Comparative Effects of Methylphenidate, Modafinil, and MDMA on Response Inhibition Neural Networks in Healthy Subjects

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Abstract

Background: Psychostimulants such as methylphenidate and modafinil are increasingly used by healthy people for cognitive enhancement purposes, whereas the acute effect of 3,4-methylenedioxymethamphetamine (ecstasy) on cognitive functioning in healthy subjects remains unclear. This study directly compared the acute effects of methylphenidate, modafinil, and 3,4-methylenedioxymethamphetamine on the neural mechanisms underlying response inhibition in healthy subjects.

Methods: Using a double-blind, within-subject, placebo-controlled, cross-over design, methylphenidate, modafinil, and 3,4-methylenedioxymethamphetamine were administrated to 21 healthy subjects while performing a go/no-go event-related functional magnetic resonance imaging task to assess brain activation during motor response inhibition.

Results: Relative to placebo, methylphenidate and modafinil but not 3,4-methylenedioxymethamphetamine improved inhibitory performance. Methylphenidate significantly increased activation in the right middle frontal gyrus, middle/superior temporal gyrus, inferior parietal lobule, presupplementary motor area, and anterior cingulate cortex compared with placebo. Methylphenidate also induced significantly higher activation in the anterior cingulate cortex and presupplementary motor area and relative to modafinil. Relative to placebo, modafinil significantly increased activation in the right middle frontal gyrus and superior/inferior parietal lobule, while 3,4-methylenedioxymethamphetamine significantly increased activation in the right middle/inferior frontal gyrus and superior parietal lobule.

Conclusions: Direct comparison of methylphenidate, modafinil, and 3,4-methylenedioxymethamphetamine revealed broad recruitment of fronto-parietal regions but specific effects of methylphenidate on middle/superior temporal gyrus, anterior cingulate cortex, and presupplementary motor area activation, suggesting dissociable modulations of response inhibition networks and potentially the superiority of methylphenidate in the enhancement of cognitive performance in healthy subjects.

Keywords: cognitive control, response inhibition, methylphenidate, modafinil, MDMA, fMRI
Significance Statement

Methylphenidate (MPH) and modafinil are being increasingly used by healthy people for cognitive enhancement purposes, whereas the acute effect of the party drug 3,4-methylenedioxymethamphetamine (MDMA, ecstasy) on cognitive functioning in healthy subjects remains unclear. This study directly compared the acute effects of MPH, modafinil, and MDMA on the behavioral and neural correlates of response inhibition in healthy subjects. Our findings reveal that MPH and modafinil but not MDMA improved inhibitory performance. MPH, modafinil, and MDMA revealed broad recruitment of fronto-parietal regions but specific effects of MPH on middle/superior temporal gyrus, anterior cingulate cortex, and presupplementary motor area activation. The additional recruitment of fronto-temporal regions might reflect the superiority of MPH for enhancing cognitive performance in healthy subjects.

Introduction

Successful response inhibition is essential to process and integrate incoming perceptual information in a flexible fashion (Bari and Robbins, 2013). Such a trial-by-trial adjustment of information processing is important for adapting ongoing behavior according to prevailing conditions in the respective environment. The inability to inhibit responses has been linked to several prevalent disorders, including obsessive-compulsive disorder, autism, and attention-deficit/hyperactivity disorder (ADHD). Response inhibition, as often operationalized by the go/no-go task (Bari and Robbins, 2013), has been associated with significant activation in a widespread network of brain regions including the superior, middle (MFG), and inferior frontal gyrus (IFG), inferior parietal lobule, striato-thalamic regions, and the presupplementary motor area (pre-SMA) / anterior cingulate cortex (ACC) (Simmonds et al., 2008; Swick et al., 2011). Although a complex neural circuit is involved in response inhibition, it is the activation in the right IFG that allows the inhibition of a prepotent motor response, a conclusion that is corroborated by connectivity and causality studies (Aron et al., 2014). Hence, the assessment of IFG activation helps to study the neuropharmacological mechanisms of response inhibition and to test the efficacy of potentially novel medications for disorders associated with impaired response inhibition.

A previous meta-analysis of stop-signal and go/no-go tasks showed that ADHD patients reveal reduced activation in the right IFG, SMA, and ACC, as well as striato-thalamic areas during response inhibition (Hart et al., 2013). It has been shown that methylphenidate (MPH), the most frequently prescribed drug for the treatment of ADHD (Briars and Todd, 2016), normalized fronto-temporal activation (including right IFG) during response inhibition in ADHD patients during the stop-signal (Rubia et al., 2014). The therapeutic effect of MPH is likely mediated through increases in extracellular levels of norepinephrine (NE) and dopamine (DA) in prefronto-striatal brain regions (Hannestad et al., 2010), both neurotransmitters critically involved in response inhibition (Bari and Robbins, 2013). Acute MPH administration also improved response inhibition during the stop-signal task (Nandam et al., 2011) and increased activation in the ACC, right IFG, superior temporal gyrus, caudate and left MFG, and left angular gyrus during the go/no-go task in healthy subjects (Nandam et al., 2014). In contrast, however, it has also been shown that MPH reduced activation in the right IFG/insula to infrequent stimuli associated with successful inhibition, failed inhibition, and attentional capture during stop-signal tasks in healthy subjects (Paula et al., 2012). Interestingly, another study found that MPH increased activation in the putamen only during inhibition errors but not during successful inhibition and only in the go/no-go but not stop-signal task (Costa et al., 2013).

