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**Role of functional imaging in the development and refinement of invasive neuromodulation for psychiatric disorders**

Williams NR *et al*. Imaging in DBS for neuropsychiatric disorders

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**Abstract**

Deep brain stimulation (DBS) is emerging as a powerful tool for the alleviation of targeted symptoms in treatment-resistant neuropsychiatric disorders. Despite the expanding use of neuropsychiatric DBS, the mechanisms responsible for its effects are only starting to be elucidated. Several modalities such as quantitative electroencephalography as well a intraoperative recordings have been utilized to attempt to understand the underpinnings of this new treatment modality, but functional imaging appears to offer several unique advantages. Functional imaging techniques like positron emission tomography, single photon emission computed tomography and functional magnetic resonance imaging have been used to examine the effects of focal DBS on activity in a distributed neural network. These investigations are critical for advancing the field of invasive neuromodulation in a safe and effective manner, particularly in terms of defining the neuroanatomical targets and refining the stimulation protocols. The purpose of this review is to summarize the current functional neuroimaging findings from neuropsychiatric DBS implantation for three disorders: treatment-resistant depression, obsessive-compulsive disorder, and Tourette syndrome. All of the major targets will be discussed (Nucleus accumbens, anterior limb of internal capsule, subcallosal cingulate, Subthalamic nucleus, Centromedial nucleus of the thalamus-Parafasicular complex, frontal pole, and dorsolateral prefrontal cortex). We will also address some apparent inconsistencies within this literature, and suggest potential future directions for this promising area.

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**Key words:** Deep brain stimulation; Functional neuroimaging; Functional magnetic resonance imaging; Functional magnetic resonance imaging; Cortical stimulation; Nuclear imaging

**Core tip:** Deep brain stimulation (DBS) is emerging as a powerful tool for the alleviation of targeted symptoms in treatment-resistant neuropsychiatric disorders. Most recently, functional magnetic resonance imaging has been used to examine the effects of focal DBS on activity in a distributed neural network. The purpose of this review is to summarize the current functional neuroimaging findings from neuropsychiatric DBS implantation and to discuss some apparent inconsistencies within this literature, and to suggest potential future directions for this promising area.

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**INTRODUCTION**

Deep brain stimulation (DBS) is a form of invasive neuromodulation in which electrodes are implanted into a neuroanatomical target (nucleus or fiber tract) in an effort to modulate activity throughout a neural network[[1](#_ENREF_1)]. Neuropsychiatric disorders can also be modulated through stimulation of cortical nodes in a neural network with similar efficacy and reduced risk[[2](#_ENREF_2),[3](#_ENREF_3)]. The cortex can be stimulated either non-invasively from outside the skull with techniques like transcranial magnetic stimulation (TMS), or by placing stimulation paddles close to the surface of the brain under the skull but outside of the dura (epidural). For the purpose of this review, these cortical stimulation studies with invasive stimulation paddles and external generators are also labeled DBS, even though the electrodes are not deep. The imaging issues with these invasive cortical stimulation methods are the same as for traditionally defined DBS. These network modulations are driven by stimulation parameters that can be optimized for maximal therapeutic benefits. DBS was first developed for movement disorders wherein high frequency electrical stimulation of the thalamus has the capacity to reduce tremors[[4](#_ENREF_4)]. Since that time, DBS has been investigated for treatment-resistant neuropsychiatric disorders[[5](#_ENREF_5),[6](#_ENREF_6)]. In this context, high frequency electrical stimulation of limbic system targets has the capacity to alter mood and alleviate affective symptoms. Similar to DBS for movement disorders[[7](#_ENREF_7)], invasive neuromodulation for neuropsychiatric disorders has similar reported open-label efficacy rates as lesion surgeries (although never directly compared) but unlike lesion surgery offers reversibility and fewer side-effects[[8](#_ENREF_8),[9](#_ENREF_9)].

Neuropsychiatric disorders are beginning to be classified based on limbic, cognitive, and motoric[[10](#_ENREF_10),[11](#_ENREF_11)] circuit dysfunction[[12](#_ENREF_12)]. DBS likely exerts its effects by directly modulating the target node(s) and indirectly modulating the regions connected to that node[[1](#_ENREF_1),[13](#_ENREF_13)]. Functional neuroimaging studies have attempted to capture these downstream network effects induced by DBS[[13](#_ENREF_13)]. While traditional nuclear medicine techniques such as positron emission tomography (PET) imaging have successfully captured gross DBS-induced changes in neural networks, functional magnetic resonance imaging (fMRI) is poised to offer more refined insights[[14](#_ENREF_14)]. Additionally, basic science techniques like optogenetics are now being used to explore the precise neural changes associated with DBS in animal models of disease[[15](#_ENREF_15)].

This review begins by describing the functional imaging techniques that have been used to examine the effects of focal DBS on activity in a distributed neural network. Thereafter the three neuropsychiatric syndromes of interest will be introduced and their DBS targets briefly identified. The subsequent sections describe the cortical and subcortical targets for each disorder in further detail, as well as discuss their imaging results and relevance to neuropsychiatric symptoms. Table 1 summarizes advantages, disadvantages and cautions for each imaging method. Tables 2-5 and Figures 1-4 are categorized by neuropsychiatric disease and illustrate the effect each DBS target has on downstream nodes.

***PET***

Positron emission tomography (PET) can safely and effectively probe the effects of DBS without disrupting the function of the device or its electrodes. PET uses short-lived radioisotopes to create a static image of averaged neural activity over time based on changes in oxygen metabolism, glucose, blood flow or neurotransmitter receptor binding[[16](#_ENREF_16),[17](#_ENREF_17)]. PET data are static in that they are averaged over time. Thus, PET cannot yield information about the immediate (*e.g.*, second to second) time course of change induced by an experimental manipulation, although oxygen PET can image activity for one minute and can thus show changes over a few minutes.

PET is the predominant functional imaging technique in the psychiatric DBS literature, in part because it was available much earlier than modalities like functional magnetic resonance imaging (fMRI), and because it is technically less difficult to combine DBS with PET[[14](#_ENREF_14)]. Although PET continues to make significant contributions to clinical Neuroscience, the technique has a number of limitations that are relevant to DBS research. First, PET has poor temporal and spatial resolution relative to fMRI. These resolution limitations might make it difficult to accurately assess DBS-induced changes in small subcortical nodes in limbic neural networks. Second, PET requires injections of radioactive tracers into patients who have no medical indication for such exposure. Furthermore, the cost and availability of these radiotracers can be prohibitive. Third, PET studies typically have methodological variations that make it difficult to compare data between studies. For example, a study that uses repeated injections of 15O-H2O PET to measure cerebral blood perfusion over 60-120 s intervals[[18](#_ENREF_18)] cannot be contextualized with a study that uses a single injection of 18F-fluorodeoxyglucose (18F-FDG) PET to measure glucose metabolism over a 60 min interval[[19](#_ENREF_19)]. Fourth, PET measures of binding capacity are not specific for receptor occupancy and could reflect altered receptor density[[20](#_ENREF_20),[21](#_ENREF_21)] . This lack of clarity might make it difficult to assess the pharmacological effects of DBS. These are just a few of the considerations that should be kept in mind when considering PET DBS research.

On the other hand, there are few if any problems with performing PET studies in patients with DBS electrodes implanted, unlike with fMRI. Also, single photon emission computed tomography (SPECT) imaging, with perfusion tracers that capture activity summed over a minute or two, offers a less expensive and readily available option[[18](#_ENREF_18),[22](#_ENREF_22),[23](#_ENREF_23)] .

***fMRI***

One of the most popular ways to assess brain function is to employ fMRI. Most fMRI studies use blood oxygen level dependent (BOLD) signals that are based on magnetization differences between oxygenated and deoxygenated hemoglobin[[24](#_ENREF_24)]. Thus, fMRI can be used to make inferences about neural activity based on time-locked alterations in neurovascular coupling. Although BOLD fMRI is widely used, it does have limitations. The BOLD hemodynamic response takes about 1-3 s to peak, which is a lifetime in terms of neuronal firing. Also, the BOLD signal reflects general vascular changes arising from new metabolic demands, and is not precise to the actual neurons driving the signal. That is, the BOLD response is anatomically imprecise or “smudged” with respect to the actual activated neurons, like “the garden being flooded for want of one thirsty flower”[[25](#_ENREF_25)]. Finally, BOLD changes occur both when a brain region is acting to excite activity, and when it is attempted to regulate of inhibit brain activity. Thus the precise interpretation of the BOLD signal direction is problematic.

In the DBS literature, fMRI is often used to examine the function of neural network(s) before and after targeted stimulation[[26](#_ENREF_26),[27](#_ENREF_27)]. Safety considerations such as MRI-induced lead migration and secondary heat lesioning have greatly slowed the utilization of this technology in patients with DBS implanted electrodes[[28](#_ENREF_28)]. There is also some concern that fMRI may deactivate implanted impulse generators (IPGs), although this phenomenon has been linked to low battery[[29](#_ENREF_29)] and more recent studies have not observed it[[30](#_ENREF_30)]. To address some of these concerns, DBS-fMRI studies have typically been conducted utilizing externalized leads connected to pulse generators that are housed in the fMRI control room. This setup may reduce the risk of hardware malfunction[[31](#_ENREF_31)]. This also means that most DBS fMRI studies are performed in humans immediately after surgical implantation and before connecting the DBS electrodes to the generator implanted in the chest wall. This makes longitudinal studies problematic.

