Identification of biomarkers that predict response to subthalamic nucleus deep brain stimulation in resistant obsessive–compulsive disorder: protocol for an open-label follow-up study

Shyam Sundar Arumugham, Dwarakanath Srinivas, Janardhanan C Narayanaswamy, TS Jaisoorya, Himani Kashyap, Philippe Domenech, Stéphane Palfi, Luc Mallet, Ganesan Venkatasubramanian, YC Janardhan Reddy

ABSTRACT

Introduction Deep brain stimulation (DBS) of bilateral anteromedial subthalamic nucleus (amSTN) has been found to be helpful in a subset of patients with severe, chronic and treatment-refractory obsessive–compulsive disorder (OCD). Biomarkers may aid in patient selection and optimisation of this invasive treatment. In this trial, we intend to evaluate neurocognitive function related to STN and related biosignatures as potential biomarkers for STN DBS in OCD.

Methods and analysis Twenty-four subjects with treatment-refractory OCD who receive treatment as usual will undergo open-label STN DBS. Structural/functional imaging, electrophysiological recording and neurocognitive assessment would be performed at baseline. The subjects would undergo a structured clinical assessment for 12 months post-surgery. A group of 24 healthy volunteers and 24 subjects with treatment-refractory OCD who receive treatment as usual would be recruited for comparison of biomarkers and treatment response, respectively. Baseline biomarkers would be evaluated as predictors of clinical response. Neuroadaptive changes would be studied through a reassessment of neurocognitive functioning, imaging and electrophysiological activity post DBS.

Ethics and dissemination The protocol has been approved by the National Institute of Mental Health and Neurosciences Ethics Committee. The study findings will be disseminated through peer-reviewed scientific journals and scientific meetings.

INTRODUCTION

Obsessive–compulsive disorder (OCD) is a common neuropsychiatric condition with a lifetime prevalence of 2%–3%. It is among the top 10 causes of neuropsychiatric disabilities worldwide. The first-line treatment of OCD includes selective serotonin reuptake inhibitors (SSRIs) and/or cognitive behaviour therapy (CBT). However, around 20%–30% of patients do not respond to standard treatment strategies. Neurosurgical interventions are considered in patients with chronic, severe and treatment-refractory OCD. Ablative neurosurgical procedures, such as gamma ventral capsulotomy, are helpful in around 45%–65% of patients with treatment-refractory OCD. Due to the potential irreversible and severe adverse effects associated with ablative procedures, there has been a surge of interest in deep brain stimulation (DBS) as an alternate treatment. DBS involves electrical stimulation of the subcortical regions through surgically implanted microelectrodes. High-frequency electrical stimulation through these electrodes modulate the activity of dysfunctional neuronal circuits. Unlike ablative procedures, the stimulation in DBS can be modulated to optimise improvement and adverse effects.

Corticostriatothalamocortical (CSTC) circuits passing through subcortical regions are implicated in the pathogenesis of OCD. Thus, several subcortical structures, including anterior limb of the internal capsule, ventral capsule/ventral striatum (VC/vs), nucleus accumbens (NAC), bed...
nucleus of stria terminalis, anteromedial subthalamic nucleus (amSTN) and inferior thalamic peduncle are targets for DBS in OCD. DBS targeting some of these structures has been found to significantly reduce OCD symptoms compared with sham stimulation. Based on evidence from a double-blinded randomised controlled trial, a recent treatment guideline recommended bilateral STN DBS for treatment-refractory OCD. Two recent randomised crossover trials found equivalent efficacy of DBS targeting amSTN compared with VC/VS, caudate nucleus and NAc. In the latter study, most patients, who were masked/blinded to the target of stimulation, preferred the amSTN stimulation based on subjective improvement. A recent systematic review also found similar efficacy for DBS of amSTN and striatal targets. Interestingly, recent evidence suggests that the different targets have similar connections along fronto/subcortical circuits and thus may be targeting the same network.

The adverse effects of DBS are frequently related to stimulation and generally mild and reversible. DBS is an invasive procedure that requires long-term close monitoring for optimisation of stimulation as well as periodic changes in battery, which adds to the treatment cost. Despite these limitations, long-term follow-up studies have shown that DBS leads to improvement in symptoms as well as the quality of life. However, DBS is helpful in only 60%–75% of patients with treatment-refractory OCD. There is a need to identify predictors of treatment response, which would assist in patient selection for this invasive and expensive treatment.

Despite decades of research, the exact mechanism of action of DBS is still not clearly understood. Recent evidence suggests that DBS has both local and distant effects with resultant amelioration of pathological network activity. The role of target nuclei such as amSTN in the pathophysiology of OCD provides useful clues to unravel the mechanism, which would help identify biopredictors of response. Further, DBS provides a unique opportunity to study the role of STN in psychopathology with spatial and temporal precision.

**Neurocircuitry of OCD**

Although the aetiology of OCD is still unknown, converging evidence from neuroimaging studies implicates the dysfunctional CSTC circuits in the pathophysiology of OCD. Particularly, the CSTC pathways involving the orbitofrontal cortex (OFC) and, to a lesser extent, the dorsal anterior cingulate cortex (dACC) are dysfunctional in OCD. Functional neuroimaging studies show that these cortical regions are hyperactive at rest, which is accentuated during symptoms provocation. Pretreatment OFC and caudate metabolism predict response to medications in OCD. Similarly, DBS over various targets modulate the activity in these circuits by increasing their connectivity. Thus, modulation of functional connectivity in the CSTC circuits is an important target for antiobsessive treatments, including DBS.

**STN in the pathophysiology of OCD**

STN is the key basal ganglia input structure of the ‘hyperdirect’ pathways. Based on hyperdirect connectivity, the STN is divided into three partially overlapping subterritories, namely the limbic, associative and motor regions. Computational models and imaging studies suggest that the ‘hyperdirect’ pathway connecting inferior frontal gyrus (IFG) and STN is involved in global response inhibition, for example, during the stop signal task (SST). Functional MRI (fMRI) studies have shown activation of STN, IFG and presupplementary motor area during SST performance. STN also plays a crucial role in decision-making and response selection in conflict situations, by setting a decision threshold that is contextually modulated. Recent evidence suggests that neurons in ventromedial STN, which is preferentially connected to dACC and OFC, are especially involved in reactive stopping and switching. Both these functions are impaired in patients with OCD.

