Transcranial direct current stimulation versus intermittent theta-burst stimulation for the improvement of working memory performance

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Abstract
Non-invasive brain stimulation (NIBS) techniques have been increasingly used over the dorsolateral prefrontal cortex (DLPFC) to enhance working memory (WM) performance. Notwithstanding, NIBS protocols have shown either small or inconclusive cognitive effects on healthy and neuropsychiatric samples. Therefore, we assessed working memory performance and safety of transcranial direct current stimulation (tDCS), intermittent theta-burst stimulation (iTBS), and both therapies combined vs placebo over the neuronavigated left DLPFC of healthy participants. Twenty-four subjects were included to randomly undergo four sessions of NIBS, once a week: tDCS alone, iTBS alone, combined protocol and placebo. The 2-back task and an adverse effect scale were applied after each NIBS session. Results revealed a significantly faster response for iTBS (b = -21.49, p = 0.04), but not for tDCS and for the interaction tDCS vs. iTBS (b = -21.49, p = 0.04).
Background

Working memory (WM) describes the temporary storage and online manipulation of the information necessary for performing cognitive tasks (Baddeley & Baddeley, 1986). WM is primarily processed in the prefrontal cortex (PFC), particularly its dorsolateral region (DLPFC) (Barbey et al., 2013). Thus, one approach to explore and manipulate WM performance is using non-invasive brain stimulation (NIBS) over this region, such as theta-burst (TBS) or direct current (tDCS) stimulation. The former uses electromagnetic pulses to trigger action potentials and produce long-term changes in cortical activity, being a novel method of transcranial magnetic stimulation (TMS) associated with stronger and faster changes in cortical excitability (Huang et al., 2005). The latter does not trigger action potentials, but rather induces either excitatory or inhibitory changes in cortical activity according to the parameters of stimulation (Nitsche & Paulus, 2000). In fact, although transcranial DCS and TBS can independently modify WM performance (Chung et al., 2018; Moreno et al., 2015), findings have been mixed, and the effects seem to be small, particularly in neuropsychiatric samples. For instance, a recent meta-analysis evaluating the WM performance of health volunteers found only a small positive effect favoring tDCS (Wischnewsk et al., 2021). Similar results were found in another meta-analysis that investigated the TBS effects over the PFC for executive functions (Lowe et al, 2018). Moreover, reviews suggest that tDCS over the PFC did not change cognitive performance in neuropsychiatric populations (Martin et al., 2018; Farhat et al., 2020).

Due to these mixed findings in the tDCS field, innovative NIBS protocols have been pursued such as combining NIBS techniques simultaneously, for achieving larger effects (Hasan et al., 2012). One strategy that has shown promising results in studies investigating the motor and visual cortex excitability was the application of tDCS to preconditioning the effects of a secondary NIBS protocol (i.e. tDCS or rTMS). Preliminary studies suggest that preconditioning the target area with tDCS may alter the effects of subsequent rTMS protocols in a manner that generates more robust outcomes (Hurley & Machado, 2017). Similar results were also observed in animal models, in which researchers found that anodal DCS augmented long-term potentiation (LTP) outcomes whereas cathodal DCS seemed to not affect it (Sharma et al., 2021). While these are promising findings, studies combining NIBS protocols in humans have focused primarily on evaluating cortical excitability, a brain outcome that is relatively easier to measure than the ones available for the DLPFC. In fact, only two trials used tDCS to precondition the DLPFC before a course of excitatory rTMS protocol for depression and stress outcomes (De Smet et al., 2021; Loo et al., 2009), but no study investigated the cognitive findings of this combination.

Therefore, here we aimed to evaluate whether working memory performance — as a proxy to DLPFC activity — is modified by application of tDCS, iTBS, or the combination of tDCS + iTBS, in a factorial, double-blinded, sham-controlled trial. Our primary hypothesis was that tDCS and iTBS combined would enhance WM performance. Our secondary hypotheses were that tDCS and iTBS would also improve cognitive performance in comparison to sham. Finally, the tolerability of the combined treatment was also investigated. Despite unveiling mechanisms of action of TBS and tDCS on the DLPFC, this would also be important from a clinical perspective; because WM impairment is associated with psychiatric disorders (Bortolato et al., 2014; Gold et al., 2019), and restoring normal WM performance might ameliorate their cognitive symptoms.

