Preoperative Electroencephalography-Based Machine Learning Predicts Cognitive Deterioration After Subthalamic Deep Brain Stimulation

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ABSTRACT: Background: Subthalamic deep brain stimulation (STN DBS) may relieve refractory motor complications in Parkinson’s disease (PD) patients. Despite careful screening, it remains difficult to determine severity of alpha-synucleinopathy involvement which influences the risk of postoperative complications including cognitive deterioration. Quantitative electroencephalography (qEEG) reflects cognitive dysfunction in PD and may provide biomarkers of postoperative cognitive decline.

Objective: To develop an automated machine learning model based on preoperative EEG data to predict cognitive deterioration 1 year after STN DBS.

Methods: Sixty DBS candidates were included; 42 patients had available preoperative EEGs to compute a fully automated machine learning model. Movement Disorder Society criteria classified patients as cognitively stable or deteriorated at 1-year follow-up. A total of 16,674 EEG-features were extracted per patient; a Boruta algorithm selected EEG-features to reflect representative neurophysiological signatures for each class. A random forest classifier with 10-fold cross-validation with Bayesian optimization provided class-differentiation.

Results: Twenty-five patients were classified as cognitively stable and 17 patients demonstrated cognitive decline. The model differentiated classes with a mean (SD) accuracy of 0.88 (0.05), with a positive predictive value of 91.4% (95% CI 82.9, 95.9) and negative predictive value of 85.0% (95% CI 81.9, 91.4). Predicted probabilities between classes were highly differential (hazard ratio 11.14 [95% CI 7.25, 17.12]); the risk of cognitive decline in patients with high probabilities of being prognosticated as cognitively stable (>0.5) was very limited.

Conclusions: Preoperative EEGs can predict cognitive deterioration after STN DBS with high accuracy. Cortical neurophysiological alterations may indicate future cognitive decline and can be used as biomarkers during the DBS screening. © 2021 The Authors. Movement Disorders published by Wiley Periodicals LLC on behalf of International Parkinson and Movement Disorder Society

Key Words: machine learning; quantitative EEG; Parkinson’s disease; deep brain stimulation; cognition; prediction; subthalamic nucleus

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E X P E R I M E N T A L A R T I C L E

EEG P R E D I C T S C O G N I T I V E D E C L I N E A F T E R S T N D B S

Methods

Consecutive patients who underwent STN DBS at the DBS Center of the Leiden University Medical Center (LUMC) and Haga Teaching Hospital between May 2017 and July 2019 were included, as part of the OPTIMIST trial (OPTIMizing patient selection for deep brain STimulation of the subthalamic nucleus) (Netherlands Trial Register NL6079). Sample size calculation was based on a different research question. All patients fulfilled MDS PD criteria. Written informed consent was obtained from all participants. The study was approved by the local medical ethics committee of the LUMC. Anonymized data may be shared upon request. The Standards for Reporting Diagnostic Accuracy (STARD) guidelines were followed during the writing process.

Procedures and Inclusion

All patients received standard questionnaires, a neuropsychological assessment, and routine EEG as part of the DBS screening procedure. After acceptance for STN DBS, patients were invited for participation in the study (1 patient declined during the inclusion period). Inclusion criteria included age >18 years, diagnosis of idiopathic PD according to established criteria, clinical indication for STN DBS, written informed consent, ability to comply with the study assessments, and ability to read and understand Dutch. Exclusion criteria included Hoehn & Yahr stage 5, Mattis Dementia Rating Scale (MDRS) scores <120, psychiatric contraindications for STN DBS, and general contraindications for stereotactic surgery. Surgery took place approximately 1–2 months after the DBS eligibility screening. Surgical procedures have been published elsewhere. Lead implantation was performed during an awake procedure, withdrawn from sedatives and dopaminergic medication. On average, 2–3 cannulas and microelectrodes were inserted simultaneously in a Ben-Gun array. Individual adjustments were made to avoid ventricles, blood vessels, and sulci. All procedures were performed bilaterally and during the same session. Permanent leads were usually positioned with the middle two contacts located at the site of best therapeutic effect. Follow-up visits were carried out 1 year after surgery (± 6 weeks) and included similar assessment of cognition and questionnaires. EEG was only performed preoperatively.

EEG Acquisition

EEG acquisition and preprocessing procedure has been detailed previously. EEGs were recorded with patients in a supine position, eyes closed, and in a state of relaxed wakefulness. Twenty-one Ag/AgCl EEG electrodes were positioned according to a standard 10–20 setup. Medication was continued according to individual schedules (ie, “ON”); dyskinesias were not observed. All data were re-referenced towards a source-derivation approaching the surface Laplacian derivation to amplify spatial resolution. Five consecutive artifact-free non-overlapping epochs (4096 points, 8.192 seconds, sampling rate 500 Hz) were manually