Following STN DBS, patients with OCD and Parkinson’s disease show decreased reaction time and performance, suggesting more ‘impulsivity’, as defined in response inhibition paradigms. STN DBS modulates uncertainty/conflict-driven decision threshold adjustment and adapting to speed/accuracy trade-offs. Impairment in task switching improves, specifically following stimulation of ventral DBS contacts in STN. Thus, DBS targeting amSTN may decrease decisional threshold towards more optimal levels, leading to a less cautious and more rapid goal-directed behaviour, which may be beneficial in patients with OCD.

STN connectivity has been implicated in the pathophysiology of OCD. Resting-state functional connectivity of STN with cortical and striatal structures has been associated with cognitive and behavioural measures of compulsivity. Tracing studies in non-human primates have found hyperdirect pathways connecting OFC, dACC and dorsolateral prefrontal cortex (DLPFC) projecting to the anteromedial STN. A prospectively acquired imaging study found that effective STN DBS targets in OCD are located in this region, that is in the anterior inferior medial border, which has direct connections with the OFC, dACC and DLPFC. Further, evidence suggests that STN DBS modulates corticostriatal connectivity during SST. STN DBS in OCD decreases metabolism in ACC and the therapeutic effects correlate with a decrease in OFC metabolism. Thus, CSTC networks connecting various prefrontal regions with STN play a role in the neurocognitive functions underlying OCD, its behavioural manifestations and may mediate the therapeutic actions of DBS in OCD. The hyperdirect pathways have especially been implicated in this regard.

Another line of evidence has used the unique opportunity provided by DBS to collect electrophysiological data directly from the nucleus. STN oscillatory activity and frontal cortico-STN coherence in β and θ frequency bands are associated with different phases of the SST, with the latter prominently seen in the ventral contacts. The medial PFC-STN θ phase coherence increases during...
high-conflict trials in the flanker task. Response inhibition and conflict monitoring, assessed through SST and flanker task, respectively, are putative endophenotypes for OCD. In OCD subjects, bursting and oscillatory activity have been localised to the associative/limbic STN, with predominant oscillatory activity in δ-band. Further, the severity of OCD is associated with low frequency oscillatory activity and burst characteristics. Low frequency activity (θ band) in the ventromedial STN has been associated with symptom provocation and cognitive/emotional functioning in OCD subjects. A recent study demonstrated that emotional images distinctively modulated STN θ band activity in OCD subjects and emotion-related θ band activity correlated with symptom severity. Thus, electrophysiological activity in STN, in particular low frequency oscillatory activity in the ventromedial region, is a putative biomarker associated with neurocognitive functions underlying OCD and its clinical manifestations.

Overall, STN-cortical connectivity (especially with the frontal cortex), scalp electroencephalographic activity in the frontal cortex, bursting/oscillatory activity in the STN and STN-related neurocognitive functioning (response inhibition) are potential biomarkers for treatment response to DBS in OCD.

The current proposal plans to identify such biomarkers with the following objectives:

**Primary objective**
1. To evaluate whether STN functional connectivity when performing response inhibition task (SST) predicts improvement in obsessive–compulsive symptoms after DBS over STN.

**Secondary objectives**
1. To evaluate whether the improvement of obsessive–compulsive symptoms following STN DBS is predicted by
   a. baseline STN resting-state functional connectivity with PFC.
   b. STN functional connectivity with PFC regions during symptom provocation paradigm.
   c. White matter structural connectivity (fractional anisotropy and other measures) of STN with cortical regions.
   d. Local field potential (LFP) time frequency amplitude in the STN at rest, during response inhibition task and symptom provocation.
   e. Frontal electroencephalogram (EEG) activity at rest, during response inhibition task and symptom provocation.
   f. Localisation of DBS electrodes within the STN.
2. To study the neuroadaptive changes following DBS by evaluating neurocognitive performance and EEG activity before and 12 months post DBS.
3. To evaluate the effect of DBS on clinical symptomatology, disability, quality of life and subjective well-being.
   We hypothesise that baseline STN prefrontal connectivity, low frequency oscillatory activity over frontal lobes/

STN and impaired performance on response inhibition task would predict improvement in obsessive–compulsive symptoms after STN DBS in OCD. The current report is prepared based on Standard Protocol Items: Recommendations for Interventional Trials checklist.

**Experimental design**
We plan to conduct a longitudinal open-label follow-up study of 24 subjects undergoing STN DBS for severe, chronic and treatment-refractory OCD to evaluate whether baseline biomarkers predict a decrease in the severity of OCD at follow-up. The DBS leads and stimulator would be implanted after baseline investigation for biomarkers. The initial 6 months of follow-up would be used for the optimisation of stimulus parameters. This would be followed by an open-label clinical follow-up of 6 months. The primary outcome would be a clinical improvement after 6 months of stable stimulation, that is, approximately 1 year after the implantation procedure. A comparison group consisting of 24 subjects with chronic, severe and treatment-refractory OCD, who would receive treatment as usual, would also be followed-up for 1 year. We would also recruit a sample of 24 healthy volunteers for comparison of biomarkers at baseline (figure 1). Subject enrolment is expected to commence in July 2021 and complete by September 2024.
Sample

DBS group (OCD-DBS)

Twenty-four subjects with severe, chronic and treatment-refractory OCD would be recruited from the OCD clinic, inpatient and outpatient services of the National Institute of Mental Health and Neurosciences (NIMHANS), Bangalore. The subjects would be screened with the following selection criteria:

Inclusion criteria

1. Age 18–60 years.
2. Primary diagnosis of OCD satisfying the Diagnostic and Statistical Manual of Mental Disorders (DSM)-5 criteria,
3. Duration of OCD ≥5 years.
4. Yale-Brown Obsessive Compulsive Scale (Y-BOCS) score of ≥28 or ≥14 if the subject has either predominantly obsessions or compulsions alone.
5. Clinical Global Impression-Severity (CGI-S) scale of ≥5.
6. Disability of ≥40% as evaluated by the Indian Disability Assessment and Assessment Scale.
7. Failure to obtain meaningful improvement in OCD despite adequate trial with standard treatment strategies, which should include:
   a. At least three trials of SRIs, one of which should be clomipramine, which was either ineffective or poorly tolerated. An adequate trial includes the recommended dose of SRIs for ≥12 weeks’ duration each.
   b. Augmentation of SRIs with at least two agents for ≥6 weeks, one of which should be either risperidone or aripiprazole.
   c. Trial of structured behaviour therapy/CBT for at least 20 sessions or demonstrated an inability to tolerate the anxiety due to therapy.