Many of the original DBS-fMRI studies were performed during the roughly 2-3-wk interval between lead implantation and IPG placement. During this time, stimulator extension wires can be accessed for externalized operation and stimulation[[32](#_ENREF_32)]. While this approach is useful for examining the short-term effects of DBS on the limbic/cognitive networks, it cannot be used to assess long-term effects that may evolve or change over time[[33](#_ENREF_33)]. Furthermore, in the weeks after surgery, a “microlesion effect” may result from mechanical manipulation at the site of lead placement. Such a lesion could itself have a unique functional imaging signature[[34](#_ENREF_34)] and therefore complicate predictions about the long-term effects of limbic DBS. These microlesion effects have also been shown to have unanticipated effects on adjacent neural circuits[[35](#_ENREF_35),[36](#_ENREF_36)]. IPG replacement surgery has been proposed as an optimal window of time during which the chronic effects of DBS may be explored[[37](#_ENREF_37)].

There are several sources of potential MRI artifacts in patients with DBS implants: 1-the skull-cap, 2- the metallic portion of the electrode contact, 3- the electrical stimulation itself, and 4-movement[[38](#_ENREF_38),[39](#_ENREF_39)]. Some of these artifacts can be reduced or eliminated. Some of these artifacts are disease specific, such as the participant having tics in the scanner[[40](#_ENREF_40)]. The artifact generated by the skull-cap, for example, can be addressed by replacing ferromagnetic screws with non-ferromagnetic options[[41](#_ENREF_41)] or by post-processing techniques normally used for sinus artifacts[[30](#_ENREF_30)]. Other artifacts, like those generated by the metallic portion of the lead during echoplanar acquisition[[29](#_ENREF_29),[42](#_ENREF_42)], are more difficult to address. The deep brain stimulation itself can also create MR artifact in the area around the tissue-lead interface. This has been addressed by turning off the device minutes before the scan[[13](#_ENREF_13)]. This artifact, if present, prevents analysis of DBS-induced activity in the area surrounding the electrode. Some authors have chosen to address these issues with strict statistical thresholding in offline analysis[[43](#_ENREF_43),[44](#_ENREF_44)].

Despite its limitations, DBS-fMRI can provide unique and valuable information regarding the neurobiological effects of DBS[[45](#_ENREF_45),[46](#_ENREF_46)]. Most psychiatric DBS-fMRI studies have only utilized the device in the OFF condition[[13](#_ENREF_13)]. Newer protocols, however, have begun to explore the possibility that DBS-fMRI could be performed in the ON condition in patients with fully-implanted DBS hardware[[30](#_ENREF_30),[43](#_ENREF_43)]. One recent study successfully imaged Parkinson’s disease patients with subthalamic nucleus (STN) DBS in the ON and OFF setting with no reported adverse effects[[43](#_ENREF_43)].This protocol utilized a 1.5 Tesla MRI scanner as well as a special transmit-receive (T/R) head coil. The specific absorption ratio (SAR) in the head was limited to under 0.1 W/kg for this experiment. This protocol was first performed in a phantom, demonstrating no significant heating with the sequences utilized in the human study[[30](#_ENREF_30)]. We encourage anyone considering such a technique to consult with a magnetic resonance physicist and take great caution as this is a risky undertaking. While this protocol appears to be safe, it is yet to be utilized in patients with DBS for psychiatric conditions.

**NEUROPSYCHIATRIC DISEASE AND THEIR DBS TARGETS**

This portion of the review will focus on the imaging of DBS in three neuropsychiatric syndromes: 1-obsessive-compulsive disorder (OCD), 2-major depressive disorder (MDD), and 3-Tourette syndrome (TS). The following sections will identify common DBS targets for each disorder and review relevant functional imaging data.

***Obsessive-compulsive disorder and its DBS targets***

Obsessive-compulsive disorder (OCD) is a neuropsychiatric condition characterized by functionally impairing obsessions and compulsions. The obsessions occur as recurrent and ego-dystonic ideas, images, or impulses. By contrast, the compulsions are stereotyped, repetitive mental acts or behaviors that are performed with the intention of reducing anxiety generally related to the obsessions. OCD affects approximately 2%-3% of the population and 20%-40% of patients with OCD remain severely disabled despite conventional treatments like exposure/response prevention as well as medications such as selective serotonin reuptake inhibitors. DBS for severe, treatment-resistant OCD was recently approved through a humanitarian device exemption[[47](#_ENREF_47)]. Several targets have been explored for this indication, including the nucleus accumbens (NAc)[[48](#_ENREF_48),[49](#_ENREF_49)], the anterior limb of the internal capsule (ALIC)[[50](#_ENREF_50)], the ventral capsule/ ventral striatum (VCVS)[[51-53](#_ENREF_51)] and the STN[[54](#_ENREF_54),[55](#_ENREF_55)]. Four of the main OCD targets that have been imaged with PET include the ALIC[[56](#_ENREF_56)], VCVS[[57](#_ENREF_57)], STN[[58](#_ENREF_58)], and NAc[[50](#_ENREF_50),[59](#_ENREF_59)](Table 2, Figure 1). Functional MRI has been performed following NAc DBS in both animals[[26](#_ENREF_26)] and more recently in humans with OCD[[13](#_ENREF_13)].

***Major depressive disorder and its DBS targets***

Depression is a psychiatric disorder characterized by extreme sadness and/or melancholia that is of sufficient length and interferes with activities of daily living as well as socialization. One in five people will experience an episode of major depression at some point in their lifetime. The World Health Organization has indicated that major depressive disorder is one of the four most disabling illnesses worldwide[[60](#_ENREF_60)]. Currently, antidepressants and psychotherapy are the primary modes of treatment, although that is evolving[[61](#_ENREF_61)]. Minimally invasive brain stimulation methods like electroconvulsive therapy (ECT) and transcranial magnetic stimulation (TMS) are typically reserved for patients with treatment-resistant depression[[62](#_ENREF_62)]. DBS is currently being utilized in a clinical research setting for patients who do not respond to the aforementioned therapies. Three of the main depression targets that have been imaged with PET include the dorsolateral prefrontal cortex[[3](#_ENREF_3)], subcallosal cingulate[[63](#_ENREF_63),[64](#_ENREF_64)], and the nucleus accumbens[[65](#_ENREF_65), [66](#_ENREF_66)] (Table 3, Figure 2). Two of the depression targets are also OCD targets have been imaged for OCD, but not depression include the anterior limb of the internal capsule[[56](#_ENREF_56)], ventral capsule/ventral striatum)[[57](#_ENREF_57)]. Functional MRI has been performed in NAc in non-depressed animals[[26](#_ENREF_26)].

***Tourette syndrome***

Tourette syndrome (TS) is a neuropsychiatric movement disorder that affects approximately 1% of the world’s population, typically in childhood and/or adolescence. TS is characterized by at least two motor tics and one or more vocal tics that persist for more than a year. Approximately 90% of individuals with TS have at least one comorbid psychiatric symptom. These symptoms include attention-deficit/hyperactivity disorder (ADHD), OCD, anxiety disorders, or impulse control disorders (ICD). DBS for TS has been shown to reduce the frequency and severity of motor tics as well as their associated psychiatric symptoms[[47](#_ENREF_47)]. Although the precise mechanism of action remains unclear, DBS for TS appears to diminish dopamine signaling in the thalamus and the striatum[[16](#_ENREF_16)]. Multiple targets have been introduced for TS DBS, including the thalamic centromedian nucleus and parafasicular complex (CM-pf)[[67](#_ENREF_67)], the globus pallidus internus (GPi)[[68](#_ENREF_68)], the ALIC, and NAc[[69](#_ENREF_69)] (Table 4, Figure 5) While three of these targets (GPi[[70](#_ENREF_70)], ALIC, and NAc) have had functional neuroimaging performed in other conditions (as described above), only the medial thalamus has been imaged in the TS population[[16](#_ENREF_16),[17](#_ENREF_17)]. In a large animal model, both the CM-Pf and GPi have been imaged using fMRI[[27](#_ENREF_27), [71](#_ENREF_71)].

**DBS TARGETS AND DOWNSTREAM NODES IN NEUROPSYCHIATRIC DBS**

***Cortical targets and nodes***

**Dorsolateral prefrontal cortex (BA 9 and BA 46):** The dorsolateral prefrontal cortex (DLPFC) (BA 9 and 46)[[72](#_ENREF_72)] is a critical node in the mesocortical system, a dopaminergic tract that modulates anticipation, goal selection, planning monitoring, and the use of feedback in task performance[[73](#_ENREF_73)]. The DLPFC is critical for working memory of both spatial and non-spatial information[[74](#_ENREF_74)]. There has been some suggestion that direct stimulation (activation) of the DLPFC may serve to modulate parietal attentional networks involved in the automatic processing of salient environmental stimuli. For example, a recent meta-analysis of 162 imaging studies revealed that co-activation of BA 9 (which spans the dorsolateral and medial prefrontal cortices) and midbrain regions like the periaqueductal gray (PAG) is essential for assigning emotional valence[[75](#_ENREF_75)]. These results may partially explain how the DLPFC plays such a critical role in mood regulation[[76](#_ENREF_76)]. In depression, the left DLPFC is hypoactive and thought to be associated with negative emotional judgment. The right DLPFC is hyperactive and this is linked to attentional modulation[[77](#_ENREF_77)]. The DLPFC has been demonstrated to have various subcortical connections, including but not limited to the mediodorsal nucleus of the thalamus[[78](#_ENREF_78)], the caudate[[79](#_ENREF_79)], hippocampus[[80](#_ENREF_80)], subcallosal cingulate[[81](#_ENREF_81)], and amygdala[[82](#_ENREF_82)] to name a few.

