0.05$. According to the FDR analysis, $p$-values $\leq 0.01$ are
significant; raw p-values between <0.01 and 0.05 will be presented here as trend-level findings that may provoke further testing.

To assess the medication effects on EEG spectral power measures, generalized linear mixed model analyses were used to compare EEG spectral power by region and frequency band at baseline (B), week 4 (W4) and week 8 (W8) for each medication group. Primary effects of interest were: 1) TIME: the main effect of time, which tests within-medication group changes in spectral power at three time points, 2) MEDICATION: the main effect of medication is a between-subjects effect that indicates whether any of the medication groups were significantly different from others across all time points, and 3) MEDICATION X TIME: interaction effect of medication group by time, which tests if spectral power changes between medication groups were significantly different over the course of the medication trial. Significant omnibus effects (p<0.01) and trends (p<0.05) were followed by individual contrasts to determine specific time point (B, W4, W8) or medication group at week 8 (GUAN, DMPH, COMB) differences.

Pearson partial correlations (with age as a covariate) between EEG variables and behavioral measures/cognitive factor scores were conducted to assess the association of medication-related changes in EEG power. Behavioral variables consisted of the ADHD-RS-IV inattentive and hyperactive-impulsive scores were used to reflect ADHD symptom severity at W8. In addition, the degree of treatment-related change in ADHD symptom severity was used in the correlation analysis. To calculate treatment change, baseline ADHD symptom severity was subtracted from Week 8 ADHD symptom severity under the assumption that there would be a reduction of ADHD symptoms through the course of treatment; thus a larger number for the treatment change variables suggests a bigger reduction in symptom severity from baseline to end of trial. The treatment change and Week 8 ADHD symptom severity variables provide different information regarding the relationship of EEG to ADHD symptomatology and consequently show different patterns of association.

RESULTS
Demographics. Of the 212 randomized participants, 182 completed the clinical trial (Week 8) and were included in the current study. Three participants were excluded based on the requirement to have at least 30 seconds of artifact EEG, leaving a final sample of 179 (GUAN n=59; DMPH n=60; COMB n=60). The 179 participants were 67% male, an average age of 10.1 years (SD=2.1), predominantly Caucasian (75%), and mean IQ of 103 (SD=14). At baseline, participants had, on average, ADHD-Inattentive symptoms= 7.9 (SD=1.3) and ADHD-Hyperactive/Impulsive symptoms=5.1 (SD=2.6). None of these variables differed significantly across medication groups. Average EEG spectral power for each medication group within each frequency is presented in Figure 2 for baseline and W8 time points. There were no significant differences in mean relative spectral power in any band or cluster across the groups at baseline (all p’s >0.2).

Within-group changes in EEG spectral power by medication group. Significant TIME effects informed whether a significant effect of medication was observed on EEG power among subjects within each medication group. Results of the linear mixed models suggested significant TIME effects on EEG spectral power for each medication condition; this is depicted by percent change in EEG power for each medication group within each frequency band by region in Figure 3. Each medication group was associated with significant changes in a different frequency band. The primary effect of COMB medications was in central and parietal theta-band power (central: F(2, 238.3)=3.0, p=0.049, and parietal F(2, 238.3)=3.4, p=0.034), which was reduced at week 8 compared to baseline and week 4. In contrast, the DMPH group exhibited increased parietal beta-band power (F(2, 238.3)=5.0, p<0.01) at Week 8 relative to baseline levels. Finally, administration of GUAN resulted in decreased global alpha-band (8-12 Hz) spectral power at Weeks 4 and 8, where significant effects of TIME emerged in all regions (frontal: F(2, 238.3)=10.0, p<0.001, central: F(2, 238.3)=14.7, p<0.001, parietal: F(2, 238.3)=4.8, p=0.008).
**Between-group changes in EEG spectral power.** Significant MEDICATION and MEDICATION x TIME interaction effects informed whether a significant medication effect was observed on EEG power relative to other medication conditions. These effects are also depicted in Fig. 2. In the beta2 band (17-21 Hz), a significant MEDICATION effect emerged in the frontal and central regions for the theta and beta2 frequency bands (central theta: F(2,172.5)=3.0, p=0.050; frontal beta2: F(2,172.5)=3.9, p=0.022; central beta2: F(2,172.5)=3.5, p=0.033). In the central region, the COMB group exhibited lower theta band power relative to the GUAN group (p<0.05), although this effect became a trend-level finding after FDR correction. DMPH also had significantly higher spectral beta band power compared to the GUAN group in frontal regions (p’s <0.01) and both DMPH and COMB trended towards higher beta2 power relative to GUAN in the central region (p<0.05).