Exclusion criteria

1. Diagnosis of bipolar disorder, a psychotic disorder of ≥3 months’ duration as assessed with MINI 7.0.2.
2. Current substance use disorder (except caffeine or nicotine use disorder) or major depressive episode or current high suicidality, as assessed with MINI 7.0.2.
3. Severe personality disorder as assessed with Structured Clinical Assessment of DSM-5-Personality Disorders.
4. Clinically significant abnormality on MRI of the brain.
5. Pregnancy, contraindication for DBS/anaesthesia/preoperative MRI and inability to comply with surgical requirements.

All potential study participants satisfying the above criteria will be assessed for suitability for DBS by two psychiatrists and a neurosurgeon from the study team. An independent review committee with members (consisting of two psychiatrists, one neurologist and one neurosurgeon) who are not a part of the study team would ratify the decision for eligibility for surgery.

OCD control group

We would recruit 24 OCD subjects fulfilling the above criteria from the same population, including those subjects who refuse consent for DBS. These subjects would receive treatment for OCD as recommended by the treating clinician.

Healthy volunteers

Twenty-four age and gender-matched healthy volunteers would be recruited from the community to compare the baseline differences in neuroimaging, neurocognitive performance and EEG with the study subjects. They would be screened for the presence of psychiatric disorder using MINI 7.0.2. Other exclusion criteria include Y-BOCS score ≥12, family history of psychiatric disorder in first-degree relatives, clinically significant neurological illness, pregnancy and contraindication for MRI.

Sample size estimation

For evaluating the abovementioned primary objective, the optimal sample size was estimated using standard principles and methods. Based on approximation and estimation from the previous data on MRI-based predictors of treatment response for other interventions in OCD (n=12, 15), a sample size of at least 24 OCD patients will be required to detect a two-tailed significant difference of α=0.05 with an estimated 90% power for an estimated correlation coefficient of 0.6 (Pearson’s) between STN-PFC connectivity and change in Y-BOCS total score with add-on DBS.

Baseline assessment

Clinical assessment

The clinical status would be assessed with the MINI 7.0.2. All OCD subjects would undergo baseline assessment with Y-BOCS, which includes a symptom checklist to assess the nature of symptoms, severity scale to assess the severity of symptoms and insight/avoidance scale to assess these domains of symptoms. The severity of illness would also be assessed with the CGI-S scale. The severity of depressive and anxiety symptoms would be assessed with the Hamilton Depression Rating Scale (HAM-D) and Hamilton Anxiety Rating Scale (HAM-A), respectively. The extent of disability, quality of life and mental well-being would be assessed with the WHO Disability Assessment Schedule 2.0–36 item version (WHODAS 2.0), WHO quality of life instrument (WHOQOL-BREF) and WHO-5 Well-Being Index (WHO-5). The healthy volunteers would undergo screening with MINI 7.0.2 and Y-BOCS.

Neurocognitive assessment

All the subjects would undergo a detailed neurocognitive evaluation consisting of the modified NIMHANS neuropsychology battery, SStask for assessing response inhibition, modified flanker task, for assessing response inhibition in a conflict situation/error monitoring, beads task for assessing decisional impulsivity and temporal discounting task for assessing reward-related decision-making.
Impairment in the above neurocognitive functions have been observed in OCD patients and they are closely associated with amSTN functioning.\(^3\)\(^7\)\(^4\)\(^3\)\(^5\)\(^6\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(^7\)\(\text{Box 1 Neurocognitive and personality assessment}\)

**Neurocognitive assessment**
1. Modified NIMHANS neuropsychology/neuropsychology battery\(^7\)\(^1\)
   - Finger tapping.
   - Bender-Gestalt test.
   - Digit symbol substitution test.
   - Colour trails 1 and 2.
   - Block design test.
   - Controlled Oral Word Association Test.
   - Animal names test.
   - Digit span.
   - Spatial span.
   - Tower of London.
   - Wisconsin Card Sorting Task.
   - Stroop test.
   - Auditory verbal learning test.
   - Rey’s complex figure test.
2. Stop signal task.\(^7\)\(^2\)
3. Modified flanker task.\(^5\)\(^5\)
4. Beads task.\(^3\)\(^7\)
5. Temporal discounting task.\(^7\)\(^4\)–\(^7\)\(^6\)

**Personality/behavioural assessment**
1. Frontotemporal Behavioural Scale.\(^7\)\(^6\)
2. Iowa Scales for Personality Change.\(^7\)\(^9\)
3. UPPS-P Impulsive Behaviour Scale.\(^8\)\(^0\)

To the best of our knowledge, ME-fMRI has not been performed under symptom provocation. Hence, it is unclear whether this paradigm would activate STN. Due to the previous data showing activation of OCD-specific circuits, activity under this paradigm will be tested as an additional predictor of treatment response.

**Electroencephalography**
All subjects would undergo an awake 64-channel EEG as per the international 10–20 EEG system electrode placement, in a comfortable resting supine position. Recordings from the frontocentral (FCz), central (Cz) and bilateral dorsolateral prefrontal (F3 and F4) montages would be the focus of analysis. The OCD subjects and healthy volunteers would undergo EEG while performing the SST and flanker tasks. The OCD subjects would also undergo EEG symptom provocation using symptom dimension-specific pictures. Artifact-free epochs of resting-state EEG epochs will be selected for power spectral analysis, which involves examining the spectral power in various frequency bands (\(\delta\)=1–3 Hz, \(\theta\)=4–7 Hz, \(\alpha\)=8–11 Hz, \(\beta\)=12–30 Hz and \(\gamma\)=30 Hz) at each electrode. For examining the EEG under symptom provocation, power spectral analysis would be conducted on 2 s epochs selected from 1 s before stimuli to 1 s after the stimuli that is, symptom dimension-specific pictures.

