Testing the therapeutic effects of transcranial direct current stimulation (tDCS) in semantic dementia: A double blind, sham controlled, randomized clinical trial

CURRENT STATUS: ACCEPTED

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Abstract

Background

Semantic dementia is a neurodegenerative disease that primarily affects the left anterior temporal lobe, resulting in a gradual loss of conceptual knowledge. There is currently no validated treatment. Transcranial stimulation has provided evidence for long-lasting language effects presumably linked to stimulation-induced neuroplasticity in post-stroke aphasia. However, studies evaluating its effects in neurodegenerative diseases as semantic dementia are still rare and evidence from double blind prospective therapeutic trials is required.

Objective

The primary objective of the present clinical trial (STIM-SD) is to evaluate the therapeutic efficacy of a multiday transcranial direct current stimulation (tDCS) regime on language impairment in patients with semantic dementia. The study also explores the time course of potential tDCS-driven improvements and uses imaging biomarkers which could reflect stimulation-induced neuroplasticity.

Methods

Double-blind sham-controlled randomized study using tDCS applied daily during 10 days, and language/semantic and imaging assessments at 4 time-points: baseline, 3 days, 2 weeks and 4 months after the 10 stimulation sessions. Language/semantic assessments will be applied at the 4 time-points. Fluorodeoxyglucose Positron Emission tomography (FDG-PET), resting-state functional Magnetic Resonance Imaging (rs-fMRI), T1-weighted images and white matter diffusion tensor imaging (DTI) will be applied at baseline and the two-weeks’ time-point. According to the principle of inter-hemispheric inhibition between left (language-related) and right homotopic regions we will use two stimulation modalities: left-anodal and right-cathodal tDCS over the anterior temporal lobes. Accordingly, the
Patient population (n=60) will be subdivided into 3 subgroups: left-anodal tDCS (n=20), right-cathodal tDCS (n=20) and sham tDCS (n=20). The stimulation duration will be sustained for 20 minutes at an intensity of 1.59 mA. It will be delivered through 25 cm² round stimulation electrodes (current density of 0.06 mA/cm²) placed over the left and right anterior temporal lobes for anodal and cathodal stimulation, respectively. A group of age, gender and education-matched healthy participants (n=20) will also be recruited and tested to provide normative values for the language/semantic tasks and imaging measures.

Discussion
The study aims at assessing the efficacy of tDCS for language/semantic disorders in semantic dementia. A potential treatment would be easily applicable, inexpensive, and renewable when therapeutic effects disappear due to disease progression.

Introduction
Semantic dementia (SD), also referred to as the semantic variant of Primary Progressive Aphasia (sv-PPA) [1], is part of the spectrum of frontotemporal lobar degeneration and constitutes one of the major clinical variants of this disorder [2]. The onset age of SD is frequently before 65 years [3] and it severely affects the ability to communicate which generates a major impact on the family and socio-professional life of patients.

SD is characterized by a gradual and severe loss of conceptual knowledge, resulting in anomia, impaired word comprehension and speech that is fluent but empty of content [2], leaving grammar and speech articulation preserved [4]. Although the most prominent deficits concern word meaning [1,4], SD might eventually cause deterioration of knowledge for all kinds of semantic concepts [5,6] impacting on face recognition [7], object feature attribution [8], sound-picture matching [9] and object-use [10]. The damage of multi-modal semantic representations, besides the verbal domain, gave birth to the
concept of "semantic dementia" [11]. It therefore appears that sv-PPA is a purely linguistic variant of SD [12] and/or that SD results from the evolving disease course of sv-PPA [13,14].

At the anatomical level SD affects the anterior temporal lobe (ATL) in both hemispheres, showing in most of the cases a left lateralization [1,4]. Correlations have been found between gray matter loss in the ATL and different semantic tasks like picture naming [15,16] and word-picture association [17]. It is associated with disruptions of functional connectivity between a broad range of brain regions across the temporal, frontal, parietal, and occipital lobes, including visual and auditory association cortices [18]. Alterations of structural connectivity such as a white matter volume reduction in the left temporal lobe, the periventricular white matter and the corpus callosum [19], and damage to white matter tracts such as the inferior longitudinal fasciculus (ILF) and the uncinate fasciculus (UF) have also been found in SD [20]. Signs of cortical hypometabolism which are a useful neuroimaging hallmark for the diagnosis have mainly been found in the ATL cortices, extending sometimes to the subgenual region and the right anterior cingulate cortex [21]. There is currently no validated treatment for SD given that speech therapy protocols have not been validated and pharmacological trials did not demonstrate significant effects [22,23,24]. In this context, new approaches based on the use of non-invasive brain stimulation and neuro-modulation [25] might represent a promising therapeutic strategy.

Two of the most common technologies for non-invasive brain stimulation are repetitive Transcranial Magnetic Stimulation (TMS) and Transcranial Direct Current Stimulation (tDCS) [26]. Repetitive TMS involves the application of a series of magnetic pulses through a stimulating coil placed in contact with an area of the scalp, which in a frequency- and pattern-dependent manner induces an intracranial electric current that subsequently modulates (inhibits or facilitates) neuronal activity [26]. Repetitive TMS is characterized
by its excellent spatial and temporal resolution and ability to make neurons discharge, features that come at the cost of low portability, high financial cost and epileptic risk. The effects of tDCS are based on a weak electric current (1-2 mA) conveyed between two electrodes (an active and a return) placed on separate locations of the scalp, with the ability to generate a polarization gradient across a large cortical area between electrodes, hence modulating cortical excitability within its boundaries [27]. Two modalities of tDCS are commonly used: anodal stimulation (the anode is placed on the target region) increases membrane resting potential in neurons between its gradient of action, hence facilitating neural activity; cathodal stimulation (the cathode is placed on the target region), which usually results in the opposite effect, decreasing membrane potential, hence inhibiting or reducing neuronal activity [27].