Modafinil, licensed for the treatment of narcolepsy, shift-work disorder, and obstructive sleep apnea (Erman et al., 2007; Sheng et al., 2013), has also emerged as a possible agent to improve cognition. Both MPH and modafinil are being increasingly used as cognitive enhancers for nonmedical reasons (Sahakian et al., 2015). Modafinil produced cognitive enhancing effects and decreased stop-signal reaction time in healthy adults (Turner et al., 2003) and improved cognitive functioning in people with ADHD (Turner et al., 2004). Acute administration of modafinil to healthy subjects improved task performance during the stop-signal task but did not modulate brain activation (Schmaal et al., 2013). Modafinil is a weak inhibitor of DA and NE transporter and has additional effects on the brain GABA, glutamate, and orexin systems (Minzenberg and Carter, 2008), although the precise neuropharmacological mode of action of modafinil remains unclear.

Response inhibition is also strongly associated with integrity of the serotonergic (5-HT) system in humans and animals (Eagle et al., 2007). A reduction of 5-HT has been linked to impulsive, suicidal, and aggressive behavior, whereas high levels of 5-HT have been shown to decrease impulsive behavior (Pattij and Vanderschuren, 2008; Fitzgerald, 2011). A previous study demonstrated improved response inhibition in a stop-signal task after acute administration of 3,4-methylenedioxymethamphetamine (MDMA, ecstasy) in healthy subjects compared with placebo (Ramaekers and Kuypers, 2006). Given that MDMA acts primarily by releasing 5-HT from presynaptic terminals (Liechti and Vollenweider, 2000; Hysek et al., 2011, 2012), the authors suggested a possible link with 5-HT neurotransmission for this effect (Ramaekers and Kuypers, 2006). However, this MDMA-induced improvement in response inhibition is in contrast to another study reporting increased reaction times in a stop-signal task after MDMA administration to healthy controls (van Wel et al., 2012).

The present study directly compares for the first time the acute effects of MPH, modafinil, and MDMA on the behavioral and neural correlates of response inhibition in healthy controls. It thereby sought to elucidate the neural mechanisms underlying MPH’s and modafinil’s cognitive enhancing effect and to provide further insights into the neuropharmacological base of response inhibition. Using a within-subject, placebo-controlled, cross-over design, MPH, modafinil, and MDMA were administered to healthy subjects while performing a fMRI go/no-go task. Our a priori hypothesis was that MPH and modafinil but not MDMA would improve inhibitory performance during the go/no-go task compared with placebo. We further predicted that MPH and modafinil but not MDMA would increase brain activation (in particular IFG activation) during response inhibition.

Methods

The study was conducted in accordance with the Declaration of Helsinki and International Conference on Harmonization...
Guidelines in Good Clinical Practice and approved by the local Ethics Committee and Swiss Agency for Therapeutic Products (Swissmedic). The administration of MDMA to healthy subjects was authorized by the Swiss Federal Office for Public Health, Bern, Switzerland. The study was registered at ClinicalTrials.gov (NCT01951508). After receiving a written and oral description of the aim of this study, all participants gave written informed consent statements before inclusion.

Participants and Study Design
We recruited a group of 24 healthy subjects by word of mouth or an advertisement placed on the web market platform of the University of Basel. Subjects were excluded due to the following reasons: (1) chronic or acute medical condition including clinically relevant abnormality in physical exam, laboratory values, or ECG (in particular: seizures or a cardiac or neurological disorder); (2) current or previous psychotic or major affective disorder; (3) prior illicit drug use more than 5 times (except occasional use of THC-containing products) or any time within the previous 2 months; (4) pregnant or nursing women; (5) participation in another clinical trial (currently or within the last 30 days); (6) use of medications that are contraindicated or otherwise interfere with the effects of the study medications (monoamine oxidase inhibitors, antidepressants, sedatives, etc.); and (7) tobacco smoking (regularly > 10 cigarettes/d). Previous drug use is reported in supplementary Table 1. Subjects were asked to abstain from any illicit drug use including cannabis during the study, and drug tests were performed during screening and randomly before test sessions. There were no positive urine tests for stimulants, opioids, THC, or hallucinogens and therefore no exclusions from the study. Subjects were also asked to abstain from excessive alcohol consumption between test sessions and in particular to limit their use to one glass on the day before the test sessions.

Using a double-blind, placebo-controlled, randomized, within-subject design, participants received MPH (60 mg), modafinil (600 mg), MDMA (125 mg), and placebo in a counter-balanced order. The wash-out period between sessions was at least 7 days. Three participants did not complete all 4 go/no-go sessions, resulting in a final sample of 21 subjects (10 men, 11 women; mean age: 23.6 ± 2.8, range: 21–30).

MDMA was administered in a single absolute dose of 125 mg corresponding to a relatively high dose of (mean ± SD) 1.9 ± 0.3 mg/kg body weight. This dose of MDMA is in the high range of the doses typically used in clinical research (Kirkpatrick et al., 2014, 2015; Kuypers et al., 2017) and is within the dose range that is used recreationally (Brunt et al., 2012). MPH was administered in a single relatively high dose of 60 mg as done in previous studies (Martin et al., 1971; Korostenskaja et al., 2008). The subjective and cardiostimulant effects of this dose have previously been assessed on the same tests (Hysek et al., 2014) and have also been statistically compared with a lower dose of 40 mg (Schmid et al., 2014). The doses of MDMA and MPH were expected to be equivalent regarding their cardiovascular stimulant effects (Hysek et al., 2014). The therapeutic starting dose of modafinil is 100 mg and common doses are 400 mg/d. In this study, modafinil was administered in a single high dose of 600 mg. The goal was to use high single doses of all substances to maximize the subjective drug effects and reach responses close to \(E_{\text{max}}\).