**Deep brain stimulation**
Following the baseline assessments, subjects in the OCD-DBS will undergo bilateral STN DBS.

**Surgical procedure**
STN would be located through direct visualisation in MR images. Quadripolar electrodes would be implanted at the boundary of limbic and associative territories of STN, bilaterally, under local anaesthesia and antiseptic precautions. The amSTN would be targeted 2 mm anterior and 1 mm medial to the motor STN target used for DBS in Parkinson’s disease.\(^1\) The targets would be refined through intraoperative microrecording along multiple trajectories.\(^8\)\(^4\) Intraoperative macrostimulation would be conducted to evaluate the acute effects of stimulation. The trajectory would be chosen based on acute adverse vis-a-vis beneficial effects. The position of the electrodes would be confirmed by postoperative MRI using a 3D atlas MRI co-registration.\(^8\)\(^5\) Electrophysiological activity from STN would be recorded perioperatively and postoperatively. The stimulator would be implanted subcutaneously 5–7 days following STN stimulation after electrophysiological recording.

**Electrophysiological recording from STN**
Perioperative single-unit recording would be recorded through microelectrodes used for STN localisation. Spiking activity would be recorded during 2 min of...
rest and symptom provocation through stereotactically lowered microelectrode leads from various regions of the STN. Symptom provocation would be performed by showing individualised symptom-provoking pictures. LFPs would be recorded through the DBS stimulating electrodes postoperatively. LFPs would be recorded through bipolar montages (to reduce volume conductance effect) from all adjacent contacts, yielding three channels/hemisphere. The electrodes would be externalised through the scalp with extension cables connected to an amplifier for LFP recording. Adequate precautions would be taken to prevent postoperative infection. The LFP recording would be performed on postoperative days 3–5 at rest, during SST, flanker task and under symptom provocation. Power spectral analysis would be performed to study oscillatory activity in various frequency bands (δ=1–3 Hz, θ=4–7 Hz, α=8–11 Hz, β=12–30 Hz and γ>30 Hz).

### Optimisation of stimulation parameters

After the implantation of the stimulator, programming would be finalised over the subsequent 6–8 months. The stimulator would be set at a frequency of 130 Hz and pulse width of 60 μs. Successive trials of monopolar stimulation from each contact would be attempted, beginning with the most ventral contact. Voltage would be increased gradually to monitor for adverse effects. Each contact would be evaluated for 6–8 weeks to evaluate therapeutic efficacy. The final contact would be chosen based on a balance between clinical efficacy and adverse effects.

### Follow-up intervention

Following the optimisation of parameters that may take up to 6 months postimplantation of stimulator and electrodes, the patient would be followed up for another 6 months, considering the latency in onset of improvement with DBS in OCD. DBS would be provided as an unblinded add-on treatment. Although the subjects chosen are treatment-refractory, medication changes and CBT would be permitted during the follow-up period due to ethical concerns as well as to improve the external validity of the study. CBT would also be permitted during follow-up as it may improve the outcome after DBS.

Subjects in the OCD-C would undergo the best alternative treatment for treatment-refractory OCD based on the standard treatment practices of OCD clinic, NIMHANS, and evidence-based clinical practice guidelines. The participants would be unblinded to the group allotment. If they wish to undergo neurosurgical interventions, including DBS for OCD during follow-up, they would be offered the same and would be removed from the control arm of the study.

### Follow-up assessments

#### Clinical assessments

Subjects from both OCD-DBS and OCD-C groups would undergo structured clinical assessment during follow-up. Assessment during follow-up would include a clinical evaluation consisting of monthly assessment of Y-BOCS, CGI-S, HAM-D and HAM-A (box 1). The subjects would do a self-assessment of their obsessive–compulsive symptoms using a smartphone-based app, WHODAS 2.0, WHOQOL-BREF and WHO-5 would be administered every 3 months. Y-BOCS scores at the end of 6 months of follow-up postoptimisation of stimulation parameters would be the primary outcome variable. The structured assessment would be performed through an in-person interview or video conferencing. Subjects in the OCD-C group would also be assessed with same baseline and follow-up clinical assessments, as specified above.

#### Neurocognitive assessment, MRI and EEG

At the end of 6 months of stable stimulation (ie, approximately 12 months postimplantation), both groups of OCD subjects would undergo a detailed neurocognitive assessment as well as EEG (using the same paradigms described above). Subjects in the OCD-DBS group would undergo follow-up EEG both in the stimulation turned ‘ON’ and ‘OFF’ phases on 2 consecutive days. The DBS artifacts in the follow-up EEG recordings during the ‘ON’ phase would be removed using recommended techniques, including oversampling, temporal low-pass filtering and frequency-domain Hampel filtering. The effect of DBS on neurocognitive and EEG markers would be evaluated and correlated with symptom change. Acquiring MRI scans with DBS electrodes in situ may carry the risk of image distortion, interference with DBS stimulator and risk of tissue damage due to overheating of the electrodes. However, recent reports suggest that MRI scans could be obtained safely under appropriate precautions. DBS manufacturers recommend precautions for acquiring MRI scans in this population, such as acquisition using 1.5T MRI. Thus, postsurgical MRI scans (including T1-weighted structural images, fMRI and DTI) would be acquired under appropriate precautions, only for subjects who provide informed consent after explaining the risks and benefits associated with the procedure. Subjects in the OCD-C group would undergo follow-up MRI, EEG and neurological assessments at the end of 12 months, using the same paradigms and protocols as in the OCD-DBS group. As the subjects would not undergo surgery, the electrophysiological recording would not be obtained from the STN. The assessment schedule for the three groups is shown in table 1.

The OCD-C subjects would serve as a comparator for the intervention group to establish the efficacy of DBS. The baseline and follow-up biomarkers would be compared between the two groups using the standard statistical techniques mentioned in the protocol. The healthy volunteer group would serve as a comparator to establish the baseline differences in MRI, EEG and neurocognitive test performances with the OCD subjects.