Patients with Parkinson’s disease (PD) may be eligible for subthalamic deep brain stimulation (STN DBS) in cases of debilitating motor complications refractory to oral therapy. STN DBS has similar, or even better, effects on motor function compared to oral therapy, reduces motor complications, and improves quality of life (QoL). However, deterioration in one or more cognitive domains may be observed after STN DBS: 34 of 59 studies on STN DBS reported deterioration in ≥1 cognitive domain, and 12 of 20 studies reported cognitive deterioration in ≥1 cognitive domain in controlled settings. Careful and accurate assessment of cognitive functioning is a crucial element during the DBS eligibility screening, but mild cognitive deterioration can occur even after rigorous patient selection. It remains difficult to determine the severity of the impact of alpha-synucleinopathy on the central nervous system, which may in turn influence the long-term impact of DBS in terms of postoperative complications, including worsening of specific cognitive functions. Biomarkers based on quantitative electroencephalography (qEEG) have been previously described to reflect cognitive functioning in both the general PD population and DBS candidates specifically. The utility of qEEG in predicting future cognitive decline has been previously described in the general PD population in studies with 3–5 years’ follow-up. Given that cognitive deterioration may already be manifest within 1 year post-surgery, we hypothesize that qEEG may have potential utility during the DBS screening to prognosticate cognitive decline after DBS.

In previous studies we demonstrated the utility of a fully automated machine learning algorithm to classify patients according to their cognitive status through EEGs. These models apply a series of automatic processes to sequentially extract a large number of EEG-features, select EEG-features to create a comprehensive EEG-signature of either class, and subsequently build and optimize a classification model without requiring arbitrary decisions by the researcher. This process is considered highly efficient and reduces the need for prior knowledge of existing EEG-features, thereby increasing the likelihood of identifying previously unknown biomarkers of cognitive decline.

The aim of this study was to develop an EEG-based fully-automated machine learning model to evaluate the utility of EEG during the DBS screening to prognosticate cognitive deterioration according to Movement Disorder Society (MDS) criteria 1 year after STN DBS.
selected for offline analysis; patients with fewer artifact-free epochs were excluded. Brainwave software was used to calculate global peak frequencies and spectral slowing ratios (relative $[\frac{8 + 0}{|a + \beta|}]$ band-power) (BrainWave version 0.9.152.12.26, C. J. Stam; available at https://home.kpn.nl/stam7883/brainwave.html).

### Used Scales and Determination of Outcome

From the neuropsychological assessment, six tests were selected to reflect cognitive functioning: Rey Auditory Verbal Learning Test (RAVLT),\textsuperscript{21} Verbal Fluency,\textsuperscript{22} Trail Making Test (TMT) B corrected for A\textsuperscript{23} Stroop Color and Word Test (Stroop) section 3 - interference,\textsuperscript{24} and Digit Cancellation Test (DCT) sections “correct” and “wrong + corrected”.\textsuperscript{25} All six scores provided a Test-score (T-score) with normally distributed population means of 50 and standard deviations (SD) of 10. A global cognitive composite score (Cog-score) was calculated by averaging scores from all six tests; a maximum of two missing subscores was allowed. Cognitive deterioration was based on MDS level I criteria, that is, presence of significant cognitive decline in $\geq 2$ tests (ie, $\geq 10$ points deterioration).\textsuperscript{16}

Secondary outcomes included motor function (Movement Disorder Society-Unified Parkinson’s Disease Rating Scale [MDS-UPDRS] III),\textsuperscript{26} motor fluctuations (MDS-UPDRS IV),\textsuperscript{26} Levodopa Equivalent Dose (LED),\textsuperscript{27} DBS impairment scale (DBS IS),\textsuperscript{28} (SENS-PD),\textsuperscript{29} Parkinson’s Disease Quality of Life scale 39 (PDQ39),\textsuperscript{30} MDRS,\textsuperscript{31} Montreal Cognitive Assessment (MoCA),\textsuperscript{32} psychiatric symptoms (depression: Becks Depression Inventory [BDI]),\textsuperscript{33} Parkinson Anxiety Scale (PAS),\textsuperscript{34} Apathy Evaluation Scale (AES),\textsuperscript{35} impulse-control disorder (Quip-RS and Quip-RS-ICD),\textsuperscript{36} autonomic symptoms (SCOPA-AUT),\textsuperscript{37} nighttime sleeping problems (SCOPA-SLEEP),\textsuperscript{37} and excessive daytime sleepiness (EDS).\textsuperscript{38}

Motor outcomes after STN DBS were assessed using a Stimulation Challenge Test (SCT) as published previously,\textsuperscript{39} in the conditions “Med-OFF-Stim-ON”, “Med-OFF-Stim-OFF”, and “Med-ON-Stim-ON”. Baseline levodopa-response was determined through a Levodopa Challenge Test (LCT) using a 120% suprathreshold dosage of the early morning LED.\textsuperscript{40}

