The Impact of High-Frequency Repetitive Transcranial Magnetic Stimulation on Fine Motor Functions in Medication-Resistant Major Depression

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Key Words
Transcranial magnetic stimulation • High-frequency repetitive transcranial magnetic stimulation • Psychomotor symptoms • Retardation • Major depressive disorder

Abstract

Objectives: Although high-frequency repetitive transcranial magnetic stimulation (HF-rTMS) over the left dorsolateral prefrontal cortex (DLPFC) has been reported to improve mood symptoms in major depressive disorder (MDD), research on its impact on psychomotor symptoms is scarce. This study assessed the psychomotor effects of 1 and 10 sessions, respectively, of HF-rTMS over the left DLPFC. Methods: Ten HF-rTMS sessions were applied in 21 medication-free MDD patients over a 2-week period. At the beginning, one placebo (sham)-controlled rTMS session was also applied in a cross-over, single-blind design. Psychomotor variables were digitally recorded during completion of a Fitts’ task, at baseline, after the first and second real/sham session and at the end point. Results: The total 10-session treatment period resulted in a decrease of depression severity. One HF-rTMS session resulted in improvements on the Fitts’ task, without a difference between active and sham stimulation, however. No further improvements occurred from session 2 to session 10. Conclusions: No evidence was provided to link the observed psychomotor improvements to HF-rTMS stimulation, as a practice effect could have impacted the significant psychomotor outcomes.

Introduction

In recent years, studies investigating the therapeutic efficacy of repetitive transcranial magnetic stimulation (rTMS) for major depressive disorder (MDD) have exponentially increased [1–3]. A growing body of evidence has been published indicating that depressed patients can be successfully treated with high-frequency (HF)-rTMS when this is administered on the left dorsolateral prefrontal cortex (DLPFC) [4–6]. Recent meta-analytic studies support the antidepressant efficacy of this technique in treatment-resistant depressed patients when stimulation periods are long enough (e.g. >2 weeks) [2, 3]. Besides the treatment duration, the frequency and which side is...
stimulated should also be taken into account when evaluating the efficacy of rTMS [3].

Most rTMS studies focus on the reduction in mood symptoms in MDD, but the effects of rTMS on psychomotor functioning have been rarely explored.

Notwithstanding, cognitive dysfunctions and psychomotor retardation have also been determined as core features of episodes of MDD [7–9]. Psychomotor slowing appears to be a strong diagnostic marker for MDD with melancholic features. As psychomotor retardation is one of the key symptoms of the melancholic subtype of depression, it would be very appropriate to investigate psychomotor changes following rTMS treatment in a cohort of melancholic patients. From a neurobiological perspective, psychomotor retardation in major depression – and especially the melancholic subtype – has been linked to a hypodopaminergic state [7, 9]. Moreover, prefrontal rTMS has been found to influence striatal dopaminergic activity [10, 11]. Furthermore, a higher level of psychomotor retardation has been associated with reduced metabolic activities in the left DLPFC suggesting an important role of this cortical area on psychomotor functioning [9]. Therefore, given the possible neurobiological and clinical implications, a thorough investigation of the psychomotor effects of rTMS in MDD is warranted.

A limited number of studies have already examined the neurocognitive effects of rTMS in MDD [12, 13], but even fewer studies have examined the impact of rTMS on psychomotor performance in MDD. Applying the motor agitation and retardation scale (MARS), Hoeppner et al. [14] observed a reduction in psychomotor retardation following 10 HF-rTMS sessions over the left DLPCF. They could only demonstrate a trend in the reduction of psychomotor agitation, but not for retardation in MDD following 15 left prefrontal HF-rTMS sessions [15]. Finally, Baeken et al. [16] demonstrated the positive effect of 10 sessions of HF-rTMS on psychomotor slowing as measured by means of the Salpêtrière Retardation Rating Scale (SRRS).

Given the limited number of studies on this subject and their divergent results, this study further investigated the psychomotor effects of HF-rTMS in MDD, applying the Fitts’ task. This computerized task is an objective and reliable method to assess fine motor activity, and is generally considered to be a rater-independent and more objective measurement method than the more subjective rating scales [9]. The Fitts’ task has been widely used in the research into psychomotor symptoms in MDD [17–19]. This fine motor task requires a precise sensorimotor programming, initiation and execution of the muscle commands [17–19]. One HF-rTMS session with parameters comparable to rTMS treatment for depression has been reported to affect this psychomotor functioning in healthy subjects [20].

