Continuous Theta-Burst Stimulation to the Right Dorsolateral Prefrontal Cortex May Increase Potentiated Startle in Healthy Individuals

Marta Teferi, Walid Makhoul, Zhi-De Deng, Desmond J. Oathes, Yvette Sheline, and Nicholas L. Balderston

ABSTRACT

BACKGROUND: Convergent neuroimaging and neuromodulation studies implicate the right dorsolateral prefrontal cortex (dPFC) as a key region involved in anxiety-cognition interactions. However, neuroimaging data are correlational, and neuromodulation studies often lack appropriate methodological controls. Accordingly, this work was designed to explore the role of right prefrontal cognitive control mechanisms in the expression/regulation of anxiety using continuous theta-burst transcranial magnetic stimulation (cTBS) and threat of unpredictable shock. Based on prior neuromodulation studies, we hypothesized that the right dPFC contributed to anxiety expression, and that cTBS should downregulate this expression.

METHODS: We measured potentiated startle and performance on the Sternberg working memory paradigm in 28 healthy participants before and after 4 sessions (600 pulses/session) of active or sham cTBS. Stimulation was individualized to the right dPFC site of maximal working memory–related activity and optimized using electric-field modeling.

RESULTS: Compared with sham cTBS, active cTBS, which is thought to induce long-term depression–like synaptic changes, increased startle during threat of shock, but the effect was similar for predictable and unpredictable threat. As a measure of target (dis)engagement, we also showed that active but not sham cTBS decreased accuracy on the Sternberg task.

CONCLUSIONS: Counter to our initial hypothesis, cTBS to the right dPFC made individuals more anxious, rather than less anxious. Although preliminary, these results are unlikely to be due to transient effects of the stimulation, because anxiety was measured 24 hours after cTBS. In addition, these results are unlikely to be due to off-target effects, because target disengagement was evident from the Sternberg performance data.

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While there are neuromodulation data targeting the right dIPFC in anxiety, these results are much less clear about whether this region positively or negatively affects anxiety expression. Repetitive transcranial magnetic stimulation (rTMS) is one common noninvasive neuromodulatory technique to causally manipulate neural activity (42). Different patterns of rTMS can be used to up- or downregulate cortical excitability and induce lasting changes in synaptic plasticity (43-46). rTMS tends to increase excitability at high frequencies (i.e., >5 Hz) and decrease excitability at low frequencies (47), although this is likely an oversimplification with exceptions depending on site, context, and other stimulation parameters. Similarly, patterned theta-burst stimulation (TBS) can induce long-term potentiation (LTP) (46) or long-term depression (LTD) (48) if delivered in an intermittent (TBS) (LTP-like changes) or continuous (cTBS) (LTD-like changes) pattern. Accordingly, if the right dIPFC contributes to the expression of anxiety, one might expect inhibitory stimulation patterns (i.e., low-frequency rTMS and cTBS) to reduce anxiety. In contrast, if the right dIPFC contributes to the regulation of anxiety, one might expect inhibitory stimulation patterns to increase anxiety. For simplicity, we define inhibitory as neuromodulation techniques that interfere with or downregulate ongoing processes in a region. However, we understand that this is an outdated and reductionistic conceptualization that equates long-term therapeutic effects to the induced transient states of neuronal excitability and suppression, which is almost certainly false. Accordingly, it is critical for studies to establish a behavioral measure of target engagement and to distinguish between acute and long-term effects.

As mentioned above, the results are mixed. Results in patients with depression with comorbid anxiety suggest that high-frequency stimulation to the left dIPFC followed by low-frequency stimulation to the right dIPFC can reduce anxiety symptoms (49), consistent with the expression hypothesis. In contrast, data in patients with posttraumatic stress disorder suggest that either 5 Hz (50) or iTBS (51) to the right dIPFC can reduce posttraumatic stress disorder symptoms, consistent with the regulation hypothesis. As for patients with generalized anxiety disorder, there have actually been very few randomized controlled trials targeting the right dIPFC with rTMS, and across studies, the data are inconclusive (42,52-54). In a previous study, we delivered within-session 10-Hz stimulation and found that this increased anxiety-potentiated startle (APS), which is consistent with the expression hypothesis, potentially offering a (preliminary) mechanistic explanation for the low-frequency results in patients with anxious depression (55). It is important to note as well that targeting approaches vary across studies, and many of these trials were based on trial and error-like modifications of depression protocols rather than mechanistic work in patients with anxiety or model systems.