Drug Administration
Each of the 4 test sessions lasted 7 hours. Subjects arrived at the laboratory at 8:45 AM. MPH, modafinil, MDMA, or placebo was administered orally at 9:45 AM. fMRI scanning was performed between 11:15 AM and 12:15 PM during the expected drug peak effects (Wong et al., 1998; Hysek et al., 2014; Schmid et al., 2014). The sessions ended at 3:45 PM. Additional study findings are reported elsewhere (Dolder et al., 2017).

Psychometric Assessment
The Adjective Mood Rating Scale (AMRS) (Janke, 1978) was used to assess subjective drug effects directly before (75 minutes posttreatment) and after (150 minutes posttreatment) the fMRI scanning took place. We averaged the values of the 75 and 150 min posttreatment assessments to best relate the subjective drug effects to the go/no-go task. In this study, we focused on the AMRS subscales related to cognitive control functioning such as activation, concentration, and performance-related activation. Performance-related activation is the sum of activation and concentration ratings.

The Go/No-Go Task
Ninety minutes after drug administration, all patients underwent an event-related go/no-go fMRI paradigm that was conducted with jittered inter-stimulus intervals (ISIs) and incorporated infrequently presented oddball stimuli to optimize statistical efficiency, that is, the accuracy with which the hemodynamic response to different stimuli can be estimated for a given amount of imaging time (Dale, 1999). The task is a well-validated paradigm used in previous fMRI studies (Rubia et al., 2006; Schmitz et al., 2006; Smith et al., 2006; Borgwardt et al., 2008; Lawrence et al., 2009; Atakan et al., 2013; Schmidt et al., 2013; Bhattacharyya et al., 2014, 2015; Daly et al., 2014), requiring either the execution or the inhibition of a motor response, depending on the visual presentation of the stimuli. The basic go task is a choice reaction time paradigm, in which arrows point either to the left or to the right side for 500 milliseconds, with a mean ISI of 1800 milliseconds (jitter range: 1600–2000 milliseconds). During go trials, subjects were instructed to press a left or right response button according to the direction of the arrow. In 11% of the trials, arrows pointing upward appeared. During these so-called no-go trials, participants were required to inhibit their motor response. During another 11% of the trials, arrows pointing left or right at a 23° angle were presented, and subjects were told to respond to these in the same way as for go stimuli (even though they pointed obliquely). These oddball stimuli were used to control for novelty effects associated with the low frequency and different orientation of the no-go relative to the go trials (stimulus-driven attention allocation). In total, there were 24 no-go, 160 go, and 24 oddball trials, with task duration of approximately 6 minutes.

Analyses of Inhibitory Performance and Subjective Feelings of Cognitive Controls
Behavioral task performance was evaluated by the probability of inhibition, correct number of go trials, and reaction time to go trials. Treatment differences in task performance and the 4 AMRS items were examined using a repeated-measures ANOVA with treatment as within-subject factor. Where the ANOVA null hypothesis of equal means was rejected, we used posthoc tests (Bonferroni).

fMRI Image Acquisition and Analysis
Scanning was performed on a 3T scanner (Siemens Magnetom Verio; Siemens Healthcare) using an interleaved T2*-weighted
echo planar imaging (EPI) sequence with 2.5-second repetition time, 28-millisecond echo time, a matrix size of 76 x 76, and 38 slices with 0.5-mm interslice gap, providing a resolution of 3 x 3 x 3 mm³ and a field of view of 228 x 228 cm². In total, 160 volumes were acquired.

EPIs were analyzed using an event-related design with SPM12 (www.fil.ion.ucl.ac.uk/spm). During preprocessing, images were realigned to the first image in the series, spatially normalized to the Montreal Neurological Institute template, and smoothed with a Gaussian kernel of 8 mm full half-width maximum. All images underwent visual inspection, and participants with a high number of severely corrupted images and/or gross artefacts were excluded (none). Additionally, all images were checked for movement artefacts, and all scans with more than 3 mm deviation from the previous scan in any dimension, resulting in corrupted volumes, were excluded and replaced with the average of the neighboring volumes (4 volumes were replaced in total, all after MDMA administration). Subjects with >10% corrupted volumes were excluded (none). There were no movement differences across treatment in any dimension (supplementary Table 2).

Voxel-wise maximum likelihood parameter estimates were calculated during the first-level analysis using the general linear model. Our design matrix included an autoregressive AR(1) model of serial correlations and a high-pass filter with a cut-off of 128 seconds. Onset times for go, no-go, and oddball trials across all 4 treatments were convolved with a canonical hemodynamic response function, and motion parameters acquired during the realignment procedure were added to the individual design matrix as multiple regressors. Five subject-specific contrast images were generated per participant: 4 images representing response inhibition controlled for the attentional oddball effect due to the low frequency occurrence of no-go trials (no-go vs. oddball trials) for each treatment (placebo, MPH, modafinil, MDMA); and 1 average image for response inhibition across all treatments (to compute the effect of task). A 1-sample t test was performed to examine whole brain activation during response inhibition across all treatments (effect of task). Treatment differences were examined using a within-subject ANOVA design. To control for drug order effects, we included drug order as a regressor of no interest into the ANOVA analysis. According to recent recommendations on cluster-extent based thresholding in fMRI analyses (Woo et al., 2014), significance was assessed at a cluster-level threshold of P < .05 family-wise error corrected across the whole brain, using an uncorrected cluster-forming threshold of P < .001 with an extent threshold of 20 voxels.