### Machine Learning Classification Algorithm

The machine learning algorithm has been previously reported in relation to EEG research.\textsuperscript{14,15} The machine learning pipeline consists of four fully automated phases: feature-extraction, feature-selection, classifier-training, and hyperparameter-optimization. EEG-features are derived from the “Time Series FeatureExtraction on basis of Scalable Hypothesis tests” (tsfresh) library,\textsuperscript{41,42} resulting in 16,674 features per patient (794 features per EEG-channel). Features from all five epochs were averaged per patient. A Boruta selection algorithm was used for feature-selection, which tests the variable importance (VIMP) against fictive features created by random shuffling of the extracted features. Multiple independent trials test whether the VIMP of the real features are consistently higher than the VIMP of the fictive features during a random forest classifier (RFC).\textsuperscript{43} After saturation of the feature-selection (ie, only “real features” are retained), a RFC was trained using the resulting feature set; a majority vote from all individual decision trees was used for binary classification.\textsuperscript{44} Hyperparameter-optimization was performed using a Bayesian optimization method (Mixed Integer Parallel Efficient Global Optimization).\textsuperscript{45,46} The robustness of the resulting model was validated using 10-fold cross-validation; the resulting T-scores were averaged to obtain cross-validated metrics. Split-sample validation was considered, but demonstrated to be inferior to our cross-validation approach.\textsuperscript{14} Each model was developed five times sequentially to evaluate its robustness. All models were built, tested, and validated without manual interference in relation to feature-extraction or feature-selection.

### Statistical Analyses

Differences between baseline and follow-up were compared using paired $t$ tests or Wilcoxon signed-rank tests and Pearson $\chi^2$ tests, if appropriate. $P$ values were reported using Monte Carlo estimation in the case of Wilcoxon signed-rank tests, and Fisher’s exact test in cases of dichotomous $\chi^2$ tests. Continuous demographic and clinical data, including change indices (ie, follow-up – baseline values), was compared among the cognitive groups in a similar fashion.

The machine learning classification algorithm was evaluated using sensitivity, specificity, accuracy, and F1 scores; receiver–operator characteristic (ROC) curves were used for visualization. Kaplan–Meier curves were used to demonstrate the differentiation in predicted probabilities between the groups, in which the predicted probability of being classified as cognitively stable was used as “time” and the label as “event”. The y-axis is the inverse likelihood of being classified (100% - % classified), separate lines show at what levels of the predicted probability of being classified as cognitively stable patients were indeed classified (ie, green: stable cognition; red: cognitive deterioration). The assumption of proportional hazards was confirmed through visual inspection of the log minus log plots. Curves were compared using Cox regression analyses.

All analyses were performed using IBM Statistical Package for the Social Sciences (SPSS) 25 Software and Python software.
Results

Seventy-three patients were originally included in this study. Two patients had missing baseline Cog-scores, and 11 patients were lost-to-follow-up (reasons for lost-to-follow-up were not systematically documented): 60 patients were included for analyses, of which 42 had usable preoperative EEG-recordings; 18 patients had EEGs with insufficient artifact-free epochs (see Fig. S1 for the STARD flowchart). Mean age at baseline was 60.3 (8.3) years, with 10.0 (5.7) years disease duration, 33% were female (n = 20) (see Table 1 for a full overview). Mean duration between screening and follow-up was 14.5 (2.0) months. Cognitive performance was significantly reduced after STN DBS, mean Cog-score reduction /C0 3.6 (4.3) points (FU-baseline) (P < 0.001). Patients without available EEG-recordings did not differ significantly from patients with EEG-recordings for any demographic or clinical characteristic.

Based on a cut-off of significant deterioration of ≥2 tests, 25/60 patients were classified as having clinically established

| TABLE 1  | Cohort description |
|-----------|--------------------|
| Clinical characteristic | Baseline | Follow-up | P value |
| Age at baseline, y | 60.0 (55.0, 67.8) | | |
| % female sex (n) | 33 (20) | | |
| Disease duration, y | 8.8 (6.6, 11.7) | | |
| % opt for DBS again (n) | 81 (46) | | |
| MDS-UPDRS III (Med-OFF [Stim-OFF]) | 43.3 (11.1) | 50.6 (13.6) | 0.001 |
| MDS-UPDRS III (Med-ON [Stim-ON]) | 19.0 (8.4) | 17.2 (9.9) | 0.140 |
| Therapy-responsiveness\[^b\,\,c\] | 56.4 (15.1) | 66.0 (15.2) | <0.001 |
| MDS-UPDRS IV\[^a\] | 9.0 (6.3, 11.0) | 1.0 (0.0, 5.0) | <0.001 |
| LEDD\[^b\] | 1129 (482) | 435 (273) | <0.001 |
| LEDD-DA\[^a\] | 80 (0, 248) | 30 (0, 120) | 0.036 |
| Cog-score | 49.2 (5.4) | 45.5 (6.2) | <0.001 |
| MDRS\[^a\] | 140.0 (137.0, 142.0) | 140.0 (138.0, 142.8) | 0.658 |
| MoCA\[^b\] | 26.4 (2.1) | 25.7 (2.5) | 0.025 |
| BDI\[^b\] | 12.2 (7.0) | 10.1 (6.3) | 0.005 |
| PAS\[^a\] | 10.0 (6.0, 18.0) | 8.0 (3.0, 14.0) | 0.017 |
| Apathy Scale\[^b\] | 10.5 (3.0) | 12.0 (3.8) | 0.014 |
| Quip-RS\[^a\] | 8.0 (2.0, 18.8) | 4.0 (0.0, 13.0) | 0.007 |
| SENS-PD\[^a\] | 11.0 (7.3, 14.0) | 11.0 (6.8, 15.0) | 0.660 |
| DBS IS\[^b\] | 17.3 (9.9) | 18.6 (12.8) | 0.568 |
| Scopa-AUT\[^b\] | 16.2 (7.1) | 13.9 (7.2) | 0.004 |
| Scopa-Sleep\[^b\] | 6.0 (3.0) | 3.7 (3.4) | <0.001 |
| Scopa-EDS\[^b\] | 4.2 (3.3) | 3.4 (2.7) | 0.021 |
| PDQ39\[^b\] | 47.4 (21.8) | 36.1 (23.4) | <0.001 |
| EQ5D\[^a\] | 11.0 (9.0, 12.0) | 8.0 (7.0, 10.0) | <0.001 |