Significant MEDICATION x TIME interaction effects emerged in the theta-, alpha-, and beta2 frequency bands. In frontal and central regions (frontal F(4,238.3)=2.9, p=0.024, central: F(4,238.3)=4.0, p=0.004; parietal F(4,238.3)=3.1, p<0.016), theta power was significantly lower in the COMB group compared to GUAN (p<0.005) across all regions and trended in the same direction relative to the DMPH group in central and parietal regions (p’s < 0.05). Significant interaction effects for alpha power in central (F(4,238.3)=4.2, p=0.002) and parietal (F(4,238.3)=4.8, p=0.001) region emerged. The GUAN group exhibited lower alpha power relative to both the COMB and DMPH group in the central region (p<0.05) and the COMB condition in the parietal region (p<0.05) only. The COMB group alone exhibited an EEG profile of reduced theta-band and higher alpha and beta band power after medication administration.

**Correlations between EEG and behavioral and cognitive variables.** Partial Pearson correlations (controlling for age) were used to assess the relationship between resting state spectral power and behavioral and cognitive functioning at the Week 8 time point. Significant between-group medication condition differences were observed in global theta, central and parietal alpha and frontal and central beta band power, therefore these EEG variables were tested for association with behavioral and cognitive measures. As seen in Table 1, severity of ADHD inattentive or hyperactive-
Impulsive symptoms at W8 were not associated with EEG spectral power beyond two trend-level correlations between theta-band power and hyperactive-impulsive symptoms.

In contrast, theta band spectral power was significantly negatively associated with treatment-related change in ADHD symptoms such that lower theta band power was correlated with a larger reduction in inattentive symptoms (r's range -0.23 to -0.29, p<0.01). In central and parietal regions, reduced theta power was also associated with larger treatment change in hyperactive-impulsive symptoms (r's range 0.23 to 0.25, p<0.01). Spectral power in alpha and beta bands was not associated with ADHD symptom severity at W8 or treatment-related change in ADHD symptomatology. The primary effect of the COMB medications was lower theta power, which was significantly associated with medication-related improvement in ADHD symptom severity.

Within the cognitive domain, the relationship between the same EEG variables and the factor scores for broad domains of cognitive functioning were tested (see Table 1). Although the correlations were modest, significant correlations emerged between resting state EEG power and cognitive performance, both of which were measured at week 8. Theta band power in the parietal region was negatively correlated at a trend level with the working memory factor (r=0.20, p=0.033) and significantly correlated with reaction time (r =0.24, p=0.010) and reaction time variability (r=0.30, p=0.002). These correlations suggest a relationship between improved cognitive performance (better working memory performance and lower reaction time variability) with lower spectral power in the theta frequency band during resting state. Coupled with the behavioral findings above, these data further strengthen the idea that reduced theta band power, which is evident in the COMB medication condition, is associated with enhanced functioning in ADHD. Alpha band power, on the other hand, was significantly negatively correlated with reaction time (central r = 0.32, p=0.001; parietal r = 0.30, p=0.001) and reaction time variability (central r =0.31, p=0.001; parietal r = 0.29, p=0.002) and trended toward significance with the Inhibition factor score (central r = -0.21, p=<0.031; parietal r = -0.22, p<0.019). Thus, the primary effect of GUAN, which was lower alpha band power, was
associated with a slower and more variable cognitive task performance and slightly poorer performance on inhibition tasks. Beta band activity was not significantly associated with any of the behavioral or cognitive variables, suggesting that resting state cortical activity in this frequency band may be a poor predictor of later behavior or cognitive performance.

DISCUSSION

The current study is the first to report the comparative medication effects of $\alpha$2A agonist, guanfacine, d-MPH, and their combination on resting state EEG spectral power among a large sample of youth with ADHD. Analyses of spectral power measures during resting EEG suggest that each medication is associated with unique effects on cortical activity: GUAN with decreased alpha band power, DMPH with increased frontal and central beta power, and COMB with decreased theta band power and focal increases in beta power. Correlations between spectral power and behavior/cognitive variables, although modest, suggest that lower theta and higher alpha at post-treatment are associated with improved behavioral and cognitive functioning. Thus, these data suggest that each medication has a distinct neural signature, which is associated with behavioral and cognitive functioning.

