Review

Efficacy of deep brain stimulation for treatment-resistant obsessive-compulsive disorder: systematic review and meta-analysis

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ABSTRACT
Deep brain stimulation (DBS) is an established and growing intervention for treatment-resistant obsessive-compulsive disorder (TROCD). We assessed current evidence on the efficacy of DBS in alleviating OCD and comorbid depressive symptoms including newly available evidence from recent trials and a deeper risk of bias analysis than previously available. PubMed and EMBASE databases were systematically queried using Preferred Reporting Items for Systematic reviews and Meta-Analyses guidelines. We included studies reporting primary data on multiple patients who received DBS therapy with outcomes reported through the Yale-Brown Obsessive-Compulsive Scale (Y-BOCS). Primary effect measures included Y-BOCS mean difference and per cent reduction as well as responder rate (≥35% Y-BOCS reduction) at last follow-up. Secondary effect measures included standardised depression scale reduction. Risk of bias assessments were performed on randomised controlled (RCTs) and non-randomised trials. Thirty-four studies from 2005 to 2021, 9 RCTs (n=97) and 25 non-RCTs (n=255), were included in systematic review and meta-analysis based on available outcome data. A random-effects model indicated a meta-analytical average 14.3 point or 47% reduction (p<0.01) in Y-BOCS scores without significant difference between RCTs and non-RCTs. At last follow-up, 66% of patients were full responders to DBS therapy. Sensitivity analyses indicated a low likelihood of small study effect bias in reported outcomes. Secondary analysis revealed a 1 standardised effect size (Hedges’ g) reduction in depressive scale symptoms. Both RCTs and non-RCTs were determined to have a predominantly low risk of bias. A strong evidence base supports DBS for TROCD in relieving both OCD and comorbid depression symptoms in appropriately selected patients.

INTRODUCTION
Obsessive-compulsive disorder (OCD) is a complex neuropsychiatric illness characterised by intrusive and persistent obsessive thoughts along with dysfunctional and ritualised behaviours. The disorder often begins in childhood, puberty, or early adulthood and thus affects a critical period of development. OCD can be a debilitating disease with many patients experiencing severe comorbid depressive and anxiety disorders as well as the inability to work or attend school. The lifetime prevalence of OCD in the general population is 1%–3%, and while 50%–70% of patients can significantly improve with conventional therapies including pharmacotherapy and cognitive–behavioural therapy with exposure and response prevention (ERP), at least 10% of patients will develop severe symptoms refractory to multimodality therapy.

Well-established surgical methods to address treatment-resistant OCD (TROCD) symptoms include various forms of ablative lesioning such as anterior cingulotomy and anterior capsulotomy and have been practised since the 1950s. Over the last two decades, deep brain stimulation (DBS) has emerged as a viable method to treat TROCD, offering an adjustable and partially reversible alternative to ablative techniques, with a similar reported efficacy. The first reported case of DBS for OCD involved targeting of the anterior limb of the internal capsule (ALIC) based on the authors’ previous experiences with anterior capsulotomies. Since the first sham-controlled randomised trial targeting the ALIC in 2005, a multitude of trials have been conducted with varying surgical targets, methodologies and reported outcomes. DBS for TROCD at the ALIC was granted a humanitarian exemption by the US Food and Drug Administration in 2009. A variety of white and grey matter areas including ALIC, subthalamic nucleus (STN), ventral capsule/ventral striatum (VC/VS), bed nucleus of stria terminalis (BNST) and nucleus accumbens (NAC) have been targeted surgically, as all are theorised to share important roles in regulating mood, reward-learning and decision-making within the hypothesised cortico-striato-thalamo-cortical (CSTC) circuit. More recently, it has been described that the various striatal areas targeted in TROCD DBS are likely modulating similar CSTC and orbitofrontal networks and that a conserved pathway is implicated in optimal symptom improvement across targets.

In the evolving field of DBS for TROCD, a better understanding of treatment efficacy across studies with varying methodological designs is desired. To date, several meta-analyses have synthesised the body of evidence on DBS for OCD. However, since the most recent publication, one novel randomised controlled trial (RCT) and at least six observational cohort studies have been published. In addition, previous studies have not attempted to quantitatively assess bias in outcomes reporting.
which may improve confidence in the reported results of variously powered and designed studies. In light of this, we present here a systematic review and meta-analysis with the objective of critically assessing the efficacy of DBS in alleviating OCD and comorbid depressive symptoms across targets in patients with TROCD.