\[^a\]Median (interquartile range), Wilcoxon signed-rank test.
\[^b\]Mean (standard deviation), paired t tests.
\[^c\]Therapy-responsiveness: percentage (%) response to dopaminergic therapy (baseline) or oral dopaminergic therapy + stimulation (follow-up) ([OFF–ON]/[OFF]).

Abbreviations: DBS, deep brain stimulation; MDS-UPDRS, Movement Disorder Society Unified Parkinson’s Disease Rating Scale; LEDD, Levodopa Equivalent Dose (DA, dopamine agonists); Cog-score, global composite cognitive scores; MDRS, Mattis Dementia Rating Scale; MoCA, Montreal Cognitive Assessment; BDI, Becks Depression Inventory; PAS, Parkinson’s Disease Anxiety Scale; SENS-PD, Severity of Predominantly Non-Dopaminergic Symptoms in Parkinson’s Disease; DBS IS, Deep Brain Stimulation Impairment Scale; Scopa, Scales for Outcomes of Parkinson’s Disease; AUT, autonomic symptoms; EDS, Excessive Daytime Sleepiness; PDQ39, Parkinson’s Disease Questionnaire 39; EQ5D, EuroQol 5 Dimensions.
| Clinical characteristic | Stable cognition | Cognitive deterioration | P value |
|-------------------------|------------------|-------------------------|---------|
| Age at baseline, ya*    | 58.5 (53.5, 68.0) | 62.0 (55.5, 67.0)      | 0.470   |
| % female sex (n)        | 41 (14)          | 20 (5)                  | 0.100   |
| Disease duration, ya*   | 8.1 (6.3, 11.4)  | 9.6 (6.6, 12.3)        | 0.342   |
| % opt for DBS again (n) | 100 (30)         | 67 (16)                | 0.030   |
| MDS-UPDRS III OFFb      | Baseline         | 41.5 (11.9)            | 0.155   |
|                         | Follow-up        | 47.6 (13.1)            | 0.068   |
|                         | Change           | 5.8 (14.9)             | 0.535   |
| MDS-UPDRS III ONb       | Baseline         | 18.4 (8.7)             | 0.565   |
|                         | Follow-up        | 15.6 (9.5)             | 0.215   |
|                         | Change           | −3.2 (9.0)             | 0.456   |
| Therapy-responsivenessb,d,e | Baseline     | 56.1 (16.0)           | 0.816   |
|                         | Follow-up        | 66.9 (14.8)           | 0.645   |
|                         | Change           | 0.1 (0.2)              | 0.611   |
| MDS-UPDRS IVa           | Baseline         | 9.0 (5.0, 12.0)        | 0.787   |
|                         | Follow-up        | 2.0 (0.0, 5.0)         | 0.853   |
|                         | Change           | −5.1 (6.6)             | 0.503   |
| LEDDd                   | Baseline         | 1009 (411)             | 0.039   |
|                         | Follow-up        | 391 (266)              | 0.140   |
|                         | Change           | −618 (437)             | 0.136   |
| LEDD-DAa                | Baseline         | 80 (0, 240)            | 0.789   |
|                         | Follow-up        | 58 (0, 120)            | 0.327   |
|                         | Change           | 22 (305)               | 0.347   |
| Cog-scored              | Baseline         | 49.3 (5.7)             | 0.994   |
|                         | Follow-up        | 47.3 (6.0)             | 0.008   |
|                         | Change           | −2.0 (3.9)             | <0.001  |
| RAVLTd                  | Baseline         | 41.9 (10.5)            | 0.381   |
|                         | Follow-up        | 41.8 (10.7)            | 0.189   |
|                         | Change           | −0.1 (8.9)             | 0.013   |
| Verbal Fluencyd         | Baseline         | 59.3 (13.5)            | 0.703   |
|                         | Follow-up        | 49.2 (11.3)            | 0.011   |
|                         | Change           | −10.1 (10.5)           | 0.005   |
| TMT B corrected for A4d | Baseline         | 46.0 (12.6)            | 0.169   |
|                         | Follow-up        | 45.1 (11.6)            | 0.708   |
|                         | Change           | −0.9 (14.8)            | 0.349   |
| Stroop 3 – interferenced| Baseline         | 52.1 (7.6)             | 0.676   |
|                         | Follow-up        | 53.2 (8.0)             | 0.040   |
|                         | Change           | 1.1 (5.6)              | 0.016   |
| DCT – total correctd    | Baseline         | 45.7 (9.4)             | 0.342   |