Materials and methods

Design

The study was conducted at the Institute of Psychiatry, University of São Paulo, Brazil, from March 2019 to March 2021. It was approved by the local and national Ethics Committee (CAAE: 89310918.8.0000.0068) and participants provided written, informed consent. We used a factorial, double-blinded, within-subjects design, in which participants received, once a week and in a randomized order: sham tDCS/sham iTBS (placebo); active tDCS/active iTBS (combined protocol), sham tDCS/active iTBS (iTBS-only); and active tDCS/sham iTBS (tDCS-only) (Razza et al., 2021) (Sup. Material - Appendix A).

Participants

We included right-handed, 18-45 years-old subjects, without previous or current neuropsychiatric disorders or clinical conditions. Participants were recruited through flyers and social media platforms. Volunteers were first screened by e-mail and those who met the inclusion criteria underwent on-site evaluation by a trained psychologist. Exclusion criteria were specific contraindications for NIBS interventions and MRI (e.g., metal implants), current smoker (>10 cigarettes/day), use of substances, pregnancy, and use of psychoactive drugs.

Procedures

The baseline assessment included a brief sociodemographic questionnaire, the Portuguese-based version of the Mini International Neuropsychiatric Interview (MINI) (Amorim, 2009), and a mood questionnaire, the Portuguese-based version of the Mini International Neuropsychiatric Interview (MINI) (Amorim, 2009), but no study investigated the cognitive findings of this combination.
the Hamilton Depression Rating Scale (HDRS) (Hamilton, 1960), the Beck Depression Inventory (BDI) (Beck et al., 1961).

After these assessments, we collected an anatomical brain scan of the participants in a 3-Tesla Magnetic Resonance Imaging (MRI) equipment (General Electric PET/MRI equipment). In the following days, a neuronavigation procedure (Brainsight, Rogue Resolutions, Inc) was performed, in which we found the left and right DLPFC according to the MNI coordinates x = -38, y = +44, z = +26, per previous studies (Blumberger et al., 2018; Fox et al., 2012).

Each session lasted twenty minutes. TDCS was performed alone for around 11 minutes and TBS was concomitantly applied for the last 8 minutes and 40 seconds. Afterwards, the n-back task was performed, and a questionnaire of adverse effects was applied (Sup. Material - Appendix B) (Aparicio et al., 2016). (Figure 1).

Interventions

The tDCS procedure lasted 20 minutes and was applied with a current of 2mA using 25cm² saline-soaked sponges. Anode and cathode were placed over the left and right DLPFCs, as described above. Sham tDCS sessions were identical, but the active current only lasted 30 seconds. The tDCS device was set to deliver either active or sham stimulation according to a randomized code (Neuroconn DC-Stimulator, Ilmenau, Germany).

Regarding iTBS, the coil was placed over the anode and positioned 45° relative to the midline. We performed 54 cycles of 10 triplet bursts with a train of 2s and an interval of 8s between trains (1620 pulses) at 110% of the resting motor threshold, corresponding to a duration of 8 minutes and 40 seconds. The coil used for iTBS has two identical sides that delivers either active or sham stimulation (Cool-B65 Active/Placebo - MagVenture), which is chosen according to a randomized code inputted in the TMS device.

2-Back task

The cognitive task was programmed in E-prime 2.0 software (Psychology Software, Tools Inc Pittsburgh, Pennsylvania, USA). The visual stimulus consisted of alphabet letters (A to Z) that appeared individually and randomly on a 15-inch screen. Three blocks of 30 letters each, displayed on the screen for 500ms, with an interstimulus interval of 3000 ms, were presented. Each block had 10 target letters – i.e., the same letter that had been presented two letters before.
Participants were instructed to identify target and not-target letters by pressing ‘2’ (‘target’) or ‘0’ (‘non-target’), respectively. A brief practice session containing 20 stimuli was performed immediately before each n-back session. The 2-back was chosen because it was previously associated with WM improvement after NIBS in healthy participants (Brunoni & Vanderhasselt, 2014).