Results

Subjective Feelings of Cognitive Performance

All stimulants overall significantly increased scores on activation as revealed by a significant main effect of treatment (F (3, 18) = 3.622, P = .033). Posthoc testing showed a significantly enhanced activation following MPH administration relative to placebo (P = .018), while no other differences between drugs were found (Figure 1). We found no main effect of treatment for concentration (F (3, 18) = 1.953, P = .157; Figure 1). Furthermore, there was a main effect of treatment for performance-related activation (F (3, 18) = 3.323, P = .043). Bonferroni posthoc testing yielded that this effect was driven by a significant enhancement of performance-related activation after MPH intake relative to placebo (P = .022) (Figure 1).

Behavioral Task Performance

We found a significant treatment effect for probability of inhibition (F (3, 18) = 4.24, P = .020), indicating a generally improved inhibitory performance after stimulant exposures. Posthoc analysis revealed that MPH (P = .012) and modafinil (P = .038) but not MDMA improved inhibitory performance relative to placebo (Figure 2A).

Stimulants significantly increased the responses to go trials (F (3, 18) = 7.30, P = .002). The number of correct go trials was higher following MPH (P = .003) and modafinil (P = .005) but not MDMA administration compared with placebo (Figure 2B). There was also a significant treatment effect for reaction times (F (3, 18) = 6.19, P = .004), indicating faster reactions in response to go trials after stimulant intake. Relative to placebo, reaction times were significantly lower after MPH exposure (P = .003) (Figure 2C).

Brain Activation during Response Inhibition

Effect of Task

Combined treatment maps revealed significant activation in frontal, parietal, temporal, striato-thalamic, and cerebellar brain regions during response inhibition (supplementary Table 3).

Treatment Effects during Response Inhibition

Brain activation during response inhibition significantly differed across treatments in the right MFG, superior/inferior parietal lobule and supramarginal gyrus, and pre-SMA (supplementary Figure 1; supplementary Table 4).

Posthoc testing revealed that MPH increased brain activation relative to placebo in the right MFG, inferior parietal lobule, supramarginal gyrus, middle/superior temporal gyrus, as well as in the right ACC and pre-SMA (Figure 3A; supplementary Table 5). Compared with placebo, modafinil increased activation in the right MFG and superior and inferior parietal lobule, whereas MDMA increased activation in the right MFG (extending to the IFG), superior parietal lobule, and angular gyrus (Figure 3B-C; supplementary Table 5). Finally, ACC and pre-SMA activation was significantly higher after MPH compared with modafinil administration (Figure 3D; supplementary Table 5).
Discussion

To the best of our knowledge, this is the first within-subject study directly comparing the acute effects of MPH, modafinil, and MDMA on the behavioral and neural correlates of response inhibition in healthy controls. We found that both MPH and modafinil enhanced inhibitory performance with a concomitant increase in fronto-parietal activation. In addition, MPH was dissociable from modafinil and placebo in terms of its modulation of ACC and pre-SMA activation. Finally, MDMA did not alter...
inhibitory performance but was associated with alteration in fronto-parietal activation relative to placebo.

Besides their use for treating symptoms in ADHD patients (Briars and Todd, 2016), psychostimulants such as MPH and modafinil are being increasingly used by healthy people for cognitive enhancement purposes mainly to produce alertness and enhance professional performance (Sahakian et al., 2015). In the present study, both MPH and modafinil improved inhibitory performance in healthy people during the go/no-go task relative to placebo, supporting the view of beneficial effect of MPH and modafinil on cognitive performance in healthy people (Sahakian and Morein-Zamir, 2015). These findings are in line with other studies in healthy subjects demonstrating improved response inhibition (stop-signal reaction time) after acute MPH (Nandam et al., 2011) and modafinil (Turner et al., 2003; Schmaal et al., 2013).

Consistent with another go/no-go study in healthy subjects (Nandam et al., 2014), we found that MPH increased activation in right MFG, superior temporal gyrus, and ACC compared with placebo. MPH also increased activation in the right supramarginal gyrus, inferior parietal lobule, middle temporal gyrus, and pre-SMA in this study. Although MPH also increases DA levels in prefrontal regions, the observed increase in prefrontal activation here is likely mediated via NE neurotransmission (Chamberlain and Sahakian, 2007), given that in frontal regions MPH increases NE more than DA via reuptake inhibition of NE transporter (Hannestad et al., 2010). This corresponds with a previous fMRI go/no-go study, which showed that atomoxetine, a selective NE reuptake inhibitor increased fronto-temporal brain activation in healthy people (Chamberlain et al., 2009).