Consequently, this study aims to further explore the psychomotor effects of HF-rTMS over the left DLPFC in a sample of medication-resistant MDD patients, applying the Fitts’ task. To evaluate the effect of a single stimulation on psychomotor functioning, we assessed the effects of one sham-controlled session of HF-rTMS delivered on the left DLPCF, in a single-blind placebo controlled cross-over design. To examine the effect of HF-rTMS treatment, the effect on the Fitts’ task was assessed after 10 such sessions spread over a period of 2 weeks of treatment. To ascertain that the melancholic depressed patients displayed decreased psychomotor speed, their baseline Fitts’ measurements were compared to an age- and gender-matched control group.

As mentioned above, few studies on psychomotor effects in MDD have been executed up to now, psychomotor retardation has been associated with DLPFC hypofunction and a hypodopaminergic state and rTMS has been supposed to exert an impact on the dopaminergic system. Together with the limited existing evidence mentioned above, all these findings led us to hypothesize that an improvement in psychomotor functioning could be expected after HF-rTMS treatment, particularly in our treatment responder group.

Methods

Patients

Our group consisted of 21 medication-free unipolar depressed patients of the melancholic subtype (female: male = 13:8; age 44.7 ± 10.3 years). Psychiatric disorders were assessed using the Mini-International Neuropsychiatric Interview [21]. Because comorbid personality disorders were not part of the exclusion criteria no formal diagnostic screening on axis II diagnosis was performed. Severity of depression was assessed with the 21-item Beck Depression Inventory (BDI) [22] and the 17-item Hamilton Depression Scale (HADS) [23]. The HADS was administered by an experienced psychiatrist, not related to the study. Eleven participants were current in-patients during HF-rTMS treatment. Treatment resistance was assessed with criteria according to Thase and Rush [24]. All were right-handed and considered at least stage-III treatment-resistant: they had had a minimum of 2 unsuccessful trials of SSRI/NSRI treatment and 1 failed trial with tricyclic antidepressants as described by Rush et al. [25]. Because concomitant antidepressant treatment can confound outcome results, all patients went through a 2-week antidepressant washout before entering the study (this was over 3 weeks if they were on fluoxetine). Where necessary, patients were kept on a steady dose of their ‘somatic’ medications. During the washout period, patients

Impact of HF-rTMS on Fine Motor Functions in MDD

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had contact with their physicians on a regular basis. Only habitual benzodiazepine agents were allowed: 1 subject took alprazolam (1 mg), 1 took clonazepam (0.5 mg), 1 took flunitrazepam (1 mg) and 1 took alprazolam (1 mg) and flurazepam (27 mg). During our stimulation protocol, all psychopharmacological changes were considered as dropout from the study. Additionally, no changes of habitual somatic treatment were allowed. All subjects underwent physical, neurological (MRI, EEG) and psychiatric examinations.

Exclusion criteria were: a current or past history of epilepsy, neurosurgery, having metal or magnetic objects in the brain and being pregnant. Patients with suicide attempts during their current depressive episode or alcohol/drug dependence and/or abuse were not included.

A group of 28 healthy controls (mean age: 40.82 ± 6.93; male:female ratio = 11/17) was included matched for sex and age with the patient group (p = 0.20 and p = 0.77, respectively). These controls were recruited in the context of a previous research project on psychomotor functioning in major depression [26]. This study was part of a larger project investigating the influence of HF-rTMS on different neurocognitive markers. The study was consistent with the latest version of the Helsinki Declaration and was approved by the ethics committee of the University Hospital (UZBrussel) of the Vrije Universiteit Brussel. All subjects gave written informed consent.

Psychomotor Assessments

For the objective psychomotor measurement, all participants carried out a Fitts’ task (see fig. 1) using a pressure-sensitive ink ballpoint pen on sheets of paper placed on a digitizer that was connected to a personal computer. The Fitts’ task is a computerized fine motor task that has been designed especially to evaluate sensorimotor programming, initiation and execution of muscle commands without requiring higher-order cognitive processes [17–19]. In this task, subjects had to connect two vertically placed circles, depicted on a normal sheet of paper, by drawing a line of about 1 cm. They were instructed to start in the middle of the top circle and to end in the middle of the lower circle. Per trial, 6 lines had to be drawn. The accuracy of movement was varied by changing the circle diameter from 0.50 cm in trials 1 and 4 (fig. 1, upper part) to 0.25 cm in trials 2 and 3 (fig. 1, lower part). Movement time was recorded, i.e. the time between the starting and completion of each separate line-drawing movement. The movement times of inaccurate line drawings, i.e. when the line was drawn from the lower to the upper circle and/or when the start/end point of the connecting line was situated outside the circle diameter, were not included in the analyses.

