2008

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Meghan C. Campbell
Washington University School of Medicine in St. Louis

Morvarid Karimi
Washington University School of Medicine in St. Louis

Patrick M. Weaver
Washington University School of Medicine in St. Louis

J. Wu
Washington University School of Medicine in St. Louis

Dana C. Perantie
Washington University School of Medicine in St. Louis

See next page for additional authors

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RUNNING HEAD: STN DBS-INDUCED COGNITIVE VARIABILITY

Neural correlates of STN DBS-induced cognitive variability in Parkinson disease

M.C. Campbell, Ph.D.¹, M. Karimi, M.D.², P.M. Weaver¹, J. Wu¹, D.C. Perantie¹, N.A. Golchin², S.D. Tabbal, M.D.², J. S. Perlmutter, M.D.²,³,⁴,⁵, and T. Hershey, Ph.D.¹,²,³

Departments of Psychiatry¹, Neurology², Radiology³, Anatomy and Neurobiology⁴ and Program in Physical Therapy⁵, Washington University School of Medicine, St Louis, MO

Correspondence to:
Tamara Hershey, Ph.D.
Campus Box 8225
Washington University School of Medicine
4525 Scott Avenue
St. Louis, MO 63110
Phone: 314-362-5593
Fax: 314-362-0168
email: tammy@wustl.edu

Disclosure: Dr. Karimi received partial fellowship funding from Medtronic, Inc., the manufacturer of the implanted stimulators; no other authors have any conflicts of interest to disclose.
Abstract

Background: Although deep brain stimulation of the subthalamic nucleus (STN DBS) in Parkinson disease (PD) improves motor function, it has variable effects on working memory (WM) and response inhibition (RI) performance. Currently, little is known about the relationship between the neurophysiological response to STN DBS and cognitive functioning. The purpose of the present study was to determine the neural correlates of STN DBS-induced variability in cognitive control performance. Methods: We measured bilateral STN DBS induced blood flow changes (PET and [15O]-water on one day) in the supplementary motor area (SMA), dorsolateral prefrontal cortex (DLPFC), anterior cingulate cortex (ACC), and right inferior frontal cortex (rIFC) as well as WM and RI changes (Spatial Delayed Response and Go-No-Go tasks on the next day) in 24 PD participants. On both days, participants withheld PD medications overnight and conditions (DBS off vs. bilateral on) were administered in a counterbalanced, double-blind manner. Results: As predicted, STN DBS-induced change in DLPFC blood flow correlated with STN DBS-induced change in WM error, but not RI performance. Furthermore, change in ACC blood flow correlated with change in RI but not WM performance. Both were inverse relationships, such that increased blood flow related to decreased cognitive performance in response to STN DBS. Conclusions: The results of the present study demonstrate that the variability in the effects of STN DBS on cognitive control performance relates to STN DBS-induced cortical blood flow changes. This relationship highlights the need to further understand the mechanism(s) of STN DBS as variability in stimulation characteristics could alter behavioral and cortical responses.
Keywords: Parkinson disease, deep brain stimulation, working memory, response inhibition, PET
Deep brain stimulation of the subthalamic nucleus (STN DBS) provides effective treatment of the motor symptoms for many individuals with Parkinson disease (PD) (Limousin et al., 1995). However, recent work suggests that STN DBS has a more variable effect on cognitive functioning (Voon, Kubu, Krack, Houeto, & Troster, 2006) and may even negatively affect select cognitive processes, especially cognitive control (Temel, Blokland, Steinbusch, & Visser-Vandewalle, 2005). In fact, a recent review of the literature revealed that approximately 41% of individuals with STN DBS experience cognitive problems (Temel et al., 2006). Cognitive dysfunction in non-demented individuals with PD is common (30-70%) (Green et al., 2002); can significantly decrease perceived quality of life (Schrag, Jahanshahi, & Quinn, 2000); can impair functional ability in work and home environments (Weintraub, Moberg, Duda, Katz, & Stern, 2004); and is a relatively common effect of STN DBS (Temel et al., 2006). To further optimize this therapeutic technique, it is important to determine the mechanisms that produce the variability in cognitive effects of STN DBS. This information may also provide new insights into basal ganglia-thalamocortical pathway involvement in cognitive control.