METHODS
A systematic review was completed using 2020 Preferred Reporting Items for Systematic reviews and Meta-Analyses (PRISMA) guidelines. PubMed and EMBASE databases were queried using advanced search strategies including Medical Subject Headings (MeSH) terms in PubMed—“Deep Brain Stimulation”[Mesh] AND “Obsessive-Compulsive Disorder”[Mesh] OR (“OCD” OR “obsessive-compulsive disorder”) AND (“DBS” OR “deep brain stimulation”)—and an exhaustive set of terms in EMBASE including (“OCD” OR ‘obsessive-compulsive disorder’/exp OR ‘obsessive-compulsive disorder’ OR ‘TROCD’) AND (“DBS” OR ‘brain depth stimulation’/exp OR ‘brain depth stimulation’ OR ‘deep brain stimulation’/exp OR ‘deep brain stimulation’). The search was completed through September 2021.

Selection criteria
Studies were included if they met the following criteria: (1) subjects were human adults (age >18 years) with a primary diagnosis of OCD according to the Diagnostic and Statistical Manual of Mental Disorders Fourth or Fifth edition (DSM-IV or DSM-V) or International Classification of Diseases criteria; (2) DBS was the primary intervention; (3) primary outcome was improvement in clinical OCD symptoms after DBS; (4) outcome was assessed using the Yale-Brown Obsessive Compulsive Scale (Y-BOCS); (5) treatment response was defined as a ≥35% reduction in Y-BOCS score; (6) published in English in peer-reviewed journals.

Studies were excluded according to several criteria: (1) reviews, meta-analyses, comments, letters and editorials lacking de novo patients; (2) single case reports; (3) studies investigating neuroimaging, neuropsychiatric, behavioural, and/or electrophysiological changes after DBS as primary outcomes; (4) animal studies; (5) studies focusing on non-OCD indications for DBS; (6) technical reports on the safety or procedural aspects of DBS for OCD.

Selection process
All search results from both PubMed and EMBASE databases were exported to a spreadsheet and duplicates were removed. Studies were excluded using automated detection tools in Excel 2016 (Microsoft Corporation, Redmond, Washington, USA). Two reviewers independently screened the title of each record retrieved. One reviewer screened the abstracts and full texts of all remaining records and reports for eligibility and final inclusion. In any case where several records reported on all or part of the same cohort of patients, the study with the most detailed dataset for the largest number of patients was selected for inclusion.

Data collection and organisation
The following data items were collected wherever available:

- General study information including study location, first author, publication year, study design, patient inclusion and exclusion criteria, sample size, treatment response criteria, response rate, and rates of complications or adverse events.
- Patient-level data including stimulation target(s), primary diagnosis, patient sex, age at onset of OCD, age at DBS surgery, comorbid psychiatric diagnoses, active medications, preoperative/baseline Y-BOCS scores, all follow-up Y-BOCS scores (with time points in months), per cent Y-BOCS score reduction at last follow-up, length of follow-up (in months), DBS stimulation parameters (amplitude, pulse width, frequency, contact configuration, polarity), quality of life outcomes, baseline depression scale (Hamilton Depression Rating Scale (HAM-D, HDRS-17, HDRS-24), Montgomery-Åsberg Depression Rating Scale (MADRS), Depression Anxiety Stress Scale-Depression (DASS-21-D) and Beck Depression Inventory (BDI), baseline anxiety scale (Hamilton Anxiety Rating Scale (HAM-A and HARS), State-Trait Anxiety Inventory (STAI-1/STAI-2/X2), DASS-Anxiety (DASS-21-A) and Beck Anxiety Inventory), and baseline Global Assessment of Functioning (GAF) scores and all follow-up scores.

If patient-level data were not available, pooled means were collected. All data were manually recorded in a single spreadsheet by two reviewers who worked in conjunction.