For several years a growing number of studies have explored the effects of non-invasive brain stimulation in patients with aphasia following left hemisphere strokes. These studies rely on the assumption that weak electrical currents can interact with neural networks subtending language and promote neural plasticity, allowing short-term modulations, and eventually clinical recovery. Within these language-related networks, left and right homotopic regions are connected via transcallosal connections [28,29] which, according to the principle of inter-hemispheric inhibition, tend to convey mutual net inhibitory influences [30]. The use of non-invasive brain stimulation in left hemisphere stroke aphasia relies on three potential mechanisms or combinations thereof [31,32]: 1) the use of left-excitatory (anodal tDCS or high-frequency TMS) stimulation on left hemisphere language systems to reactivate language processes implemented by peri-lesional regions; 2) the delivery of right-inhibitory (cathodal tDCS or low-frequency TMS) stimulation to reduce the inhibition that right hemisphere systems exert on the left dominant language network; 3) the delivery of right-excitatory stimulation to activate potential language
contributions of right hemisphere networks. The most promising clinical outcomes on language disabilities have been achieved from studies adopting the first two approaches using either TMS [e.g., 33-36] or tDCS [e.g. 37-39]. Beneficial effects have been shown either transiently with single session applications, or as longer lasting impacts (> 6 months) following periodical stimulation sessions across several days, probably related to stimulation-induced neuroplasticity [34,40].

Non-invasive brain stimulation has also been used with relative efficacy in Alzheimer's disease by targeting the left and/or right dorsolateral prefrontal cortex [41,42]. In PPA, including SD (or sv-PPA), two small-cohort studies have suggested encouraging results with both TMS and tDCS [43,44]. However, the authors did not target language-specific brain regions and the small number of patients precluded a counterbalanced study design. In addition, left-excitatory vs. right-inhibitory TMS or tDCS have not been systematically evaluated to reveal the most efficient strategy in aphasia of degenerative origin.

The purpose of this article is to present a clinical protocol (PHRC ‘STIM-SD’) for implementing a multi-day tDCS regime in a large population of SD patients (n=60) specifically targeting the left ATL (anodal stimulation) or the right ATL (cathodal stimulation) to provide evidence for potential therapeutic effects and brain plasticity outlasting the duration of the treatment. This intervention builds on a previous pre-therapeutic double-blind, sham-controlled study by our team using a single tDCS session applied to the left and right ATL in SD patients [45]. This approach allowed to compare left anodal (excitatory) stimulation to right cathodal (inhibitory) stimulation and showed that a single session of both left anodal and right cathodal tDCS resulted in transient but highly significant intra-semantic effects. [45]. The goals of the present study include the evaluation of the potential therapeutic efficacy of repetitive tDCS over the ATL during 10 days on language/semantic performance in patients with SD, the assessment of the time
course of potential improvements, the exploration of potential effects on brain plasticity using functional connectivity measures (resting-state fMRI) and cortical metabolism (FDG-PET), the identification of the most efficient stimulation modality (left-anodal versus right-cathodal), and the identification of biomarkers such ATL atrophy levels which could be individually indicative of an efficient tDCS impact.

Methods

Study design

The STIM-SD protocol is a double-blind, sham-controlled, randomized study testing the efficiency of periodical tDCS sessions during 10 days (Monday to Friday for two weeks) using language/semantic assessments, and PET and MRI-based neuroimaging at 4 time-points: baseline, 3 days, 2 weeks and 4 months following the end of the tDCS sessions (Figure 1A and B). The patient population (n=60) is randomly assigned to three subgroups each receiving a different treatment over the ATL: left-anodal tDCS (n=20), right-cathodal tDCS (n=20) and sham tDCS (n=20).

Language/semantic assessments are applied at the 4 time-points by a set of computer-based tasks. For each task, we have developed two equivalent versions matched on various linguistic variables, which are used alternatively either baseline or during follow up evaluations, in a counterbalanced order across patient subgroups to avoid test/re-test confounds. Neuroimaging acquisitions (structural MRI, resting-state fMRI, FDG-PET) are performed at baseline and the ‘2-weeks’ time-point. We also acquire resting-state EEG recordings at baseline and after the end of tDCS sessions (‘3-days’ time point) given that SD patients might have an altered pattern of resting state neuronal synchronizations [46].

To ensure the double-blindness a first investigator performs and supervises the tDCS sessions whereas a second investigator conducts the language/semantic tasks, blinded to the stimulation condition (anodal, cathodal or sham). During sham stimulation, tDCS
current is ramped up and down along 30 seconds respectively during the initial and final phases of the session to emulate the transient skin-itching sensations characterizing active anodal or cathodal stimulation. Unnoticed by the patients, the stimulation unit is turned off during the 20 minutes of the sham tDCS session.

Stimulation sessions are applied daily for 10 days (Monday to Friday for two weeks). Each tDCS session lasts for 20 minutes. The direct current has an intensity of 1.59 mA (25 cm$^2$ round electrodes, current density of 0.06 mA/cm$^2$). A group of 20 healthy subjects are also evaluated at baseline to provide normative values for the language/semantic tasks and for the imaging measures. Healthy participants do not undergo tDCS treatment and are only assessed once with the different language/semantic tasks and neuroimaging explorations.