Although the role of the prefrontal cortex (PFC) in cognitive control processes has been emphasized, the STN also may contribute to cognitive control systems (Baunez et al., 2001; Chudasama, Baunez, & Robbins, 2003; Nakano, Kayahara, Tsutsumi, & Ushiro, 2000) through its connections to the PFC (Alexander, Crutcher, & DeLong, 1990; Alexander, DeLong, & Strick, 1986). For example, working memory (WM) processes rely on the dorsolateral prefrontal cortex (DLPFC) (Braver et al., 1997; Goldman-Rakic, 1990; Wager & Smith, 2003), whereas performance on response inhibition (RI) tasks is often associated with
the anterior cingulate cortex (ACC) and the right inferior frontal cortex (rIFC) (Aron, Robbins, & Poldrack, 2004; Barch et al., 2001; Botvinick, Nystrom, Fissell, Carter, & Cohen, 1999; Braver, Barch, Gray, Molfese, & Snyder, 2001; Konishi et al., 1999; Menon, Adleman, White, Glover, & Reiss, 2001; Nee, Wager, & Jonides, 2007; Wager et al., 2005). According to the most commonly accepted models of frontal-striatal circuitry (Alexander et al., 1990; Alexander et al., 1986; Middleton & Strick, 2000), basal ganglia output directly targets prefrontal cortex including DLPFC and ACC, and the STN plays an important role in these circuits (Temel et al., 2005). Recent evidence also suggests that rIFC connects to STN via the “hyperdirect” pathway (Aron, Behrens, Smith, Frank, & Poldrack, 2007). A functional role of the STN in these cognitive control processes is supported by the effects of lesions of the STN in animals, specifically demonstrating poor cognitive control and impaired ability to inhibit responses under conditions of strong conflict (Baunez et al., 2001; Baunez, Nicoullon, & Amalric, 1995; Baunez & Robbins, 1997; Temel et al., 2005).

However, STN DBS has variable effects on tasks that rely on cognitive control (Voon et al., 2006; Temel et al., 2005) whereas those that do not depend heavily on cognitive control (e.g., non-declarative memory, decision-making, visuomotor sequencing, and language) appear to be relatively unaffected by STN DBS (Funkiewiez et al., 2004; Halbig et al., 2004; Morrison et al., 2004a). Two aspects of cognitive control that have contradictory findings across studies are WM and RI performance. STN DBS has been demonstrated to improve performance (Pillon et al., 2000a; van den Wildenberg et al., 2006), impair performance (Dujardin, Defebvre, Krystkowiak, Blond, & Destee, 2001; Hershey et al., 2004; Witt et al.,
The variability in STN DBS responses across studies may, in part, reflect methodological differences. The most obvious difference across studies is the use of different cognitive tasks. However, other methodological limitations also may contribute to these discrepancies. For example, small sample size limits confidence in some findings (N < 15; c.f., Jahanshahi et al., 2000; Schroeder et al., 2002; Schroeder et al., 2003). Medication (i.e., levodopa) effects may have confounded several previous investigations (c.f., Pillon et al., 2000b; Trepanier et al., 2000; Witt et al., 2004) since levodopa has its own variable effects on cognition (Cools, 2006), making it difficult to isolate the specific effects and mechanisms of STN stimulation. Other studies focused on the surgical effects of STN DBS rather than the direct manipulation of only stimulation (c.f., Saint-Cyr, Trepanier, Kumar, Lozano, & Lang, 2000; Trepanier, Kumar, Lozano, Lang, & Saint-Cyr, 2000).