Unilateral stimulation of the DLPFC has bilateral effects[[83](#_ENREF_83)] as long as the corpus callosum is intact. While there have been varied downstream neural network changes in response to non-invasive magnetic stimulation of the DLPFC, it is clear that the effects of DLPFC stimulation are not isolated to a single node in the network. There is some suggestion that the antidepressant efficacy of non-invasive, repetitive transcranial magnetic stimulation of the left DLPFC sites is related to the anticorrelation of the subgenual cingulate, further supporting that the effects of DLPFC stimulation modulated deep structures in the neural network[[81](#_ENREF_81)]. It is also important to note that there are individual differences in DLPFC connectivity and these difference appear to affect efficacy of stimulation at this node[[84](#_ENREF_84)].

Left as well as bilateral epidural DLPFC stimulation has been shown to have a positive effect on depression[[2](#_ENREF_2),[3](#_ENREF_3)], and when combined with bilateral epidural frontopolar epidural cortical stimulation (EpCS) appears to be a durable therapy for treatment–resistant depression[[85](#_ENREF_85)]. Left DLPFC EpCS causes contralateral activation of the right DLPFC as well as superior frontal gyrus, cuneus, and posterior cingulate. In the Van Laere study, chronic, bilateral ALIC stimulation reduced right DLPFC activity on PET[[56](#_ENREF_56)]. Conversely, in the Sturm study, DBS of the right NAc was shown to increase the activity of the right DLPFC in one patient on PET[[59](#_ENREF_59)]. Subcallosal cingulate gyrus (SCG) DBS has been demonstrated to increase metabolism in bilateral DLPFC[[63](#_ENREF_63)] in an earlier study, then decrease DLPFC bilaterally in a later study[[64](#_ENREF_64)] at essentially the same time points of 3 and 6 mo[[63](#_ENREF_63),[64](#_ENREF_64)]. An increase in activity of the DLPFC was also seen with fMRI of NAc DBS in large animals[[26](#_ENREF_26)]. In a functional connectivity analysis of NAc DBS, the efficacy of the stimulation was related to reduced hyperconnectivity between the lateral (and medial) prefrontal cortex and the NAc [[13](#_ENREF_13)].

**Frontopolar cortex (BA 10):** Frontopolar cortex (FPC) is the most uniquely human area in the hominid brain and is the only part of the prefrontal cortex (PFC) that has no direct inputs from the sensory cortex[[86](#_ENREF_86),[87](#_ENREF_87)]. Several studies have demonstrated the frontal pole has a significant role in self-reflection, long-term goals, past or future events, or hypothetical scenarios[[87-89](#_ENREF_87)]. Damage to this area impairs an individual’s ability to multitask[[90](#_ENREF_90)]. The FPC functions to consider events beyond the present moment[[87](#_ENREF_87)]. Pathological patterns of rumination and self-reflection are core features of depression. The FPC is being more and more implicated in the pathogenesis of depression as well as in its recovery[[91](#_ENREF_91),[92](#_ENREF_92)]. A recent neuroimaging meta-analysis demonstrated a consistent finding of increased resting-state activity in the frontal pole (BA10) in patients with depression[[93](#_ENREF_93)].

Invasive neuromodulation over the FPC (along with DLPFC) bilaterally has been implanted in 5 individuals with treatment-resistant depression, termed epidural cortical stimulation[[2](#_ENREF_2)]. Furthermore, subgenual cingulate DBS appears to only be effective when it contacts the white matter tracts that cause downstream changes in the frontal polar cortex[[47](#_ENREF_47),[94](#_ENREF_94),[95](#_ENREF_95)]. The tractographic connection to the frontal pole was the invasive neuromodulation target common between the ALIC target and the SCG target[[2](#_ENREF_2),[96](#_ENREF_96)]. Chronic deep brain stimulation of NAc reduces activity in the gyri that terminate in the area of the frontal polar cortex[[65](#_ENREF_65)]. The therapeutic effects of NAc DBS for OCD are related to reducing functional connectivity between the NAc and the FPC[13](#_ENREF_13)].

The frontal pole can be divided into a medial (ventromedial prefrontal cortex (vmPFC) part that is seen in all primates and a lateral part (the lateral frontopolar cortex (lFPC)) that is seen only in humans[[86](#_ENREF_86)]. Activity of the vmPFC has been shown to be increased with NAc deep brain stimulation in an animal model, but these animals were not noted to be depressed[[26](#_ENREF_26)]. Furthermore, it is unclear the function of the vmPFC in non-primate mammals, although it has been consistently demonstrated to have connections with the NAc[[97](#_ENREF_97)]. vmPFC has been implicated in the pathogenesis of obsessive-compulsive disorder through its neural network connections to nodes involved in affective and reward processing[[13](#_ENREF_13),[98](#_ENREF_98)]. A PET study investigating the effects of STN DBS for OCD demonstrated that a decrease in Yale-Brown Obsessive Compulsive Scale (Y-BOCS) score, a 10-item scale designed to both determine severity of OCD and to monitor improvement during treatment, was correlated with a decrease of metabolic activity in the vmPFC[[58](#_ENREF_58)].

**Orbitofrontal cortex (BA 11 and BA 12/47):** The orbitofrontal prefrontal cortex (OFC) includes BA 12 caudally (or more recently 47[[99](#_ENREF_99)] and BA 11 anteriorly[[100](#_ENREF_100)]. Orbitofrontal hyperactivity/hyperconnectivity has been demonstrated on numerous functional imaging studies of obsessive-compulsive disorder[[101](#_ENREF_101)] and Tourette[[102](#_ENREF_102)]. Numerous neuroimaging studies in idiopathic depression and Parkinson’s-related depression have also implicated this area[[93](#_ENREF_93),[103](#_ENREF_103)]. Reduction of OFC activity is correlated with symptom improvement in medication and psychotherapy trials[[104](#_ENREF_104),[105](#_ENREF_105)]. With DBS to the inferior thalamic peduncle (connection from OFC to thalamus), it has been demonstrated to have significant reduction in OCD and depression symptoms[[106](#_ENREF_106)]. OFC activity has been demonstrated to be reduced in chronic NAc stimulation[[65](#_ENREF_65)]. In ALIC DBS for OCD, reduction in OFC activity was associated with clinical response in one study[[50](#_ENREF_50)] and regardless of clinical response in another[[107](#_ENREF_107)], as well as in larger studies of VCVS DBS[[57](#_ENREF_57)], ALIC DBS[[56](#_ENREF_56)], and capsulotomy[[108](#_ENREF_108)]. It is also reduced in SCG stimulation[[63](#_ENREF_63),[64](#_ENREF_64)], potentially explaining its anti-obsessional effects[[109](#_ENREF_109)]. A PET study investigating the effects of STN DBS for OCD revealed that the decrease in Y-BOCS score was correlated with a decrease of metabolic activity in the orbitofrontal cortex[[58](#_ENREF_58)]. Given that decreased orbitofrontal metabolism is associated with Parkinson’s related depression, this may explain why STN is associated with increased depressive symptoms[[47](#_ENREF_47)], although that is likely not the whole story[[110](#_ENREF_110)]. Reduction in the activity of the OFC appears to be clearly linked to improvements from OCD surgery regardless of the target or methodology chosen[[57](#_ENREF_57),[58](#_ENREF_58), [108](#_ENREF_108)].

**Cingulate cortex (BA 23, BA 24, BA 25, BA 31, BA 32):** The SCG is the portion of the cingulum that lies ventral to the corpus callosum[[111](#_ENREF_111),[112](#_ENREF_112)]. The SCG has been recognized for more than two decades to be an important node in the neural networks that contribute to mood regulation. This network includes cortical structures such as the DLPFC and the FPC as well as the subcortical structures such as the limbic system, thalamus, hypothalamus, and brainstem nuclei[[111](#_ENREF_111)]. It has been demonstrated that activity in the SCG can reflect antidepressant efficacy, regardless of the nature of the intervention[[81](#_ENREF_81),[113-115](#_ENREF_113)]. The SCG has reciprocal connections to the orbitofrontal, FPC, anterior cingulate cortex (ACC), intralaminar thalamus, hypothalamus and amygdala. The SCG also has afferents from BA 9/46 and efferents to numerous brainstem nuclei[[111](#_ENREF_111), [116](#_ENREF_116), [117](#_ENREF_117)]. The SCG was the first target chosen because of functional neuroimaging data and not as the result of lesioning studies[[63](#_ENREF_63)]. SCG was first targeted for depression DBS (unipolar and bipolar)[[63](#_ENREF_63),[64](#_ENREF_64),[118](#_ENREF_118)], but has also been targeted for anorexia[[109](#_ENREF_109)]. The gray matter of the subgenual cingulate has reduced activity with chronic NAc DBS[[65](#_ENREF_65)], ALIC DBS[[56](#_ENREF_56)] and SCG (white matter) DBS[[63](#_ENREF_63)]. The white matter of the subgenual cingulate has increased activity with SCG DBS[[63](#_ENREF_63)] and VCVS DBS[[57](#_ENREF_57)]. It appears that in order to achieve an antidepressant response, the active contact has to interface with the white matter tracts to the ventral pallidum, the mediodorsal (MD) thalamus, and to the FPC[[47](#_ENREF_47), [94](#_ENREF_94)].