Risk of bias assessment
Two reviewers completed all risk of bias (RoB) assessments independently. For RCTs, the revised Cochrane tool for assessing risk of bias in randomised trials\(^{24}\) was used to critically evaluate six domains of bias: randomisation, period/carryover (for crossover trials), assignment to intervention, missing outcome, outcome measurement and selection of reported results. Non-RCTs were assessed using Cochrane’s Risk Of Bias In Non-randomized Studies-of Interventions\(^{25}\) which examines seven domains of bias: confounding, selection of participants, classification of interventions, deviations from intended interventions, missing data, measurement of outcomes and selection of the reported results. All bias assessments were performed by two authors who were blinded to each other’s ratings. Following individual assessment, incongruencies were mediated and results were aggregated and visualised using the RoB visualisation web app robvis.

Meta-analytical methods
Meta-analysis was performed with the meta package (RRID: SCR_019055) using R statistical computing software (R Core Team, 2021). Primary outcome measures were (1) Y-BOCS mean difference at last follow-up compared with baseline, (2) treatment effect, or Y-BOCS per cent reduction, at last follow-up, and (3) responder rate at last follow-up. Secondary outcomes included (1) standardised mean difference (SMD) in depression scale scores and (2) depression scale responder rate, defined as >50% reduction in depression scores.\(^{26}\) As different studies used varying scales (ie, MADRS, HDRS, BDI) to assess depression preoperatively and postoperatively, SMD was calculated using the Hedges’ g statistic, which is a commonly used method to compare depression scales.\(^{26,27}\) To determine a treatment effect size corrected by pooled sample variance across studies. A g of <0.2 indicates a small effect, 0.5–0.8 indicates a medium effect and >0.8 indicates a statistically large effect.\(^{27}\) Both fixed and random-effects models were used to synthesise primary and secondary outcomes and assess the relative effects of smaller studies compared with larger ones. The DerSimonian-Laird method was used to assess between-study variance (\(τ^2\)) and I\(^2\) was used to determine the fraction of the variance due to various forms of study heterogeneity (eg, statistical, clinical, methodological).\(^{28}\) Funnel plot analysis was performed to evaluate small
study effect biases and symmetry was quantitatively assessed using the Thompson-Sharps test for asymmetry.28 Subgroup analysis was performed on RCTs and non-RCTs to determine possible causes of outcome heterogeneity.28 Furthermore, meta-regression was performed on primary outcome measures using the model sum of square statistic (Q_M) covaried by reported duration and one used the DASS-D46 to assess depression preoperatively and postoperatively. Of the 16 studies that reported anxiety scores, 7 used a Hamilton anxiety scale (9 HAM-A, 2 HARS),9 22 23 30 32 33 35–37 39–41 47 50 51 seven studies used STAI-A,2 21 22 29 54 and I used DASS-A.46 Seventeen studies reported GAF scores; however, seven of these studies only reported baseline scores.

Study characteristics: safety
Approximately 70% of studies (24 of 34; n=249) reported complete data on serious adverse events (SAEs), including, but not limited to: hardware-related complications, infections, seizures, suicide attempts, intracranial haemorrhage (ICH) and the development of de novo obsessions associated with stimulation. Overall, ~31% of patients (n=78) experienced at least one SAE. Incidence of device-related complications, that is, lead damage or malposition, was ~3% (n=20). There were 11 cases of postoperative infection (~2.4%)—of which 6 required explantation and/or replacement of a pulse generator—and 9 instances of postoperative seizure (~3.6%). One patient (included in both infection and seizure groups) experienced several SAEs, including a generalised tonic–clonic seizure, intracranial infection, shock and a pharmacologically induced coma.48 Additionally, there were six cases of attempted suicide (~3.6%). Six studies reported five cases of postoperative ICH (~1.6%); of which one resulted in prolonged finger palsy25 and another resulted in prolonged dystonia.35 Finally, in two cases (0.8%), DBS therapy itself became the source of a new obsession (eg, checking settings and battery life), which contributed to worsening OCD.40 47 For a
full list of reported complications and adverse events, see online supplemental file A.

**RoB within studies**

In RCTs, RoB in randomisation, missing outcomes and selection of reported results was found to be low in all studies. RoB in assignment to intervention and outcome measurement was low in most studies but uncertain in several due to unblinding of investigators secondary to adverse events during the trial period. Three of the nine RCTs included dedicated washout periods from stimulation ON to stimulation OFF arms and thus incurred a low RoB in this domain.

