Intermittent Theta-Burst Stimulation Transcranial Magnetic Stimulation Increases GABA in the Medial Prefrontal Cortex: A Preliminary Sham-Controlled Magnetic Resonance Spectroscopy Study in Acute Bipolar Depression

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Background: Magnetic resonance spectroscopy (MRS) has been used to identify gamma-aminobutyric acid (GABA) alterations in mood disorders, particularly in the medial prefrontal cortex (mPFC) where decreased concentrations have been associated with anhedonia. In major depressive disorder (MDD), prior work suggests that repetitive transcranial magnetic stimulation (rTMS) increases mPFC GABA concentrations proportional to antidepressant response. To our knowledge, this has not been examined in acute bipolar depression.

Methods: As part of a multicentre 4-week randomized, double-blind, sham-controlled trial using intermittent theta-burst stimulation (iTBS) of the left dorsolateral prefrontal cortex (DLPFC) in individuals with acute bipolar depression, we quantified mPFC GABA and Glx (glutamate+glutamine) concentrations using a 3T MRS scan at baseline and after the intervention. Depressive symptoms were measured using the Montgomery-Asberg Depression Rating Scale (MADRS) and the Hamilton Depression Rating Scale-17 (HRDS-17), and anhedonia was measured using the Snaith-Hamilton Pleasure Scale (SHAPS).

Results: The trial was terminated for futility and magnetic resonance spectroscopy data was acquired for 18 participants. At baseline, there were no associations between GABA or Glx concentrations and anhedonia, however GABA was negative correlated with depressive symptom severity on the HRDS-17. Compared to the sham-iTBS group, participants receiving active-iTBS had a significant increase in mPFC GABA concentrations. This was unrelated to antidepressant outcomes or improvements in anhedonia.
**INTRODUCTION**

Bipolar disorder (BD) is a leading cause of global disability that is defined by episodes of mania or hypomania (1). Yet, patients with BD type I spend as much as 70% of the time of the symptomatic periods experiencing syndromic or sub-syndromic depressive symptoms, while this proportion is over 80% in patients with BD type II (2). There are currently only four treatments that have been approved by the US Food and Drug Administration for the treatment of acute bipolar depression, namely olanzapine and fluoxetine combination, quetiapine, lurasidone, and cariprazine, and limited alternatives for those who do not respond to or tolerate these treatments (3).

Non-invasive neurostimulation techniques, such as transcranial magnetic stimulation (TMS), are efficacious in the treatment of Major Depressive Disorder (MDD). Accordingly, they are often considered in acute bipolar depression, however the literature in BD is small (4, 5). Newer TMS protocols, such as intermittent theta-burst stimulation (iTBS), are efficacious in MDD (6, 7) but have not demonstrated efficacy in BD to date (8, 9). Given lack of efficacy in BD, this may provide an opportunity to dissociate physiological effects that are and are not treatment mechanisms.

A growing body of research highlights alterations in brain gamma-aminobutyric acid (GABA) and glutamate systems in mood disorders (10, 11). $^1$H-MRS studies report decreases in cortical GABA in unipolar depression that correlate inversely with glutamate (12–15). Lower GABA levels have been repeatedly seen in the medial prefrontal cortex (mPFC) in depression, and this has been associated with anhedonia (16–18). Importantly, mPFC GABA levels increase with successful pharmacological treatment of MDD (19), as well as with repetitive transcranial magnetic stimulation (rTMS) treatment (20).

Several lines of evidence highlight the importance of the mPFC, and the anterior cingulate cortex more specifically, in the therapeutic effects of rTMS when the target is the dorsolateral prefrontal cortex (DLPFC). Indeed, a single treatment to the left DLPFC is associated with sustained changes in fMRI BOLD response and the GABA/Glx ratio in the anterior cingulate (21). Hyperactivity of the anterior cingulate predicts successful rTMS treatment (22), and clinical response to rTMS is associated with the degree of anticorrelated activity between the precise DLPFC target and the anterior cingulate cortex (23–25). In light of these findings, GABA concentrations in the medial prefrontal cortex (mPFC) may represent a physiological marker associated with large scale network changes in depression (26) that are amenable to rTMS.