(Continues)
| Clinical characteristic | Stable cognition | Cognitive deterioration | \( P \) value |
|-------------------------|-----------------|-------------------------|--------------|
| **Follow-up**           | 44.1 (10.8)     | 37.7 (9.1)              | 0.021        |
| **Change**              | −0.9 (6.8)      | −4.4 (6.2)              | 0.058        |
| **DCT – total wrong and corrected** |                   |                         |              |
| **Baseline**            | 50.9 (9.2)      | 52.6 (8.4)              | 0.484        |
| **Follow-up**           | 50.6 (10.4)     | 49.4 (9.1)              | 0.648        |
| **Change**              | −0.3 (8.6)      | −3.2 (8.1)              | 0.218        |
| **MDRS**                |                 |                         |              |
| **Baseline**            | 140.0 (138.5, 142.5) | 139.0 (136, 141.8) | 0.294        |
| **Follow-up**           | 140.4 (138.0, 143.0) | 140.0 (136, 142.0) | 0.548        |
| **Change**              | −0.2 (3.0)      | 0.7 (3.7)               | 0.323        |
| **MoCA**                |                 |                         |              |
| **Baseline**            | 26.6 (2.3)      | 26.2 (1.7)              | 0.442        |
| **Follow-up**           | 25.9 (2.7)      | 25.6 (2.3)              | 0.655        |
| **Change**              | −0.7 (2.3)      | −0.6 (2.1)              | 0.762        |
| **BDI**                 |                 |                         |              |
| **Baseline**            | 12.8 (7.2)      | 11.4 (6.9)              | 0.452        |
| **Follow-up**           | 9.7 (5.5)       | 10.2 (7.0)              | 0.730        |
| **Change**              | −3.1 (6.3)      | −1.1 (4.2)              | 0.181        |
| **PAS**                 |                 |                         |              |
| **Baseline**            | 11.0 (6.0, 18.0) | 10.0 (8.0, 17.5)        | 0.730        |
| **Follow-up**           | 5.5 (3.0, 15.5) | 10.0 (4.5, 14.0)        | 0.549        |
| **Change**              | −2.6 (7.7)      | −1.3 (6.1)              | 0.510        |
| **Apathy Scale**        |                 |                         |              |
| **Baseline**            | 10.4 (3.4)      | 10.4 (2.4)              | 0.948        |
| **Follow-up**           | 12.2 (3.4)      | 11.9 (4.4)              | 0.806        |
| **Change**              | 2.0 (3.7)       | 1.5 (3.5)               | 0.710        |
| **Quip-RS**             |                 |                         |              |
| **Baseline**            | 9.0 (1.5, 18.8) | 7.0 (2.0, 23.0)         | 0.849        |
| **Follow-up**           | 4.5 (0.0, 12.8) | 2.5 (0.0, 13.3)         | 0.614        |
| **Change**              | −3.8 (8.0)      | −4.2 (17.6)             | 0.922        |
| **SENS-PD**             |                 |                         |              |
| **Baseline**            | 9.0 (6.8, 13.0) | 12.0 (8.5, 15.0)        | 0.162        |
| **Follow-up**           | 11.0 (6.0, 15.0) | 11.0 (7.3, 15.0)        | 0.666        |
| **Change**              | 0.1 (4.2)       | −0.4 (3.7)              | 0.665        |
| **DBS IS**              |                 |                         |              |
| **Baseline**            | 18.3 (10.2)     | 15.8 (9.5)              | 0.350        |
| **Follow-up**           | 18.3 (11.9)     | 18.8 (14.3)             | 0.875        |
| **Change**              | −0.8 (10.6)     | 3.0 (13.0)              | 0.224        |
| **Scopa-AUT**           |                 |                         |              |
| **Baseline**            | 14.6 (6.6)      | 17.9 (7.2)              | 0.077        |
| **Follow-up**           | 13.3 (7.0)      | 14.6 (7.7)              | 0.505        |
| **Change**              | −1.3 (5.7)      | −3.2 (5.9)              | 0.199        |
| **Scopa-Sleep**         |                 |                         |              |
| **Baseline**            | 6.0 (2.9)       | 5.9 (2.9)               | 0.906        |
| **Follow-up**           | 3.7 (2.9)       | 3.3 (3.2)               | 0.598        |
| **Change**              | −2.3 (3.2)      | −2.6 (4.2)              | 0.729        |
| **Scopa-EDS**           |                 |                         |              |
| **Baseline**            | 4.3 (3.4)       | 3.9 (3.0)               | 0.639        |
| **Follow-up**           | 3.7 (2.7)       | 2.8 (2.2)               | 0.223        |
| **Change**              | −0.7 (2.8)      | −1.1 (2.4)              | 0.563        |
cognitive decline (of which 17 had EEG-recordings), and 34 patients were classiﬁed as cognitively stable (of which 25 had EEG-recordings) (1 patient was missing due to missing data on two follow-up cognitive tests, such that the criteria for cognitive decline could not be assessed).