Although methodological differences may have contributed to some of the discrepancies across studies, other factors must explain variability of STN DBS-induced cognitive effects across individuals within a study. Clarification of these other factors requires an understanding of the neurophysiological mechanisms of STN DBS which still remain unclear. The most commonly proposed mechanisms of STN DBS are that it 1) blocks local neuronal activity (“conduction block”), 2) excites local inhibitory afferent neurons to reduce neuronal output, or 3) directly excites output neurons (Montgomery, Jr. & Baker, 2000; Perlmutter & Mink, 2006). In support of the latter mechanism, PET studies have revealed 2004), or to have no significant effect on cognitive control performance (Morrison et al., 2004).
that STN stimulation increases subcortical while decreasing cortical blood flow (Hershey et al., 2003; Schroeder et al., 2003), suggesting that stimulation increases STN output leading to increased thalamic inhibition of cortical activity (Hershey et al., 2003). Furthermore, PET studies have demonstrated that STN DBS increased blood flow or glucose metabolism in both the thalamus as well as areas of the frontal, temporal, and parietal cortex (Hilker et al., 2004; Trost et al., 2006). Additional PET studies have also been conducted with the inclusion of motor or cognitive tasks during PET scanning (Schroeder et al., 2002; Strafella, Dagher, & Sadikot, 2003). Interestingly, STN stimulation was reported to decrease activation in the ACC during a response inhibition (Stroop) task and the decreased activity in the ACC correlated with decreases in Stroop interference performance (Schroeder et al., 2002). However, because these studies assessed the effects of stimulation on blood flow changes during a cognitive task, it is impossible to separate the effects of stimulation on blood flow from the effects of the cognitive task on blood flow (Schroeder et al., 2002; Strafella et al., 2003). Thus, these studies have limited ability to determine the neurophysiologic underpinnings of STN DBS’s effects on cognitive control.

Therefore, the purpose of the present study was to investigate the possible neural correlates of STN DBS induced variability in cognitive control, focusing on the relationships between STN DBS blood flow responses in cortical areas and WM and RI performance. To address this issue, we correlated regional PET blood flow change with WM and RI change induced by STN DBS in people with PD. We predicted that DBS-induced blood flow changes in the DLPFC would be associated with DBS-induced changes in WM performance, whereas DBS-induced blood flow changes in the ACC would be related to RI performance. Based on the
growing evidence that the rIFC is important for successful RI performance, we also explored DBS-induced blood flow changes in the rIFC as another possible neural correlate of STN DBS induced changes in RI performance. Finally, we also examined stimulation-induced blood flow changes in the supplementary motor area (SMA). Although we expected stimulation-induced changes in blood flow in the SMA, we did not expect any specific relationship between blood flow in the SMA and cognitive performance. Thus, the SMA region served as a “control” to guard against the possibility of a more global relationship between blood flow and cognitive performance. Importantly, we assessed the effects of stimulation on blood flow and cognitive functioning separately, without the confound of medication, thus focusing exclusively on the effects of stimulation.

Materials and Methods

Participants. Twenty-nine individuals with PD and previously implanted bilateral STN stimulators were studied. Each met the diagnostic criteria for clinically definite PD (Racette, Rundle, Parsian, & Perlmutter, 1999). Exclusionary criteria included a history of neurological events or diagnoses other than PD, or dementia on clinical exam prior to surgery. The surgical implantation of stimulators (Medtronic model 3389 DBS leads) targeted STN with a technique that combines conventional stereotactic planning using formulas with reference to the anterior-posterior commissural line, visual targeting on T2 weighted magnetic resonance imaging (MRI), frame-based targeting using computerized methods (Medtronic STEALTH STATION, Framelink IV) and microelectrode recording (Tabbal et al., 2007). The degree of subsequent clinical benefit achieved by stimulation, as measured by change in UPDRS motor subscore 3 (mean improvement in total UPDRS motor
scores was 52%, (SD = 14); paired t-test OFF versus ON, t(22)= 9.8, p < .001), is comparable to other centers. Participants had to be at least 2 months post-STN stimulator implantation to allow time for programming to achieve optimal clinical benefit for each individual. All participants except 3 were taking levodopa/carbidopa daily and 19 were taking other PD medications (e.g. amantadine, pergolide, pramipexole, or entacapone). All participants except two were right handed. This study was approved by the Institutional Review Board at Washington University School of Medicine and all participants gave informed consent.

Data from the 29 PD participants who had adequate PET scans were examined. However, one participant was excluded for being an outlier (>2.5 SDs from the mean) on blood flow change in the ACC region, and 4 participants were excluded for having invalid GNG data due to tremor. On average, the remaining 24 participants with valid PET and cognitive data (10 female, 14 male) were 61.7 years old (SD = 9.2), had been diagnosed with PD for 12.8 years (SD = 4.7) and were tested 8.7 months (SD = 5.9) following STN stimulator implantation.