**Conclusion:** Our data suggests that iTBS targeting the DLPFC is associated with physiological changes in the mPFC. In acute bipolar depression, our preliminary data suggests that mPFC GABA is dissociated from antidepressant iTBS treatment outcomes and anhedonia.

**Keywords:** repetitive transcranial magnetic stimulation, intermittent theta burst stimulation, bipolar disorder, bipolar depression, magnetic resonance spectroscopy, gamma-aminobutyric acid

To our knowledge, mPFC GABA concentrations during rTMS treatment in acute bipolar depression have not been measured. Therefore, in a randomized sham-controlled trial that examined the efficacy of iTBS in comparison to sham-iTBS (9), we used magnetic resonance spectroscopy (MRS) to assess mPFC GABA in the study patients. We hypothesized that mPFC GABA would be related to anhedonia, would increase with active-iTBS but not sham-iTBS, and that the change would be related to antidepressant response.

**METHODS**

We conducted a double-blind randomized controlled study in two Canadian centers (the University of British Columbia and University of Calgary) between October 2016 and March 2020. The study was registered with ClinicalTrials.gov (NCT02749006) and was approved by the clinical ethical review boards of the University of British Columbia and conjoint health ethical review board of the University of Calgary. The University of Calgary site, but not the University of British Columbia site, acquired GABA-edited MEGA-PRESS data in the mPFC.

The primary outcomes are described elsewhere (9). Briefly, the trial was terminated for futility after 37 participants were randomized due to the absence of an efficacy signal and overall low rates of clinical response (15.7% sham-iTBS vs. 16.6% active-iTBS). This low rate of clinical response was also observed after a 4-week open-label continuation for those that had not responded in the double-blind phase (23.8%).

**Participants**

Participants were recruited by referral or by online and community advertisements. Participants provided written informed consent. Eligibility criteria included: males and females 18–70 years of age with a primary diagnosis of Bipolar I Disorder or Bipolar II (BD1 or BD2), experiencing a major depressive episode with ≥15 on the 17-item Hamilton Depression Rating Scale (HRDS-17) (27), and having failed to achieve clinical response with CANMAT first-line recommended treatments (lithium, lamotrigine, quetiapine, lurasidone with or without concurrent lithium or valproate) for an acute major depressive episode (3). Participants were required to have been on a stable pharmacological regimen for 2 weeks prior to screening that had to include a mood stabilizer (lithium >0.6 mEq/L or valproate >350 mM/L), an atypical antipsychotic, or a combination of a mood stabilizer and an atypical antipsychotic. For participants with BD2 only, lamotrigine monotherapy was acceptable if the dose was >100 mg daily.
Clinical Measures and Self-Reports
Participants were assessed by an independent evaluator (AM) blinded to treatment conditions. The diagnosis of BD was confirmed with the MINI International Neuropsychiatric Interview 7.0. Clinician rated instruments included the Hamilton Rating Depression Scale 17-item (HRDS-17), the Montgomery-Asberg Depression Rating Scale (MADRS) (30), the Young Mania Rating Scale (YMRS) (31), and the Clinical Global Impression (CGI) subscales for severity and improvement (32) at baseline, after 2 weeks, and after 4 weeks of double-blind treatment. Clinical response was defined as a reduction of 50% or more in MADRS score, and clinical remission was defined as a MADRS score of 12 or less. To measure anhedonia, participants completed the Snaith-Hamilton Pleasure Scale (SHAPS) (33) at baseline and at the conclusion of the double-blind phase.

GABA-MRS
Imaging was performed at 3 T (MR750, General Electric Healthcare, USA). Because the GABA peaks are overlapped by more abundant metabolite peaks, advanced spectroscopy methods are required for its accurate quantification; the most common being J-difference editing (34). For voxel localization and subsequent segmentation, first a T1-weighted acquisition was performed (TR/TE = 8.3/3.2 ms, 1mm isotropic voxels, aligned to the AC-PC line). Gamma-aminobutyric acid data were acquired from a 30 × 30 × 30 mm³ voxel in the mPFC using a GABA-edited MEGA-PRESS acquisition (TR/TE = 2 s/68 ms, 320 averages, 14 ms editing pulses alternating between 1.9 and 7.46 ppm every two averages, and eight unsuppressed averages for metabolite quantification). Voxels were then placed parallel to the AC-PC line anterior to the genu of the corpus callosum (Figure 1). Gamma-aminobutyric acid and the co-edited Glx (glutamate+glutamine) peak were analyzed in Gannet (35) including voxel localization and segmentation with SPM12, tissue correction for tissue specific T1 and T2 relaxation, and water visibilities (36, 37). Gamma-aminobutyric acid data also included tissue correction for concentration differences of GABA in white and gray matter, assuming the concentration of GABA in gray matter is twice that of white matter (38).