The primary endpoint of the STIM-SD protocol is to evaluate the potential therapeutic efficacy of multiday tDCS (10 days) on language/semantic performance in SD patients (see below the section ‘Language/semantic Tasks’ – the semantic association task). In addition, we will also: i) assess the time course of potential language/semantic improvement through the application of four follow-up time-points; ii) assess imaging biomarkers that could reflect stimulation-induced neuroplasticity and response to stimulation; iii) compare the effects of left-anodal and right-cathodal tDCS to define the most efficient stimulation modality; iv) identify biomarkers which could predict individually an efficient tDCS impact; and v) improve the understanding of the semantic roles of the left and right ATL and their potential anatomical connectivity, contributing to the definition of anatomo-functional models of semantics.

The protocol has been approved by the local Ethics Committee and is registered in ClinicalTrials.gov with the identifier NCT03481933 and the study title ‘Evaluation of a Transcranial Stimulation with Direct Current on Language Disorders in Semantic Dementia

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Written informed consent is obtained from all the patients and healthy subjects before the onset of any of the study procedures. All research visits of the protocol take place at the same site, the Pitié-Salpêtrière Hospital. Two additional centers contribute to the protocol by recruiting SD patients.

Participants

SD patients are recruited in the ‘National Reference Center for Rare or Early Onset Dementias’ at the Pitié-Salpêtrière Hospital, at the Rothschild Ophthalmologic Foundation and at the Léopold Bellan Hospital, in Paris. Patients are recruited based on the following inclusion criteria: (1) diagnosis of SD based on current research criteria [1] comprehending progressive language impairment, single-word comprehension deficits and anomia, without sentence repetition impairment, agrammatism or motor speech disorders; (2) age > 18 years old; and (3) affiliation to a social security regime.

Non-inclusion criteria are the following: (1) psychiatric disorders or neurologic diseases other than SD; (2) contraindication for MRI, PET or tDCS such as carrying intracranial ferromagnetic devices, scalp or skull lesions or epilepsy; (3) MRI images revealing pathological processes other than those associated with SD; (4) high aphasia severity (severity score < 3 in the Boston Diagnostic Aphasia Evaluation-BDAE [47]); (5) Mini Mental State Examination (MMSE) [48] score < 15; (6) Frontal Assessment Battery (FAB) [49] score < 10; (7) Montgomery Asberg Depression Rating Scale [50] (MADRS) score ≥ 20 indicating a major depressive disorder; (8) no French mother tongue; (9) non-right-handed patients; (10) curatorship or tutorship. Prior to the inclusion in the protocol all patients undergo a neuropsychological and speech therapist evaluation to check for inclusion/non-inclusion criteria, and to characterize global cognitive/language/semantic capacities on the basis of several published standard tests.
Healthy participants are recruited among hospital staff, caregivers and among patients’ relatives via announcements in the neurology department. They are matched for sex, age, handedness and number of years of education with the SD patients. Healthy participants non-inclusion criteria are the following: (1) neurological or psychiatric disorders or physical deficits that can interfere with cognitive function; (2) contraindications to MRI or PET; (3) no French mother tongue.

Randomization

A computer-generated block randomization list has been prepared by the statistician of the Clinical Research Unit of the Pitié-Salpêtrière - Charles Foix Hospital group. The randomization list is incorporated into the eCRF and a randomization number assigning patients to one of the 3 stimulation conditions (anodal, cathodal or sham tDCS) is attributed automatically upon completion of the inclusion/non-inclusion criteria. The randomization is not stratified by center of inclusion since all stimulation sessions and time-point evaluations are performed at the Pitié Salpêtrière Hospital.

Stimulation administration

The stimulation procedure (electrode montage and stimulation parameters) is the same as the procedure used in the above-mentioned pre-therapeutic study of our group [45]. During the baseline visit patients undergo an FDG-PET and MRI including anatomical 3D T1-weighted images. The T1-weighted images are registered in standardized Montreal Neurological Institute (MNI) space and the left and right ATL are identified and labeled with a 5mm sphere centered on the coordinates [x=-52, y=2, z=-28] and [x=53, y=4, z=-32], respectively [51] using custom made SPM8 (Statistical Parametric Mapping, Matlab Mathworks) procedure. Images are then denormalized in each patient’s native space. The
day of the stimulation a scalp inspection is performed to check for the absence of skin
lesions. Before the placement of the tDCS electrodes a carefully cleaning of the scalp
using an abrasive paste is realized to limit impedance losses between the skin and the
electrodes. Two round sponge electrodes, one acting as the active electrode (anodal or
cathode) and the other as return (Sponstim®, 5.65cm diameter, 25cm² surface,
NEuroelectrics, Barcelona, Spain) are placed under MRI guidance using a stereotactic
neuronavigation system (Brainsight®, Rogue System®, Montreal, Canada). This procedure
minimizes the distance between the labeled cortical target on a 3D reconstruction of each
patient’s MRI and its closest (shortest Euclidian path) scalp location (Figure 2).
Stimulation is delivered using a wireless hybrid EEG/tDCS 8-channel neurostimulator
(Starstim, NEuroelectrics, Barcelona, Spain). For left-anodal tDCS the active electrode is
placed on the left ATL (between 10-20 EEG coordinates ~FT7 and FT9), while the cathodal
tDCS targets the right ATL (between ~FT8 and FT10). For sham tDCS stimulation, the
active electrode is placed over the same MNI coordinates as left-anodal tDCS. The return
electrode is placed on the contralateral supra-orbital region with regards to the active
electrode location (10-20 EEG coordinates AF8 for left anodal tDCS and AF7 for right
cathodal tDCS). Additionally, 6 EEG scalp electrodes (NG Geltrode® Ag/AgCl, 1.0 cm²)
provided by the same tDCS device, which allow for continuous monitoring of brain activity
during the session, are placed in 10-20 EEG system locations F4, F3, C4, C3 and P4, P3.
During anodal or cathodal tDCS, current intensity is linearly increased during 30 seconds
to reach a maximum of 1.59mA. This level of tDCS intensity has been chosen to ensure
similar levels of current density (0.06 mA/cm²) with our 25cm² electrodes as those applied
in previous post-stroke aphasia or PPA studies with larger leads [37,45,52]. Current is kept
on at this intensity during 20 minutes before being ramped down along 30 seconds at the
end of the tDCS session. During sham stimulation tDCS current is ramped up and down along 30 seconds at the initial and final phase of the session but is turned off during the 20 minutes of the session. This process makes active (anodal or cathodal) and sham stimulations similar (same somatosensory skin sensations) as required by a double-blind design. During each stimulation session, values of mean voltage (V), mean current intensity (uA) and mean impedance (Ohm), directly provided by the stimulation software, are recorded for means of tDCS verification intents. Treatment duration has been chosen based on previous studies showing beneficial tDCS effects with the same amount of time [e.g., 44,45]. During tDCS sessions patients perform a language-neutral visuo-motor task on a laptop screen consisting in pressing the space bar of a computer keyboard every time a slowly moving dot contacts the edge of a surrounding rectangle. This task is intended to limit variability in neural activity states across sessions and patients (forcing patients to maintain vigilance) but without interfering with the language processes and tasks. At each session, performances (number of trials, number of successes and number of errors) in the visuo-motor task are recorded for patient resting-state standardization purposes during stimulation.