In the present study, acute modafinil administration also increased activation in the right MFG and superior/ inferior parietal lobule compared with placebo. Modafinil is a non-ephedrine psychostimulant that elevates synaptic NE and DA levels in prefrontal regions (de Saint Hilaire et al., 2001). Many of modafinil’s cognitive and behavioral effects are mediated by adrenergic receptors (Minzenberg and Carter, 2008). A previous fMRI study in healthy humans reported that modafinil increased activation in the locus coeruleus and prefrontal cortex and the functional coupling between these regions during a cognitive control task (Minzenberg et al., 2008). Another study in rats demonstrated that response inhibition in the stop-signal paradigm improved after modafinil and MPH administration, and these effects were not blocked by concurrent DA receptor antagonism, nor was response inhibition affected by DA receptor antagonism per se (Eagle et al., 2007). Furthermore, direct infusion of the alpha-2 adrenoceptor antagonist yohimbine into the prefrontal cortex of nonhuman primates impaired inhibitory control on a go/no-go paradigm and was associated with increased locomotor hyperactivity (Ma et al., 2005). Together, these findings suggest that increased prefrontal activation after modafinil and MPH administration is more likely mediated by increased levels of NE than DA. Such an interpretation resonates with previous works suggesting a key role for prefrontal NE neurotransmission in the inhibition of an already initiated response, whereas DA appears to modulate motor readiness for both inhibition and activation, potentially at the level of the striatum (Chamberlain and Sahakian, 2007; Bari and Robbins, 2013).

Party drugs such as MDMA are consumed recreationally for their acute mood- and social-enhancing effects. It has been shown that the acute psychological and physiological effects of MDMA in humans are mediated via an increase in 5-HT (Liechti et al., 2000, 2001; Liechti and Vollenweider, 2000). Acute MDMA administration has been shown to increase impulse control when 5-HT levels are high (Ramaekers and Kuypers, 2006). In the present study, acute MDMA administration did not affect inhibitory performance relative to placebo during the go/no-go task. This finding supports previous evidence that pharmacological manipulations of the 5-HT system have no detectable behavioral effects on response inhibition (Chamberlain and Sahakian, 2007). Although behavioral no-go effects of 5-HT interventions are often mild or absent in humans, neuroimaging has revealed altered activation in frontal regions. For instance, reduced IFG activation during the go/no-go task has been found after acute tryptophan depletion in healthy volunteers (Rubia et al., 2005), while the selective 5-HT reuptake inhibitor escitalopram increased activation in the ACC and middle frontal and temporal gyrus during successful inhibition during a stop-change paradigm (extension of stop-signal task) (Druiske et al., 2013). Along this line, citalopram also increased activation in the right dorsolateral prefrontal cortex and middle frontal gyrus in healthy subjects during the go/no-go task (Del-Ben et al., 2005). Consistent with these findings, we found increased activation in the right MFG/IFG after MDMA administration compared with placebo and also in the superior parietal lobule and angular gyrus. However, we cannot be certain whether this effect is mediated directly via a MDMA-induced increase in 5-HT or rather NE (Hysek et al., 2012). This interpretation fits with findings from a recent study showing that the psychotrophic effects of MDMA are not only mediated through 5-HT but also NE release (Hysek et al., 2011, 2012). However, although there were indications of improvements in all behavioral measures after MDMA intake in this study, the MDMA-induced increase in right IFG/MFG and inferior parietal lobule activation may not have been sufficient to improve inhibitory performance. The lack of effect on inhibitory performance following MDMA administration despite neural changes is intriguing. It supports previous studies using acute tryptophan depletion in healthy controls that failed to find effects on inhibitory control during the go/no-go task in healthy subjects (Rubia et al., 2005; Lamar et al., 2009). We can speculate that potential noradrenergic MDMA effects on inhibitory performance are offset by its serotonergic effects.

Finally, we found that acute MPH administration increased ACC and pre-SMA activation compared with modafinil. It has been proposed that the ACC is functionally interconnected with the basal ganglia and prefrontal cortex to form a key node of the response inhibition network (Aron et al., 2014). While the prefrontal cortex maintains goals and the basal ganglia suppress irrelevant motor responses, the ACC may detect response conflict (Aron et al., 2014). Therefore, ACC activation during response inhibition might reflect the conflict that occurs when 2 incompatible responses, such as whether to go or stop, are both compelling (MacDonald et al., 2000). Consistent with such an interpretation, a previous work showed that MPH acutely increased activation in the ACC and superior frontal gyrus for failed inhibitions during a stop-signal task, but only after controlling for attentional capture (Pauls et al., 2012). Furthermore, MPH also increased activation in the putamen during inhibition errors in the go/no-go task (Costa et al., 2013). Based on these findings and having in mind that we also controlled our imaging results for effects of stimulus-driven attention allocation, our finding suggests that MPH induced higher ACC activation in response to failed inhibitions than modafinil. ACC functioning has been associated with DA (Jocham and Ullasperger, 2009; Ko et al., 2009) and NE signalling (Aston-Jones et al., 2010; Aston-Jones and Cohen, 2005). Therefore, the MPH effect on ACC activation relative to modafinil is perhaps caused by differential effects of dopamine and noradrenaline, but future research is needed to explore the underlying neurochemical mechanisms.
modulation of DA pathways that project from the ventral tegmental (Williams and Goldman-Rakic, 1998) and of NE projections from the locus coeruleus (Aston-Jones et al., 2000). We might speculate that increased prefrontal activation after MPH is due to a higher blockade of DA and NE transporter relative to modafinil, leading to increased levels of DA and NE. In other words, the MPH-induced increase in prefrontal activation relative to modafinil is probably mediated through differential dynamic effects of MPH and modafinil on DA and NE in the prefrontal cortex (Rowley et al., 2014).