Data quality was visually assessed, which included assessing the GABA peak and resolution of subtraction artifact following spectral alignment, confirming the stability of water frequency (indicative of movement or scanner drift), confirming the quality of creatine alignment and fit of the spectrum and residuals. The included MEGA-PRESS acquisitions had a GABA fit error of 7.74 ± 3.28, full-width half-maximum of 22.03 ± 4.48 Hz, and signal to noise ratio of 11.97 ± 3.05.

Voxel Placement Consistency
The T1-weighted image from the follow-up session was registered to the baseline acquisition [FSL FLIRT (39, 40)] and the transformation matrix for that registration was then applied to the follow-up voxel such that both voxels for each subject could be visualized in the same space to ensure voxel placement consistency. In addition to visual confirmation of voxel placement consistency across sessions, the dice coefficient...
of the voxel overlap was calculated. The average dice coefficient was 72% (standard deviation = 10%).

**Statistical Analysis**

Statistical analysis was performed in SPSS (IBM SPSS Statistics Version 26, IBM). Data normality was confirmed using the Shapiro-Wilk test and outliers were identified using Tukey’s Fences (41). Cross-sectional analyses of continuous data were performed using Student’s t-test or Mann Whitney U Test. Chi-square was used for dichotomized variables. One way ANOVA and linear regression was used to explore relationships between mPFC GABA and clinical characteristics. Change in mPFC GABA was quantified with repeated measures ANOVA. Two tailed significance was set at $\alpha < 0.05$.

**RESULTS**

The progress of participants through the University of Calgary site within this multisite study is described in the CONSORT flow diagram (Figure 2). Twenty-one participants were randomized between sham-iTBS ($n = 10$) and active iTBS-rTMS groups ($n = 11$). Three individuals did not tolerate the MRI, and therefore we acquired valid GABA data at baseline from 18 participants ($n = 7$ sham-iTBS and $n = 11$ active-iTBS). Sociodemographic and clinical characteristics were comparable between groups at baseline, and treatment outcomes also did not differ (Table 1).

mPFC GABA concentrations at baseline were skewed with one notable outlier >3 standard deviations from the mean. When this participant was removed, this data was normally distributed. There were no significant baseline differences in GABA levels between sham-iTBS and active-iTBS groups [$n = 7$ sham-iTBS $2.13 \pm 0.36$ mmol vs. $n = 10$ active-iTBS $1.96 \pm 0.45$ mmol $t_{(15)} = -0.84, p = 0.41$]. mPFC Glx concentrations were normally distributed and there was a trend toward higher Glx levels at baseline in the sham-iTBS group [$n = 7$ sham-iTBS $11.20 \pm 2.53$ mmol/kg vs. $n = 10$ active-iTBS $9.41 \pm 0.94$ mmol $t_{(15)} = -2.06, p = 0.057$].

At baseline, there were no significant associations between GABA and SHAPS score [$F_{(1,15)} = 0.07, p = 0.79$], medication regimen [$F_{(2,16)} = 0.02, p = 0.97$], MADRS score [$F_{(1,16)} = 2.41, p = 0.14$], or CGI-severity score [$F_{(1,15)} = 1.53, p = 0.23$], however there was a significant association with HRDS-17 scores [Figure 3A; $F_{(1,16)} = 5.18, p = 0.038$, Standardized $B = -0.50$]. mPFC Glx did not significantly correlate with any clinical characteristic ($p > 0.18$).
TABLE 1 | Sample characteristics and clinical outcomes.