One study, which did not have a washout period in its design, tested specifically for carryover effects and found none, thus incurring a low RoB.

Four others, which included neither washout periods nor any structured analyses testing for carryover effects, prompted an unclear RoB designation. The most recent RCT was a parallel-arm non-crossover study.

Of the non-RCTs, nearly all studies demonstrated low-moderate RoB in participant selection, intervention classification, intended intervention deviation, missing data, outcome measurement and selection of reported results. Approximately one-third of studies demonstrated a low RoB due to confounding, while the other two-thirds had a moderate-high RoB due to confounding. This risk was most often related to within-study variability in target implanted, length of follow-up or baseline OCD severity/phenotype. See figure 2 for full RoB assessments of RCTs and non-RCTs.

**Synthesis of results**

The meta-analytical Y-BOCS mean difference (MD) at last follow-up was 14.28 (95% CI 12.51 to 16.05) points across 345 patients pooled from 31 studies. Three of the 34 studies were excluded from MD meta-analysis as they did not report individual Y-BOCS scores both preoperatively and postoperatively.

Using available pre-disambiguated and post-disambiguated data, the meta-analytical treatment effect (TE) was found to be a 47% (95% CI 40% to 53%) reduction in Y-BOCS scores at last follow-up across 249 patients from 28 studies in which precision estimates could be gathered or measured. The responder rate (RR) at last follow-up was found to be 66% (95% CI 57% to 74%). Both Y-BOCS MD and TE demonstrated pooled statistical significance ($p<0.01$). There was, however, statistically significant between-study variance in both measures with between 70% and 84% of the variance arising due to study heterogeneity.

Funnel plot analysis was performed on TE data. The relative symmetry of the dispersion of SE plotted against treatment effect along with the finding that larger studies tended toward lower errors while smaller studies tended toward larger errors suggests low risk for small study effects bias. Twenty-seven of 28 studies achieved statistically significant TE ($p<0.05$). The lone study that reported non-significant effects ($p>0.1$) was the only one to target MD/VA thalamus. Furthermore, a Thompson-Sharps test for funnel plot asymmetry demonstrated random dispersion of TE relative to SE indicating a low

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**Figure 2** Risk of bias assessments. (A) Individual RCT Cochrane RoB 2 breakdown and (B) summary of RCT assessments. (C) Individual Cochrane ROBINS breakdown for non-RCTs and (D) summary of non-RCT assessments. RCT, randomised controlled trial; RoB 2, Cochrane tool for assessing risk of bias in randomised trials; ROBINS, Risk Of Bias In Non-randomized Studies.
risk for between-study heterogeneity due to small study effects bias (t=1.11, df=19, p=0.2814) (online supplemental file B). Subgroup analysis by methodology demonstrated no significant differences between RCTs and non-RCTs in any primary outcome measure by random-effects modelling (MD p=0.87, TE p=0.67, RR p=0.61). Meta-regression of Y-BOCS outcomes by striatal versus non-striatal targets (Q M(df=2)=1.92, p=0.385), as well as across various reported targets (Q M(df=13)=19.99, p=0.095), demonstrated no significant difference. Interestingly, studies explicitly targeting NAc (adjusted p<0.05) and MD/VA (adjusted p<0.01) did report Y-BOCS effects of lower magnitude than other reported targets.

Raw mean reduction in depressive symptoms was 7.15 points (95% CI 4.82 to 9.47) or 40% (n=140) with HDRS, 11.89 points (95% CI 2.66 to 21.13) or 41% (n=68) with MADRS, and 8.43 points (95% CI 4.80 to 12.05) or 32% (n=57) with BDI. Using pooled mean averages and variances, overall SMD calculation yielded an approximate 1 standardised effect size (Hedges’ g) reduction in depressive scale scores (95% CI 0.67 to 1.32) (figure 4). Of note, there was a significant difference between studies using HDRS or MADRS compared with those using BDI, whereby those using the latter reported a less significant effect size (χ2=8.89, df=2, p=0.01). Using available reported data at last follow-up, we found that 47% of patients were considered full responders relative to their preoperative/baseline comorbid depression. An additional 16% of patients were considered partial responders (30%–49% reduction in pre/post-treatment depressive symptoms) and 37% were considered non-responders (<30% reduction).