The ACC has been heavily implicated in the neural circuitry of mood and anxiety regulation as well as the pathogenesis of TS[[119](#_ENREF_119)]. In depression, the ACC is underactive while the SCG is overactive[[120](#_ENREF_120)]. Conversely, the ACC is overactive in OCD during error processing[[121](#_ENREF_121),[122](#_ENREF_122)]. The ACC is involved in self-referential processing. It is an important area for determining treatment response[[120](#_ENREF_120)]. The higher the activity in the ACC, the higher likelihood of depression remission[[123](#_ENREF_123)]. Anterior cingulotomy has been demonstrated to be quite effective in the treatment of OCD and to a lesser degree depression, but with significant side effects[[9](#_ENREF_9),[124](#_ENREF_124)]. Success with anterior capsulotomy has been associated with significant reductions in dorsal anterior cingulate[[108](#_ENREF_108)]. DBS of the right NAc has been shown to increase the activity of the right ACC[[59](#_ENREF_59)] and animal studies have confirmed the laterality of these findings in a porcine model[[26](#_ENREF_26)]. VCVS DBS increases activity of ACC acutely[[57](#_ENREF_57)]. Conversely, DBS of the SCG demonstrated decreases in ACC metabolism[[63](#_ENREF_63),[64](#_ENREF_64)]. Interestingly, limbic STN DBS in appears to reduce activity in the ACC[[71](#_ENREF_71)]. The ACC has significant connections to the posterior cingulate cortex (PCC) and the medial PFC, another target for invasive neuromodulation for depression[[2](#_ENREF_2)].

The PCC is a key node in the default mode network. It has been hypothesized that the posterior cingulate cortex has a role in supporting internally-directed cognition and increases in activity when individuals retrieve autobiographical memories or plan for the future[[125](#_ENREF_125)]. The PCC has been shown to be decreased with depression[[126](#_ENREF_126)] and obsessive-compulsive disorder[[127](#_ENREF_127)] and is associated with treatment response to multiple modalities[[128](#_ENREF_128),[129](#_ENREF_129)]. PCC activity is increased in the neural activity of the PCC is correlated with good response in DBS, but additionally in medication trials[[125](#_ENREF_125)]. Cingulate activity was increased with NAc DBS in a large animal model[[26](#_ENREF_26)] and increased in another case with OCD[[59](#_ENREF_59)], but appears to decrease after chronic NAc DBS[[65](#_ENREF_65)]. YBOCS scores for ALIC DBS were negatively correlated with PCC activity on PET[[56](#_ENREF_56)]. Activity of the PCC increases with SCG stimulation[[63](#_ENREF_63),[64](#_ENREF_64)]. The activity of SCG was reduced with limbic STN DBS[[58](#_ENREF_58)]. The PCC has been demonstrated to have reduced activity with chronic NAc stimulation for depression[[65](#_ENREF_65)], but this may reflect homeostatic mechanisms during chronic stimulation.

**Insula:** The anterior insula appears to be an important node in mood and anxiety regulation networks. A recent study demonstrated that relative hypo- or hyperactivity of the anterior insula is correlated with response to depression treatment whether cognitive behavioral therapy (CBT) or medication[[130](#_ENREF_130)]. Disgust associated with OCD is thought to be mediated by the insula[[131](#_ENREF_131)]. The insula has also been implicated in tic generation in TS[[132](#_ENREF_132)]. The metabolism of the insula has been demonstrated to have a significant correlation preoperative Y-BOCS score in patients undergoing capsulotomy[[108](#_ENREF_108)].Anterior insular activity is reduced with ALIC DBS stimulation [[56](#_ENREF_56)] and NAc DBS stimulation[[66](#_ENREF_66)]. It is similarly reduced in SCG stimulation[[63](#_ENREF_63),[64](#_ENREF_64)] as well as in STN DBS (limbic)[[58](#_ENREF_58)]. Conversely, the insular signal has been demonstrated to also be decreased with a model of porcine CM-Pf DBS[[27](#_ENREF_27)], but increased in porcine NAc DBS[[26](#_ENREF_26)]. It should be noted that both porcine DBS studies utilized non-depressed animal subjects[[26](#_ENREF_26),[27](#_ENREF_27)].

**Pre-supplementary motor area and supplementary motor area (BA 6):** While the supplementary motor area (SMA) is an area of motor planning, the pre-SMA is an area related to response inhibition[[133](#_ENREF_133)]. Lesions to the SMA cause reduction of spontaneous movements and difficulty in performing voluntary motor acts[[134](#_ENREF_134)]. Inhibitory transcranial magnetic stimulation over the SMA and pre-SMA can reduce motor tics and OCD symptoms, respectively[[10](#_ENREF_10),[11](#_ENREF_11)]. The effects of ALIC DBS and anterior capsulotomy have been demonstrated to spread to the pre-SMA/SMA[[56](#_ENREF_56),[108](#_ENREF_108)]. SCG DBS has been shown to have effects on BA 6[[63](#_ENREF_63), [64](#_ENREF_64)].

**SUBCORTICAL TARGETS**

***Striatum (nucleus accumbens, caudate and putamen)***

The NAc is an important node in the human reward system. It can be divided into two principal parts; the core and the shell[[135](#_ENREF_135)]. The NAc core projects to the pallidal and nigral complexes and the NAc shell projects to the lateral hypothalamic areas, dopaminergic cell groups, and caudal mesencephallic areas[[136](#_ENREF_136)]. Hyperactivity of the NAc has been shown to have a negative correlation with recovery from depression[[123](#_ENREF_123)]. DBS has been implanted in the NAc not only for depression and OCD[[59](#_ENREF_59)], but also TS[[137](#_ENREF_137)], addiction[[138-142](#_ENREF_138)], eating disorders[[143](#_ENREF_143)], and even suggested for pain[[144](#_ENREF_144)].

DBS of the ALIC normalized pre-op hyperactivity of the NAc[[56](#_ENREF_56)]. The caudate has been shown to be involved in reward processing and hypoactive caudate responses to reward may be the functional explanation for anhedonia. The caudate has been shown to have reduced activity with chronic NAc stimulation[[65](#_ENREF_65)]. Furthermore, it has been implicated in the pathophysiology of OCD where specific symptom profiles can be mapped to subregions within the caudate nucleus[[145](#_ENREF_145)]. The caudate has been targeted for OCD DBS with apparent efficacy[[145](#_ENREF_145)]. VCVS has been shown to activate striatum[[57](#_ENREF_57)]. Right NAc inhibited the activity of the right dorsolateral rostral putamen[[59](#_ENREF_59)]. Recently, the white matter tract afferent from the ventral tegmental area, medial forebrain bundle, to the NAc was targeted for treatment-resistant depression with marked success[[146](#_ENREF_146)].

***Amygdala and extended amygdalar complex***

The amygdala is a brain nucleus that responds to novelty, salience and a variety of emotional stimuli. The amygdala has been implicated as an important component of the neural network that underlies social behavior[[147](#_ENREF_147)]. The most commonly referenced role of the amygdala is in mediating fear and anxiety[[148](#_ENREF_148)]. The amygdala has been shown to be overactive in patients with both MDD and Bipolar Disorder[[149](#_ENREF_149)] and its activity is elevated in anticipatory anxiety in patients with generalized anxiety disorder[[150](#_ENREF_150)] and in contamination fear with OCD[[151](#_ENREF_151)]. Patients with Tourette appear to have altered functional connectivity of the amygdala[[152](#_ENREF_152)] as well as amygdalar response to negative face pictures[[153](#_ENREF_153),[154](#_ENREF_154)]. Furthermore, the mean amygdala metabolism decreases in antidepressant treatment responders for depression, but persistence of elevated amygdala metabolism following remission is associated with a high risk for depressive relapse[[155](#_ENREF_155)].

DBS stimulation of the ALIC was shown to normalize baseline hyperactivity of the amygdala and trended towards significant correlation with reductions in YBOCS[[56](#_ENREF_56)], while this is not seen after anterior capsulotomy[[108](#_ENREF_108)]. Acutely (1 wk), there is an increase in amygdalar activity with NAc DBS[[66](#_ENREF_66)], which may explain the acute panic that has been observed[[156](#_ENREF_156)]. Reduction in amygdalar hyperactivity was correlated with response to chronic NAs DBS for depression and anxiety[[65](#_ENREF_65)]. In a functional connectivity analysis of NAc DBS, positive coupling was found with the amygdala[[13](#_ENREF_13)]. CM-Pf DBS in a large animal model demonstrated reductions in amydalar signal which may explain some of the positive psychiatric improvements from CM-Pf DBS for TS[[27](#_ENREF_27)].