The statistical analysis would be conducted using the SPSS software (SPSS, Chicago, Illinois, USA). The fMRI processing and analysis would be performed using the Statistical Parametric Mapping (SPM) software (http://www.fil.ion.ucl.ac.uk/spm/). Functional connectivity...
professionals. If subjects from the OCD-monitored for adverse effects by trained mental health
OCD—benefits of participating in the study. Subjects in the
obtaining written informed consent explaining the risks
pre-
16 July 2019). The trial has been registered in the Clin-
Identification
WHOQOL-BREF, WHO quality of life instrument; 69
WHOQOL-BREF, WHO quality of life instrument; 69
-analysis would be conducted using SPM toolboxes and
FMRI Software Library (FSL) (https://fsl.fmrib.ox.ac.
uk/fsl/fslwiki/). Resting-state fMRI would be analysed
using SPM, FSL and Data Processing & Analysis for Brain
Imaging (DPABI) toolbox (http://rfmri.org/dpabi)
according to established processing pipelines. DBS elec-
trodes would be localised from postoperative and preop-
erative images following standardised pipelines using
appropriate toolboxes (eg, lead DBS). 92 The structural
connectivity of volume of tissue activated and location of
active electrode within STN would be evaluated as a poten-
tial predictor of outcome. EEG analysis and electrophys-
iological analysis would be conducted using appropriate
software. The demographic, clinical (including the prin-
cipal symptom dimension and age-at-onset), neurocog-
nitive and electrophysiological data will be examined for
normality. Correlational followed by multiple regression
analysis would be used to assess whether the biomarkers
predict clinical response (Y-BOCS score) at the end of 6
months of stable stimulation. Similar analyses would be
carried out to test other study objectives.

Ethical considerations
The study proposal and the informed consent forms
have been approved by NIMHANS Ethics Committee
(No.NIMHANS/EC(BEH.SC.DIV; 19th meeting dated
17 June 2019). The trial has been registered in the Clin-
cical Trial Registry of India (CTRI/2020/05/024131)-
pre-results stage. The subjects would be recruited after
obtaining written informed consent explaining the risks
and benefits of participating in the study. Subjects in the
OCD-DBS group would be diligently educated about and
monitored for adverse effects by trained mental health
professionals. If subjects from the OCD-C group wish to
undergo neurosurgical interventions for OCD during
follow-up, they would be offered the same and would be
removed from the control arm. Postoperative MRI would
be conducted only in consenting subjects from the OCD
(intervention) group after explaining the risks and ben-
efits. The subjects in the intervention group would be
protected by an insurance plan funded by the research
proposal during the process of the study. A data safety
monitoring board would be constituted to periodically
review the progress of the study to assess safety issues and
recommend continuation, modification and termination
of the trial as required. All OCD subjects can continue
to receive treatment from the OCD clinic, NIMHANS,
following the study proposal irrespective of whether they
complete the study.

Dissemination
The data will be anonymised, curated and stored in a
computer hard disk, with appropriate backup. In keeping
with the policies of the funding agency, the data and meta-
data would be shared with the wider research community
through appropriate platforms. The study findings would
be disseminated in peer-reviewed scientific publications
and scientific conferences, irrespective of outcome.

Expected outcome
Identification of biomarkers of treatment response to
bilateral STN DBS in OCD would pave way for person-
alised medicine by assisting patient selection for this
invasive and expensive treatment. Furthermore, this would
help understand the mechanisms of action of treatment,
which would assist in future refinement/optimisation of
treatment and open the scope for innovations, including
closed-loop stimulation.

| Table 1  | Assessment schedule |
|---------|---------------------|
|         | OCD intervention group | OCD control group | Healthy volunteers |
| MINI 7.0.2 | Baseline | Baseline | Baseline |
| Y-BOCS | Baseline every month | Baseline every month | Baseline |
| CGI-S | Baseline every month | Baseline every month | Baseline |
| HAM-A | Baseline every month | Baseline every month | Baseline |
| HAM-D | Baseline every month | Baseline every month | Baseline |
| WHOQOL-BREF | Baseline every 3 months | Baseline every 3 months | Baseline |
| WHODAS 2.0 | Baseline every 3 months | Baseline every 3 months | Baseline |
| WHO-5 | Baseline every 3 months | Baseline every 3 months | Baseline |
| Neurocognitive testing | Baseline end of 12 months | Baseline end of 12 months | Baseline |
| MRI | Baseline end of 12 months* | Baseline end of 12 months | Baseline |
| EEG | Baseline end of 12 months | Baseline end of 12 months | Baseline |
| Subthalamic nucleus recording | Baseline | x | x |

* Would be acquired after obtaining separate informed consent

CGI-S, Clinical Global Impressions—Severity; 64 EEG, electroencephalography; HAM-A, Hamilton Anxiety Rating Scale; 67 HAM-D, Hamilton Depression Rating Scale; 66 MINI 7.0.2, 65 Mini International Neuropsychiatric Interview; OCD, obsessive–compulsive disorder; WHO-5, WHO-5Five Well-Being Index; 60 WHODAS 2.0, WHO Disability Assessment Schedule; 69 WHOQOL-BREF, WHO quality of life instrument; 69 Y-BOCS, Yale-Brown Obsessive Compulsive Scale. 59
REFERENCES