**Overview of protocol.** Participants were assessed on two consecutive days. On each study day, participants refrained from taking any PD medications for at least 12 hours prior to testing and were tested with both stimulators off (OFF condition) and with both stimulators on (ON condition). Order of stimulation condition was counterbalanced across participants and both participants and examiners were blind to condition. On the first study day participants had PET scans with STN DBS either ON or OFF and on the second day had cognitive testing and UPDRS motor ratings under the same STN DBS conditions.
**PET Scans.** PET scans were performed at least 42 minutes after change of the stimulator settings. Participants were scanned on the Siemens/CTI ECAT EXACT HR 47 tomograph (Wienhard et al., 1994) in 2D acquisition mode with a 15 cm axial field of view, 3.125 mm center to center slice separation and simultaneous collection of 47 slices. A 20 gauge plastic catheter was placed in an antecubital vein for $^{15}$O-water injection. The participant was positioned in the PET scanner using cross laser lines and a Polyform mask. The mask was marked on the participant’s face to detect any change in the head position relative to the mask. Attenuation was measured using three rotating rod sources of $^{68}$Ge/$^{68}$Ga. During each scan lights were dimmed and the room was quiet while participants kept their eyes closed. About 50 mCi of $^{15}$O-water was injected intravenously as a bolus followed by two minutes of data acquisition. Each participant had as many as six separate PET scans in each study block. The scans were 14 minutes apart to permit adequate radioactive decay. After the first block, the participant was removed from the scanner to change settings. A second attenuation scan and a second set of six PET scans were done at least 42 minutes after the settings change.

The entire study was recorded with two video cameras; one camera recorded the head and the other recorded the body during the PET scans. Videos were reviewed to exclude possible movement during data acquisition. In addition, surface EMG electrodes over bilateral biceps, wrist flexors, quads and gastroc muscles were used to detect any muscle activation without visible movement. EMG signals were amplified at a gain of 2000 and filtered on line with a band pass of 10Hz-1000 kHz. The EMG signals were monitored on line and stored on
computer using a CED micro 1401 interface (Cambridge Electronic Design). Codes were inserted into the data stream to coincide with the onset and offset of each scan. Any observed movement, loud noise, etc was remarked with a separate code.

Cognitive Testing

Spatial Delayed Response (SDR) Task. The SDR task is an experimentally derived working memory task (Hershey, Craft, Glauser, & Hale, 1998) that has been closely linked to lateral prefrontal cortex functioning in animals and humans (Funahashi, Bruce, & Goldman-Rakic, 1989; Funahashi, Bruce, & Goldman-Rakic, 1993; Funahashi, Chafee, & Goldman-Rakic, 1993; Goldman-Rakic, Funahashi, & Bruce, 1990; Luciana, Depue, Arbisi, & Leon, 1992). A central fixation cross appeared on a computer screen placed approximately 40 cm away from the participant. While fixated, either one or two cues (each 1 cm in diameter) appeared for 150 msec in any of 32 possible unmarked locations at an 11.5 cm radius from the central fixation. A delay period (5 or 15 sec) was then imposed. During the delay, participants performed a continuous performance task (CPT) in which a series of geometric shapes (triangle, square and diamond) appeared in place of the fixation cross (1000 msec duration, 750-1250 msec inter-trial interval). Participants pressed the spacebar whenever the diamond shape appeared. After the delay, the fixation cue returned, and participants pointed on the computer screen where they remembered seeing the cue(s). Responses were measured in X and Y coordinates and compared to the actual location of the cue. Delay trials and trials with no mnemonic load (cue-present trials) were presented in random order. On the cue-present trials the cue (dot) was present during the response phase. This set of trials gave an indication of participants' pointing accuracy. Mean error in mm (distance between recall and
actual target) was calculated for each participant for each type of trial. There were either one or two cues to be remembered on each trial. In the two-cue condition, both locations were presented simultaneously, and in the recall phase, participants pointed to both locations, in any order desired. Forty experimental trials were presented, 20 with only one cue presented and 20 with two cues presented. Trials were blocked by number of cues and the order of blocks was counterbalanced across participants. Participants performed 4 cue-present trials and 8 test trials per delay (i.e., 16 cue-absent trials) for each block.