|                        | Sham-iTBS          | Active-iTBS        | Statistic    | p    |
|------------------------|--------------------|--------------------|--------------|------|
| n (%) or M ± SD        |                    |                    |              |      |
| **Age**                | 42.20 ± 13.32      | 45.27 ± 14.52      | t(19) = 0.50 | 0.62 |
| Female sex             | 5 (50.0%)          | 5 (45.5%)          | χ² = 0.04    | 0.83 |
| Education (years)      | 16.00 ± 2.16       | 16.64 ± 3.72       | t(19) = 0.47 | 0.64 |
| Diagnosis              |                    |                    | χ² = 0.11    | 0.73 |
| BD1                    | 2 (28.6%)          | 4 (38.4%)          |              |      |
| BD2                    | 5 (71.4%)          | 7 (83.6%)          |              |      |
| Medication stratification |                |                    | χ² = 0.06    | 0.96 |
| Mood stabilizer        | 4 (40.0%)          | 5 (45.5%)          |              |      |
| Atypical antipsychotic | 1 (10.0%)          | 1 (9.1%)           |              |      |
| Mood stabilizer + atypical antipsychotic | 5 (50.0%) | 5 (45.5%) |              |      |
| **Baseline characteristics** |            |                    |              |      |
| HRDS-17                | 20.50 ± 3.30       | 22.54 ± 3.88       | t(19) = 1.29 | 0.21 |
| MADRS                  | 31.60 ± 3.89       | 32.90 ± 4.30       | t(19) = 0.72 | 0.47 |
| CGI-Severity           | 4.20 ± 0.42        | 4.70 ± 0.94        | t(19) = 1.52 | 0.14 |
| SHAPS                  | 11.00 ± 3.09       | 10.30 ± 4.00       | t(18) = −0.43 | 0.66 |
| **Treatment outcomes** |                    |                    |              |      |
| Percent MADRS improvement at 4 weeks | 40.33 ± 34.93 | 24.69 ± 36.48 | t(15) = −0.90 | 0.38 |
| Clinical response      | 3 (33.3%)          | 1 (12.5%)          | χ² = 1.02    | 0.31 |
| Clinical remission     | 3 (33.3%)          | 1 (12.5%)          | χ² = 1.02    | 0.31 |

**FIGURE 3** | (A) Baseline mPFC GABA concentrations were negative associated with HRDS-17 scores. (B) Active-iTBS, but not sham-iTBS, was associated with an increase in mPFC GABA concentrations after 4 weeks of treatment. *time interaction p < 0.05.

At the conclusion of double-blind treatment, paired pre-post MRS data were available for n = 6 sham-iTBS and n = 6 active-iTBS participants. We observed a significant increase in mPFC GABA in the active-iTBS group, but not the sham-iTBS group [Figure 3B; Time F(1, 10) = 4.65, p = 0.056, Time * Group F(1, 10) = 6.99, p = 0.025]. This significant effect persisted after controlling for percent change in MADRS scores [Time F(1, 9) = 0.69, p = 0.42, Time * %MADRS F(1, 9) = 1.51, p = 0.24, Time * Group F(1, 9) = 8.68, p = 0.016]. We found no evidence for mPFC Glx time or treatment group effects [Time F(1, 9) = 0.00, p = 0.99, Time * Group F(1, 9) = 0.00, p = 0.98].

**DISCUSSION**

To our knowledge, this is the first study to use GABA-edited MRS to investigate mPFC metabolite changes during TMS therapy in acute bipolar depression. Though preliminary, our data suggests iTBS results in increased GABA in the mPFC, consistent with previous data in MDD samples for the mPFC (20) and in the DLPFC (42). However, contrary to both of these rTMS studies, and others using pharmacological interventions (19), the GABA effect we observed was dissociated from antidepressant outcomes. We also failed to observe an association between mPFC GABA and anhedonia.
This data is interesting because it confirms a common physiological effect of DLPFC TMS stimulation in different pathologies. Yet, the lack of antidepressant effect in this subsample or in the larger multisite trial from which it is drawn highlights that not all physiological effects of therapeutic TMS are causally associated with improvement in depressive symptoms. Moreover, we did not observe associations between mPFC GABA and clinical features such as anhedonia, a replicated finding in MDD (18). While it is possible that the physiological effect we observe precedes symptomatic improvement, the 4 week open label extension of this trial for non-responders also had a low rate of clinical response (9).