1 Stein DJ, Costa DLC, Lochner C. Obsessive–compulsive disorder. Nat Rev Dis Primers 2019;5:1–21.
2 Collins PY, Patel V, Joestl SS, et al. Grand challenges in global mental health. Nature 2011;475:27–30.
3 Arumugham SS, Reddy YCJ. Commonly asked questions in the treatment of obsessive-compulsive disorder. Expert Rev Neurother 2014;14:151–63.
4 Fineberg NA, Hollander E, Pallanti S, et al. Clinical advances in obsessive-compulsive disorder: a position statement by the International College of obsessive-compulsive spectrum disorders. Int Clin Psychopharmacol 2020;35:173–93.
5 Bear RE, Fitzgerald P, Rosenfeld JV, et al. Neurosurgery for obsessive-compulsive disorder: contemporary approaches. J Clin Neurosci 2010;17:1–5.
6 Janardhan Reddy YC, Sundar AS, Narayanaswamy JC, et al. Clinical practice guidelines for obsessive-compulsive disorder. Indian J Psychiatry 2017;59:74–90.
7 Kohl S, Schönher DM, Luigjes J, et al. Deep brain stimulation for treatment-refractory obsessive compulsive disorder: a systematic review. BMC Psychiatry 2014;14:214.
8 Vicheva P, Butler M, Shotbolt P. Deep brain stimulation for obsessive–compulsive disorder: a systematic review of randomised controlled trials. Neurosci Biobehav Rev 2020;109:129–38.
9 Kumar KK, Appelboom G, Lamsam L, et al. Comparative effectiveness of neuroablation and deep brain stimulation for treatment-resistant obsessive-compulsive disorder: a meta-analytic study. J Neurol Neurosurg Psychiatry 2019;90:469–73.
10 Balachander S, Arumugham SS, Srinivas D. Ablative neurosurgery and deep brain stimulation for obsessive-compulsive disorder. Indian J Psychiatry 2019;61:577–84.
11 van den Heuvel OA, van Wingen G, Soriano-Mas C, et al. Brain circuitry of compulsivity. Eur Neuropsychopharmacol 2016;26:810–27.
12 Doughtery DD, Brennan BP, Stewart SE, et al. Neuroscientifically informed formulation and treatment planning for patients with obsessive-compulsive disorder: a review. JAMA Psychiatry 2018;75:1081–7.
13 Martinho FP, Duarte GS, Couto FSdo, do CFS. Efficacy, effect on mood symptoms, and safety of deep brain stimulation in refractory obsessive-compulsive disorder: a systematic review and meta-analysis. J Clin Psychiatry 2020;81. doi:10.4088/JCP.19dr12821. [Epub ahead of print: 26 05 2020].
14 Mallet L, Polosan M, Jaafar N, et al. Subthalamic nucleus stimulation in severe obsessive-compulsive disorder. N Engl J Med 2008;359:2121–34.
15 Staudt MD, Pouratian N, Miller JP, et al. Congress of neurological surgeons systematic review and evidence-based guidelines for deep brain stimulations for obsessive-compulsive disorders: update of the 2014 guidelines. Neurosurgery 2020;87:10–27.
16 Tyagi H, Apergis-Schoume AT, Akram H, et al. A randomized trial directly comparing ventral capsule and anteroomedial subthalamic nucleus stimulation in obsessive-compulsive disorder: clinical and imaging evidence for dissociable effects. Biol Psychiatry 2019;85:726–34.
17 Welter M-L, Alves Dos Santos JF, Clair A-H, et al. Deep brain stimulation of the subthalamic, acumbens, or caudate nuclei for patients with severe obsessive-compulsive disorder: a randomized crossover controlled study. Biol Psychiatry 2020;10;Li N, Balderrmann JC, Kibbel A, et al. A unified connectomic target for deep brain stimulation in obsessive-compulsive disorder. Nat Commun 2020;11:3364.
18 Haber SN, Vendli A, Jabadi S. Four deep brain stimulation targets for obsessive-compulsive disorder: are they different? Biological Psychiatry.
19 Alonso P, Cuadras D, Gabriëls L, et al. Deep brain stimulation for obsessive-compulsive disorder: a meta-analysis of treatment outcome and predictors of response. PLoS One 2015;10:e0133591.
20 Ooms P, Mantione M, Fige M, et al. Deep brain stimulation for obsessive-compulsive disorders: long-term analysis of quality of life. J Neurol Neurosurg Psychiatry 2014;85:153–8.
21 Mallet L, Du Montcel ST, Clair A-H, et al. Long-term effects of subthalamic stimulation in obsessive-compulsive disorder: follow-up of a randomized controlled trial. Brain Stimul 2019;12:1080–2.
22 van Westen M, Rietveld E, Fige M, et al. Clinical outcome and mechanisms of deep brain stimulation for obsessive-compulsive disorder. Curr Behav Neurosci Rep 2015;2:41–8.
23 Saxena S, Rauch SL. Functional neuroimaging and the neuroanatomy of obsessive-compulsive disorder. Psychiatr Clin North Am 2000;23:563–86.
24 Burgièire E, Monteiro P, Mallet L, et al. Striatal circuits, habits, and implications for obsessive-compulsive disorder. Curr Opin Neurol Biol 2015;30:59–65.
25 Burgièire E, Monteiro P, Feng G, et al. Optogenetic stimulation of lateral orbitofronto-striatal pathway suppresses compulsive behaviors. Science 2013;340:1243–6.
26 Robbins TW, Vaghi MM, Banca P. Obsessive–Compulsive Disorder: puzzles and prospects. Neuron 2019;102:27–47.
27 Ahmari SE, Spellman T, Douglass NL, et al. Repeated cortico-striatal stimulation generates persistent OCD-like behavior. Science 2013;340:1234–9.
28 Rotge J-Y, Guell D, Diliarreugy B, et al. Provocation of obsessive-compulsive symptoms: a quantitative voxel-based meta-analysis of functional neuroimaging studies. J Psychiatry Neurosci 2008;33:405–12.
30 Saxena S, Brody AL, Ho ML, et al. Differential brain metabolic predictors of response to paroxetine in obsessive-compulsive disorder versus major depression. *Am J Psychiatry* 2003;160:522–32.
31 Saxena S, Brody AL, Maitem KM, et al. Localized orbitofrontal and subcortical metabolic changes as predictors of response to paroxetine treatment in obsessive-compulsive disorder. *Neuropsychopharmacology* 1999;21:683–93.
32 Figeer M, Luigjes J, Smolders R, et al. Deep brain stimulation restores frontostriatal network activity in obsessive-compulsive disorder. *Nat Neurosci* 2013;16:386–7.
33 Millet B, Jaafar N, Polosan M, et al. Limbic versus cognitive target for deep brain stimulation in treatment-resistant depression: accumbens more promising than caudate. *Eur Neuropsychopharmacol* 2014;24:1229–39.
34 Mulders AEP, Mennes M, BR, Schruers K, et al. Deep brain stimulation of the subthalamic nucleus in obsessive-compulsive disorder: neuroanatomical and pathophysiological considerations. *Eur Neuropsychopharmacol* 2016;26:1909–19.
35 Lambert C, Zinno Z, Nagi Z, et al. Functionality of functional zones within the human subthalamic nucleus: patterns of connectivity and sub-parcellation using diffusion weighted imaging. *Neuroimage* 2012;60:83–94.
36 Haynes WIA, Haber SN. The organization of prefrontal-subthalamic inputs in primates provides an anatomical substrate for both functional specificity and integration: implications for basal ganglia models and deep brain stimulation. *J Neurosci* 2013;33:4804–14.
37 Voon V, Droux F, Morris L, et al. Decisional impulsivity and the associative-limbic subthalamic nucleus in obsessive-compulsive disorder: stimulation and connectivity. *Brain* 2017;140:442–56.
38 Wiesciak TV, Frank MJ. A computational model of motor control in frontal cortex and basal ganglia. *Psychol Rev* 2013;120:329–55.
39 Aron AR, Behrens TE, Smith S, et al. Triangulating a control network using diffusion-weighted magnetic resonance imaging (MRI) and functional MRI. *J Neurosci* 2007;27:3743–52.
40 Aron AR, Poldrack RA. Cortical and subcortical contributions to stop signal response inhibition: role of the subthalamic nucleus. *J Neurosci* 2006;26:2424–33.