**Go-No-Go (GNG) Task.** The GNG task assessed the ability to inhibit a prepotent response under conditions of low or high prepotent response strength (Barch et al., 2001; Braver et al., 2001; Casey et al., 1997), and requires active cognitive control processes such as conflict monitoring. This task involved monitoring a visual display while single uppercase letters were presented one at a time interspersed with the number “5” (250 msec duration, 1000 msec intertrial interval). In this task, participants were instructed to push a target response button at the occurrence of every letter but to withhold a response when the number “5” was presented. Target frequency (percent of trials where a button press was required, e.g. letters) was manipulated in a blocked fashion. There were two levels of target frequency (medium = 50%; high = 83%). The high target frequency block is designed to produce a strong prepotent response (e.g. press the button) so is more challenging for response inhibition skills. One block at each frequency level was performed with the order randomly determined for each participant. Each block contained 150 trials. Reaction times and accuracy rates were recorded.

**Analyses**
PET Scans. PET emission scans were reconstructed using filtered back projection and measured attenuation. Only the initial 40 seconds of the data after the arrival of the radioactive water in the brain were analyzed for each scan. Images were smoothed with a three dimensional Gaussian filter to a final resolution of 16 mm full width at half maximum. All images were coregistered to the initial emission image (Woods, Cherry, & Mazziota, 1992). The two transmission images from each participant were coregistered to each other and averaged. This average attenuation image was resliced to match each emission image and new attenuation corrections were forward projected. All emission scans were reconstructed a final time using these coregistered, averaged attenuation corrections. The new PET images were coregistered and resliced to a standard mean blood flow image in Talairach atlas (Talairach & Tournoux, 1988) space with 12-parameter fit. Individual images were normalized using mean whole brain counts and masked to include only the voxels in common among all scans. Scans were excluded from further analyses if the patient had tremor, other movements or sustained substantial EMG activity above background noise within the 40 seconds of data acquisition. Participants’ data were retained for analysis if they had at least one usable scan per condition.

Based on the functional neuroimaging evidence that the DLPFC is involved in working memory processes and that the ACC and rIFC are involved in response inhibition, these regions of interest (ROIs) were specifically chosen. The ROIs were defined on the basis of previous fMRI results with similar versions of the SDR (Leung, Gore, & Goldman-Rakic, 2002) and GNG (Braver et al., 2001) tasks. A 10.4 mm diameter (equivalent to the width of five voxels) sphere was placed on coordinates for left and right DLPFC (Talairach coordinates of -34, 44, 27 and 34, 44, 27, respectively; (Leung et al., 2002)) a midline
anterior cingulate cortex region (Talairach coordinates of 3, 19, 35; (Braver et al., 2001)), and the right IFC (Talairach coordinates of 41, 16, 19 (Konishi et al., 1999)). See Figure 1 for illustration of the location of these coordinates. Mean blood flow within each sphere was obtained for each participant’s OFF and ON average blood flow image, averaging across left and right DLPFC. In addition, as a “control” region, we also extracted the mean blood flow in the SMA for each participant’s OFF and ON average blood flow image. Percent change in blood flow from OFF to ON conditions was calculated as the dependent measure.

**Effects of stimulation.** To determine the effects of STN stimulation on blood flow and cognitive control, one sample t-tests were performed for change in blood flow and change in performance on the SDR and GNG tasks. To simplify analyses and thus reduce the number of correlations performed, correlational analyses focused on the more difficult conditions of both tasks previously found to be more sensitive to the effects of STN DBS (Hershey et al., 2004). For the SDR, this was the 2 cue condition; for the GNG, this was a discriminability measure (Pr; [target accuracy – (1 – nontarget accuracy)]) from the high demand condition. Of note, both of these measures are based on performance accuracy and are not dependent on motor speed, thus reducing the potential influence of motor symptoms (e.g., bradykinesia) on cognitive performance. Significance level was set at $p < .05$.