To address these gaps in the literature, this work is designed to explore the role of right prefrontal cognitive control mechanisms in the expression/regulation of anxiety using TBS that should be observable across sessions. We targeted right dIPFC control circuits using the Sternberg WM paradigm (22,25,57,58). In addition, we measured performance on this task before and after stimulation as an index of target engagement. According to the expression hypothesis and the assumption that cTBS induces LTD-like changes in synaptic plasticity, we expected to see reductions in APS following active but not sham stimulation.

METHODS AND MATERIALS

Participants

A total of 34 right-handed participants between the ages of 18 and 50 were recruited from the Philadelphia metropolitan area to take part in this study. Exclusion criteria included current or past Axis I psychiatric disorder(s) as identified with the Structured Clinical Interview for DSM-IV, nonpatient edition (2), use of psychoactive medications, any significant medical or neurologic problems (e.g., cardiovascular illness, respiratory illness, neurologic illness, seizure), and any magnetic resonance imaging (MRI)/TMS contraindications (e.g., implanted metal, history of epilepsy or seizure). For a complete list, see: http://www.clinicaltrial.gov (Identifier: NCT03993509).

A total of 28 participants completed the study (21 females, 7 males, mean age = 26.61 years, SD = 7.04). Six consented subjects were excluded from the final sample (2 screen failures, 1 pilot subject, 3 subjects withdrew [2 due to scheduling, 1 withdrew during consent]). All participants signed an informed consent form, and the protocol was approved by the Institutional Review Board for human subject research at the University of Pennsylvania. The authors assert that all procedures contributing to this work comply with the ethical standards of the relevant national and institutional committees on human experimentation and with the Helsinki Declaration of 1975, as revised in 2008.

General Procedure

The basic procedure can be seen in Figure 1A. Subjects completed 8 study visits over the course of 4 weeks. During week 1, subjects completed an intake/pretest visit that included the consent, screening questionnaires, the no-shock, predictable-shock, unpredictable-shock (NPU) task, and the Sternberg task. They also completed a targeting session in the MRI scanner that included structural, resting-state, and task functional MRI runs. During weeks 2 and 4, subjects completed 2 days (2 sessions per day) of either active or sham cTBS. The order of the visits was counterbalanced across subjects. They also completed a post-cTBS testing session 24 hours after the final cTBS session that included the NPU and Sternberg WM tasks.

Consent Visit

Subjects began by completing the informed consent form. They then completed the MRI safety form, the TMS adult safety screen (59), a medical history questionnaire, a demographics questionnaire, the State-Trait Anxiety Inventory (60), the Beck Anxiety Inventory (61), the Montgomery–Åsberg Depression Rating Scale (62), and an eligibility checklist. Afterward, the study coordinator administered the Structured
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Clinical Interview for DSM-IV (3). Participants that met screening criteria then completed the prestimulation test visit procedure.

Test Visits

Test Visit Procedure. The coordinator began the test visit by cleaning and preparing the skin for electrode placement. Then, electrodes for the blink recording, electrodermal activity recording, and shock delivery were attached and tested. Next a startle habituation task was completed, followed by a shock workup procedure. Once this initial setup was complete, the subjects completed 2 runs of the NPU threat task and 2 runs of the Sternberg+threat WM task (additional methods in the Supplement).

NPU Task. During each test visit, subjects had 2 runs of the NPU task (Figure 1B). Each run consisted of alternating blocks of neutral (no shock), predictable (at risk for shock only during cue), and unpredictable (at risk for shock throughout) conditions (20,55,56). Predictable and unpredictable blocks were always separated by a neutral block to yield the following 2 block orders: NPNUNNP and NUNPNPNU. Subjects were informed of the contingencies before the task, and the block type was displayed at the top of the screen. Each block contained cue and intertrial interval (ITI) trials where a white noise probe was presented during the presence or absence of a visual cue. Cues were (8 s) simple colored (orange, teal, and purple) shapes (triangle, square, and pentagon), and the color and shape were varied across conditions. Each of the 4 neutral blocks had 2 trials per condition, while predictable (×2) and unpredictable (×2) blocks had 4 trials per condition, for a total of 8 trials per condition per run. Three shocks were presented during each run at random points during either the cue (predictable condition) or the ITI (unpredictable condition). Subjects rated their anxiety from 0 (not anxious) to 10 (extremely anxious) throughout the task using an onscreen numerical scale.