**DISCUSSION**

We report the largest meta-analysis of efficacy and mood response data concerning the use of DBS for TROCD to date. Our results indicate that patients with severe TROCD experience a near 50% reduction in their OCD symptoms by a median follow-up of approximately 24 months. Sixty-six per cent of patients in well-designed studies achieved response to DBS therapy, which compares with or outperforms recent estimates of treatment response with lesional procedures (36%–59%). A relatively recent review compiled data in order to compare the outcomes and complications of anterior capsulotomy versus DBS of the VC/VS or NAc for OCD and found that both procedures carry similar risk–benefit profiles. We found a strong effect of DBS for TROCD on comorbid depression, with nearly half of reported patients attaining a complete response and an additional 16% at least partially responding to therapy.

A previous meta-analysis of a smaller set of DBS for TROCD data found a correlative effect between Y-BOCS and depression response. A potential and intuitive explanation for the strong co-therapeutic effect seen in our and previous results is that the more a patient’s OCD symptoms improved, the more positive the effect was on their mood. The same study found that illness severity at baseline was a negative predictor of treatment response at last follow-up. Their results give credence to the idea that studies that enroll patients with overly variable illness severities assume a potential risk of variability in results that can be avoided by further narrowing severity inclusion criteria. Additionally, our finding that NAc targeting resulted in lower magnitude effects supports the idea that the optimal target is not
grey matter (NAc/VS) but rather white matter. This observation reinforces the developing idea that optimal targeting requires engaging a network of regions via cortical white matter hubs.11,58

Of the nine RCTs included, only three had a washout period. In a crossover study, the washout period should be of sufficient duration to eliminate carryover effects (ie, the treatment effects from a stimulation ON period carrying over into a stimulation OFF period, resulting in an underestimation of the overall treatment effect). The absence of a washout period is thus a potentially significant source of bias. RCTs included in our meta-analysis had stimulation ON periods ranging from 2 weeks to 3 months. In the case of the study using 2-week crossover periods (ON/OFF), despite the lack of washout, the investigators found no significant carryover effects.32 One explanation for this is that 2 weeks of stimulation is likely not long enough to significantly alter the circuits implicated in OCD. On the other hand, for the four RCTs with a 3-month crossover design, the likelihood of carryover effects is much higher after a 3-month stimulation period, and so the washout period omission should be considered when interpreting results.

For non-RCTs, two-thirds of those included in our meta-analysis carried a moderate-high RoB due to confounding. The most prominent confounder in this group of studies was the deviation of inclusion criteria from strict, commonly accepted standards. For example, two non-RCTs included patients with hoarding disorder.51,52 Hoarding is a ‘phenotype’ that has lacked diagnostic standards. For example, two non-RCTs included patients with hoarding disorder.51,52 Hoarding is a ‘phenotype’ that has lacked diagnostic standards. For example, two non-RCTs included patients with hoarding disorder.51,52 Hoarding is a ‘phenotype’ that has lacked diagnostic standards. For example, two non-RCTs included patients with hoarding disorder.51,52 Hoarding is a ‘phenotype’ that has lacked diagnostic standards. For example, two non-RCTs included patients with hoarding disorder.51,52 Hoarding is a ‘phenotype’ that has lacked diagnostic standards. For example, two non-RCTs included patients with hoarding disorder.51,52 Hoarding is a ‘phenotype’ that has lacked diagnostic standards. For example, two non-RCTs included patients with hoarding disorder.51,52 Hoarding is a ‘phenotype’ that has lacked diagnostic standards. For example, two non-RCTs included patients with hoarding disorder.51,52 Hoarding is a ‘phenotype’ that has lacked diagnostic standards. For example, two non-RCTs included patients with hoarding disorder.51,52 Hoarding is a ‘phenotype’ that has lacked diagnostic standards. For example, two non-RCTs included patients with hoarding disorder.51,52 Hoarding is a ‘phenotype’ that has lacked diagnostic standards. For example, two non-RCTs included patients with hoarding disorder.51,52 Hoarding is a ‘phenotype’ that has lacked diagnostic standards. For example, two non-RCTs included patients with hoarding disorder.51,52 Hoarding is a ‘phenotype’ that has lacked diagnostic standards. For example, two non-RCTs included patients with hoarding disorder.51,52 Hoarding is a ‘phenotype’ that has lacked diagnostic standards. For example, two non-RCTs included patients with hoarding disorder.51,52 Hoarding is a ‘phenotype’ that has lacked diagnostic standards. For example, two non-RCTs included patients with hoarding disorder.51,52 Hoarding is a ‘phenotype’ that has lacked diagnostic standards. For example, two non-RCTs included patients with hoarding disorder.51,52 Hoarding is a ‘phenotype’ that has lacked diagnostic standards. For example, two non-RCTs Included patients with a host of typically excluded conditions, when designing studies with the goal of reporting TE, it would be useful to reduce or control for these sources of biases. In addition, patients in non-RCTs had more variable follow-up periods (range 6–72 months) with many patients having less than 12–18 months of follow-up. Patients undergoing DBS for OCD have been shown to derive maximal symptom improvement with prolonged follow-up and this may also confound the pooled TE between studies.53 A recent follow-up study by the pioneering group from Amsterdam recapitulates this point by reporting a highly maintained response for both OCD and mood symptoms at prolonged mean follow-up (6.8±3 years) with improvement in functioning and overall well-being.60