Statistical analysis

Statistical analyses were performed using RStudio Version 4.1.2 (Team & Others, 2013). Accuracy (binary outcome) and reaction time (RT, in milliseconds (ms)) of the target stimuli were the dependent variables. Missed responses were considered errors. Also, RTs < 200ms and >2500ms were considered not genuine responses and were excluded (Whelan, 2008). Due to the non-normal distribution of RT responses, generalized linear mixed models (GLMM; using the ‘lme4’ package) were calculated to select the statistical model with the best fit. The final model for our analysis was chosen by using the Akaike Information Criterion (AIC) and checking the normality of residuals (See Sup. Material - Appendix C). We employed the inverse-gaussian and binomial distributions for the analyses of RT and accuracy, respectively.

RT or accuracy were the dependent variables, while the factors tDCS (i.e. tDCS yes / tDCS no) and iTBS (i.e. iTBS yes / iTBS no), the variable session and their interaction (tDCS*iTBS*Session) were added as fixed effects (Sup. Material - Appendix C). The variable subject was included as a random intercept. Our primary hypothesis would be confirmed if the interaction between tDCS and iTBS were significant, indicating synergistic effects of these techniques. Linear mixed models were conducted to evaluate adverse effects among groups. Pairwise comparisons were post hoc and performed using the R Package ‘emmeans’. All results were considered significant at a p-value < 0.05. More information regarding sample size calculation can be found elsewhere (Razza et al., 2021).

Results

Out of 211 volunteers, 56 of them underwent onsite or online screening and 24 were included in our study (Sup. Material - Appendix E). Their mean age was 28.7 (standard deviation (SD) = 6.95) years, and most of them were women with university degrees. Twenty-three participants completed all NIBS sessions, with a total of 93 NIBS sessions being performed. The mean accuracy and RT for each group is displayed in Table 1.

Regarding our primary hypothesis, the factorial analysis revealed no significant effects for the interaction iTBS vs. tDCS for either RT (Figure 2A) or accuracy (Figure 2B) in the 2-back task (Table 2). Analyses “at the margins” revealed that iTBS (iTBS and sham tDCS, and iTBS and active tDCS) presented a faster response than no iTBS (sham iTBS and sham tDCS, and sham iTBS and active tDCS). No effects “at the margins” were found for tDCS.

Moreover, a significant effect of “session” on reaction time ($t = -7.53, CI = -88.7; -61.8, t = -5.45, p < 0.001$) (Figure 3A) and accuracy ($t = 0.56, CI = -0.29; 0.83, p = 0.04$) (Figure 3B) was found, showing cognitive performance increasing over time. (Sup. Material - Appendix F).

Regarding tolerability, no serious adverse effects, such as seizures, were reported. The combined intervention group presented higher rates of headache compared to placebo and tDCS, but not to iTBS. Patients receiving tDCS presented higher rates of skin redness. Finally, facial twitches were more common in the iTBS group (Sup. Material - Appendix G).

Discussion

The present study was the first to evaluate the cognitive effects of combined and standalone protocols of tDCS and iTBS on the DLPFC in healthy participants. Here, we could not confirm our hypothesis that tDCS and iTBS combined could increase WM performance. Interestingly, we found that iTBS resulted in faster reaction times. Additionally, headache, skin redness, and facial twitches differed among protocols.

Interestingly, previous studies found that the combination of tDCS and iTBS over the motor cortex could enhance motor cortical excitability results (Rubi-Fessen et al., 2015). However, the combination of different NIBS interventions over the DLPFC has been less investigated. In this context, our findings are in line with the results of a recent study that showed no superiority of the combination tDCS-iTBS, compared to iTBS alone, over the DLPFC, on stress response (De Smet et al., 2021). In turn, synergistic effects between tDCS and neurocognitive techniques (Dedoncker et al., 2021) have been previously reported. For instance, the combination of tDCS over the DLPFC with cognitive control training (CCT) and psychotherapy have shown larger effects compared with tDCS alone (Nasiri et al., 2020; Segrave et al., 2014). Differently from NIBS, psychological interventions can engage cortical target areas (Linden, 2006) via behavioral methods. These methods, applied together with tDCS, seem to modulate neural activity in a way that boosts the overall effects more easily (Dedoncker et al., 2021).