**Cognitive and PET correlations.** Percent change in SDR and GNG performance from bilateral OFF to bilateral ON conditions was calculated for each measure and correlated (Pearson r) with blood flow change in DLPFC, ACC, rIFC, and SMA. To determine the strength of these correlations, potential confounding variables were included as covariates in
hierarchical linear regression models. For these models, change in task performance was the dependent variable, and age, disease severity (measured by off medication, off stimulation UPDRS score), and finally change in regional blood flow were entered in a hierarchical fashion. The unique relationship between blood flow and task performance change ($R^2$ change), after effects of age and disease severity were removed, was tested for significance. Significance for all analyses was set at $p < .05$.

**Results**

**Cognitive Performance.** For the SDR task of working memory, there was a significant main effect of task difficulty ($F(1,22) = 32.77$, $p < .001$) but not a significant main effect of stimulation ($p = .67$) or interaction between task difficulty and stimulation ($p = .84$). Similarly, for the GNG task of response inhibition, there was a main effect of task difficulty ($F(1,23) = 35.61$, $p < .001$), but no significant effect of stimulation ($p = .30$) or interaction between task difficulty and stimulation condition ($p = .13$). See Table 1 for descriptive statistics of task performance. Based on the effects of task difficulty and previous evidence that the more difficult conditions of both of these tasks are most likely to be affected by stimulation (Hershey et al., 2004), subsequent analyses focused on the more difficult condition of each task.

**Effects of stimulation.** As expected, stimulation significantly improved motor symptoms (UPDRS: $t(23) = 8.76$, $p < .001$). In contrast, there were no significant differences in blood flow in the selected regions or cognitive performance between the ON and OFF stimulation conditions. Changes in blood flow (DLPFC: -1.49 (SD = 9.55); ACC: 0.07 (SD = 2.94);
rIFC: -0.62 (SD = 3.72); SMA: -0.56 (SD = 2.53)) and changes in cognitive performance (SDR: 4.92 (SD = 20.37); GNG: -5.32 (SD = 22.19)) were not significantly different from zero (all $p$s > .24). However, the trend was for impaired performance on both cognitive tasks with STN DBS ON.

**Cognitive and PET correlations.** There were no significant relationships between stimulation-induced changes in regional blood flow; blood flow changes in the SMA was not related to blood flow changes in the DLPFC, ACC, or rIFC (all $p$s > .17) and there were no significant relationship among blood flow changes in the DLPFC, ACC, or rIFC (all $p$s > .10). Similarly, stimulation-induced changes on SDR performance did not correlate with changes in GNG performance ($p = .30$).

In support of our hypothesis, change in DLFPC blood flow correlated with change in SDR performance ($r = .52, p = .009$), but not GNG performance ($r = -.16, p = .45$). Also as predicted, change in ACC blood flow correlated with change in GNG performance ($r = -.44, p = .03$), but not SDR performance ($r = .14, p = .52$). Both of these significant relationships indicated that impaired performance was related to increases in blood flow with stimulation, whereas improved performance was related to decreases in blood flow with stimulation (See Figure 2 A & B). The correlations between blood flow changes in DLPFC and ACC and changes in cognitive performance remained significant even after covarying change in UPDRS motor score, age and disease severity (partial correlations, $p$s <.038). Furthermore, hierarchical linear regression analyses demonstrated that, after controlling for UPDRS motor score, age, and disease severity, stimulation-induced blood flow changes significantly
predicted stimulation-induced cognitive performance changes (SDR and DLPFC, $R^2 = .24$, $F(1,20) = 6.6$, $p = .018$; GNG and ACC change, $R^2 = .20$, $F(1,20) = 5.1$, $p = .035$).

Although our primary hypotheses were supported, our secondary analyses revealed that there was no relationship between stimulation-induced changes in rIFC and stimulation-induced changes in cognitive control performance (SDR: $p = .81$; GNG: $p = .65$). As anticipated, stimulation-induced changes in SMA blood flow were not correlated with stimulation-induced changes in cognitive performance (SDR: $p = .14$; GNG: $p = .94$).