The GUAN-related decrease in alpha band power is especially interesting to consider due to a seemingly paradoxical effect. Behaviorally, GUAN leads to improvement in ADHD symptoms\textsuperscript{6,11,12}, however neurophysiologically, guanfacine appeared to lower alpha power, which was subsequently associated with slower reaction time, higher reaction time variability, and a trend towards lower performance on tasks measuring inhibition. This result is similar to the only previous study examining EEG correlates of guanfacine among humans\textsuperscript{22}. Among 10 healthy adults, guanfacine resulted in alpha power decrease with concomitant subject report of significantly reduced alertness, both of which were interpreted as being consistent with a central nervous system depressant effect. Resting alpha power has been negatively associated with vigilance\textsuperscript{23} and arousal regulation\textsuperscript{24}, potentially arising from idling of thalamocortical circuits\textsuperscript{25}. Recent studies suggest that resting state EEG alpha
power is negatively correlated with functional connectivity (as measured by resting state functional magnetic resonance [rs-fMRI]) across a broad range of regions\textsuperscript{26}, particularly within the visual network\textsuperscript{27}. In addition, researchers have also found an association between alpha power and the degree of anti-correlation between the default network and task-positive attention networks\textsuperscript{28}, which has also found to be aberrant in ADHD\textsuperscript{29}. Finally, alpha power has been associated with affective dysregulation\textsuperscript{30} and recent studies suggest that guanfacine may modulate the influence of emotional cues on cognitive control by altering fronto-limbic connectivity\textsuperscript{31}. These associations suggest that guanfacine’s neural effect may be to decrease thalamo-cortical arousal or affective modulation of cognitive processing, which leads to slower and more variable reaction time. Thus, guanfacine may be most effective for those individuals with ADHD who have high arousal levels or affective dysregulation, perhaps in line with the reported secondary effect of decreasing emotional lability and irritability\textsuperscript{42}.

Repeated administration of DMPH leads to increased beta power, which is consistent with previous findings on psychostimulant effects on EEG in ADHD\textsuperscript{16,17}. During EC resting, beta power is associated with deactivation of the fronto-parietal attention network as well as sensory cortices\textsuperscript{32}. These associations, however, exhibited strong developmental trends and were weakest among children, perhaps explaining the lack of correlation with behavioral and cognitive measures in the current study. It is likely that measures of event-related beta band activity would show stronger association with specific aspects of cognitive performance given its association with attention and concentration in previous studies\textsuperscript{33}.

Finally, the effects of COMB medication on EEG spectral power were most prominent and resulted in ~20\% decrease in theta band power. In our study, lower theta band power was associated with greater treatment change in ADHD behaviors as well with improved cognitive functioning, such as fewer inattentive errors and lower reaction time variability. Elevated resting state theta power has been widely reported as a marker for ADHD; that the COMB group was had the lowest theta power relative to the other medication groups suggests a selective and more robust
targeted treatment effect moving towards normalization. In concurrent EEG-fMRI studies, eyes closed theta band power has been negatively correlated with default mode network activity\textsuperscript{26,32} and has been implicated in the balance of cortical excitation/inhibition\textsuperscript{34}. Aberrant connectivity within the DMN as well as weaker anti-correlation between the DMN and other task-positive networks have been associated with greater reaction time variability in ADHD\textsuperscript{35,36}, thus supporting the relationships found here among theta band power, DMN, and reaction time variability. Thus the combination of psychostimulant and $\alpha_2A$ agonist may have a synergistic effect of strengthening the DMN at rest, which has been reported to have weaker connectivity in ADHD\textsuperscript{37,38}.