Sternberg+Threat WM Task. Following the NPU task, subjects completed 2 runs of the Sternberg+threat WM task (Figure 1C). The task consisted of a series of WM trials presented during safe (no shock) and threat (shock at any time) conditions. Each trial started with an instruction keyword to indicate the trial type. Next, subjects viewed a series of 5 letters presented sequentially. They retained them in WM for a brief interval and then gave a forced choice response during a subsequent response prompt. In maintain trials, subjects rehearsed the letters in the order that they were presented. In sort trials, subjects rearranged the letters in alphabetical order. When prompted with a letter/number combination, subjects indicated with a button press whether the position of the letter in the series matched the number.

Targeting Visit

Targeting Visit Procedure. Subjects arrived at the scanner and were cleared by the scanning technician or principal investigator to enter the scan room. They were given ear plugs, a button box, an emergency squeeze ball, and padding to minimize head movement. A pulse oximeter and respiration belt were also attached. Once setup was complete, structural scanning was completed from start to finish without intervention. Subjects then completed 1 run of the Sternberg WM task, followed by 2 resting-state runs (additional methods in the Supplement).

TMS Visits

TMS Visit Procedure. Subjects began the TMS visit by affirming their previous answers to the TMS adult safety screen and acknowledging any potential changes. The coordinator then secured the neuronavigation sensors using a swimcap and attached the e-stim electrodes. The subject was then registered to their MRI in Brainsight. On the first TMS visit, the subject’s resting motor threshold was obtained (specifications below). Next, the subject completed the remaining TMS visit procedures in the following order: Sternberg WM task (before stim run), cTBS (specifications below), and Sternberg WM task (after stim run). The subjects were given a 30-minute break and the TMS visit procedures were repeated. We chose to administer 2 sessions per day for 2 days based on evidence that multiple spaced cTBS applications lead to more robust changes in plasticity that are less susceptible to de-depression (63). We also chose not to deliver a longer cTBS session to avoid...
potential excitatory effects (64). We gave participants a 30-minute break between the sessions to yield a net ~50- to 60-minute temporal gap (gap = break + setup + 2 × Sternberg runs) between the cTBS trains, which has been shown to be optimal for inducing metaplasitcacy (65) (additional methods in the Supplement).

**Testing Session: NPU Task Anxiety Ratings and Startle.** Anxiety ratings at the time of each white noise presentation were extracted and averaged across trials. Likewise, electromyography data were processed, and startle magnitude was averaged across trials. For both ratings and startle, difference scores were calculated to correspond to fear (fear-potentiated startle [FPS]; predictable cue – predictable ITI), anxiety during the ITI (APS_ITI; unpredictable ITI – neutral ITI), and anxiety during the cue (APS_cue: unpredictable cue – neutral cue). A 2 (coil: active vs. sham) × 3 (trial type: FPS vs. APS_ITI vs. APS_cue) repeated-measures analysis of variance was conducted on these values.

**Testing Session: Sternberg Threat WM Performance.** Percent correct and reaction time were calculated for the sort and maintain trials during safe and threat blocks. WM-related effects were calculated by creating WM-related difference scores (sort – maintain). A 2 (coil: active vs. sham) × 2 (condition: safe vs. threat) repeated-measures analysis of variance was conducted on these difference scores. For all measures, outliers (i.e., values greater than 2 × SD) were truncated to 2 standard deviations from the mean (i.e., \(x|\mu ± 2 × SD| = M ± 2 × SD\)). Significant two-way interactions and multilevel one-way main effects were probed using post hoc paired-sample t tests.

**RESULTS**

**Targeting Session: Whole-Brain Blood Oxygen Level–Dependent Data**

Figure 2 shows the data used to individualize targeting across subjects. The sort > maintain contrast yielded activations in previously identified task-positive regions of dorsal attention and cognitive control regions including the dorsomedial prefrontal cortex, bilateral dlPFC, anterior insula, and posterior parietal cortex/intraparietal sulcus (Figure 3A, Table S1; see Table S2 for dlPFC target coordinates). It also yielded deactivations in previously identified task-negative regions of the default mode network including the ventromedial prefrontal cortex, hippocampus, and posterior cingulate cortex. These results are largely replications of established WM manipulation findings.

