Deep brain stimulation (DBS) of the nucleus accumbens in treatment-refractory patients with obsessive-compulsive disorder (OCD).

No registrations found.

Ethical review Positive opinion

Status Recruitment stopped

Health condition type -

Study type Interventional

Summary

ID

NL-OMON20428

Source

Nationaal Trial Register

Brief title

N/A

Health condition

Obsessive-compulsive disorder (OCD)

Sponsors and support

Primary sponsor: Implantation equipment is provided by Medtronic Europe, Tolochenaz, Switzerland.

No personal funding of research personell.

Source(s) of monetary or material Support: No public funding.

Intervention

Outcome measures

Primary outcome

1 - Deep brain stimulation (DBS) of the nucleus accumbens in treatment-refractory pa ... 22-06-2025

- 1. Change on the Y-BOCS;
- 2. Number of responders, defined as a decrease on the Y-BOCS >35%.

Secondary outcome

- 1. Hamilton Depression Rating Scale (HDRS-17);
- 2. Hamilton Anxiety Scale (HAS);
- 3. Symptom Checklist 90 (SCL-90);
- 4. Quality of life enjoyment and satisfaction questionnaire;
- 5. Sheehan Disability Scale (SDS);
- 6. Clinical Global Impre3ssion (CGI);
- 7. Y-BOCS checklist.

Study description

Background summary

Objective of the study is to test the hypothesis that bilateral DBS in the nucleus accumbens of patients with severe treatment-refractory OCD can lead to long-term improvement of OCD symptoms and functioning, without unacceptable side-effects.

The study design is a double-blind cross-over trial in which sixteen patients are to be included.

Selected patients are reviewed by an independent approval-board. After electrode implantation an optimisation period is used to test stimulation parameter settings and check for side-effects of stimulation. In the ensuing cross-over period of six weeks without and six weeks with stimulation, the order being determined by randomization, patients are followed closely on an outpatient-basis. Thereafter the study continues with stimulation on in all patients.

Ethical review boards of both hospital have approved the study. An independent safety-committee is informed of all surgeries being performed and all events encountered in the study.

Study objective

DBS in the nucleus accumbens can lead to long-term improvement of obsessive-compulsive symptoms and funtioning, without unacceptable side-effects.

2 - Deep brain stimulation (DBS) of the nucleus accumbens in treatment-refractory pa ... 22-06-2025

Study design

N/A

Intervention

Stereotactic implantation of bilateral DBS electrodes in the nucleus accumbens, placebo: no stimulation.

Contacts

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Eligibility criteria

Inclusion criteria

- 1. Primary diagnosis: OCD (300.3) according to DSM-IV criteria using the MINI Plus-interview as diagnostic instrument;
- 2. Illness duration > 5 years;
- 3. Yale-Brown Obsessive-Compulsive Scale (Y-BOCS) total > 27, measured twice at least two weeks apart;
- 4. Disabling severity with substantial funtional impairment according to the DSM-IV criterion
 - 3 Deep brain stimulation (DBS) of the nucleus accumbens in treatment-refractory pa ... 22-06-2025

C and a Global Assessment of Function (GAF) score of <45;

- 5. Age 18 65 years;
- 6. Written informed consent;
- 7. Able to fully understand the consequences of the procedure (IQ>80);
- 8. Dutch speaking and able to answer all study questions;
- 9. Capable to make his or her own choice without coercion;
- 10. Treatment refractory is defined as no or insufficient response (still fulfilling the inclusion criteria) following:
- a. Two treatments with a SSRI at maximum dose for and least 12 weeks, and
- b. One treatment with clomipramine at the maximum dose for at least 12 weeks, with assessment of clomipramine/desmethylclomipramine plasma levels to control for sufficient bioavailability, and
- c. At least one augmentation trial with an atypical antipsychotic for 8 weeks in combination with a SSRI, and
- d. At least one (cognitive) behaviour therapy trial for 16 weeks in combination with an effective drug for the treatment of OCD.

Exclusion criteria

Any of the following: unstable physical condition, Parkinson's disease, dementia, epilepsy, schizophrenia or history of psychosis, alcohol or substance abuse during last 6 months, current tic disorder, antisocial personality disorder, body dismorphic disorder, pregnancy, use of psychiatric medication other than: stable use of one SSRI or clomipramine, one benzodiazepine, one atypical antipsychotic.

Study design

Design

Study type: Interventional

Intervention model: Crossover

4 - Deep brain stimulation (DBS) of the nucleus accumbens in treatment-refractory pa ... 22-06-2025

Allocation: Randomized controlled trial

Masking: Double blinded (masking used)

Control: Placebo

Recruitment

NL

Recruitment status: Recruitment stopped

Start date (anticipated): 27-03-2006

Enrollment: 16

Type: Actual

Ethics review

Positive opinion

Date: 10-03-2006

Application type: First submission

Study registrations

Followed up by the following (possibly more current) registration

No registrations found.

Other (possibly less up-to-date) registrations in this register

No registrations found.

In other registers

Register ID
NTR-new NL

NTR-new NL565 NTR-old NTR621 Other : N/A

ISRCTN ISRCTN23255677

Study results

Summary results

N/A