In addition, visual analog scales (VAS) were used to examine subjective mood changes: subjects were asked to rate their mood on 5 horizontal 100-mm VAS in order to detect subtle changes in feelings of ‘depression’, ‘fatigue’, ‘tension’, ‘anger’ and ‘vigor’. The minimum score on each VAS subscale is 0 and the maximum score is 100. Right-handedness was assessed with the Van Strien questionnaire [27].

Design and rTMS Procedure

Patients underwent 10 sessions of HF-rTMS on the left DLPFC within a period of 2 weeks. At the beginning of this open-treatment trial, each subject also received 1 placebo (sham) HF-rTMS stimulation session, separated from the first active stimulation session by 1 day. This phase was a placebo-controlled cross-over, single-blind design allowing examination of single session, specific rTMS effects in MDD patients.

Potential mood changes were assessed before (Tpre), immediately after (Tpost) and 30 min after (Tpost30) terminating the first rTMS (real/sham) session, using VAS scales.

The antidepressant effects of 2 weeks of rTMS treatment were investigated by assessing the HDRS and BDI at baseline and at end point, i.e. after the eleventh stimulation session (1 of which was a placebo session). The Fitts’ task was applied before (Tpre) immediately after (Tpost) terminating the first and second rTMS (real/sham) session, and at the end of the rTMS treatment period (Tpost-treatment i.e. after the last session).

Because this study was part of a larger project investigating the influence of HF-rTMS on different neurocognitive markers, additional tasks were also administered that were not used for the current study purposes. Measures were always presented in the same order for all participants.

Patients were kept unaware of the type of stimulation; they wore earplugs and were blindfolded. Importantly, all patients

![Fig. 1. Fitts’ task.](image-url)
were stimulated on all occasions within the same time schedule, i.e. between 10 am and noon.

For the application of rTMS, we used a Magstim high-speed magnetic stimulator (Magstim Company Limited, Wales, UK), connected to a figure-of-8-shaped coil. Before each application, the motor threshold of each individual was determined using electromyography. A stimulation intensity of 110% of the subject’s motor threshold of the right abductor pollicis brevis muscle was used. In order to accurately target the left DLPFC (Brodman area 9/46), the precise stimulation site and position of the coil were determined using MRI nonstereotactic guidance (Philips Intera, Best, The Netherlands). Perpendicular to this point, the precise stimulation site on the skull was marked and stimulated [28]. In each HF (10 Hz) stimulation session, subjects received 40 trains of 3.9-second duration, separated by an intertrain interval of 26.1 s. Each session, therefore, lasted 20 min (1,560 pulses per session). For the sham condition, the coil was held at an angle of 90°, with one edge only resting on the scalp. The International Society of Transcranial Magnetic Stimulation safety guidelines were followed [29, 30].

Statistics
All results were analyzed using the SPSS for Windows 16.0 software package. Statistical analyses were performed using ANOVAs. The significance level was set at $p < 0.05$ for all analyses. Baseline and end-point psychomotor outcomes in the patient group were compared with the outcomes of an age- and gender-matched control group that participated in a previous research project in which psychomotor measurements were registered [26]. Separate analyses were conducted to investigate the single-session effect (1 active/sham session) and the treatment effect (10 active sessions) of HF-rTMS.

Regarding the single-session effects, we used separate $3 \times 2$ ANOVAs with the VAS mood scales as dependent variables and Session (Tpre, Tpost or Tpost30) and Stimulation (active, sham) as within-subject factors. With regard to the analyses for the single-session effects on the Fitts’ task, a $2 \times 2 \times 2$ factorial design was used with Stimulation (active, sham), Session (Tpre, Tpost) and Complexity (small circles, large circles) as within-subject factors.

Subsequently, overall improvement in HDRS and BDI scores between the baseline and final assessment were analyzed using paired Student t tests over the whole group. Clinical response was defined as a 50% reduction of the baseline HDRS score.