**Discussion**

Despite the beneficial effects of STN DBS on the motor symptoms of PD, the effects on cognition are highly variable. The present study demonstrates that STN DBS-induced change in WM performance is associated with change in regional blood flow in the DLPFC, while change in RI task performance is associated with change in regional blood flow in the ACC. These correlations were specific and as predicted; they are consistent with frontal-striatal circuitry and the neurophysiological effects of STN DBS as well as the functional role of these areas in cognitive control. These relationships were not due to participant characteristics such as age, motor symptom severity, or motor benefit from STN DBS. In addition, the results indicate an inverse relationship between regional blood flow and cognitive control as measured by both tasks; STN DBS-induced *increases* in regional blood flow were associated with *decreased* cognitive control, whereas STN DBS-induced *decreases* in regional blood flow were associated with *increased* cognitive control performance. However, the factor(s) contributing to the variability in responses to STN
stimulation, which may mediate the relationship between blood flow changes and cognitive control, are yet to be determined.

The key finding of the present study is that the stimulation-induced change in cognitive control performance was inversely related to the stimulation-induced change in regional blood flow in the DLPFC and ACC. Importantly, the relationship between cognitive control and regional blood flow changes was not limited to just the degree of change, but also the direction. Participants with stimulation-induced decline in cognitive control performance demonstrated stimulation-induced increased blood flow in the relevant cortical regions whereas participants with improved cognitive control performance had reduced rCBF in these same regions (see Figure 2). The specific relationships between blood flow change in the DLPFC with WM and blood flow change in ACC with RI performance further support the notion that these areas are important for cognitive control (Braver et al., 2001; Leung et al., 2002).

Despite the evidence from other studies demonstrating that rIFC is involved in response inhibition, our exploratory analysis of DBS-induced blood flow responses in this area did not indicate any relationship with stimulation-induced changes in GNG performance. Several possible reasons may explain this negative finding. First, methodological differences may account for our seemingly discrepant findings. For example, the majority of the research demonstrating involvement of the rIFC in response inhibition has utilized the Stop Signal Task (c.f., Aron & Poldrack, 2006), which requires the inhibition of an already initiated response, whereas the GNG task in the present study requires the inhibition of a prepotent
(but not yet initiated) response. In fact, there is even evidence for differences in cortical activation based on the specific GNG task that is used (Simmonds, Pekar, & Mostofsky, 2008). A second possibility may relate to individual or group differences. The participants in the current study were primarily older adults, who may rely on slightly different functional neuroanatomy for response inhibition (c.f., Nielson, Langenecker, & Garavan, 2002).

Finally, and most likely, it is highly probable that rIFC is critical for response inhibition, but that it simply is not significantly affected by STN stimulation. The DLPFC and ACC, however, are specific targets of frontal-striatal circuits (Alexander et al., 1990; Alexander et al., 1986; Middleton et al., 2000), demonstrate altered rCBF with STN stimulation (Sestini et al., 2002), and are also involved in WM and RI performance.

Several caveats regarding this study should be noted. Our main findings are based upon correlational analyses and do not prove causal relationships between the STN DBS-induced changes in blood flow and cognitive control performance. However, this type of analysis does provide meaningful information. If the STN DBS-induced behavioral change depends on functional modifications of basal ganglia-prefrontal circuits that can be measured by cortical blood flow changes, then these changes should be correlated. Our findings support this interpretation and are consistent with a potential causal relationship. Although we failed to confirm a significant STN DBS-induced impairment of cognitive control, these results fit with the high degree of variability in cognitive and blood flow responses to STN DBS across studies (Burn & Troster, 2004; Takeshita et al., 2005; Temel et al., 2005; Voon et al., 2006). In this study, the mean change in cognitive response to STN stimulation was in the direction of impairment but did not reach statistical significance due to the high variability across
participants. In fact, this variability across participants permitted us to identify significant correlations between changes in behavioral performance and blood flow response to STN DBS. The factors driving these differing individual responses have not yet been identified nor have their mechanisms of action been delineated, thus requiring further investigation.