To ensure safety and comfort and to assess the tolerance of patients to stimulation, immediately after each tDCS session patients are asked to fill a ‘tDCS adverse effects questionnaire’ [53] that measures, through a rating scale, patients’ sensations in a set of the most frequent adverse effects reported in tDCS studies such as itching, tingling, burning sensations, skin redness or sleepiness.

Language/semantic tasks

A set of computer-based language/semantic tasks is used to assess the potential effectiveness of tDCS. These tasks are applied at 4 time-points: pre-tDCS (baseline), 3
days, 2 weeks and 4 months following the end of the 10 days tDCS regime. The 5
language/semantic tasks are the same as the tasks elaborated and used in the pre-
therapeutic tDCS study on SD of our team [45]:

Semantic association task (SA)
Picture naming task (NAME)
Reading task (READ)
Letter and category fluency task (FLU)
Category judgment task (CJ)

Two additional tasks evaluating tDCS impacts on other cognitive functions are also
applied:

Executive function task (EXE)
« Recognition of famous faces » subtest of the French Batterie Imagerie-Perception (BIP).

The tasks are computer-programmed using the E-Prime software (E-Prime®, Psychology
Software Tools, Sharpsburg, PA, USA). Participants are comfortably seated in front of a
laptop computer screen (HP EliteBook 8770w, USA) and a response box recording the
responses. Performance accuracy, reaction times and voice-records are automatically
registered by the software. The sessions are carried out in the presence of an investigator
immediately after familiarization/training blocks made of 5 trials for each task.

Semantic association task (SA): This task assesses semantic capacities and provides the
primary endpoint criterion. It is based on the principle of the Pyramid Palm Trees Test
[54]. The material includes 78 French words, which are grouped into 26 trials. Each trial
includes three words, two of them are semantically related (the target item and test item),
while the third word is a semantically unrelated distractor. The 3 words are displayed on
the computer screen for 8 seconds and the participant has to decide, using two response
buttons, if the target item (on top of the screen) is associated with the test item (on the
bottom of the screen, either left or right) or the distractor item (on the bottom of the screen, either left or right, opposite to the test item). Thirteen test items appear on the left and 13 on the right. The lack of response during this 8-second time interval is recorded as an error. The task assesses two category dimensions using the contrast between trials containing only living items (n=13) and trials containing only non-living items (n=13). Two modalities of this task are applied: a *verbal modality* (described above) and a *picture modality* that uses pictures instead of words. The stimuli of “living” and “non-living” trials are matched for (1) lexical frequency, (2) number of letters, (3) familiarity of words and pictures and (4) visual complexity of the pictures. Stimuli of Version 1 and Version 2 of the test are also matched for these 4 variables. Outcome measures will focus on performance (number of correct responses), and reaction times in milliseconds. Figure 3 illustrates different trials of the task.

*Picture naming task* (NAME): this task evaluates lexical and semantic abilities. The material includes 40 pictures, derived from two picture-naming databases [55,56]. Each picture is displayed on the computer screen for 8 seconds and subjects are asked to name it aloud. Responses are voice-recorded and notified. The lack of a response during this time interval is recorded as an error. Stimuli of Version 1 and Version 2 of the test are matched for (1) lexical frequency of words, (2) familiarity of words and pictures and (3) visual complexity of the pictures. The task will allow for assessing (1) the number of correct responses, (2) the number of non-responses, (3) the number of semantic paraphasias. The number of correct responses and non-reponses are markers of lexico-semantic abilities, and the number of semantic paraphasias is an additional marker of semantic abilities.
The reading task (READ): The task provides a semantic marker in written language. During reading the phonological pathway allows for mapping each letter (grapheme) to a phoneme while the lexical-semantic pathway allows for ‘whole word reading’ that depends on knowledge of written words [57]. The lexical-semantic route is therefore critical for reading irregular words where the grapheme-phoneme correspondence is not transparent (e.g. bear). In contrast, the phonological route is essential to read unknown or non-words. Regular words can be read through both the lexical-semantic and the phonological route. It has been shown that SD patients have difficulties reading irregular words linked to their semantic impairment [e.g., 58] and performance with such irregular items will therefore provide a semantic marker. The task contains 45 stimuli: 15 irregular words, 15 regular words and 15 non-words that are matched for the number of graphemes. Each word appears on the computer screen for 8 seconds and subjects are asked to read them aloud. Responses are voice-recorded and notified. The lack of response during this interval is recorded as an error. Stimuli of Version 1 and Version 2 of the test are also matched for the number of graphemes.