Clinical Differences Between Cognitive Classes

Both cognitive classes were not signiﬁcantly different in any baseline characteristic, including baseline individual cognitive T-scores (see Table 2). Of note, mean baseline Cog-score was 49.3 points in both groups, relatively similar to the population average of 50.0.

At follow-up, groups differed in terms of Cog-score (47.3 (6.0) points versus 43.0 (5.7) points, \(P = 0.008\); patients classiﬁed as having cognitive decline at 1 year follow-up had similarly signiﬁcantly lower T-scores for the Verbal Fluency, Stroop 3, and DCT – total correct tests. Patients with stable cognition were unanimous in choosing DBS again if given the opportunity, however only two-thirds of patients with cognitive deterioration would opt for DBS again (\(P = 0.030\)), despite no signiﬁcant difference in QoL. Apart from cognitive performance, the groups did not differ at 1 year follow-up or in change index in any other domain. Both groups did not differ in spectral EEG band-powers (see Table S1).

Machine Learning Performance

Eighteen EEG-features were selected by the machine learning pipeline during each of the ﬁve cross-validation runs. None of the features were retained in each of the individual runs; one feature was selected four times, two features were selected three times, and six features were selected twice (see Fig. S2 for an overview of each of the relative feature-importances; the cerebral location of each feature is shown in Figure S3; see Table S2 for a brief explanation of the mathematical background).

The machine learning algorithm differentiated patients with stable cognition at 1 year follow-up from patients with cognitive deterioration with a mean (SD) accuracy of 0.88 (0.05) in the test-set (see Table 3), with a positive predictive value (PPV) (ie, likelihood of stable cognition 1 year after surgery) of

| TABLE 2   |         | Stable cognition | Cognitive deterioration | \(P\) value |
|-----------|---------|------------------|-------------------------|-------------|
| Clinical characteristic |     |                  |                         |             |
| PDQ39\(^a\) | Baseline | 47.4 (24.1)      | 46.9 (18.9)             | 0.931       |
|           | Follow-up | 33.1 (20.4)      | 38.1 (25.4)             | 0.400       |
|           | Change\(^c\) | −14.3 (22.2)     | −8.8 (21.4)             | 0.339       |
| EQ5D\(^a\) | Baseline | 10.5 (9.0, 12.0) | 11.0 (9.0, 12.0)        | 0.852       |
|           | Follow-up | 8.0 (7.0, 10.0)  | 8.0 (7.0, 10.5)         | 0.913       |
|           | Change\(^c\) | −1.7 (2.6)        | −1.5 (2.2)              | 0.739       |

\(^a\)Median (interquartile range), Mann–Whitney U tests.

\(^b\)Follow-up: MDS-UPDRS III Med-OFF-Stim-OFF or Med-ON-Stim-ON.

\(^c\)Follow-up – baseline (mean [standard deviation], Student t tests).

\(^d\)Therapy-responsiveness: percentage (%) response to dopaminergic therapy (baseline) or oral dopaminergic therapy + stimulation (follow-up) ([OFF–ON]/[OFF]).

Abbreviations: DBS, deep brain stimulation; MDS-UPDRS, Movement Disorder Society Uniﬁed Parkinson’s Disease Rating Scale; LEDD, Levodopa Equivalent Dose (DA, dopamine agonists); Cog-score, global composite cognitive scores; RAVLT, Rey Auditory Verbal Learning Test; TMT, Trail Making Test; Stroop 3, Stroop Color and Word Test section 3; DCT, Digit Cancellation Test; MDRS, Mattis Dementia Rating Scale; MoCA, Montreal Cognitive Assessment; BDI, Becks Depression Inventory; PAS, Parkinson’s Disease Anxiety Scale; SENS-PD, Severity of Predominantly Non-Dopaminergic Symptoms in Parkinson’s Disease; DBS IS, Deep Brain Stimulation Impairment Scale; Scopa, Scales for Outcomes of Parkinson’s Disease; AUT, autonomic symptoms; EDS, Excessive Daytime Sleepiness; PDQ39, Parkinson’s Disease Questionnaire 39; EQ5D, EuroQol 5 Dimensions.

| TABLE 3   |         |       |       |     |
|-----------|---------|-------|-------|-----|
| Dataset   | Accuracy | Sensitivity | Specificity | F1  |
| Training-set\(^a\) | 0.97 (0.02) | 0.99 (0.01) | 0.93 (0.05) | 0.97 (0.02) |
| Validation results\(^a\) | 0.88 (0.05) | 0.95 (0.04) | 0.76 (0.07) | 0.84 (0.06) |

\(^a\)Mean (standard deviation); all metrics were averaged over ﬁve separate computation runs.