A change in blood flow could reflect alterations in interneuronal activity within these regions (DLPFC or ACC), changes in input from distant pathways such as the basal ganglia-thalamo-cortical circuits, or both, since regional blood flow changes reflect neuronal activity in target synaptic fields. Therefore, measurements of stimulation-induced blood flow changes permit insight into possible underlying mechanisms (Hershey & Mink, 2006). Some have hypothesized that STN DBS forces “regularization” of irregular STN output leading to improved motor performance in people with PD (Vitek, 2002). Although regularization of STN output may improve motor function, a forced regular rate of firing may interfere with the phasic burst firing related to cognitive control processes (Funahashi et al., 1989; Kropotov & Etlinger, 1999; Schultz, 1997). Our findings support this idea. Stimulation-induced increased input or interneuronal activity in DLPFC or ACC could override the firing patterns that support optimal cognitive control functioning. Likewise, reduced input or interneuronal activity in these prefrontal regions could reflect decreased competition or noise thereby permitting firing patterns underlying cognitive function to operate more optimally. The crucial next step is to identify the factor(s) that determine the neurophysiological response to STN DBS as this may also mediate the cognitive response to STN stimulation.
It is possible that stimulation variables, such as the precise location of the active electrode contact, the extent of the field of stimulation (Morrison et al., 2004b; Smeding et al., 2007; Temel et al., 2006) or patient variables, such as degree of dopaminergic denervation (Foster, Black, Antenor-Dorsey, Perlmutter, & Hershey, 2007; Hershey et al., 2007) could modulate cognitive control as well as the direction and degree of change in associated prefrontal cortical blood flow. For example, without direct visual identification of electrode contacts within the brain, uncertainty remains regarding their precise location and the spatial extent of the effects of stimulation, both of which may contribute to STN DBS effects (Temel et al., 2005). The frequency, voltage, and amplitude of STN stimulation also could influence cortical functioning (Strafella et al., 2003; Temel et al., 2005). Future studies that incorporate the exact location of contacts as well as stimulation variables, including the degree and strength of current spread, may be useful in understanding the physiological characteristics of the anatomical pathways underlying the cognitive effects of stimulation.
Acknowledgments

Supported by: the Greater St. Louis Chapter of the American Parkinson Disease Association (APDA), NIH (NS41248; NS41509; DA07261; NS057105), APDA Advanced Center for PD Research at Washington University, and the Barnes-Jewish Hospital Foundation (Elliot Stein Family Fund and Jack Buck Fund for PD Research).
Reference List


Table 1

Cognitive control performance across stimulation conditions (OFF v. ON).

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<thead>
<tr>
<th>Stimulation Condition</th>
<th>Bilateral OFF</th>
<th>Bilateral ON</th>
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<td><strong>SDR Task</strong></td>
<td></td>
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</tr>
<tr>
<td>One Cue Error</td>
<td>18.06 (6.77)</td>
<td>18.02 (5.02)</td>
</tr>
<tr>
<td>Two Cue Error</td>
<td>23.05 (5.85)</td>
<td>23.28 (5.19)</td>
</tr>
<tr>
<td><strong>GNG Task</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Medium Frequency RT</td>
<td>525.31 (100.37)</td>
<td>498.43 (67.86)</td>
</tr>
<tr>
<td>High Frequency RT</td>
<td>486.09 (98.31)</td>
<td>439.50 (80.87)</td>
</tr>
<tr>
<td>Medium Frequency Pr</td>
<td>0.84 (0.17)</td>
<td>0.84 (0.17)</td>
</tr>
<tr>
<td>High Frequency Pr</td>
<td>0.75 (0.18)</td>
<td>0.71 (0.20)</td>
</tr>
</tbody>
</table>

Note. Values shown as mean (SD). SDR error in mm. Pr = discriminability index. RT = response time in msec.
**Figure 1.** Locations of the (A) anterior cingulate cortex (ACC), (B) right inferior frontal cortex (rIFC), and (C) dorsolateral prefrontal cortex (DLPFC) regions of interest. A 10.4mm sphere was placed on these locations.
Figure 2. Scatterplots showing the significant correlations between (A) change in DLPFC blood flow and change in WM, as indicated by SDR errors (larger numbers reflect *decreased* performance with stimulation), and (B) change in ACC blood flow and change in RI, as indicated by GNG discriminability (larger numbers reflect *improved* performance with stimulation). Both correlations indicate that stimulation-induced decreased cognitive control performance is associated with stimulation-induced increased blood flow responses.