**Targeting Session: Performance and dlPFC Blood Oxygen Level–Dependent Data**

Results from the targeting session showed no significant difference in accuracy \((t_{27} = 1.89; p = .071; d = 0.38)\) (Figure 3B) or reaction time \((t_{27} = 0.48; p = .635; d = 0.1)\) (Figure 3C). However, consistent with our previous studies, accuracy was marginally better on maintenance trials than on sort trials. In contrast, sort trials evoked significantly greater blood oxygen level–dependent responses in the dlPFC target mask (\(t_{27} = 5.04; p < .001; d = 0.96\)) (Figure 3D), validating the current targeting approach.

**Testing Session: NPU Anxiety Ratings and Startle**

For ratings during the NPU task (Figure 4A; Table S3), there was a significant main effect for trial type \((F_{2,54} = 64.35; p < .001; \eta^2 = 0.7)\), but no main effect for coil (sham vs. active; \(F_{1,27} = 1.75; p = .2; \eta^2 = 0.06\)) and no coil × trial type interaction \((F_{2,54} = 1.47; p = .24; \eta^2 = 0.05\)). These effects are comparable when order of stimulation is included as a fixed factor. To characterize the significant main effect, we conducted pairwise post hoc t tests for the levels of trial type. FPS was significantly reduced compared with APS_cue \((t_{27} = -7.86; p < .001; Cohen’s d = -1.49)\) and APS_ITI \((t_{27} = 8.32; p < .001; Cohen’s d = 1.57)\), but APS_cue and APS_ITI were not significantly different from one another \((t_{27} = 0.07; p = .94; Cohen’s d = -0.01)\).

In contrast, for startle during the NPU task (Figure 4B and Table S3), there was a significant main effect for coil (active > sham; \(F_{1,27} = 7.08; p = .01; \eta^2 = 0.21\)) and trial type \((F_{2,54} = 18.49; p = 0; \eta^2 = 0.41)\), but no coil × trial type interaction \((F_{2,54} = 0.1; p = .9; \eta^2 = 0)\). These effects are comparable when order of stimulation is included as a fixed factor. To characterize the significant main effect of trial type, we conducted pairwise post hoc t tests for the levels of trial type. FPS was significantly larger compared with APS_cue \((t_{27} = 4.17; p < .001; Cohen’s d = 0.79)\) and APS_ITI \((t_{27} = -5.67; p < .001; Cohen’s d = -1.07)\), but APS_cue and APS_ITI were not significantly different from one another \((t_{27} = -1.39; p = .18; Cohen’s d = -0.26)\).
It should also be noted that when counterbalance was included in the model, there was a counterbalance × stimulation interaction ($F_{1,26} = 5.14; p = .032$). When probed further with paired-sample t tests, we found that the group that received active stimulation first showed a significantly larger startle response for the active than the sham condition ($t_{14} = 3.89; p = .002$; Cohen’s $d = 1.00$). However, this effect was not significant for the group that received sham stimulation first ($t_{12} = 0.269; p = .793$; Cohen’s $d = 0.07$). This pattern seems to be inconsistent with carryover effects, which would likely lead to a larger active versus sham effect for the group receiving sham stimulation first. The most likely explanation for these findings is that there was a tendency for startle to habituate over time. In the active-first counterbalance, this effect was additive with the stimulation effect. However, in the sham-first counterbalance, the stimulation and habituation effects were counter to each other.

Testing Session: Sternberg Threat WM Performance

For accuracy during the Sternberg threat WM sessions (Figure 5A), there was a significant main effect for threat ($F_{1,27} = 4.53; p = .04; \eta^2 = 0.14$), with larger WM-related differences (i.e., sort > maintain) in safe compared with threat conditions. There was also a significant main effect for coil ($F_{1,27} = 4.22; p = .05; \eta^2 = 0.14$), with larger WM-related differences following active compared with sham stimulation, suggesting that active stimulation reduced accuracy on sort trials (Table S4). However, there was no threat × coil interaction ($F_{1,27} = 0.73; p = .4; \eta^2 = 0.03$). These effects are comparable when order of stimulation is included as a fixed factor.

For reaction time (Figure 5B), there was a significant main effect for coil ($F_{1,27} = 5.37; p = .03; \eta^2 = 0.17$), with greater WM-related differences following sham compared with active stimulation. However, there was no significant threat main
effect ($F_{1,27} = 0.68; p = .42; \eta^{2} = 0.02$) or threat × coil interaction ($F_{1,27} = 0.01; p = .93; \eta^{2} = 0$). These effects are comparable when order of stimulation is included as a fixed factor.