FIG. 1. Receiver–operator characteristic (ROC) curve of all individual cross-validation runs combined. Individual ROC curves of the cross-validation runs separately are shown in Figure S2.
For high predicted probabilities of being classified as cognitively stable (ie, >0.5), the likelihood of cognitive deterioration appears slim. Kaplan–Meier curves for all cross-validation runs separately are shown in Figure S3. Green: cognitively stable patients; red: patients with cognitive deterioration; dotted line: 95% CI. [Color figure can be viewed at wileonlinelibrary.com]

91.4% (95% CI 82.9, 95.9) and a negative predictive value (NPV) (ie, likelihood of cognitive deterioration) of 85.0% (95% CI 81.9%, 91.4%). Performance metrics of all five individual model-runs, as well as overall, are shown in Figure S4. A ROC curve based on the predicted probabilities of all the individual model-runs combined is shown in Fig. 1; ROC curves for the individual runs are shown in Figure S5; the robustness of the cross-validation runs indicates no large effect of overfitting. A Kaplan–Meier curve (Fig. 2) using the pertaining class as “event” and the predicted probability of being classified as cognitively stable as “time” shows the differentiation between the predicted probabilities of the two classes, combined over all cross-validation runs, with a hazard ratio (HR) 11.14 (95% CI 7.25, 17.12) (individual Kaplan–Meier curves for each of the individual cross-validation runs are shown in Fig. S6). For high predicted probabilities of being classified as cognitively stable (ie, >0.5), the likelihood of cognitive deterioration approached zero.

Discussion

The aim of this study was to develop a fully automated machine learning model based on preoperative EEGs to predict cognitive deterioration 1 year after STN DBS. Our model differentiates patients with stable cognition with 88% accuracy from patients with cognitive decline according to MDS criteria, with a PPV of 91.4% and a NPV of 85.0%. The robustness of this approach is demonstrated by relatively stable performance-metrics for each of the individual cross-validation runs, and the stable difference in predicted class-probabilities between the groups. Survival analyses demonstrate that the cognitive classes had highly differential predicted probabilities; the chance of suffering from cognitive decline 1 year after STN DBS was particularly small in cases of probabilities >0.5.

To our knowledge, no previous study has investigated the actual model-performance of EEG to predict cognitive decline after STN DBS. Although the exact retest reliability of the individual tests used to compose the Cog-score in patients both with and without neuromodulation is unknown, previous literature reported cognitive test performance as relatively stable in PD patients without DBS. A previous paper investigated the potential of qEEG to predict cognitive decline after STN DBS in 17 patients by evaluating the predictive potential of relative 0 power in predicting cognitive decline. Although results indicated the utility of qEEG to predict postoperative decline, only one spectral metric was studied and correlated with only 3/13 cognitive tests, whereas its predictive potential was not discussed. Another paper described Grand Total EEG (GTE) scores to predict cognitive decline after STN DBS in 30 patients, and reported an odds ratio of 2.78 (95% CI 1.25–6.18) in predicting cognitive decline, although the minority class (cognitive decline based on Mini-Mental State Examination [MMSE]/DemTect scores) consisted of only six patients and model performance was not reported. Moreover, GTE scores are semiquantitative in nature and still require expert knowledge of EEGs, limiting generalizability compared to automated machine learning.

Eighteen features were retained in total during cross-validation runs, none of which were selected in all five models. This suggests that there is not one representative neurophysiological signature of cognitive deterioration, but that several patterns of preoperative cognitive (dys)function exist with roughly equal potential to reflect postoperative cognitive decline. Eleven of 18 features were Fast Fourier Transformation (FFT)-based, although baseline spectral features that are more routinely studied in PD did not differ between the two groups. We hypothesize that our more in-depth spectral features are more suitable as novel biomarkers reflecting neurophysiological signaling speed than conventional band-powers. Three other metrics were related to the mass quantile (ie, calculation of a relative index in which q% of the mass of the time series is left of the index). To our knowledge, quantile functions have not been studied previously in relation to qEEG in PD, or in cognitive research, and represent a novel mathematical approach in this setting. The retained EEG-features for modeling cognitive state in a cross-sectional model (using the same methodology and feature-library) are different from those retained for modeling future cognitive deterioration, however both congregate by prioritizing FFT-features. We speculate that different FFT-aspects underlie either current
cognitive functioning or future cognitive dysfunction. Conventional EEG band-powers are probably too inter-related to optimally reflect this distinction. It should be noted that our model pertains to cognitive decline after STN DBS specifically and has not been validated for use in prognosticating cognition without DBS. Future research should evaluate its utility in non-surgical cohorts as well, and investigate cognitive functioning after a longer follow-up. However, including a non-surgical cohort would have introduced substantial bias as ineligibility for surgery is often based on poor cognitive functioning, which would have set these patients apart from the groups characterized here.