The bed nucleus of the stria terminalis (BNST) is part of the extended amygdaloid complex and a relay/integral regulator of the hypothalamic-pituitary-adrenal stress axis[[157](#_ENREF_157),[158](#_ENREF_158)]. The BNST is the major output of the basolateral (BL) amygdala and has been associated with sustained fear[[159](#_ENREF_159)]. Some have suggested that the bed nucleus of the stria terminalis has been responsible for the positive effects of DBS that was targeted at the VCVS[[160](#_ENREF_160)]. In a recent world-wide analysis, it was noted that when the “VCVS target” was shifted posterior (*i.e.*, closer to the BNST), lower voltages were required to achieve efficacy[[51](#_ENREF_51)].

The amygdala has been implicated in the pathogenesis of autism[[161](#_ENREF_161)] and its hyperarousal has been seen on fMRI[[162](#_ENREF_162)]. There is one case of BL amygdalar DBS, which was performed for self–injurious behavior in autism. In addition to improving self injurious behavior, DBS of the BL amygdala was found to improve the core symptoms of the autism spectrum in the emotional, social, and cognitive domains[[163](#_ENREF_163)].

***Subthalamic nucleus***

The sensorimotor STN has been demonstrated to be an efficacious target for the treatment of Parkinson’s disease (PD), although stimulation in this node can cause a range of neuropsychiatric symptoms[[164](#_ENREF_164)]. In addition to the sensorimotor region, the STN also has limbic and executive regions with segregated circuitry involved in affective and cognitive processes[[164](#_ENREF_164)]. Several PET studies have identified activity changes in limbic and executive regions during STN-DBS in Parkinson’s patients, including the dorsolateral prefrontal cortex and cingulate gyrus[[70](#_ENREF_70),[165](#_ENREF_165),[166](#_ENREF_166)]. The ventromedial or limbic STN has been demonstrated to have profound anti-OCD effects through two serendipitous findings where STN DBS treated both PD and OCD[[47](#_ENREF_47)].

Limbic STN DBS has been utilized successfully utilized for OCD[[54](#_ENREF_54),[55](#_ENREF_55)] and TS[[167](#_ENREF_167)]. Neuroimaging studies of the limbic and cognitive STN circuits have proven valuable, as these help with the STN DBS target refinement. Several of the STN DBS OCD patients underwent PET imaging[[58](#_ENREF_58)]. When compared to OFF stimulation, DBS ON resulted in reduction in activity of the cingulate, orbitofrontal cortex, and supplementary motor areas (BA 6). The cingulate and orbitofrontal cortex are major nodes in the emotional and executive networks and the SMA is intimately involved in OCD/TS pathophysiology as was described above. The cingulate, SMA, and orbitofrontal cortex are hyperfunctional in untreated OCD, suggesting that a possible mechanism of DBS action may be the normalization of the activity of the SMA, OFC and cingulate[[98](#_ENREF_98),[133](#_ENREF_133)[168](#_ENREF_168),[169](#_ENREF_169)].

***Pallidum (globus pallidus interna, globus pallidus externa, and ventral pallidum)***

The pallidal areas have been demonstrated to be immensely important in the treatment of movement disorders, such as Parkinson’s disease. This target has been associated with much better neuropsychiatric side effect profiles for PD DBS than STN[[164](#_ENREF_164)]. The GPi has been implicated in the pathophysiology of TS[[170](#_ENREF_170)] and OCD[[171](#_ENREF_171)]. While the GPi has been implicated as a motor target, it is clear that the anterio-medial portions of the GPi are limbic. In fact the ventral pallidum has been implicated to be the limbic pleasure generator[[172](#_ENREF_172)]. Both TS and OCD are successfully treated with GPi DBS[[68](#_ENREF_68),[173](#_ENREF_173)] and there has even been some report of mood improvement with GPi DBS[[174](#_ENREF_174)]. The globus pallidus externa has also been demonstrated to treat TS[[175](#_ENREF_175)]. Surprisingly, many of the functional imaging studies do not demonstrate changes in the pallidum with DBS[[26](#_ENREF_26),[50](#_ENREF_50),[63](#_ENREF_63)]. The exception is PET of patients with VCVS DBS[[57](#_ENREF_57)]. The efficacy of SCG DBS is optimized when the active contact interfaces with the white matter tract to the ventral pallidum[[47](#_ENREF_47),[94](#_ENREF_94)].

***Thalamus***

**Centromedian nucleus of the thalamus and parafascicular complex:** The CM-Pf nuclei complex is located in the posterior part of the intralaminar thalamus and serves as the main basal ganglia input station. Additionally, the CM-Pf controls striatal dopamine function and provides a majority of the thalamic input to the striatum[[176-178](#_ENREF_176)]. CM-Pf lesions have been shown to cause complex attention deficits. This has been hypothesized to be due to the role of the CM-Pf in directing attention to motivationally relevant stimuli[[179](#_ENREF_179)]. The CM-Pf DBS target has been utilized in TS, but has also been shown to have positive effects on mood, anxiety, and OCD[[180](#_ENREF_180)].

In addition to the striatum, the CM-Pf has been demonstrated to have connections with the amygdala[[181](#_ENREF_181)], hippocampus[[182](#_ENREF_182)], globus pallidus[[183](#_ENREF_183)], nucleus accumbens[[184](#_ENREF_184)], insular cortex, and anterior cingulate cortex[[176](#_ENREF_176)]. A recent large animal study looking at DBS-fMRI in the CM-Pf study was recently completed[[27](#_ENREF_27)]. In this study, DBS at the Pf was shown to have negative BOLD downstream effects on the hippocampus as well as the dorsal anterior cingulate cortex and posterior cingulate cortex. DBS at the CM was shown to have negative BOLD downstream effects on the sensorimotor areas and the associative area. The observed differences in structural connections of CM versus Pf in non-human primate topographical analyses were verified by this large animal CM-Pf DBS-fMRI study[[27](#_ENREF_27)]. The Pf has been demonstrated to innervate associative and limbic striatal areas[[185-187](#_ENREF_185)]. This large animal study also demonstrated a primarily limbic downstream effect with high frequency stimulation[[27](#_ENREF_27)]. Similarly, the CM innervates the sensorimotor striatal area[[185](#_ENREF_185),[188](#_ENREF_188),[189](#_ENREF_189)]. The large animal study also confirmed that the CM is has primarily sensorimotor downstream effects with high frequency stimulation[[27](#_ENREF_27)].

***Medial dorsal thalamus***

The limbic nucleus of the thalamus, the medial dorsal nucleus of the thalamus, is an important node in the limbic circuitry. Using high-resolution 7T fMRI, Metzger, et.al. demonstrated general emotional arousal is localized to the mediodorsal nucleus while preceding attention and expectancy were localized to the intralaminar centromedian/parafascicular complex[[190](#_ENREF_190)]. A component of several of the frontal-striatal circuitry, the MD thalamus has been shown to be hyperactive in depression. In the depression associated with Parkinson’s, volumetric studies have shown that the MD nucleus volume is larger than non-depressed controls, but hypoactive on the left in this population suggesting that the increase in volume is compensatory[[191](#_ENREF_191)]. There has been four cases where the MD was targeted for TS[[16](#_ENREF_16),[17](#_ENREF_17)]. In addition to improvement in TS symptoms, there was a hint of improvement in mood and obsessive thinking for at least one of these individuals[[16](#_ENREF_16)]. During chronic NAc stimulation, MD nucleus activity is reduced[[65](#_ENREF_65)]. This reduction in MD activity is also observed with ALIC stimulation[[56](#_ENREF_56)]. In large animal studies looking at porcine NAc DBS stim, similar reductions were seen[[26](#_ENREF_26)]. In a recent analysis of SCG DBS, tractography interfacing with the active contact was compared to presence or absence of remission and only individuals with white matter projections extending to the MD thalamus were able to achieve remission[[47](#_ENREF_47)]. It appears that the MD thalamus is a critical node in mood/affect regulation as well as mediating anxiety disorders.

**OTHER MAGNETIC RESONANCE IMAGING TECHNIQUES**

***White matter tracts, tractography, and correct positioning***

It is possible to non-invasively determine the anatomical connection pathways using diffusion tractography imaging (DTI). These pathways are assumed to be consistent with known anatomy[[192](#_ENREF_192),[193](#_ENREF_193)]. This MRI technology has been utilized in determining the elements that contact and are stimulated with DBS[[194](#_ENREF_194)]. The benefits observed in patients with DBS appear to heavily depend on exact electrode position, which can only be truly verified with sophisticated tractography techniques[[94](#_ENREF_94),[95](#_ENREF_95),146,[194-196](#_ENREF_194)]. In the case of medial forebrain bundle DBS, the target can only be identified with sophisticated tractography imaging as one cannot visualize this target on conventional imaging[[146](#_ENREF_197)]. Slight alterations in the position of the electrode in the SCG appear to change whether this target is efficacious or not[[95](#_ENREF_95)]. The white matter tracts have to be completely intact for DBS to be efficacious where any alteration in the macroanatomy of the limbic white matter tracts can cause the stimulation to be ineffective[197-[199](#_ENREF_199)].