A similar $2 \times 2 \times 2$ design was applied for the psychomotor outcomes related to the 10-session treatment with Session (Tpre, Tpost-treatment) and Complexity (small circles, large circles) as within-subject factors, and Treatment Response (responders, non-responders) as a between-group factor.

Results

Baseline Psychomotor Outcomes
A comparison of the baseline Fitts’ outcomes of the patient group with those of an age- and gender-matched control group [mean movement time (MT) 255 ms and standard deviation (SD) 0.25] pointed to a significantly slower performance of the patients (mean MT 321 ms, SD 0.12, $t = 2.21$ and $p < 0.05$).

Single Session Effects
Mood. VAS analyses were conducted on 20 patients because 1 subject had numerous missing values. The separate ANOVAs did not reveal any significant effect, neither for the main effect of Stimulation (all F values <1.33) or Session (all F values <2.43), nor for the Stimulation by Session interaction (all F values <1.09). The only exception was the VAS depression subscale that showed a main effect of Session [$F(2, 18) = 8.38, p < 0.01$] with the scores slightly decreasing from Tpre (6.46) over Tpost (5.12) to Tpost30 (5.74), without a significant Stimulation by Session interaction [$F < 1$], however.

Psychomotor Variables
As demonstrated in previous studies, a smaller diameter of the circles resulted in significantly higher MTs [$F(1, 20) = 94.6, p < 0.001$]. Moreover, a significant main effect of Session was found with the MTs improving from prestimulation to poststimulation [$F(1, 20) = 5.68, p < 0.05$; Tpre: 309 ms, Tpost: 284 ms]. However, neither the Stimulation by Session interaction nor any of the other interactions was significant [all Fs<1], with the exception of the Complexity by Session interaction [$F(1, 20) = 3.91, p < 0.1$]. These results indicate that the first rTMS session did improve fine motor performance but no difference could be demonstrated between active (Tpre: 306 ms; Tpost: 279 ms) or sham (Tpre: 311 ms; Tpost: 287 ms) stimulation within that time frame. It should be noted that patients remained slower than healthy controls after the first stimulation session ($F = 3.14, p < 0.1$).
**Treatment Effects**

**Mood.** Mean HDRS scores before entering the study were 25.24 (SD = 3.9) and mean BDI scores were 33.82 (SD = 12.19), indicating severe depression. The overall patient group demonstrated a significant improvement in HDRS and BDI scores between the baseline and final assessment (final HDRS: 15.35, t = 5.61, p < 0.001; final BDI: 25.27, t = 2.76, p < 0.05). Eleven patients (52%) were considered clinical nonresponders and the other 10 (48%) considered clinical responders.

**Psychomotor Variables**

A significant psychomotor improvement was observed from baseline to end point [Tpre: 321 ms, Tpost-treatment: 276 ms; F(1, 19) = 4.22, p < 0.05; see fig. 2], as well as the well known effect of Complexity [F(1, 19) = 80.88, p < 0.001]. Neither the main effect of, nor any interactions with Treatment Response were significant [Fs < 2.12].

A comparison of the end-point Fitts’ outcomes of the patient group (mean MT: 276 ms) with the baseline data of the control group (mean MT: 255 ms) revealed no significant differences in psychomotor performance (t = 0.78, p = 0.44).

Note that no further substantial psychomotor improvements occurred between session 2 (which was an HF-rTMS or a sham session) and session 10 (last HF-rTMS session): F < 1, p = 0.36. This could imply that the observed psychomotor improvements are obtained between the first and the second psychomotor assessment (following 1 sham/ rTMS stimulation), which further underscores the previously mentioned impact of there being a practice effect.

Additional correlational analyses were conducted to further investigate the association between the clinical and psychomotor variables. No significant correlations could be observed, either between the absolute difference of the baseline and end-point rTMS psychomotor variable and the absolute difference in HDRS/BDI scores, or between the proportional changes of mentioned variables (all r values <0.34, all p values >0.15). In this context, it should also be mentioned that the Fitts’ outcomes at baseline and end point did not correlate with the duration of the current episode or the stage of treatment resistance (all r values <0.19 and all p values >0.4).

**Discussion**

This study explored the clinical and psychomotor effects of 1 sham-controlled and 10 active HF-rTMS sessions, respectively, in treatment-resistant MDD applying clinical rating scales and an objective psychomotor assessment method, i.e. the Fitts’ task.