Verbal fluency task (FLU): This task assesses language fluidity and access to lexical-semantic representations of words. It has two modalities: (1) in the "letter fluency" subtask participants are asked to produce, during 1 minute, a maximum of words beginning with a particular letter displayed in the center of the screen; (2) in the "category fluency" subtask participants are asked to produce, during 1 minute, a maximum of words belonging to a given semantic category displayed in the center of the computer screen. For the “letter fluency” subtask, stimuli of Version 1 and Version 2 are matched for the number of existing words starting with the given letters and also their cumulative frequencies. For the “category fluency” subtask stimuli of Version 1 and
Version 2 are matched for the number of existing items within that category. The measured variables are the number of items produced per minute in each of the two tasks. The subjects' responses are voice-recorded and quantified.

*Category judgment task (CJ):* The task assesses semantic capacities in the verbal modality. The material includes 40 French words, 20 of which representing “living items” and 20 representing “non-living items”. Words representing “living” and “non-living” items, and words of both Versions of the task, are matched for lexical frequency, number of letters and number of phonemes. Each word stimulus is displayed in the center of a computer screen for 8 seconds. Subjects are asked to judge whether a given word item belongs to a “living” or to a “non-living” semantic category and answer by pressing the correspondent buttons of the ‘response box’.

*Executive function task (EXE):* This task is used as a control task to assess whether tDCS over anterior temporal regions has semantic-specific effects or whether it might impact on executive functioning which may indirectly modulate semantic performance. Using a similar task design and procedure as in the SA task the EXE task assesses executive/attention and decision-making abilities without the influence of semantics. As in the SA task, the task contains a verbal and a picture modality. For the verbal modality, the material includes 78 French words which are grouped into 26 trials. Each trial includes three words, two of them have the same initial and final letters (the target item and test item), while the third word is a distractor sharing only the initial or the final letter with the target item. The three words presented in each trial are semantically unrelated. The picture modality of the test uses pictures instead of words. The pictures represent colored geometrical shapes. The material includes 78 pictures, which are grouped into 26 trials.
Each of the 26 trials includes three pictures, two of them representing the same geometrical shape or color (the target item and test item), while the third picture represents a distractor not sharing any of these features with the target item. The items of the verbal modality are matched for the number of letters with the word items of the SA task. Stimuli of Version 1 and Version 2 are matched for the number of letters. The 3 items are displayed on the computer screen for 8 seconds and subjects decide (using two response buttons) if the target item (on top of the screen) is related to the test item (on the bottom of the screen, either left or right) or the distractor item (on the bottom of the screen, either left or right, opposite to the test item). Thirteen test items appear on a left and 13 on a right bottom location. The lack of response during this 8-second time interval is recorded as an error. Figure 4 illustrates two trials of the test.

Recognition of famous faces subtest: This test is adapted from the French Batterie Imagerie-Perception (BIP) [59]. It is used to detect eventual negative effects of tDCS decreasing the activity of the right ATL which has been shown to play an important role in the recognition of known faces [7]. The material includes 28 pictures of faces of famous people. Each picture is displayed on the computer screen for 8 seconds. Participants have to decide if the face corresponds to one of the four professional categories (‘politician’, ‘actor’, ‘singer’ or ‘TV presenter’) by pressing one of 4 associated keys in the ‘response box’. The lack of response during this time interval is recorded as an error.

Ecological evaluation

Additionally, a semi-quantitative daily life communication questionnaire – Echelle de Communication Verbale de Bordeaux [60], which assesses the effectiveness of communication of patients with aphasia in everyday situations, is applied at baseline and
at the ‘2-week’ time-point. This questionnaire was included in the protocol to have an ecological measure of the possible language improvements after tDCS and to assess its impact in patients’ day-to-day life.

PET-MRI neuroimaging and EEG recording

Neuroimaging data (PET, MRI) are collected before the tDCS sessions (baseline) and two weeks after the tDCS sessions. These acquisitions provide all the sequences underlying the following explorations: cortical metabolism, gray matter thickness measures, white matter fiber tracking and functional connectivity. The exam is performed on a hybrid PET-MRI scanner (Signa 3T GE Healthcare, USA). The injection of FDG (Fluoro-Deoxy-Glucose 2-18F: MÉTATRACE®, half-life of 109.77 minutes, or GLUSCAN®, half-life of 110 minutes) is performed if the blood glucose checked prior to injection is equal or less than 1.5 g/l. Participants are lying in neurosensory rest in a quiet and unlit room for at least 30 minutes post-injection, prior to image acquisition.

The acquisition of brain images with PET is conducted in _list mode_. It begins 30 to 40 minutes after injection of the radiopharmaceutical FDG tracer and lasts for 20 minutes (3 x 5 minutes). Images are reconstructed and corrected for physical phenomena. They are expressed in Standard Uptake Value (SUV). Magnetic Resonance Images are acquired simultaneously and include: Two anatomical sequences (3D-T1, 3D-FLAIR), a functional imaging sequence (resting state fMRI) and a diffusion tensor imaging (DTI) sequence.