41 Frank MJ. Hold your horses: a dynamic computational role for the subthalamic nucleus in decision making. *Neural Netw* 2006;19:1120–33.
42 Herz DM, Little S, Pedrosa DJ, et al. Mechanisms underlying decision-making as revealed by deep-brain stimulation in patients with Parkinson’s disease. *Curr Biol* 2018;28:1169–78.
43 Pasquevau B, Turner RS. A selective role for ventromedial subthalamic nucleus in inhibitory control. *Elife* 2017;6:e31627.
44 Chamberlain SR, Fineberg NA, Nutt DJ, et al. Motor inhibition and cognitive flexibility in obsessive-compulsive disorder and trichotillomania. *Am J Psychiatry* 2006;163:1282–4.
45 Gu B-M, Park J-Y, Kang D-H, et al. Neural correlates of cognitive inflexibility during go/no-go switching in obsessive-compulsive disorder. *Brain* 2008;131:155–64.
46 Ballanger B, van Iemen T, Moro E, et al. Stimulation of the subthalamic nucleus and impulsivity: release your horses. *Ann Neurol* 2009;66:917–24.
47 Green N, Bogacz R, Huelb J, et al. Reduction of influence of task difficulty on perceptual decision making by STN deep brain stimulation. *Curr Biol* 2013;23:1681–4.
48 Greenhouse I, Gould S, Houser M. Stimulation of contacts in ventral but not dorsal subthalamic nucleus normalizes response switching in Parkinson’s disease. *Neuropsychologia* 2013;51:1302–9.
49 Kibielur A, Gras-Combe G, Benis D, et al. Modulation of motor inhibition by subthalamic stimulation in obsessive-compulsive disorder. *Transl Psychiatry* 2016;6:e922.
50 Morris LS, Baek E, Voon V. Distinct cortico-striatal connections with subthalamic nucleus underlie facets of compulsivity. *Cortex* 2017;88:143–50.
51 Le Jeune F, Virin M, ND’Iaye K, ND’Iaye K, et al. Decrease of prefrontal metabolism after subthalamic stimulation in obsessive-compulsive disorder: a positron emission tomography study. *Biol Psychiatry* 2010;68:1016–22.
52 Senova S, Clair A-H, Palfi S, et al. Deep brain stimulation for refractory obsessive-compulsive disorder: towards an individualized approach. *Front Psychiatry* 2019;10:905.
53 Alegre M, Lopez-Azarate J, Obeso J, et al. The subthalamic nucleus is involved in successful inhibition in the stop-signal task: a local field potential study in Parkinson’s disease. *Exp Neurol* 2013;239:1–12.
54 Zavala B, Tan H, Ashkan K, et al. Human subthalamic nucleus-medial frontal cortex theta phase coherence is involved in conflict and error related cortico-thalamic interactions. *Alzheimers Dement* 2017;13:179–87.
55 Riesel A, Kathmann N, Endrass T. Overactive performance monitoring in obsessive-compulsive disorder is independent of symptom expression. *Eur Arch Psychiatry Clin Neurosci* 2014;264:707–17.
56 Chamberlain SR, Fineberg NA, Menzies LA, et al. Impaired cognitive flexibility and motor inhibition in unaffected first-degree relatives of patients with obsessive-compulsive disorder. *Am J Psychiatry* 2007;164:335–8.
57 Welter M-L, Burbaud P, Fernandez-Vidal S, et al. Basal ganglia dysfunction in OCD: subthalamic neuronal activity correlates with symptoms severity and predicts high-frequency stimulation efficacy. *Transl Psychiatry* 2011;1:e6.
58 Rappel P, Marmor O, Bick AS, et al. Subthalamic theta activity: a novel human subcortical biomarker for obsessive compulsive disorder. *Transl Psychiatry* 2018;8:1–11.
59 Buot A, Karachi C, Lau B. Emotions Modulate Subthalamic Nucleus Activity: New Evidence in Obsessive-Compulsive Disorder and Parkinson’s Disease Patients. Biological Psychiatry: Cognitive Neuroscience and Neuroimaging. (Published Online First: 11 August 2020).
60 Chan A-W, Tetzlaff JM, Gotzsche PC, et al. Spirit 2013 explanation and elaboration: guidance for protocols of clinical trials. *BMJ* 2013:346:e7586.
61 American Psychiatric Association. *Diagnostic and Statistical Manual of Mental Disorders*. American Psychiatric Association, 2013. Available: http://dx.doi.org/10.1176/appi.books.9780890425596.ch1.
62 Sheehan DV, Lecrubier Y, Sheehan KH, et al. The Mini-International Neuropsychiatric Interview (M.I.N.I.): the development and validation of a structured diagnostic psychiatric interview for DSM-IV and ICD-10. *J Clin Psychiatry* 1998;59 Suppl 20:22–33.
63 Goodman WK, Priche P. The Yale-Brown obsessive compulsive scale. I. Development, use, and reliability. *Arch Gen Psychiatry* 1989;46:1006–11.
64 Guy W. *ECDEU Assessment Manual for Psychopharmacology* — Revised. Rockville, MD: U.S. Department of Health, Education, and Welfare, Public Health Service, Alcohol, Drug Abuse, and Mental Health Administration, NIMH Psychopharmacology Research Branch, Division of Extramural Research Programs 1976:218–22.
65 The Rehabilitation Committee of the Indian Psychiatric Society. *Ideas (Indian disability evaluation and assessment scale) — a scale for measuring and quantifying disability in mental disorders.* Gurgaon, India: Indian Psychiatric Society.
66 Hamilton M. A rating scale for depression. *J Neurol Neurosurg Psychiatry* 1960;23:56–62.
67 Hamilton M. The assessment of anxiety states by rating. *Br J Med Psychol* 1959;32:50–5.
68 Who | who disability assessment schedule 2.0 (WHODAS 2.0). who. Available: http://www.who.int/classifications/icf/who/ disabilities/en/ [Accessed 16 May 2018].
69 Skevington SM, Lofy M, O’Connell KA, et al. The world Health organization’s WHOQOL BREF quality of life assessment: psychometric properties and results of the International field trial. A report from the WHOQOL group. *Qual Life Res* 2004;13:299–310.
70 Bech P, Olsen LR, Kjoller M, et al. Measuring well-being rather than the absence of distress symptoms: a comparison of the SF-36 mental health subscale and the WHO-Five Well-being scale. *Int J Methods Psychiatr Res* 2003;12:85–91.
71 Rao S, Subbakrishna DK, Gopukumar K. NHMANS neuropsychological battery. Bangalore, India: National Institute of Mental Health and Neurosciences, 2004.
72 Logan GD, Cowan WB, Davis KA. On the ability to inhibit simple and choice reaction time responses: a model and a method. *J Exp Psychol Hum Percept Perform* 1984;10:276–91.
73 Jacoby RJ, Abramowitz JS, Buck BE, et al. How is the beads task related to intolerance of uncertainty in anxiety disorders? *J Anxiety Disord* 2014;28:495–503.
74 Kable JW, Gliewer PW. The neural correlates of subjective value during intertemporal choice. *Nat Neurosci* 2007;10:1625–33.
75 Figner B, Knoch D, Johnson EJ, et al. Lateral prefrontal cortex and self-control in intertemporal choice. *Nat Neurosci* 2010;13:538–9.
76 Steinglass JE, Lempert KM, Choo T-H, et al. Temporal discounting across three psychiatric disorders: anorexia nervosa, obsessive compulsive disorder, and social anxiety disorder. *Depress Anxiety* 2017;34:463–70.
77 Norman LJ, Carlisi CO, Christakou A, et al. Neural dysfunction during temporal discounting in paediatic attention-deficit/hyperactivity disorder and obsessive-compulsive disorder. *Psychiatry Res Neuroimaging* 2017;269:97–105.
78 Lebert F, Pasquier F, Soulez L, et al. Frontotemporal behavioral scales. *Alzheimer Dis Assoc Disord* 1998;12:335–9.
79 Barrash J, Anderson S. The Iowa Scales for Personality Change. Iowa City: University of Iowa, Department of Neurology, 1997.
80 Whiteside SP, Lynam DR. The five factor model and impulsivity: using a structural model of personality to understand impulsivity. *Pers Individ Dif* 2001;30:669–89.