**DISCUSSION**

Here, we examined anxiety after 4 sessions (600 pulses/session) of active or sham cTBS to the right dPFC. We used functional MRI during the Sternberg WM task to identify subject-specific stimulation sites (55), and electric-field modeling to optimize stimulation at those individualized sites (66). We found that active cTBS increased potentiated startle during both predictable and unpredictable threat compared with sham cTBS. However, there was no effect of stimulation type on anxiety ratings during the task, suggesting that these results were not driven by explicit expectations for how TMS should affect anxiety (i.e., placebo effects) (55). In addition, we have concurrent evidence from the Sternberg WM paradigm that suggests that performance during the sort trials of this task (i.e., WM manipulation trials) was reduced following active cTBS compared with sham cTBS, consistent with the hypothesis that cTBS induces LTD-like effects at the stimulation site (46). These results suggest that applying cTBS to the right dPFC can lead to increases in anxiety expression, which was counter to our initial hypothesis.

These results leave open two questions. The first question pertains to the role of the right dPFC in the expression/regulation of anxiety. While it is clear from neuroimaging data that both the left and right dPFC are activated during the manipulation of items in WM (11,25,67), it is unclear how this executive control relates to emotion regulation. In a recent study, we showed that left dPFC deficits in WM manipulation exhibited by patients with anxiety could be rescued by recruiting the right dPFC (25). Accordingly, this led to the hypothesis that both the left and right dPFC were specialized executive control centers, but that the domain of function differed across the hemispheres. While the left dPFC was specialized for verbal information (11,25,67), the right dPFC may be specialized for emotional content (9,15–19). In other words, our hypothesis is that the primary domain of function of the right dPFC is the flexible manipulation of emotional content in WM.

The second question pertains to the short- and long-term effects of different TMS protocols on right dPFC cognitive control circuits. It is hypothesized that cTBS induces LTD-like metaplastic effects at the site of stimulation (46) [see (68) for within-session effects]. Consistent with this hypothesis, we observed a performance deficit following active cTBS compared with sham cTBS on our Sternberg WM task. Accordingly, our results suggest that inducing LTD-like processes in right dPFC control circuits that are important for WM manipulation (10–14), which can make people more anxious.

Assuming that low-frequency rTMS should have a similar effect (47), we might expect low-frequency rTMS to increase anxiety rather than decrease anxiety. However, there are some inconsistencies in the literature suggesting that excitatory stimulation to the right dPFC can increase anxiety symptoms, while inhibitory stimulation to the right dPFC can decrease anxiety symptoms, supported by early research into the effects of high (assumed to be excitatory) and low (assumed to be inhibitory) frequency rTMS on mood/anxiety (69). While there have been some preliminary studies in both patients with primary generalized anxiety disorder (70–73) and patients with generalized anxiety disorder/major depressive disorder (49) to support the low-frequency hypothesis, these studies have small sample sizes and lack of adequate control conditions (74,75). In addition, there are counterexamples suggesting that high-frequency right dPFC stimulation can reduce anxiety symptoms as well (76), and there is no mechanistic explanation that can sufficiently explain this pattern of results. While we are not questioning the efficacy of 1-Hz stimulation, we instead suggest that the mechanism of action is unlikely due to downregulation of plasticity at the stimulation site. Indeed, there is some evidence to suggest that low-frequency stimulation is not sufficient to induce observable metaplastic effects outside the window for transient effects on excitability (77). While it may be tempting to equate cTBS with low-frequency stimulation and begin using right dPFC cTBS in place of the longer 1-Hz protocol to treat anxiety, current results suggest that such an approach might not yield favorable clinical outcomes. However, because this study was an exploratory, preclinical study on a small group of healthy volunteers, it would be premature to use the findings to anticipate clinical outcomes in patients with anxiety.

These results are also seemingly inconsistent with our previous work showing that 10 Hz to the stimulation site increases anxiety within session. Aside from the stimulation pattern, the primary difference between these studies is the interval between the stimulation and the poststimulation anxiety test. In the 10-Hz study, we measured anxiety immediately following a single course of 10-Hz stimulation (55). In contrast, in this cTBS study, we measured anxiety 24 hours following...
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four 600-pulse sessions of stimulation. Critically, this difference means that in the cTBS study, our results are unlikely to be due to acute, transient fluctuations in excitability. Instead, these results are likely driven by long-term changes in synaptic plasticity (46). Likewise, our Sternberg results showing decreased WM manipulation performance following active compared with sham stimulation confirm the predicted cTBS target (dis)engagement, potentially ruling out the possibility of paradoxical excitatory cTBS effects.