A limitation of many machine learning models is the “black-box-problem”, in which it is unclear how the model reaches a decision or what these decisions are based on. As we minimized arbitrary choices in our model, the resulting EEG-features can be considered novel biomarkers in relation to EEG research in PD, and results can be traced back to pathophysiological EEG alterations (eg, EEG-slowing, mass quantile alterations, etc.). However, we realize that the exact mathematical background of the resulting features may be relatively unclear and, as a consequence, their pathophysiological substrate. In terms of etiology, we cannot readily explain why some features were selected and others were not; however, in terms of implications for clinical practice the prognostic performance of the selected features currently supersedes their content. Despite our relatively small sample size, we have demonstrated good accuracy in differentiating cognitively stable patients from those with cognitive decline. We demonstrate that overfitting did not play a major role in our models despite an approximate drop in accuracy of 9% relative to the training-set, and differentiation of the groups based on predicted probability was stable.

Our results further demonstrate that a standard 21-channel EEG probably suffices to predict cognitive decline after STN DBS. Further insight into pathophysiological alterations underlyiing cognitive decline may be provided through source localization to more closely pinpoint anatomical substrates, especially when combined with high-density EEG setups or pre–post EEG settings. However, these setups are less applicable in a clinical prediction setting which would have limited its current utility.

In this study, cognitive deterioration was classified based on MDS criteria for cognitive decline, which increases the generalizability of our results. MDS level I criteria for cognitive decline (ie, ≥2 tests showing significant deterioration) are less strict than those for PD dementia (PDD), which is the likely reason for our observed high incidence of cognitive decline. Similar to previous studies, the observed cognitive decline is still within the range of normal cognitive functioning. Our classification therefore reflects “deterioration” rather than actual clinical impairment. Although class-transition to mild cognitive impairment or PDD would have been clinically more relevant, this transition within 1 year is most likely rare after DBS and would require larger sample sizes than currently available, as well as introducing a significant class imbalance. Whether the observed cognitive deterioration is a result of disease progression, a differential impact of stimulation, or a postoperative hypodopaminergic state cannot be determined. However, since the groups did not differ in any baseline characteristic, our results indicate the prognostic utility of EEG regardless of the mechanism behind cognitive deterioration.

Strikingly, generic tests such as the MDRS and MoCA did not show the same deterioration as the Cog-score, even in patients classified as cognitively deteriorated. This is in line with previous literature, as 27/29 studies reported no significant cognitive decline in these global tests after STN DBS. Both generic tests are subject to confounding such as educational level. We hypothesize that T-score-based cognitive tests resulted in more sensitive metrics to reflect cognitive functioning. Our observed cognitive deterioration was predominantly fueled by deterioration in the domains “executive functioning” and “language” in line with previous research. As there are multiple possible cognitive tests to quantify cognition in PD, the ideal classification would be based on ≥2 tests per cognitive domain according to Diagnostic and Statistical Manual of Mental Disorders (DSM-5) and MDS level II criteria to provide a more complete overview of cognitive functioning.

Our results imply that EEG may be used to prognosticate cognitive decline after STN DBS in a clinical setting. A risk of (mild) cognitive decline does not necessarily equal ineligibility for surgery, but improved prognostication may contribute to better patient education and informed consent. Moreover, a risk of postoperative decline as indicated by our machine learning algorithm could skew clinicians towards targets with possible less impact on cognition such as pallidal DBS, although our model has not been tested for prognostication in these settings. Strikingly, patients classified as cognitively stable were unanimous in choosing DBS again if given the opportunity, whereas this was significantly lower in patients with cognitive deterioration despite no differences in any other characteristic. This underscores the importance of cognitive function for patient satisfaction after DBS. Our results indicate that EEG-based prognostication could be included in the DBS screening to improve shared-decision-making.

Strengths of this study include using a fully automated machine learning algorithm which enhances the generalizability and potential clinical utility of our results. Based on the nature of the Boruta selection algorithm which uses multiple independent trials during feature-selection, we are confident that the resulting
model provides an accurate and robust electrophysiological signature underlying the cognitive classes, despite a relatively small sample size. A limitation of our study is a possibly selective loss-to-follow-up as several patients that declined the follow-up assessments reported that cognitive assessments would be burdensome for them at this stage, which may result in an underrepresentation of cognitively impaired patients in the entire cohort, although this would not affect the prediction analyses. Although patients without EEG-recordings did not differ clinically from patients with available recordings, the utility of EEG may be limited due to unavailability of artifact-free epochs. The amount of missing data due to EEG-artifacts is similar in previous literature.\textsuperscript{10} Given our lack of external validation, our results are best interpreted as proof-of-concept that machine learning algorithms can accurately predict cognitive deterioration after STN DBS.

Our results indicate that cortical neurophysiological alterations prior to STN DBS may reflect a risk of postoperative cognitive deterioration, and the use of algorithms such as those detailed here may be helpful during the DBS eligibility screening to directly improve clinical practice. Automated analyses such as these have the ability to identify new biomarkers. Further exploration of the features selected by our model may ultimately provide greater insight into pathophysiological processes and the mechanism behind the association between EEG and cognitive decline.

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**Supporting Data**

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