Additionally, prior and following stimulation, the tDCS device (_Startsim, NEuroelectrics, Barcelona_) automatically records brain activity through EEG scalp electrodes (sampling at 1500 Hz, low pass high pass filter 4 to 40 Hz). Resting-state EEG recordings are obtained from 10-20 EEG system locations F4, F3, C4, C3, P4, P3 and T8, T7.
Computational models of tDCS current magnitude and distribution

There is evidence that the thickness of the skull, the volume of the cerebrospinal fluid in the subdural space and the distance of the targeted region in the brain to the tDCS anode or cathode account for up to 50% of the spatial variation of the electric field strength [62, 63, 64]. Kim et al. [65] found that performance improvement in a working memory task correlated with the simulated current magnitude, suggesting that inconsistent behavioral outcomes of tDCS might be partly due to individual anatomical differences.

A computational approach of modeling should allow for defining if and how tDCS current magnitude and distribution in an individual head can differ and how these differences will affect clinical outcomes. Individual computational models will be produced using the open-source tool ROAST [66] to simulate tDCS current magnitude and distribution using the anatomical 3D T1-weighted images of each patient (Figure 5). Then, model-estimated data for current magnitude values will be correlated with clinical data, specifically with the changes in scores of the SA task from baseline to post-stimulation. A region of interest (ROI) will be defined and the mean of the 5% highest electric field values in this region will be obtained for each patient. More specifically, we will define the left anterior/middle temporal lobe as our region of interest, since this region is primarily damaged in SD and it is the targeted region during the stimulation. Finally, measures of cortical thickness of the different tissue layers (skin, skull, cerebrospinal fluid volume, grey matter and white matter) and distances between stimulation target (ATL) and the stimulating electrode will be obtained and regression models will be run to identify specific features that most influence changes in current magnitudes. This will allow for identifying the characteristics that might influence patients’ response to tDCS and help predict if an individual patient might benefit from a tDCS treatment.

Data management
Data collection is performed via an electronic case report form (eCRF) that was developed at the outset of the study. All clinical and language/semantic task information required by the protocol is entered in the eCRF. The data are collected as and when they are obtained, and any missing data are clearly coded. Every investigator participating in the protocol has access to the eCRF via a web-based password-protected data collection system. All data are collected in the same center, the Pitié-Salpêtrière Hospital. Investigators have been given instructions for using this tool prior to the beginning of the protocol. Regular monitoring by the promoter of the study (Assistance Public-Hôpitaux de Paris) will be done to ensure the accuracy and quality of all the data, and to detect and address any issues related to the implementation.

All data collected regarding participants are anonymized through a specific number attributed to the participant at inclusion and his/her respective last and first name initials. Data collected in this study include both quantitative (neuropsychological and language standard tests scores, study task scores) and qualitative data (clinical history, neurological information). A print-out, authenticated by the principal investigator of the protocol, will be requested at the end of the research by the promoter of the study and the investigator will archive a copy of the authenticated document that was issued to the promoter. Given the high-volume and complexity of the pre-processing and analyses of PET-MRI and EEG data, the outcome measures cannot be added to the eCRF and will be treated by the expert investigators.

Data analyses

We will accept a significance level of 5% (p < 0.05) corrected for multiple comparisons when needed. To assess our primary endpoint criterion, i.e. performance changes on the SA task, baseline and 2-week time-point outcomes will be compared with a two-way
analysis of variance with ‘time’ (baseline, ‘2-week’ post-stimulation) and ‘group’ (anodal, cathodal, sham) as factors. Since two comparisons will be made (sham vs left-anodal and sham vs right-cathodal tDCS) Dunnett’s two-tailed t-test will be used to handle with multiple comparisons. In case the data do not meet the normal distribution assumption, the non-parametric Mann-Whitney test will be used to compare performance improvements between sham vs left-anodal and between sham vs right-cathodal groups.

To assess the time course of potential improvements and test if these potential improvements differ between the treatment groups, linear mixed models will be used with ‘time’ (the 4 time-points), ‘group’ (anodal, cathodal and sham) and group-by-time interactions as fixed effects. The Benjamini and Hochberg method will be used for controlling for false discovery rate.

Regarding the study of imaging biomarkers for stimulation-induced neuroplasticity, we will analyze cortical metabolism and resting-state functional connectivity. For cortical metabolism, PET images will be used to analyze tissue metabolic activity, regional glucose uptake, in the brain. A voxel-based analysis, using the Statistical Parametric Mapping software [67], will be implemented to highlight regional differences of metabolism between the images obtained at baseline and post-stimulation. The resting-state fMRI sequence will be explored to study the status of functional connectivity within the language/semantic network and possible alterations within it between baseline and post-stimulation images. Quantification of functional connectivity will be based on integration measures in specific networks established from the known anatomy, on indices derived from the graph theory or the theory of information. Cortical regions of interest will be obtained automatically by using SPM toolboxes. More specifically, the language/semantic network will include: the middle and anterior portions of the lateral ATL, the inferior frontal gyrus, the dorsomedial and ventromedial prefrontal cortex and the inferior parietal
lobe. EEG resting state data will be analyzed for day 1 (pre-stimulation) and 3 days after the end of the tDCS sessions allowing for studying the impact of tDCS on specific brain networks. Data will be filtered (4 to 40 Hz) and power distribution and local synchrony calculated across frequency bands and compared. Functional connectivity maps across the 8 electrodes will be estimated by calculating the phase-locking value between the 8 electrodes and compared across.