Conversely, verification of correct position with EpCS has been shown to only require a postoperative computed tomography (CT) scan coregistered with the presurgical MRI scan. This method will allow for all four contact electrodes on each paddle lead to be identified. In all 5 subjects of the Nahas study, contact electrodes were over the bilateral frontal pole and dorsolateral prefrontal cortex[[2](#_ENREF_2)]. In the Kopell study, the responders were noted to have the contacts of the electrodes placed at or anterior to the target site (DLPFC). Non-responders or partial responders had their electrodes placed posterior to the DLPFC target site[[3](#_ENREF_3)].

***Functional connectivity analysis***

Recent developments in noninvasive functional connectivity analysis demonstrate that this allows for a powerful network-level view of the neuropsychiatric disease[[200](#_ENREF_200)] and the effects DBS on that network[[13](#_ENREF_13)]. In this technique, the spontaneous fluctuations in BOLD are identified as the intrinsic marker for functional connectivity. A seed region is then identified and correlated with nodes in the connected neural network in an effort to determine areas where there is functional connectivity[[201](#_ENREF_201)]. The utility of this approach in DBS research comes from its ability to not only identify the aberrant functional connectivity of a particular neuropsychiatric disorder, but also normalize this aberrant functional connectivity through neuromodulation. This technique has been utilized to probe the resting state connectivity changes in patients receiving NAc DBS for OCD[[13](#_ENREF_13)]. In this study, the NAc DBS was shown to normalize aberrant hyperconnectivity between the NAc and lateral/medial prefrontal cortices. This reduction in connectivity correlated with the reductions in the YBOCS. While NAc is also a target for treatment-resistant depression (TRD) and TS, there have been no functional connectivity analysis studies in TRD and TS to date.

**CONCLUSION**

Invasive neuromodulation holds promise as a tool for treatment-resistant neuropsychiatric conditions[[47](#_ENREF_47)]. DBS resides at an interface between neurophysiology, Neurosurgery, movement disorders Neurology, and interventional psychiatry[[202](#_ENREF_202)]. DBS has been demonstrated to be an effective therapy for essential tremor, dystonia and Parkinson’s disease, and is being investigated in many other neuropsychiatric diseases[[47](#_ENREF_47)]. Advanced neuroimaging techniques aid in targeting, investigate the effects of, and further refine targeting of DBS all in a non-invasive manner. Although many of the functional imaging studies to date are limited to small case series and have had conflicting results, a growing trend is emerging. For most neuropsychiatric DBS imaging studies, the effects of DBS appears to have downstream effects on both cortical areas (DLPFC, FP, SCG, ACC, PCC, insula, SMA) as well as subcortical areas (thalamus, pallidum, STN, amygdala, striatum). These findings affirm that the therapeutic properties of DBS reside not only in local effects, but also downstream effects on the distributed neural network.

The effects and apparent efficacy of DBS for OCD appears to be related its downstream effects on the prefrontal cortex[[13](#_ENREF_13)], more specifically orbitofrontal cortex[[50](#_ENREF_50),[58](#_ENREF_58)]. This also appears to be true for invasive neuromodulation in treatment-resistant depression where the tractography and the functional imaging of depression DBS targets all converge on PFC[[13](#_ENREF_13)], more specifically Brodmann 10[[47](#_ENREF_47),[94-96](#_ENREF_94)]. Recent optogenetics findings have confirmed the critical role of the medial prefrontal cortex in mood regulation[[92](#_ENREF_92)]. While there has only been one study investigating the effects of direct BA 10 stimulation [[2](#_ENREF_2)], this technique appears remarkably durable[[85](#_ENREF_85)]. The critical role of the frontopolar cortex in mood regulation was further demonstrated when its activation was demonstrated to be associated with acute, stimulation-induced depressive symptoms in a patient with Parkinson’s and a mispositioned lead[[31](#_ENREF_31)]. Furthermore, this cortical stimulation approach seems to avoid many of the methodological challenges seen with other techniques such as the need for advanced tractography[[95,146](#_ENREF_197)] and the risk of cerebral hemorrhage[[199](#_ENREF_199)]. The therapeutic effects of TS DBS appears to be correlated with the therapy’s ability to reduce dopamine levels in various downstream targets [[17](#_ENREF_17)].

While the apparent efficacy of DBS for neuropsychiatric disease is promising, there are several remaining questions. Are cortical or subcortical targets better for invasive neuromodulation? Will sophisticated tractography techniques be essential to the widespread implementation of this therapy for TRD[[95](#_ENREF_95),[196](#_ENREF_196),203,[204](#_ENREF_204)]? How do symptom clusters of a given neuropsychiatric disease correlate with the pre-operative functional imaging and the DBS-induced functional activation? Will imaging eventually dictate the target with the best chance of efficacy? Will DBS move beyond its current role to affect plasticity[[205](#_ENREF_205)]? Will DBS confirm findings that the basal ganglia may in fact be organized in a gradient of medial to lateral where emotional/reward (most medial) –cognitive—and motor subcomponents of the nuclei are in fact highly interconnected.

As this field develops both scientifically and technically, we anticipate that larger studies will be conducted and our collective knowledge will converge. In addition to extending ongoing functional imaging studies in this area, future investigations will likely temporally pair functional MRI with information on structural integrity (including diffusion tensor imaging and voxel based morphometry) which will enable us to have a more comprehensive understanding of the effects of DBS on neural network integrity among DBS patients. Some disease specific issues to consider in future studies include inherent microstructural pathology in a given disease may influence the effect of DBS outcomes[[206](#_ENREF_206),[207](#_ENREF_207)], the role of acute *vs* chronic stimulation on neural network plasticity[[13](#_ENREF_13)], and baseline firing rates in areas downstream from the DBS target[[208](#_ENREF_208)]. DBS is an extremely nonselective method of stimulation and this non-specific nature may be harmful in one disorder[[209](#_ENREF_209)], while that same neural element being efficacious in treating another[[146](#_ENREF_146)]. Currently, imaging studies are unable to distinguishing between circuits of interest and those being activated unintentionally. Current steering technologies coupled with functional imaging will potentially allow for isolation of a single neural element and the ability to image that neural element[[210](#_ENREF_210)]. Such an approach when coupled with the selective circuit manipulation of optogenetics[[15](#_ENREF_15)] may allow for further verification of findings. It is truly an exciting time to be embarking on a journey into the interface between advanced imaging and invasive neuromodulation.

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**Table 1 Advantages and disadvantages of imaging methods**

|  |  |  |
| --- | --- | --- |
| **Modality** | **Advantages** | **Disadvantages** |
| PET | Unlikely to disrupt implant placement or  function Widely available Technically simple to combine with DBS (relative to fMRI) Customized radiotracers to measure binding capacity | Cost, availability and health risks of radiotracers Static data (averaged over set time interval) Poor temporal and spatial resolution (relative to fMRI) Difficult to interpret data (receptor binding *vs* density) |
| MRI | Time-locked data High temporal and spatial resolution (relative to PET) Flexible data acquisition, processing and analysis  Possible analysis of ON and OFF settings | BOLD signal interpretation (temporal lag, anatomical imprecision) DBS safety considerations (lead migration, secondary lesioning) Possible deactivation of implanted impulse generators Potential DBS-related artifacts on images |

PET: Positron emission tomography; DBS: Deep brain stimulation; fMRI: Functional magnetic resonance imaging; BOLD: Blood oxygen level dependent.

**Table 2 Obsessive compulsive disorder deep brain stimulation targets: Activation/deactivation of cortical and subcortical targets**

|  |  |  |  |
| --- | --- | --- | --- |
|  | **STN\_DBS** | **ALIC\_DBS** | **VCVS\_DBS** |
| **Frontal cortex** |  | **↑Nuttin1a** |  |
| **Medial frontal gyrus** | **↓Le Jeune1f** |  |  |
| **Inferior frontal gyrus** |  | **↓Zuo1b** |  |
| **Supplementary motor area** | **↑Min1b,5** | **↓Zuo2** |  |
| **Prefrontal cortex** |  |  | **↑ Knight1d,7** |
| **Lateral PFC** |  |  | **↓Figee1b,2,↓ Knight** |
| **Dorsolateral PFC** | **↑Min1b,5** |  | **↑Sturm, ↑ Knight** |
| **Medial PFC** | **↓Le Jeune2** |  | **↓Figee1a,2 , ↑ Knight** |
| **Cingulate** |  |  | **↑Sturm; ↑ Knight1d,7** |
| **Anterior cingulate cortex** | **↑Min1b; ↓Le Jeune1f** | **↓Zuo1b,2** | **↑ Knight1d;↑Rauch1f** |
| **Orbitofrontal cortex** | **↓Le Jeune2** | **↓Zuo1b; ↓Abelson3** | **↑Rauch1f** |
| **Insula** | **↑Min1b** |  | **↑ Figee; ↑Knight1d,6** |
| **Striatum** |  | **↑ Nuttin1a** |  |
| **Caudate** | **↑Min1b** | **↓Zuo1b** | **↑Figee** |
| **Putamen** | **↑Min1b** |  | **↑Figee; ↓Sturm;↑Rauch1f** |
| **Nucleus accumbens** |  |  | **↑Figee1c** |
| **Temporal cortex** | **↑ Min1b,4** |  |  |
| **Superior temporal gyrus** |  | **↓Zuo1b ;↑Nuttin1a** |  |
| **Medial temporal gyrus** |  | **↑Nuttin1a** |  |
| **Thalamus** | **↑ Min1b,4** | **↓Zuo1b** | **↑Figee; ↓Knight1d,6;↑Rauch1f** |
| **Pons** |  |  | **↑Nuttin1a** |
| **Hippocampus/parahippocampus** | **↑ Min1b,4** |  | **↑ Knight1d,6** |
| **Globus pallidus** |  |  | **↑ Rauch1f** |

1Statistically significant, aP < 0.05, bP < 0.001, dP < 0.0001, fP < 0.005, cP = 0.031 *vs* scans; 2Statistically correlated with treatment response; 3Decreased activity for 2 of 3 patients; 4Significant activation to STN *vs* Gpi; 5Significant at 2 V; 6As the stimulation voltage increased from 3 V to 5V, the region of BOLD signal modulation increased; 7As the stimulation voltage increased from 3 V to 5V, the region of BOLD signal modulation area decreased for *n* = 2 pigs. STN: Subthalamic nucleus; ALIC: Anterior capsulotomy of anterior limb of internal capsules; DBS: Deep brain stimulation; VCVS: Ventral (anterior internal) capsule/ventral striatum; BOLD: Blood oxygen level dependent.