Given the small sample and inference from a null finding, further research is required to determine whether our findings represent a physiological difference between acute bipolar depression and unipolar depression. It is possible that MRS quantified GABA in the mPFC is not as tightly linked to mood state in BD, given that previous studies in BD have not found differences compared to healthy controls in either depression (43) or euthymia (44, 45), and this has also been suggested in meta-analyses (46). Post-mortem staining for GABAergic subpopulations in the anterior cingulate cortex indicate heterogeneity between BD and MDD, however this is principally for calbindin positive neurons and appears layer specific and therefore cannot be resolved with MRS methods (47). Alternatively, concomitant medications indicated for acute and prophylactic treatment in BD may confound MRS GABA measures. Indeed, mood stabilizers modulate GABA (48–50) and lithium modulates Glx (51), and all but one participant was treated with these medications.

**LIMITATIONS**

Our preliminary data comes from a single site from a randomized controlled trial that was terminated for futility. As a result, the sample size is small, and the availability of paired pre-post MRS data is further reduced. As such, the magnitude of change seen in GABA concentrations in the active-iTBS group may be inflated due to small sample size bias. Nevertheless, this increase is not observed in the sham-iTBS group, and therefore is likely to be a physiological effect of iTBS. Larger studies are required to more reliably determine the effect size, and independence from clinical improvements in acute bipolar depression. Participants in this trial were required to have stable dosing of an antimanic agent, which resulted in 20 of the 21 participants using medications known to influence GABA (48–50). While an unmedicated sample would resolve this limitation, it represents a safety concern as even with antimanic agents in this sample a treatment emergent affective switch occurred after a single active-iTBS treatment, and another participant experienced a treatment emergent affective switch within eight treatments after transitioning from sham-iTBS to open-label iTBS.

There is an inherent challenge in resolving GABA at 3 T due to its low concentration and overlapped, more abundant metabolites. The use of editing techniques, such as the MEGA-PRESS sequence used in this study, is preferable to the use of short-echo point resolved spectroscopy sequences for measuring GABA, particularly at 3 T (52–54). Nonetheless, there are some limitations and caveats worthy of discussion. Given the low signal and the J-difference editing approach used in MEGA-PRESS, a large voxel and many signal averages are required to obtain high signal-to-noise data for a reliable GABA measurements (53). The voxel used in the present study was 27 cm$^3$, placed in the mPFC. This large voxel limits the regional specificity of our results as it encompasses multiple functionally or anatomically distinct structures. Secondly, the measured GABA signal at 3 ppm includes a macromolecule contribution due to the limited specificity of the editing pulses applied at 1.9 ppm (55, 56). Macromolecules are typically considered functionally irrelevant, though the macromolecular signal is variable between subjects (53, 57). Finally, while frequency drift is known to affect efficiency of the editing pulses, the level of macromolecular signal contribution and can cause subtraction artifacts (58), the scanner used in this study is highly stable and no included data showed a frequency drift of >0.1 ppm.

**CONCLUSION**

Our sham-controlled data in acute bipolar depression suggests that iTBS targeting the DLPFC is associated with physiological changes in the mPFC. Unlike previous reports in MDD, in acute bipolar depression mPFC GABA concentrations appear to be dissociated from antidepressant treatment outcomes and anhedonia or have a much smaller effect. Future research is needed to better understand the clinical implications and network level changes associated with neural metabolites in acute bipolar depression.

**DATA AVAILABILITY STATEMENT**

The original contributions presented in the study are included in the article/supplementary material, further inquiries can be directed to the corresponding author/s.

**ETHICS STATEMENT**

The studies involving human participants were reviewed and approved by University of Calgary Conjoint Health Research Ethics Board. The patients/participants provided their written informed consent to participate in this study.

**AUTHOR CONTRIBUTIONS**

Study conception: AM, IY, and AH; Securing study funding: IY; Data collection: JC and AM; Data analyses: CD, MD, AH, and AM; Figure generation: AH and AM; Drafting manuscript: CD; Editing and approval of final manuscript: MD, JC, LY, AH, and AM. All authors contributed to the article and approved the submitted version.

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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