At baseline, MDD patients performed significantly slower than an age- and gender-matched control group, which has also been demonstrated repeatedly in previous studies from our research group [9]. The total patient sample manifested a clear improvement in depression severity following 10 active rTMS sessions with approximately half of the patient sample manifesting a clear clinical response, as determined with a 50% decrease of the initial HDRS scores, which is in line with previous HF-rTMS treatment studies in MDD [31]. Regarding the VAS, one HF-rTMS session did decrease the depression subscale scores, irrespective of the stimulation type (active or sham), however. Scores on the other VAS subscales did not significantly change following one session.

Focussing on the effects of one HF-rTMS-session, psychomotor improvements were observed on the Fitts’ task, but did not reveal a difference between active and sham stimulation. The analyses investigating the total treatment period of 10 HF-rTMS sessions did point to a better end-point performance on the Fitts’ task, but these psychomotor improvements are likely due to a practice effect on the task. Indeed, the psychomotor improvements observed after session one emerged irrespective of sham or active HF-rTMS stimulation, and no further improvements occurred from session 2 to session 10. Notwithstanding, it should be mentioned that for every assessment session, patients were given the opportunity to practice with and get used to the task, before the proper recordings started.

Besides the mentioned practice effect, it might be possible that due to certain nonspecific aspects related to the rTMS procedure – such as its impressive name, its discomfort and its sophisticated-looking equipment – a placebo effect could have influenced the one-session outcomes [32]. In addition, we cannot totally rule out the possible impact of our sham control condition: although this was performed at a 90° angle, ensuring minimal stimulation of the DLPFC, it could still be possible that a partially active placebo was used [33].

Whereas the current study mainly focused on fine motor performance, it would also be interesting to obtain more knowledge on the effects of rTMS on gross motor performance. However, the relationship between gross and fine motor performance in MDD has not yet been elucidated and it is not clear whether MDD patients with fine motor dysfunction are affected to the same degree in their gross motor performance [9]. Therefore, in order to investigate the impact of rTMS on gross motor perfor-
The psychomotor effects of rTMS treatment applying clinical rating scales. Hoeppner et al. [14] reported a significant improvement of baseline MARS-rated psychomotor retardation after 10 HF-rTMS sessions spread over 2 weeks, for both a group treated with 20 Hz HF-rTMS and a group treated with 1 Hz HF-rTMS, whereas a sham group did not manifest any psychomotor improvement. In another study, however, Hoeppner et al. [15] could not demonstrate a beneficial effect of left prefrontal 10 Hz HF-rTMS treatment during 15 days on MARS-assessed psychomotor retardation. Instead, they found nearly significant reductions in the agitation symptoms. Very recently, Baeken et al. [16] reported a decrease in SRRS scores after 10 sessions of 10 Hz HF-rTMS over the left DLPFC. In this context, it needs to be mentioned that rating scales and the applied psychomotor assessment method substantially differ in their duration of observation: rating scales are based on prolonged clinical observations whereas experimental tasks only capture fine motor performance during task execution [9]. Moreover, rating scales can be rater-dependent whereas the currently used assessment method does not depend on the rater [9].

A limitation of this study could be the number of HF-rTMS sessions. Despite rTMS treatment parameters being quite intense, the duration of 2 weeks might be considered as rather short. Indeed, current HF-rTMS treatment protocols stimulate patients daily for 3–6 weeks [6, 34]. In addition, as a higher level of treatment resistance in the current depressive episode might be inversely related to clinical outcome, this might to some extent have impacted our fine motor task results [35].

A major strength of this study is that all patients had a sufficient washout period from their antidepressants, whereas in several other studies patients were still taking their current psychotropic medication. Moreover, during the stimulation protocol, no changes to patients’ habitual somatic treatment were allowed. On the other hand, although all included patients in our analysis continued with exactly the same benzodiazepine concentrations, the use of benzodiazepines in our sample might have been a confounding variable [19].

Future studies might do well to further objectively investigate fine motor functioning in depressed cases in association with rTMS treatment. More intensive placebo-controlled rTMS studies are needed to further disentangle the effect of HF-rTMS on the psychomotor system as clinical improvement with these kinds of techniques are reported.

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Impact of HF-rTMS on Fine Motor Functions in MDD

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