**Table 3 Treatment-resistant depression deep brain stimulation targets: Activation/deactivation of cortical and subcortical targets**

|  |  |  |  |
| --- | --- | --- | --- |
|  | **NAc\_DBS** | **SCG\_DBS** | **DLFPC\_ EpCS** |
| **Frontal cortex** |  |  |  |
| **Ventral superior frontal sulcus** | **↓Bewernick1c** |  |  |
| **Dorsal superior frontal sulcus** | **↓Bewernick1c** |  |  |
| **Medial frontal gyrus** | **↑Schlaepfer;↓Bewernick1c** |  |  |
| **Superior frontal gyrus** | **↑Schlaepfer** |  | **↓Kopell2** |
| **Inferior frontal gyrus** | **↓Schlaepfer1a** |  |  |
| **Prefrontal cortex (PFC)** | **↑Knight1d,11** |  |  |
| **Dorsomedial PFC** | **↑Schlaepfer1a** | **↓Lozano4** |  |
| **Dorsolateral PFC** | **↑Schlaepfer1a** | **↑Mayberg8; ↓ Lozano4,6f** | **↑Kopell1f;↓Kopell2** |
| **Ventral prefrontal** |  |  |  |
| **Ventrolateral prefrontal** | **↓Schlaepfer1a;↑Knight** | **↑Lozano6f** |  |
| **Ventromedial prefrontal** | **↓Schlaepfer1a** | **↑Lozano4** |  |
| **Medial prefrontal** |  | **↓Mayberg8** |  |
| **Dorsal frontal pole** |  | **↓Lozano4** |  |
| **Ventral frontal pole** |  | **↓Lozano4** |  |
| **Cingulate** | **↑Schlaepfer1a ;↑Knight1d,11** |  |  |
| **Posterior cingulate** | **↓Bewernick1c** | **↑Mayberg7b; ↑Lozano5** | **↓Kopell2** |
| **Subgenual cingulate** | **↓Bewernick1c,2** | **↓Mayberg7b; ↑Lozano4** |  |
| **Anterior cingulate** | **↓Schlaepfer1a;↑Knight1d** | **↑Mayberg8; ↓Lozano6f** |  |
| **Orbitofrontal cortex** | **↓Bewernick1e** | **↓Mayberg9b** | **↑Kopell1f** |
| **Lateral orbitofrontal** |  | **↑Lozano4** |  |
| **Medial orbitofrontal** |  | **↓Lozano6f** |  |
| **Premotor cortex** |  | **↑Mayberg7b; ↑Lozano3** |  |
| **Dorsolateral premotor** |  | **↓Lozano3** |  |
| **Striatum** |  |  |  |
| **Ventral striatum** | **↑Schlaepfer1a; ↑Knight** |  |  |
| **Caudate** | **↓Bewernick1c;↑Schlaepfer1c** | **↑(rostral) ↓(caudal)Lozano4** |  |
| **Putamen** | **↑Schlaepfer** |  |  |
| **Insula** | **↓Schlaepfer1a;↑Knight1d** | **↓Mayberg7b; ↑Lozano4** |  |
| **Amygdala** | **↑Schlaepfer1a;↑Bewernick1c,2** |  |  |
| **Pons** |  | **↓Lozano5** |  |
| **Thalamus** | **↓Bewernick1c; ↓Schlaepfer1a  ↓Knight1d,10** |  |  |
| **Hypothalamus** |  | **↓Mayberg7b** |  |
| **Hippocampus/ parahippocampus** | **↑Schlaepfer; ↑Knight1d** | **↑Lozano4,5** |  |

1Statisically significant, b*P* < 0.001, d*P* < 0.0001, a*P* < 0.05, f*P* < 0.005, c*P* = 0.05, e*P* = 0.038 *vs* scans; 2Statistically correlated with change in depression scale; 3de/activation at 3 mo only; 4de/activation at 3 and 6 mo; 5de/activation at 6 mo only; 6significant between scans at 6 mo only; 7significant between scans at 3 and 6 mo; 8statistically correlated at 3 and 6 mo; 9significant between scans at 3 mo only; 10as the stimulation voltage increased from 3 V to 5V, the region of BOLD signal modulation increased; 11as the stimulation voltage increased from 3 V to 5V, the region of BOLD signal modulation area decreased for *n* = 2 pigs; NAc: Nucleus accumbens; DBS: Deep brain stimulation; SCG: Subcallosal cingulate gyrus; DLFPC: Dorsolateral prefrontal cortex; EpCS: Epidural cortical stimulation; BOLD: Blood oxygen level dependent.

**Table 4 Tourette syndrome deep brain stimulation targets: Activation/deactivation of cortical and subcortical targets**

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
|  | **Gpi\_DBS** | **Thalamic\_DBS** | **Nac-ALIC\_DBS** | **Pf\_DBS** | **CM\_DBS** |
| **Prefrontal cortex** | **↓Min6** |  | **↑Knight1d,5** | **↑Kim7,8b,9,10b** | **↑Kim7,8a,9,10a** |
| **Dorsolateral prefrontal cortex** | **↑Min1b** |  |  |  |  |
| **Cingulate** |  |  |  |  |  |
| **Anterior cingulate cortex** | **↑Min1b** |  |  |  |  |
| **Dorsalanterior cingulate cortex** |  |  | **↑Knight4** | **↓Kim7,8,10** | **↓Kim7a,8,9,10** |
| **Dorsoposterior cingulate cortex** |  |  |  | **↓Kim7,8a,10** | **↓Kim7d,8,9,10** |
| **Motor cortex** |  |  |  |  |  |
| **Primary motor cortex** | **↑Min1b** |  |  | **↓Kim7,8a ,9,10a** | **↓Kim7,8b,9,10a** |
| **Premotor cortex** | **↑Min1b** |  |  | **↓Kim7,8a,9,10a** | **↓Kim7a,e,f, 8ba,9a,10a** |
| **Primary somatosensory cortex** | **↑Min1b** |  |  | **↓Kim7,8a,9,10a** | **↓Kim7,8,9,10a** |
| **Insula** | **↑Min1b** |  | **↑Knight1d,4** | **↓Kim7,9↑ 8a,10a** | **↓Kim7,8,9,10** |
| **Temporal lobe** |  |  |  |  |  |
| **Inferior temporal gyrus** |  | **↓Kuhn** |  | **↓Kim7↑8a,10** | **↓Kim7,9↑8,10** |
| **Striatum** |  |  |  |  |  |
| **Caudate** | **↑Min1b** | **↓Kuhn ↓Vernaleken** |  | **↓Kim8,10** | **↑Kim7↓8,10** |
| **Putamen** |  | **↑Kuhn  ↑Vernaleken** |  | **↓Kim8,10** | **↑Kim7↓8,10** |
| **Thalamus** |  | **↓Kuhn ↓Vernaleken** | **↓Knight1d,4** |  |  |
| **Right thalamus** |  | **↑ Kuhn2** |  |  |  |
| **Left thalamus** |  | **↓Kuhn3** |  |  |  |
| **Central thalamic nucleus** |  |  |  |  | **↑Kim8** |
| **Hippocampus** |  |  |  | **↓Kim7,8,10** | **↓Kim8** |
| **Parahippocampal cortex** |  |  | **↑Knight1d,4** | **↓Kim7,8,10** | **↓Kim7,8,10** |

1Statistically significant scans; b*P* < 0.001, d*P* < 0.0001, a*P* < 0.05 *vs* scans, eResponse at 130Hz 3V was significant *vs* Pf contact site at *P* < 0.05, fresponse at 60Hz 3V was significant *vs* Pf; 2Patient #3, unilateral had reduction of putamen activity; 3Only in unilateral patient; 4As the stimulation voltage increased from 3V to 5V, the region of BOLD signal modulation increased; 5As the stimulation voltage increased from 3V to 5V, the region of BOLD signal modulation area decreased for n=2 pigs; 6Activation decreased at 2V; 7Response @ 130 Hz 3V; 8Response @ 130 Hz 5V; 9Response @60Hz 3V; 10Response @ 60Hz 5V; Gpi: Globus pallidus interna; DBS: Deep brain stimulation; NAc-ALIC: Nucleus accumbens-anterior limb of internal capsule; CM-Pf: Centromedian-parafascicular nuclei complex; BOLD: Blood oxygen level dependent.