To our knowledge, this study is the first to report medication effects of GUAN, DMPH, and COMB on cortical activation among a large sample of children with ADHD. Limitations of this study include that these results represent treatment effects that do not necessarily inform what occurs with long-term therapy. In addition, the participants were a selected population of youth with ADHD who were primarily Caucasian and relatively free of psychiatric co-morbidities. Caution should be used when generalizing these results to older individuals as well as non-Caucasian youth with ADHD and those with greater psychiatric co-morbidity. In addition, considerable variability in oscillatory activity within all of the EEG frequency bands have been noted, particularly between arousal states (i.e., eyes closed versus eyes open). Thus, EEG spectral power should not be used for clinical purposes such as making ADHD diagnosis or determining medication type or dosages. Further study is needed to examine medication effects in other resting and cognitive activation conditions to examine whether parallel effects are observed.

In conclusion, this study is the first to examine the medication effects of GUAN, DMPH, and COMB on cortical activation among youth with ADHD. Although all three medication conditions resulted in improvement of ADHD behavioral symptoms, differential medication effects on cortical activation emerged, suggesting different underlying neural mechanisms for each medication condition. Participants in the COMB condition exhibited decreased theta band and increased beta band power, which in turn was associated with better treatment response and improved cognitive
Participants in the GU AN condition had reduced alpha band power, which was associated with a slower and more variable reaction time. These results are consistent with previous studies on EEG correlates of medication response and suggest distinct neural effects of the medications singly and in combination predict later cognitive functioning. Whether or not the greater effects of COMB on these previously identified ADHD-related EEG correlates are durable over time and translate to longer-term benefits on clinical functioning should compel additional research.

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Drs. Loo, Cowen, Walshaw Welker, Levitt, Del’Homme, Mr. Cho, and Ms. Sturm report no biomedical financial interests or potential conflicts of interest.
REFERENCES


Table 1. Correlations between Week 8 EEG spectral power and behavioral characteristics and cognitive performance.

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<td>-0.10</td>
<td>0.11</td>
<td>0.02</td>
</tr>
<tr>
<td>Tx chg IN</td>
<td>-0.26**</td>
<td>-0.29***</td>
<td>-0.23**</td>
<td>0.14</td>
<td>0.12</td>
<td>-0.02</td>
</tr>
<tr>
<td>Tx chg HI</td>
<td>-0.22*</td>
<td>-0.25**</td>
<td>-0.23**</td>
<td>0.06</td>
<td>0.06</td>
<td>-0.08</td>
</tr>
<tr>
<td>WM factor</td>
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<td>-0.17</td>
<td>-0.20*</td>
<td>0.06</td>
<td>0.04</td>
<td>0.02</td>
</tr>
<tr>
<td>IN factor</td>
<td>-0.02</td>
<td>0.06</td>
<td>0.13</td>
<td>-0.21*</td>
<td>-0.22*</td>
<td>0.01</td>
</tr>
<tr>
<td>RT factor</td>
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<td>0.19*</td>
<td>0.24**</td>
<td>-0.32***</td>
<td>-0.30***</td>
<td>-0.16</td>
</tr>
<tr>
<td>RTV factor</td>
<td>0.27***</td>
<td>0.29***</td>
<td>0.30***</td>
<td>-0.31***</td>
<td>-0.29***</td>
<td>-0.07</td>
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Note: INsx=Number of inattentive symptoms at week 8, HI sx=Number of hyperactive-impulsive symptoms at week 8, Tx chg=treatment change (Baseline-Week 8), WM=working memory, IN=inhibition, RT=reaction time, RTV=reaction time variability. Arrows next to cognitive factors indicate the direction of score indicating better performance (i.e., higher scores on WM and IN indicate better performance; lower scores on RT and RTV suggest quicker reaction time and lower reaction time variability. * p<.05, ** p<.01, *** p<.005. Significant correlations are in bold, trend level findings are in italics.

Figure 1. Medication titration schedule. 8-week randomized 1:1:1, comparative parallel-group fixed-flexible dosing study of three treatments. GUAN=guanfacine, DMPH= d-methylphenidate, COMB=combination.
Figure 2. Resting state EEG spectral power topographs. Relative power for each medication group by frequency band and study visit. There were no significant differences in mean relative spectral power in any frequency band or region across the groups at baseline (all p’s >0.2).
Figure 3. Change in EEG spectral power from baseline by medication group. Each figure depicts regional change in relative power in each frequency band from baseline to week 8 for each medication group. Significant medication group differences are represented by * p<.05, ** p<.01 and significant time effects are represented by † p<.05, ††† p<.01, †††† p<.001. GUAN=guanfacine, DMPH=d-methylphenidate, COMB=combination.