While most therapeutic clinical neuromodulation trials measure their effects at longer intervals after repeated sessions (offline stimulation), many mechanistic studies measure their effects at shorter intervals during a single session (online stimulation). Accordingly, it may be possible to explain both the 1-Hz and the 10-Hz findings based on distinct effects of online and offline stimulation. In the case of the 1-Hz stimulation, we have already shown that patients with anxiety have deficits in WM manipulation processing in the left dlPFC (25). Although pure speculation, perhaps temporarily decreasing excitability at the stimulation site is non-specific to the left dlPFC to compensate, leading to increased processing efficiency over time. Consistent with this hypothesis, 1-Hz stimulation to the right dlPFC seems to be most effective as an add-on to 10-Hz stimulation to the left dlPFC (49,70). Based on this hypothesis, one might also expect improved WM performance and increased WM manipulation-related left dlPFC activity following a therapeutic course of 1-Hz stimulation to the right dlPFC. In the case of the 10-Hz stimulation (55), it is possible that the increase in excitability at the stimulation site is non-specific. According to our right dlPFC emotional WM hypothesis, this non-specific increase in cortical excitability may actually interfere with the pattern-specific activity needed to flexibly manipulate emotional content in WM (10–14). If this is the case, one might expect different effects for online and offline stimulation. Specifically, one might hypothesize offline 10-Hz stimulation to strengthen right dlPFC cognitive control circuits, which would lead to better regulation of, and thus decreases in, anxiety. In contrast, one might expect non-specific transient increases in excitability induced by online 10-Hz stimulation to potentially interfere with the specific patterns of right dlPFC activity needed for manipulation of emotional content in WM, which would lead to transient impairments in regulation and transient increases in anxiety similar to the ones we observed in our previous work.

Broader Implications

Although it is common practice in the neuromodulation literature to base a study’s rationale, hypotheses, and design on hypotheses about the excitatory and inhibitory properties of rTMS/TBS, much of the data supporting these assumptions were derived from motor cortical conditioning studies using motor-evoked potentials as a stand-in for excitability (78–82). There have been few neuromodulatory studies showing a clear generalization of these properties specifically to prefrontal areas. Accordingly, the generalization of assumptions from motor cortex to prefrontal cortex could be questionable and should be critically evaluated and discussed if it forms the basis of a study. In addition, studies should be designed with reliable behavioral indicators of target engagement whenever possible.

Likewise, many of these cortical conditioning studies were measured within session (78), with little data to suggest that these transient states of neuronal excitability and suppression can account for the longer-term changes in synaptic plasticity driving the therapeutic effects of most neuromodulatory treatments. We believe that synaptic plasticity is the key to understanding the long-term effects, and thus the therapeutic impact, of neuromodulatory treatments. There is an extensive literature showing that changes at the synaptic level undergo an active consolidation process that includes the synthesis of new proteins (83), degradation of old proteins (84), and remodeling of the synapse (85), a collection of processes that can last several hours (86). Accordingly, we believe that it is critical to evaluate the performance of a potential neuromodulatory treatment at intervals outside the window for transient increases in excitability.

Limitations

Despite the strengths of the study (see Supplemental Discussion), the following limitations should be noted. First, the results were counter to our hypotheses. Although not technically a limitation, these data need to be replicated in an independent sample. Another limitation is that we included a single baseline visit rather than a within-week baseline visit for the NPU paradigm, which would have provided a more flexible baseline that could have potentially accounted for any plasticity effects related to order of administration. Although counterbalancing should control for this, it could be argued that a baseline closer in temporal proximity to the cTBS/sham would have been preferable.

Conclusions

Here, we measured fear and anxiety following active or sham cTBS to the right dlPFC and found that active cTBS increases both fear and anxiety. Results are consistent with a role for the right dlPFC in anxiety regulation but require replication. This is important because it is a potential first step toward understanding the mechanism of action of neuromodulatory treatments for anxiety aimed at the prefrontal cortex. Future research should examine how other types of stimulation paradigms (high-frequency rTMS, low-frequency rTMS, iTBS, sequential bilateral cTBS + iTBS, etc.) affect fear and anxiety. In addition, despite the translational nature of the threat paradigm used (87–93), these stimulation paradigms should be tested in patients with clinical anxiety. Finally, our results highlight the importance of studying TMS-related effects outside of the acute administration window.

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