To identify biomarkers which could predict individually an efficient tDCS impact, 3D-T1 and DTI sequences will be explored and individual computational models of tDCS current will be produced. Structural 3D-T1 MRI data will be studied with surface-based cortical thickness analysis. A region of interest will be defined in the ATL and cortical thickness values at baseline between a group of eventual ‘responders’ (mean improvement of 15% of correct responses in the AS test) and ‘non-responders’ to the stimulation will be compared using a general linear model. Diffusion MRI data will be studied using region of interest analysis of DTI metrics. Fractional Anisotropy (FA) and Mean Diffusivity (MD) maps will be calculated and the integrity of a set of anatomical white matter tracts will be assessed (FA and MD measures in each tract). The tracts analyzed will include bilaterally the inferior longitudinal fasciculus, the uncinate fasciculus, the superior longitudinal fasciculus, the inferior frontal-occipital fasciculus and different corpus callosum tracts connecting the left and right temporal lobes. This will allow for performing a detailed anatomical analysis of white matter fiber tracts and compare their status between a possible group of ‘responders’ and ‘non-responders’. The individual computational models of tDCS current magnitude will be used to identify which anatomical characteristics influence the amount of current reaching the target in the brain and if the amount of current reaching the target can help predict the response to stimulation. Finally, the analyses on DTI data will also allow to explore connectivity between left and right ATL to
improve the understanding of the semantic roles of each of these regions while contributing to the definition of anatomo-functional models of semantics.

Sample size
The primary evaluation criterion is the change over two weeks in performance on the SA task. Performance is measured by the percentage of correct responses. The trial will be considered as positive if performances after left-anodal or right-cathodal tDCS are shown to be significantly superior to the sham stimulation. The sample size is based on the results of a preliminary study comparing sham stimulation to left-anodal and right-cathodal stimulation in 12 patients [45]. In this pre-therapeutic study of our group, the mean difference in performance improvement between sham and both anodal and cathodal stimulation was about 15% with a standard deviation of 16% and an effect size of 1.464. The mean change difference between sham stimulation and each of the two other groups is then expected to be 15% of correct responses. The inclusion of 20 patients in each group will provide a statistical power of 80%.

Discussion
Language disorders in frontotemporal lobar degeneration, and particularly in SD, are a disabling feature representing an important medical problem. They are also a relevant issue for public health because most patients have symptoms prior to retirement causing substantial health costs. Given this context our project bears major importance because it could potentially provide evidence for the validity of a novel therapy strategy improving language/semantic capacities while diminishing the functional handicap and, eventually, health care expenditure.

The STIM-SD protocol proposes the first large-scale exploration of tDCS as a potential therapy approach for language/semantic impairment in SD for which no treatment is
currently available. Contrary to most of the studies using transcranial brain stimulation in neurodegenerative diseases affecting language [43,44,61,68], our study targets sites that have been selected considering the localization of anatomical damage and related contralateral regions to optimize language and semantic recovery.

We apply a double-blind, sham-controlled design in which the investigators and the patients are blinded for the type of stimulation used, reducing any source of potential bias. This design will also enable a comparison between two different stimulation strategies (left-anodal versus right-cathodal) and will contribute to identify the most beneficial strategy. To our knowledge, not a single study applying tDCS to improve cognitive deficits in neurodegenerative diseases employed supportive neuroimaging (PET, fMRI) and only one study used neurophysiological measures (EEG) [69] to explore stimulation impact on relevant brain networks or demonstrate neuroplasticity effects. In the present study, we will generate both language/semantic data and a set of neuroimaging and resting-state EEG measures which will allow for studying the impact of tDCS on language networks and neuroplasticity, to understand if this potential impact also subtends language/semantic improvements, and to better understand tDCS mechanisms.

In addition, baseline structural data (3D-T1 cortical thickness, DTI fiber tract analyses) and individualized models of tDCS current magnitude and distribution will be fundamental for understanding which anatomical features in our cohort most influence treatment efficacy, how the amount and spreading of electric current in the brain will impact individual clinical outcomes, and possibly define a profile of patients that in the future would most benefit from tDCS.

Within neurodegenerative diseases specifically affecting language, SD is the most frequent variant of primary progressive aphasia [2], which contributes to the feasibility of the project in terms of patient recruitment. In the same vein, our ‘National Reference
Center for Rare or Early Onset Dementias’, provides the unique opportunity to recruit a large and homogeneous SD patient cohort. The active patient files of early stage SD patients in the three recruiting centers contained more than 20 SD patients per year in 2013, 2014, 2015. This fact shows that the total number of 60 SD patients in the current project will be reached.

A potential tDCS treatment would be easily applicable, inexpensive, and renewable when therapeutic effects disappear due to disease evolution. Significant effects of repetitive tDCS on language/semantics would also open an avenue for future tDCS trials targeting language non-related cortical regions such as areas subsuming episodic memory which is damaged for example in Alzheimer’s disease. More generally, the protocol might improve the understanding of neuroplasticity and its modulation through inhibitory and/or excitatory tDCS-driven cortical impact while providing a rationale for appropriate stimulation modalities and for identifying brain regions likely to demonstrate relevant plasticity. Such insights could prove important for both tDCS and TMS trials, and their combination with behavioral language rehabilitation strategies which might further enhance plasticity-related modulation.

Conclusions

The STIM-SD protocol aims at implementing a novel therapeutic tDCS approach to language/semantic deficits in patients with SD for whom no treatment is available. If found to be efficient, this strategy could be regularly implemented since it is easily applicable and with low costs, and larger trials could be extended to other neurodegenerative diseases to check for efficiency on language and other cognitive functions.