Some limitations of our study merit comment. Although we used a well-established paradigm from previous fMRI studies (Rubia et al., 2006; Schmitz et al., 2006; Borgwardt et al., 2008; Lawrence et al., 2009; Schmidt et al., 2013; Bhattacharyya et al., 2014, 2015; Daly et al., 2014), we were not able to disentangle neural activation in response to successful vs. failed inhibitions in the present study due to the modest number of no-go trials. The small number of inhibition trials (i.e., no-go trials) also limits the functional relevance of our behavioral results, albeit MPH and modafinil significantly increased the probability of inhibition. In this regard, it is also possible that the modest number of inhibition trials may explain the lack of alteration after MDMA administration. Future studies with a higher number of no-go trials are required to address these points. Furthermore, we cannot exclude effects on cerebral vasoactivity induced by the drugs (Honey and Bullmore, 2004), which might have confounded our fMRI results. For instance, it is possible that the effect of modafinil and MPH on MFG, ACC, and pre-SMA activation might be driven by their effects on regional cerebral blood flow in the same regions (Udo de Haes et al., 2007; Joo et al., 2008).

In conclusion, this study shows a common recruitment of fronto-parietal regions after MPH, modafinil, and MDMA but specific effects of MPH on middle/superior temporal gyrus, anterior cingulate cortex, and presurgical motor area activation, suggesting dissociable modulations of response inhibition networks and potentially the superiority of MPH in the enhancement of cognitive performance in healthy subjects. These effects are likely mediated via increased extracellular concentrations of NE, which may have reached the highest levels after MPH administration.

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Statement of Interest
None.

References
Aron AR, Robbins TW, Poldrack RA (2014) Inhibition and the right inferior frontal cortex: one decade on. Trends Cogn Sci 18:177–185.
Aston-Jones G, Cohen JD (2005) An integrative theory of locus coeruleus-norepinephrine function: adaptive gain and optimal performance. Annu Rev Neurosci 28:403–450.
Dolder PC, Müller F, Schmid Y, Borgwardt SJ, Liechti ME (2017) Direct comparison of the acute subjective, emotional, autonomic, and endocrine effects of MDMA, methylphenidate, and modafinil in healthy subjects. Psychopharmacology. doi:10.1007/s00213-017-4650-5.

Dreuke B, Schlaegel SM, Seifert A, Moeller O, Gründer G, Gauggel S, Boecker M (2013) The role of 5-HT in response inhibition and re-engagement. Eur Neuropsychopharmacol 23:830–841.

Eagle DM, Tufft MR, Goodchild HL, Robbins TW (2007) Differential effects of modafinil and methylphenidate on stop-signal reaction time task performance in the rat, and interactions with the dopamine receptor antagonist cis-flupenthixol. Psychopharmacology (Berl) 192:193–206.

Erman MK, Rosenberg R, Modafinil Shift Work Sleep Disorder Study Group (2007) Modafinil for excessive sleepiness associated with chronic shift work sleep disorder: effects on patient functioning and health-related quality of life. Prim Care Companion J Clin Psychiatry 9:188–194.

Fitzgerald PJ (2011) A neurochemical yin and yang: does serotonin activate and norepinephrine deactivate the prefrontal cortex? Psychopharmacology (Berl) 213:171–182.

Hannestad J, Gallezot JD, Planeta-Wilson B, Lin SF, Williams WA, van Dyck CH, Malison RT, Carson RE, Ding YS (2010) Clinically relevant doses of methylphenidate significantly occupy nor-epinephrine transporters in humans in vivo. Biol Psychiatry 68:854–860.

Hart H, Radua J, Nakao T, Mataix-Cols D, Rubia K (2013) Meta-analysis of functional magnetic resonance imaging studies of inhibition and attention in attention-deficit/hyperactivity disorder: exploring task-specific, stimulant medication, and age effects. JAMA Psychiatry 70:185–198.

Honey G, Bullmore E (2004) Human pharmacological MRI. Trends Pharmacol Sci 25:366–374.

Hysek CM, Simmler LD, Ineichen M, Grouzmann E, Hoener MC, Brenneisen R, Huwyler J, Liechti ME (2011) The norepinephrine transporter inhibitor reboxetine reduces stimulant effects of MDMA (“ecstasy”) in humans. Clin Pharmacol Ther 90:246–255.

Hysek CM, Simmler LD, Nicola VG, Vischer N, Donzelli M, Krähenbühl S, Grouzmann E, Huwyler J, Hoener MC, Liechti ME (2012) Duloxetine inhibits effects of MDMA (“ecstasy”) in vitro and in humans in a randomized placebo-controlled laboratory study. PLoS One 7:e36476.

Hysek CM, Simmler LD, Schilling N, Meyer N, Schmid Y, Donzelli M, Grouzmann E, Liechti ME (2014) Pharmacokinetic and pharmacodynamic effects of methylphenidate and MDMA administered alone and in combination. Int J Neuropsychopharmacol 17:371–381.

Janke W, Debus G (1978) Die Eigenschaftswörterliste EWL: Eine mehrdimensionale Methode zur Beschreibung von Aspekten des Befindens [The adjective word list EWL: A multidimensional method for describing aspects of the actual state]. Göttingen, Germany: Hogrefe.

Jocham G, Ullsperger M (2009) Neuropharmacology of performance monitoring. Neurosci Biobehav Rev 33:48–60.

Joo EY, Tae WS, Jung KY, Hong SB (2008) Cerebral blood flow changes in man by wake-promoting drug, modafinil: a randomized double blind study. J Sleep Res 17:82–88.

Kirkpatrick M, Delton AW, Robertson TE, de Wit H (2015) Prosocial effects of MDMA: A measure of generosity. J Psychopharmacol 29:661–668.