| 2-Back | Placebo | Combined Interventions | iTBS-only | tDCS-only |
|--------|---------|------------------------|-----------|-----------|
| % Accuracy | 0.85 (0.36) | 0.83 (0.38) | 0.82 (0.38) | 0.84 (0.37) |
| Reaction Time (in ms) | 638 (299) | 665 (308) | 618 (261) | 673 (323) |

iTBS: Intermittent theta-burst stimulation; tDCS: transcranial direct current stimulation.
Although iTBS and tDCS interventions were applied over the same target, the electric fields generated by each technique present particularities, such as different orientation and strength, which may result in the activation of different cortical areas (Suen et al., 2020; Zaidi et al., 2021). Moreover, we cannot exclude that other intervention protocols—such as cathodal tDCS or continuous TBS—could better improve the effects of each other. For instance, a previous study applying tDCS+iTBS over the motor cortex has shown that preconditioning the brain with tDCS can increase iTBS effects. However, while anodal tDCS significantly reversed the effects of cTBS towards a facilitation increase, it did not affect the iTBS effects (Hasan et al., 2012). Although this could be a rationale behind the non-significant effects of our study, future trials are warranted to investigate the neurobiological effects of this combination over the prefrontal cortex.

Our findings showed that iTBS, but not tDCS, led to significantly faster reaction time. Previous study has already suggested that iTBS can improve working memory performance of healthy volunteers (Hoy et al., 2016). However, to our knowledge, no study compared tDCS and TBS effects within the same design, although a meta-analysis previously suggested an overall greater effect for rTMS than tDCS in improving working memory performance (Brunoni & Vanderschuren, 2014). Possibly, iTBS enhances endogenous neuronal firing (Suppa et al., 2016) and produces more robust cortical excitability changes in the DLPFC and networks associated with cognition. Moreover, the target method location may have influenced the cognitive effects of iTBS, as recent meta-analyses showed that studies using neuronavigation were associated with higher cognitive performance (Beynel et al., 2019; Pabst et al., 2021).

In contrast, tDCS did not change WM performance in our study. In fact, tDCS findings on cognition have been mixed. For instance, Horvath et al. showed no significant cognitive effects for a single tDCS session vs. placebo in healthy

| Table 2 | Main results. |
|---------|---------------|
|         | Beta CI t p-value |
| Reaction Time |                    |
| tDCS vs. iTBS | 40.5 -9.45 71.5 1.27 0.2 |
| iTBS      | -21.5 -32.1 -10.9 -1.97 0.04 |
| tDCS      | 13.7 -1.8 25.5 1.1 0.26 |
| Accuracy |                    |
| tDCS vs. iTBS | 0.18 0.1 0.46 0.62 0.53 |
| iTBS      | -0.38 -0.58 -0.17 -1.81 0.06 |
| tDCS      | -0.28 -0.5 -0.1 -1.26 0.2 |

CI: Confidence interval (95%); iTBS: Intermittent theta-burst stimulation; tDCS: transcranial direct current stimulation.
subjects (Horvath et al., 2015). In a recent umbrella review, we found that tDCS could improve cognitive performance, but the quality of the included studies was poor (Farhat et al., 2020). Additionally, tDCS effects over the DLPFC might be non-linear. For instance, a study found that a current intensity of 1 mA but not 2 mA improved cognitive performance (Weller et al., 2020). Another issue is the electrode positioning and the corresponding induced electric field over brain areas. For instance, prefrontal, bilateral tDCS montages produce stronger electric fields in the medial prefrontal cortex than the commonly targeted DLPFC (Suen et al., 2020). In this context, it has been proposed that tDCS parameters should be personalized according to the targeted brain region to maximize the induced electric field (Wischnewski et al., 2019). Therefore, further studies investigating optimal tDCS parameters (i.e., position and size of electrodes, current intensity, and others) to increase working memory performance are encouraged.

Figure 3 Influence of session on the (A) reaction time and (B) accuracy performance. Note: Bars represent 95% Confidence Intervals based on 1000 bootstrap samples.