**Table 5 Parkinson’s disease deep brain stimulation targets-limbic effects**

|  |  |
| --- | --- |
|  | **STN\_DBS** |
| **Prefrontal** |  |
| **Dorsolateral prefrontal cortex** | **↑Hilker1b** |
| **Lateral prefrontal cortex** | **↑Stefurak1b** |
| **Cingulate** |  |
| **Posterior cingulate** | **↓Stefurak1b;↑Hilker1b** |
| **Anterior cingulate cortex** | **↑Stefurak1b;↑Hilker1b** |
| **Dorsal anterior cingulate cortex** | **↓Stefurak1b** |
| **Frontal cortex** | **↑Stefurak** |
| **Medial frontal cortex** | **↓Stefurak1b** |
| **Medial superior frontal** | **↑Stefurak1b** |
| **Orbitofrontal cortex** | **↑Hilker1b** |
| **Parietal Inferior lobule** | **↑Hilker1b** |
| **Frontal motor cortex** | **↑Stefurak1b** |
| **Premotor cortex** | **↑Stefurak1b** |
| **Primary motor cortex** | **↑Stefurak1b** |
| **Medial supplementary motor area** | **↓Stefurak1b** |
| **Ipsilateral parietal lobe** | **↓Stefurak1b** |
| **Insula** | **↑Stefurak1b** |
| **Superior temporal gyrus** | **↑Stefurak1b** |
| **Middle temporal gyrus** | **↓Hilker1b** |
| **Inferior temporal gyrus** | **↑Stefurak1b** |
| **Striatum** |  |
| **Caudate** | **↓Stefurak1b** |
| **Putamen** | **↑Stefurak1b** |
| **Lentiform nucleus** | **↑Hilker1b** |
| **Thalamus** |  |
| **Ventrolateral thalamus** | **↑Stefurak;↑Hilker1a** |
| **Ventroanterior thalamus** | **↑Stefurak1b** |
| **Dorsomedial nuclei of Thalamus** | **↑Stefurak1b** |
| **Pons** | **↑Stefurak1b** |
| **Midbrain** | **↑Stefurak1b** |
| **Cerebellum** | **↑Stefurak1b** |
| **Right Cerebellar posterior lobe** | **↑Hilker1a** |
| **Left Cerebellar posterior lobe** | **↓Hilker1b** |
| **Parahippocampal cortex** | **↑Stefurak1b** |
| **Occipital lobe** | **↓Stefurak1b** |

1Statistically significant, b*P* < 0.001, a*P* = 0.002 *vs* scans. STN: Subthalamic nucleus; DBS: Deep brain stimulation.

**TEMPORAL CORTEX**

**Superior Temporal Gyrus**

**Medial Temporal Gyrus**

**THALAMUS**

**HIPPOCAMPUS/**

**PARAHIPPOCAMPUS**

**ORBITAL FRONTAL CORTEX**

**Medial Frontal Gyrus**

**Inferior Frontal Gyrus**

**SMA**

**FRONTAL CORTEX**

**CINGULATE**

**Anterior Cingulate**

**INSULA**

**STRIATUM**

**Putamen**

**Caudate**

**NAc**

**PREFRONTAL CORTEX**

**Lateral PFC**

**Dorsolateral PFC**

**MEDIAL PFC**

**STN DBS**

1) Min, 2012

2) Le Jeune, 2010

**ALIC DBS**

3) Nuttin, 2003

4) Zuo, 2013

5) Abelson, 2005

**VCVS DBS**

6) Knight, 2013

7) Figee, 2013

8) Sturm, 2003

9) Rauch, 2006

**Figure 1 Obsessive compulsive disorder deep brain stimulation.** aAs stimulation voltage increased from 3V 130 Hz to 5V 130 Hz, the region of BOLD signal modulation increased; bAs stimulation voltage increased from 3V 130 Hz to 5V 130 Hz, the region of BOLD signal modulation decreased for *n* = 2 pigs. SMA: Supplementary motor area; NAc: Nucleus accumbens; PFC: Prefrontal cortex; BOLD: Blood oxygen level dependent.

**Ventral Superior frontal sulcus**

**Dorsal Superior frontal sulcus**

**Medial Frontal Gyrus**

**Superior Frontal Gyrus**

**Inferior Frontal Gyrus**

**FRONTAL CORTEX**

**Dorsomedial PFC**

**PREFRONTAL CORTEX (PFC)**

**Dorsolateral PFC**

**MEDIAL PFC**

**Ventral Frontal Pole**

**Dorsal Frontal Pole**

**CINGULATE**

**INSULA**

**THALAMUS**

**AMYGDALA**

**PREMOTOR CORTEX**

**Dorsolateral Premotor**

**Ventral Striatum**

**Caudate**

**Putamen**

**STRIATUM**

**Ventrolateral PFC**

**Ventromedial PFC**

**VENTRAL PREFRONTAL**

**ORBITOFRONTAL CORTEX**

**Medial Orbitofrontal**

**Lateral Orbitofrontal**

**HYPOTHALAMUS**

**HIPPOCAMPUS/ PARAHIPPOCAMPUS**

\*Increase in regional cerebral metabolic rate of glucose (rCMRG), positive correlation (decrease) between rCMRG and subsequent clinical response; \*\*rostral caudate increased, caudal caudate decreased.

**NAc\_DBS**

1) Bewernick, 2010

2) Schlaepfer, 2008

3) Knight, 2013

**DLPFC EpCS**

4) Kopell, 2011

**SCG DBS**

5) Lozano, 2008

6) Mayberg, 2005

nucleus accumbens, NAc; dorsolateral prefrontal cortex,DLPFC; epidural cortical stimulation, EpCS; subcallosal cingulate gyrus, SCG

**Anterior Cingulate**

**Subgenual Cingulate**

**Posterior Cingulate**

**Figure 2 Treatment-resistant depression deep brain stimulation.**

**STN DBS**

1) Hilker, 2003

2) Stefurak, 2003

**DLPFC**

**Lateral PFC**

**PREFRONTAL CORTEX**

**Posterior Cingulate**

**ACC**

**DACC**

**CINGULATE**

**FRONTAL CORTEX**

**Medial Superior Frontal**

**Orbitofrtonal Cortex**

**Parietal Inferior Lobule**

**Medial Frontal Cortex**

**FRONTAL MOTOR CORTEX**

**Premotor Cortex**

**Primary Motor Cortex**

**Medial SMA**

**IPSILATERAL PARIETAL LOBE**

**PONS**

**MIDBRAIN**

**OCCIPITAL LOBE**

**CEREBELLUM**

**Right Cerebellar Posterior Lobe**

**Left Cerebellar Posterior Lobe**

**Caudate**

**Putamen**

**Lentiform Nucleus**

**STRIATUM**

**PARAHIPPOCAMPAL CORTEX**

**INSULA**

**Superior Temporal Gyrus**

**Medial Temporal Gyrus**

**Inferior Temporal Gyrus**

**Ventrolateral Thalamus**

**Ventroanterior Thalamus**

**Dorsomedial Thalamus**

**THALAMUS**

**Figure 3 Parkinson’s disease deep brain stimulation.** DLPFC: Dorsolateral prefrontal cortex; PFC: Prefrontal cortex; ACC: Anterior cingulate cortex; DACC: Dorsoanterior cingulate cortex; SMA: Supplementary motor area.

**Figure 4 Tourette syndrome deep brain stimulation.** aResponse at 130 Hz 3V AND 60 Hz 3V decreased, but increased at 130 Hz 5V and 60 Hz 5V; bResponse decreased at 130 Hz 3V, but increased at 130 Hz 5V and 60 Hz 5V; cResponse decreased at 130 Hz 3v and 60 Hz 3V, but increased at 130 Hz 5V and 60 Hz 5V; dResponse increased at 130 Hz 3V, but decreased at 130 Hz 5V and 60 Hz 5V. GPi: Globus pallidus internus; NAc: Nucleus Accumbens; ALIC: Anterior limb of internal capsule; CM-PF: Centromedian-parafascicular nuclei complex; DLPFC: Dorsolateral prefrontal cortex; ACC: Anterior cingulate cortex; DACC: Dorsolanterior cingulate cortex.

**DLPFC**

**PREFRONTAL CORTEX**

**ACC**

**DACC**

**Dorsoposterior Cingulate**

**CINGULATE**

**Primary Motor Cortex**

**Premotor Cortex**

**Primary Somatosensory Cortex**

**MOTOR CORTEX**

**THALAMUS**

**Left Thalamus**

**Right Thalamus**

**Central Thalamic Nucleus**

**HIPPOCAMPUS**

**Parahippocampal Cortex**

**INSULA**

**Inferior Temporal Gyrus**

**TEMPORAL LOBE**

**Caudate**

**Putamen**

**STRIATUM**

**Gpi DBS**

1) Min, 2012

**Thalamic DBS**

2) Kuhn, 2012

3) Vernaleken, 2009

**NAc-ALIC DBS**

4) Knight, 2013

**CM-PF DBS**

5) Kim, 2013

A) PF

B) CM