Trial Status
The protocol version number is P160937J first published in clinicaltrials.gov on March 29 2018. Patient recruitment began in June 2018. Sixteen patients have been screened for participation between June 2018 and March 2019. Of these, 13 patients were included in the study and randomized. Seven of those patients have already completed their participation in the study with the 4-month follow-up. From the 3 patients screened but not included, 2 of them did not meet all inclusion criteria while another one was not able to undergo the planned neuroimaging exams. Five healthy participants were also included in the protocol. Recruitment is expected to be completed by June 2021.

Declarations

Ethics approval and consent to participate

The protocol has been approved by the local Ethics Committee – Comité de Protection de Personnes (CPP) Ile de France IV, Hôpital Saint-Louis. All patients and healthy subjects will be required to provide written informed consent.

Consent for publication

MRI datasets, images or any other clinical details of the participants will not compromise their anonymity.

Availability of data and material

Not applicable. No data is available at this point because authors are still in the process of gathering such.

Competing interests

The authors declare that they have no competing interests.

Funding

Assistance Public des Hôpitaux de Paris – Protocole Hospitalier de Recherche Clinique
Régional. The funding body will only be involved in the monitoring process of the protocol.

Authors’ contributions

MT and AV-C provided the idea, designed the protocol and will coordinate organizational and scientific aspects of the clinical trial, with an emphasis on data interpretation and a main role in manuscript writing. RL, SB, LVG, BD and MT will be in charge of patient clinical characterization, screening and recruitment. CS and AV-C will be responsible for carrying our tDCS stimulation, stimulation coupled EEG procedures and biophysical modeling of current distribution. MOH, SS, AuK, CS, AnK, NP and RM will be responsible for imaging acquisitions. MOH, LM, NP and SS will define the neuroimaging analysis strategies with input from MT and AV-C. CS, AnK, RM, NP and MOH will be responsible for imaging analyses. AnK, CS and MT will be responsible for the evaluation and interpretation of language outcomes. CS, AB and MT elaborated language and executive evaluation tasks for this study. AV-C and CS were in charge of defining the tDCS stimulation and EEG recording strategies CS, MOH, SS, NP and RM have been in charge of designing, optimizing and will provide quality checks for neuroimaging sequences and datasets. CS, AV-C and MT were responsible for the preparation of the current manuscript. AV-C and MT were directly responsible for IRB protocol writing.

All authors have read and approved the final version of the manuscript.

Acknowledgments

The work of Ms. Clara Sanches is funded by a PhD Fellowship by the Fondation pour la Recherche sur l’Alzheimer (FRA). Dr. Marc Teichmann has received support from the ‘PSP-France’ and is currently funded by the PHRC Regional “STIM-SD”. The activities of the group of Dr. Valero-Cabré are supported by research grants IHU-A-ICM-Translationnel,
Agence National de la Recherche (ANR), project Générique “OSCILOSCOPUS”, and eraFlag JTC 2017- “CAUSALTOMICS”, the PHRC Regional “NEGLECT” and the PHRC Regional “STIM-SD”. The authors thank the FRA foundation for logistic and equipment support and the Naturalia & Biologia Foundation for financial assistance for traveling and attendance to meetings.

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Figures
Figure 1

A. Flow diagram, from patients’ selection to the end of their participation in the study. B. Standard protocol items: Recommendations for Interventional Trials (SPIRIT) Checklist.
Figure 2

The upper panels show, respectively, a coronal, axial and sagittal slice of a SD patient MRI with the crosshair in the target region (left ATL). The first bottom panel shows the Brainsight stereotactic navigation system. The second and third bottom panels show a 3D brain curvilinear and skin reconstructions from a patient MRI, performed with Brainsight, with a sphere indicating the target region to stimulate.
Two trials of the SA test in their verbal and picture modalities. A. Trial with living items for the verbal modality (‘chenille’ [caterpillar] - target item, ‘scarabée’ [beetle] - distractor, ‘papillon’ [butterfly] - test item). B. Trial with living items for the picture modality using the same items in a picture format. C. Trial with non-living items for the verbal modality (‘ceinture’ [belt] - target item, ‘pantalon’ [trousers] - test item, ‘gilet’ [vest] - distractor). D. Trial with non-living items for the picture modality using the same items in a picture format. For each trial in each modality subjects have to decide, by pressing one of two buttons in the ‘response box’, which of the items presented on the bottom of the screen is associated with the target item, on the top of the screen.
Two trials of the EXE test. A. Trial in the verbal modality. For each trial subjects have to decide which of the test items presented on the bottom of the screen begins and ends by the same letter as the target item (‘abeille’ [bee] - target item, ‘abysse’ [abyss] - test item, ‘abri’ [shelter] - distractor). B. Trial in the picture modality. For each trial subjects have to decide which of the tests items on the bottom of the screen has the same color or the same geometric form as the target item.
Example of a computational model produced with the open-source tool ROAST for the simulation of tDCS current magnitude and distribution using a 3D T1-weighted image of one patient. A. Illustration of the montage for the left-anodal stimulation. The blue disc represents the cathode, placed over the right supra-orbital region [10-20 EEG coordinates AF8], and the red disc represents the anode, placed over the left ATL [MNI coordinates: x=-52, y=2, z =-28]. B. Coronal slice view of the electric field with current flow direction represented by the black arrows. C. Electric field distribution on cortical surface - right hemisphere view, left hemisphere view, upper view and frontal view. The color-bar represents the magnitude of the electric field, in Volts per meter, in different regions of the brain (for B and C panels).

Supplementary Files

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