Kirkpatrick MG, Bagott MJ, Mendelson JE, Galloway GP, Liechti ME, Hysek CM, de Wit H (2014) MDMA effects consistent across laboratories. Psychopharmacology (Berl) 231:3899–3905.

Ko JH, Pitto A, Monchi O, Cho SS, Van Eimeren T, Pellecchia G, Ballyinger B, Rusjan P, Houle S, Strafella AP (2009) Increased dopamine release in the right anterior cingulate cortex during the performance of a sorting task: a [11C]FLB 457 PET study. Neurolmage 46:516–521.

Korostenskaja M, Kicić D, Kähkönen S (2008) The effect of methylphenidate on auditory information processing in healthy volunteers: a combined EEG/MEG study. Psychopharmacology (Berl) 197:475–486.

Kuyppers KP, Dolder PC, Ramaekers JG, Liechti ME (2017) Multifaceted evaluation of healthy volunteers after single doses of MDMA: a pooled sample of placebo-controlled studies. J Psychopharmacology 26:9981117699617.

Lamar M, Cutter WJ, Rubia K, Brammer M, Daly EM, Craig MC, Cleare AJ, Murphy DG (2009) 5-HT, prefrontal function and aging: fMRI of inhibition and acute tryptophan depletion. Neurobiol Aging 30:1038–1050.

Lawrence EJ, Rubia K, Murray RM, McGuire PK, Walshe M, Allin M, Giampietro V, Rifkin I, Williams SC, Nosarti C (2009) The neural basis of response inhibition and attention allocation as mediated by gestational age. Hum Brain Mapp 30:1038–1050.

Liechti ME, Baumann C, Gamma A, Vollenweider FX (2000) Acute psychological effects of 3,4-methylenedioxymethamphetamine (MDMA, “Ecstasy”) are attenuated by the serotonin uptake inhibitor citalopram. Neuropsychopharmacology 22:513–521.

Liechti ME, Geyer MA, Hell D, Vollenweider FX (2001) Effects of MDMA (ecstasy) on prepulse inhibition and habituation of startle in humans after pretreatment with citalopram, haloperidol, or ketanserin. Neuropsychopharmacology 24:240–252.

Liechti ME, Vollenweider FX (2000) The serotonin uptake inhibitor citalopram reduces acute cardiovascular and vegetative effects of 3,4-methylenedioxymethamphetamine (“Ecstasy”) in healthy volunteers. J Psychopharmacol 14:269–274.

Ma CL, Arnsten AF, Li BM (2005) Locomotor hyperactivity induced by blockade of prefrontal cortical alpha2-adrenoceptors in monkeys. Biol Psychiatry 57:192–195.

MacDonald AW, Cohen JD, Stenger VA, Carter CS (2000) Dissociating the role of the dorsolateral prefrontal and anterior cingulate cortex in cognitive control. Science 288:1835–1838.

Martin WR, Sloan JW, Sapira JD, Jasinski DR (1971) Physiologic, subjective, and behavioral effects of amphetamine, methamphetamine, ephedrine, phenmetrazine, and methylphenidate in man. Clin Pharmacol Ther 12:245–258.

Minzenberg MJ, Carter CS (2008) Modafinil: a review of neurochemical actions and effects on cognition. Neuropsychopharmacology 33:1477–1502.

Minzenberg MJ, Watrous AJ, Yoon JH, Ursu S, Carter CS (2008) Modafinil shifts human locus coeruleus to low-tonic, high-phasic activity during functional MRI. Science 322:1702–1702.

Nandam LS, Hester J, Bellgrove MA (2014) Dissociable and common effects of methylphenidate, atomoxetine and citalopram on response inhibition neural networks. Neuropsychologia 56:263–270.

Nandam LS, Hester R, Wagner J, Cummins TD, Garner K, Dean AJ, Kim BN, Nathan PJ, Mattingley JB, Bellgrove MA (2011) Methylphenidate but not atomoxetine or citalopram modulates inhibition effects. J Psychopharmacol 25:366–374.

Chemical actions and effects on cognition. Neuropsychopharmacology 33:1477–1502.

MacDonald AW, Cohen JD, Stenger VA, Carter CS (2000) Dissociating the role of the dorsolateral prefrontal and anterior cingulate cortex in cognitive control. Science 288:1835–1838.

Martin WR, Sloan JW, Sapira JD, Jassi DR (1971) Physiologic, subjective, and behavioral effects of amphetamine, methamphetamine, ephedrine, phenmetrazine, and methylphenidate in man. Clin Pharmacol Ther 12:245–258.

Minzenberg MJ, Carter CS (2008) Modafinil: a review of neurochemical actions and effects on cognition. Neuropsychopharmacology 33:1477–1502.

Minzenberg MJ, Watrous AJ, Yoon JH, Ursu S, Carter CS (2008) Modafinil shifts human locus coeruleus to low-tonic, high-phasic activity during functional MRI. Science 322:1702–1702.

Nandam LS, Hester R, Bellgrove MA (2014) Dissociable and common effects of methylphenidate, atomoxetine and citalopram on response inhibition neural networks. Neuropsychologia 56:263–270.

Pattij T, Vanderschuren LJ (2008) The neuropharmacology of impulsive behaviour. Trends Pharmacol Sci 29:192–199.

Pauls AM, O’Daly OG, Rubia K, Riedel WJ, Williams SC, Mehta MA (2012) Methylphenidate effects on prefrontal functioning...
during attentional-capture and response inhibition. Biol Psychiatry 72:142–149.