Small study effects is a term for the phenomenon that smaller studies sometimes show different, often more pronounced, TEs than larger ones.61 One well-known potential reason for this is publication bias in which the chances of a smaller study being published increase if that study shows a stronger effect.62 A significant limitation of prior meta-analyses of the DBS for OCD literature is that none have attempted to assess these types of biases, which are known to lead to caution in interpreting medical literature.63 Other possible causes for small study effects include outcome reporting bias and clinical heterogeneity between patients in large and small studies. Our sensitivity analyses show that small study effects do not significantly impact the aggregate TE across studies and centres. A lack of significant differences between fixed and random-effects estimates of outcome measures is another encouraging sign that small study effects do not significantly bias the reported results.63 As most prior studies of DBS for TROCD have involved smaller sample sizes, such factors merit consideration when interpreting published outcomes available in the literature.
While these results are encouraging, it is important to remember that DBS is not without its limitations. First and foremost, it requires chronic implantation of hardware and carries the associated risk of complications. Our study shows a rate of hardware-related complications of ~8%. Additionally, we report a pooled infection rate of ~4.4%, a finding which aligns closely with a recent meta-analysis showing a surgical site infection rate of 5% in DBS all-comers and 4.5% specifically in DBS for OCD.44 Furthermore, although we report a less than 1% incidence of de novo obsessions involving the DBS patient programmer or the device itself, it remains a significant barrier to the effective implementation of DBS for OCD in certain patients; one which future studies could investigate further to define predictors of such behaviour. In addition, there is currently a need for pulse generator replacement after approximately 18 months for non-rechargeable models or up to 9 years for rechargeable models.32 Finally, successful application of DBS requires a close therapeutic alliance between patient, neurosurgical and expert psychiatrist teams in centres that specialise in implantation and programming of the device.

In sum, our findings support DBS as an effective treatment for TROCD, and the average appropriately selected patient will experience OCD symptom reduction of about 50%. Two-thirds of patients will achieve at least full response to DBS therapy with sustained follow-up. Stimulation of current limbic and non-limbic targets can provide substantial relief of comorbid depressive symptoms in TROCD. The growing evidence base reporting DBS for OCD outcomes demonstrates a predominantly low RoB across studies. Future crossover RCTs should aim to consistently include washout periods between active and sham stimulation periods, while observational and open-label clinical studies should aim to minimise potential confounders of treatment response and maintain longer follow-up protocols.

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Contributors RG, RN and SAS conceived the manuscript. RG, RN and SH acquired the data. RG and RN analysed the data. RG and RN drafted the manuscript. RG, RN, AA, ES, WKG, BS and SAS critically revised the manuscript. RG and SAS approved the final version of the manuscript.

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REFERENCES