81 Simon D, Adler N, Kaufmann C, et al. Amygdala hyperactivation during symptom provocation in obsessive-compulsive disorder and its modulation by distraction. *Neuroimage Clin* 2014;4:549–57.

82 Sanematsu H, Nakao T, Yoshiura T, et al. Predictors of treatment response to fluvoxamine in obsessive-compulsive disorder: an fMRI study. *J Psychiatr Res* 2010;44:193–200.

83 Olatunji BO, Davis ML, Powers MB, et al. Cognitive-Behavioral therapy for obsessive-compulsive disorder: a meta-analysis of treatment outcome and moderators. *J Psychiatr Res* 2013;47:33–41.

84 Hutchison WD, Allan RJ, Opitz H, et al. Neurophysiological identification of the subthalamic nucleus in surgery for Parkinson’s disease. *Ann Neurol* 1998;44:622–8.

85 Yelnik J, Damier P, Demeret S, et al. Localization of stimulating electrodes in patients with Parkinson disease by using a three-dimensional atlas-magnetic resonance imaging coregistration method. *J Neurosurg* 2003;99:89–99.

86 Mantione M, Nieman DH, Figee M, et al. Cognitive-Behavioural therapy augments the effects of deep brain stimulation in obsessive-compulsive disorder. *Psychol Med* 2014;44:3515–22.

87 Lio G, Thobois S, Ballanger B, et al. Removing deep brain stimulation artifacts from the electroencephalogram: issues, recommendations and an open-source toolbox. *Clin Neurophysiol* 2018;129:2170–85.

88 Carmichael DW, Pinto S, Limousin-Dowsey P, et al. Functional MRI with active, fully implanted, deep brain stimulation systems: safety and experimental confounds. *Neuroimage* 2007;37:508–17.

89 Bronstein JM, Tagliati M, Alterman RL, et al. Deep brain stimulation for Parkinson disease: an expert consensus and review of key issues. *Arch Neurol* 2011;68:165. doi:10.1001/archneurol.2010.260

90 Zrinzo L, Yoshida F, Hariz MI, et al. Clinical safety of brain magnetic resonance imaging with implanted deep brain stimulation hardware: large case series and review of the literature. *World Neurosurg* 2011;76:164–72. doi:10.1016/j.wneu.2011.02.029

91 Saleh C, Dooms G, Berthold C, et al. Post-Operative imaging in deep brain stimulation: a controversial issue. *Neuroradiol J* 2016;29:244–9. doi:10.1177/1971400916639960

92 Horn A, Li N, Dembek TA, et al. Lead-DBS V2: towards a comprehensive pipeline for deep brain stimulation imaging. *Neuroimage* 2019;184:293–316. doi:10.1016/j.neuroimage.2018.08.068