Ramaekers JG, Kuypers KP (2006) Acute effects of 3,4-methylenedioxymethamphetamine (MDMA) on behavioral measures of impulsivity: alone and in combination with alcohol. Neuropsychopharmacology 31:1048–1055.

Rowley HL, Kulkarni RS, Gosden J, Brammer RJ, Hackett D, Heal DJ (2014) Differences in the neurochemical and behavioral profiles of lisdexamfetamine methylphenidate and modafinil revealed by simultaneous dual-probe microdialysis and locomotor activity measurements in freely-moving rats. J Psychopharmacol 28:254–269.

Rubia K, Lee F, Cleare AJ, Tunstall N, Fu CH, Brammer M, McGuire P (2005) Tryptophan depletion reduces right inferior prefrontal activation during response inhibition in fast, event-related fMRI. Psychopharmacology (Berl) 179:791–803.

Rubia K, Smith AB, Woolley J, Nosarti C, Heyman I, Taylor E, Brammer M (2006) Progressive increase of frontal striatal brain activation from childhood to adulthood during event-related tasks of cognitive control. Hum Brain Mapp 27:973–993.

Rubia K, Alegría AA, Cubillo AI, Smith AB, Brammer MJ, Radua J (2014) Effects of stimulants on brain function in attention-deficit/hyperactivity disorder: a systematic review and meta-analysis. Biol Psychiatry 76:616–628.

Sahakian BJ, Bruhl AB, Cook J, Killikelly C, Savulich G, Pieryc T, Hafizi S, Perez J, Fernandez-Egea E, Suckling J, Jones PB (2015) The impact of neuroscience on society: cognitive enhancement in neuropsychiatric disorders and in healthy people. Philos Trans R Soc Lond B Biol Sci 370:20140214.

Sahakian BJ, Morein-Zamir S (2015) Pharmacological cognitive enhancement: treatment of neuropsychiatric disorders and lifestyle use by healthy people. Lancet Psychiatry 2:357–362.

Schmaal L, Joos L, Koelman M, Veltman DJ, van den Brink W, Goudriaan AE (2013) Effects of modafinil on neural correlates of response inhibition in alcohol-dependent patients. Biol Psychiatry 73:211–218.

Schmid Y, Hysek CM, Simmler LD, Crockett MJ, Quednow BB, Liechti ME (2014) Differential effects of MDMA and methylphenidate on social cognition. J Psychopharmacol 28:847–856.

Schmidt A, Walter M, Gerber H, Schmid Q, Smieskova R, Bendfeldt K, Wiesbeck GA, Riecher-Rössler A, Lang UE, Rubia K, McGuire P, Borgeardt S (2013) Inferior frontal cortex modulation with an acute dose of heroin during cognitive control. Neuropsychopharmacology 38:2231–2239.

Schmitz N, Rubia K, Daly E, Smith A, Williams S, Murphy DG (2006) Neural correlates of executive function in autistic spectrum disorders. Biol Psychiatry 59:7–16.

Sheng P, Hou L, Wang X, Huang C, Yu M, Han X, Dong Y (2013) Efficacy of modafinil on fatigue and excessive daytime sleepiness associated with neurological disorders: a systematic review and meta-analysis. PLoS One 8:e81802.

Simmonds DJ, Pekar JJ, Mostofsky SH (2008) Meta-analysis of Go/No-go tasks demonstrating that fMRI activation associated with response inhibition is task-dependent. Neuropsychologia 46:224–232.

Smith AB, Taylor E, Brammer M, Toone B, Rubia K (2006) Task-specific hypoactivation in prefrontal and temporoparietal brain regions during motor inhibition and task switching in medication-naïve children and adolescents with attention deficit hyperactivity disorder. Am J Psychiatry 163:1044–1051.

Swick D, Ashley V, Turken U (2011) Are the neural correlates of stopping and not going identical? Quantitative meta-analysis of two response inhibition tasks. Neuroimage 56:1655–1665.

Turner DC, Clark L, Dowson J, Robbins TW, Sahakian BJ (2004) Modafinil improves cognition and response inhibition in adult attention-deficit/hyperactivity disorder. Biol Psychiatry 55:1031–1040.

Turner DC, Robbins TW, Clark L, Aron AR, Dowson J, Sahakian BJ (2003) Cognitive enhancing effects of modafinil in healthy volunteers. Psychopharmacology 165:260–269.

Udo de Haes JI, Maguire RP, Jager PL, Paans AM, den Boer JA (2007) Methylphenidate-induced activation of the anterior cingulate but not the striatum: a [15O]H2O PET study in healthy volunteers. Hum Brain Mapp 28:625–635.

van Wel JH, Kuypers KP, Theunissen EL, Bosker WM, Bakker K, Ramaekers JG (2012) Effects of acute MDMA intoxication on mood and impulsivity: role of the 5-HT, and 5-HT, receptors. PLoS one 7:e40187.

Williams SM, Goldman-Rakic PS (1998) Widespread origin of the primate mesofrontal dopamine system. Cereb Cortex 8:321–345.

Wong YM, King SP, Laughton WB, McCormick GC, Grebow PE (1998) Single-dose pharmacokinetics of modafinil and methylphenidate given alone or in combination in healthy male volunteers. Journal of clinical pharmacology 38:276–282.

Woo CW, Krishnan A, Wager TD (2014) Cluster-extent based thresholding in fMRI analyses: pitfalls and recommendations. Neuroimage 91:412–419.