Table 3 Frequency of the adverse effects in each group.

| Adverse Effects                | Placebo (Sessions= 23) | Combined Interventions (Sessions = 23) | iTBS-only (Sessions = 23) | tDCS-only (Sessions = 24) | p-value |
|-------------------------------|------------------------|----------------------------------------|---------------------------|---------------------------|---------|
| Headache                      | 0 (0%)                 | 13 (10.9%)                             | 10 (8.7%)                 | 0 (0%)                    | 0.02    |
| Neck Pain                     | 2 (1.7%)               | 0 (0%)                                 | 3 (2.6%)                  | 4 (3.3%)                  | 0.75    |
| Local Pain                    | 0 (0%)                 | 0 (0%)                                 | 0 (0%)                    | 0 (0%)                    | -       |
| Itching                       | 29 (25.2%)             | 31 (25.8%)                             | 28 (24.3%)                | 31 (25.8%)                | 0.99    |
| Tingling                      | 29 (25.2%)             | 31 (25.8%)                             | 28 (24.3%)                | 31 (25.8%)                | 0.99    |
| Burning                       | 2 (1.7%)               | 8 (6.7%)                               | 2 (1.7%)                  | 6 (5.2%)                  | 0.47    |
| Skin Redness                  | 23 (20%)               | 42 (35%)                               | 20 (17.4%)                | 45 (37.5%)                | < 0.001 |
| Somnolence                    | 16 (13.9%)             | 12 (10%)                               | 10 (8.7%)                 | 25 (20.8%)                | 0.24    |
| Cognition Changes             | 3 (2.6%)               | 2 (1.7%)                               | 0 (0%)                    | 4 (3.3%)                  | 0.75    |
| Mood Changes                  | 3 (2.6%)               | 5 (4.2%)                               | 3 (2.6%)                  | 6 (5.2%)                  | 0.92    |
| Fatigue                       | 0 (0%)                 | 0 (0%)                                 | 0 (0%)                    | 0 (0%)                    | -       |
| Nausea                        | 0 (0%)                 | 0 (0%)                                 | 0 (0%)                    | 0 (0%)                    | -       |
| Facial Contraction            | 8 (6.9%)               | 23 (19.2%)                             | 29 (25.2%)                | 6 (5.2%)                  | 0.01    |
| Dizziness                     | 0 (0%)                 | 0 (0%)                                 | 0 (0%)                    | 0 (0%)                    | -       |

iTBS: Intermittent theta-burst stimulation; tDCS: transcranial direct current stimulation.
In our study, learning effects have probably occurred, as we found reaction time decreasing and accuracy increasing over time. This effect has been described in previous studies in the field (Curtin et al., 2019). Nonetheless, our randomized design and statistical analysis might have mitigated this issue.

Although tDCS and iTBS were not associated with any serious adverse event, we observed higher rates of skin redness, headache, and facial twitching after tDCS, the combined treatment, and iTBS, respectively. Such adverse events are consonant with previous studies in the field (Sudbrack-Oliveira et al., 2021; Oberman et al., 2011).

Some limitations should be underscored. First, the sample size was small, which might have increased the rate of false positive and false negative results. Second, accuracy rates were high. As our sample was composed by healthy, educated, and young participants, we probably should have used more challenging versions of the n-back test. Third, as different parameters of iTBS and tDCS produce distinct effects (Brunoni et al., 2019) our findings are limited to those used in our study. Fourth, blinding integrity was not assessed. Finally, in our statistical analysis, RT residuals were not normally distributed, an issue that might have decreased the power of some statistical analyses.

Conclusion

The combination of tDCS and iTBS over the DLPFC of healthy participants was tolerable and did not lead to additive or synergistic effects. Also, effects of iTBS, but not tDCS, indicated an improvement in reaction time, but not accuracy. Our findings encourage future trials evaluating other NIBS parameters and distinct outcomes, (Table 3).

Conflict of interest

The other authors have no conflicts of interest to disclose.

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Supplementary materials

Supplementary material associated with this article can be found, in the online version, at doi:10.1016/j.ijchp.2022.100334.

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