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Clinical and electrophysiological effect of right and left repetitive transcranial magnetic stimulation in patients with major depressive disorder

Maria García-Anaya,1 Jorge González-Olvera,1 Josefina Ricardo-Garcell,2,3 Gabriela Armas,2 Edgar Miranda,2 Ernesto Reyes,2 Gloria Adelina Otero4

SUMMARY

Major Depressive Disorder (MDD) is a common psychiatric disorder that represents one of the main public health problems worldwide. It has been projected that for 2020 it will be the second cause of disability-adjusted life years just below ischemic heart disease.

Quantitative electroencephalogram provides the opportunity to study cortical oscillatory activity across the different frequency bands. It constitutes an accessible tool to explore the clinical and neurophysiologic correlates underlying psychiatric disorders as well as the effect of diverse therapeutic options and the performance through cognitive tasks.

Repetitive transcranial magnetic stimulation is a technique that allows the stimulation of the cerebral cortex noninvasively, relatively painlessly and with fairly few side effects.

The vast majority of rTMS studies target left dorsolateral prefrontal cortex (DLPFC) based on imaging studies showing that left prefrontal cortex dysfunction is pathophysiologically linked to depression. However, there is some evidence implicating right PFC in the pathophysiology of depression.

Comparison of antidepressant efficacy of diverse stimulation frequencies is relevant since a main concern around rTMS is its potential to induce seizures; hence we consider that frequency of stimulation is an important aspect to be studied.

For this study we aimed to elucidate the clinical efficacy of rTMS comparing two groups of depressed patients stimulated over DLPFC, one over the left (at 5 Hz) and other over the right (at 1 Hz). We also meant to know if there were clinical and electroencephalographic differential long-term after-effects between those groups of treatment.

We included twenty right-handed patients with a DSM-IVR diagnosis of MDD. They were assigned into two groups of treatment.

Group 1 received 5 Hz rTMS over the left DLPFC. Group 2 received 1 Hz rTMS over the right DLPFC.

We obtained two EEG measurements in order to analyze Z score of broad-band spectral parameters and cross-spectral.

No statistical differences among groups were found in response to treatment after weekly comparisons of clinimetric scores and significant differences between baseline and final assessment by HDRS, MADRS, BDI and HARS.

The major rTMS effect on EEG was observed in the group that received 1 Hz over the right DLPFC and no significant effects were observed for the group that received 5 Hz over the left DLPFC.

Our results propose that administration of 15 sessions on either left (5 Hz) or right (1 Hz) rTMS over DLPFC is sufficient to reach response to treatment, assessed by HDRS, MADRS, BDI and HARS in subjects with MDD. Moreover, in both cases rTMS was able to induce an equivalent antidepressant effect.

The major effect of rTMS on EEG was observed in the right 1 Hz rTMS group where changes were elicited mainly over frontal, central and temporal regions on alpha and particularly beta frequency bands. In a lesser extent for left 5 Hz rTMS group the main effect was observed on anterior regions for beta and particularly alpha frequency bands.

We believe it is pertinent to continue exploring the therapeutic potential of lower stimulation frequencies, for what further research including larger samples is still necessary to confirm these trends.

Key words: Major depressive disorder, rTMS, EEG, laterality, 5Hz, 1Hz.

RESUMEN

El trastorno depresivo mayor es una entidad psiquiátrica que representa uno de los principales problemas de salud pública a nivel mundial. Se ha proyectado que para el año 2020 será la segunda causa de discapacidad únicamente por debajo de la cardiopatía isquémica.

La utilización del electroencefalograma cuantitativo ofrece la oportunidad de estudiar la actividad oscilatoria cortical a través de las diferentes bandas de frecuencias.

Éste constituye una herramienta para explorar las características clínicas y neurofisiológicas que subyacen a los trastornos psiquiátricos, así como un instrumento para evaluar el efecto de diversas opciones terapéuticas y el desempeño de los sujetos durante la realización de tareas cognitivas.

La estimulación magnética transcraneal repetitiva (EMTr) es una técnica que permite la estimulación de la corteza cerebral de manera no invasiva, relativamente sin dolor y con pocos efectos secundarios.

Con base en los estudios de neuroimagen que vinculan la fisiopatología de la depresión con disfunción en la corteza prefrontal...
INTRODUCTION

Major depressive disorder is a common psychiatric disorder that represents one of the main public health problems worldwide. It has been projected that for 2020 it will be the second cause of disability-adjusted life years just below ischemic heart disease. MDD has a lifetime prevalence estimated at 5.8% in women and 2.5% in men in Mexican population, while in the USA the lifetime prevalence of CIDI/DSM-IV MDD for adult population is 16.2%.3

In a real life scenario, these numbers represent a huge social and economical burden, therefore, research focused on successful diagnostic and therapeutic tools should be a priority of mental health systems.

In that sense, quantitative electroencephalogram (QEEG) provides the opportunity to study cortical oscillatory activity across the different frequency bands. It constitutes an accessible tool to explore the clinical and neurophysiologic correlates underlying psychiatric disorders as well as the effect of diverse therapeutic options and the performance through cognitive tasks.5

On the other hand, currently, the pharmacological approach still constitutes the first election treatment for MDD. However, it is not exempt from having a significant percentage of failures and an important amount of drop outs is explained in part by adverse and side effects of pharmacological antidepressants.7

Repetitive transcranial magnetic stimulation (rTMS) has emerged as an alternative to standard pharmacological therapies. Its antidepressant efficacy has been examined by several meta-analyses which in general have shown evidence of statistical clinical benefit that has importantly improved from the first studies until the most recent ones. These findings remarkably point out how the more the rTMS therapeutic potential is studied the better results are achieved. In relation to this, it has been demonstrated how the larger amount of total pulses is administered, the stronger antidepressant effect is achieved.13

Repetitive transcranial magnetic stimulation is a technique that allows the stimulation of the cerebral cortex noninvasively, relatively painlessly and with fairly few side effects. The focused magnetic field over the surface of the head induces electrical currents in the brain that as a result depolarizes the underlying superficial neurons. Even when rTMS effects vary depending on the stimulus frequency, intensity and duration, as well as on the number of sessions, it is generally accepted that rTMS involves a wide range of excitatory, inhibitory and plastic neuronal processes.

One pathophysiological hypothesis of MDD stands on evidence from imaging studies showing that decreased left prefrontal cortex (PFC) function with respect to the right is linked to depression. On that basis, stimulation parameters chosen as treatment are selected from evidence of how high frequency rTMS (1 Hz) increases excitability below the underlying cortex just as low frequency rTMS (≤1 Hz) does it so.

Thus, the vast majority of rTMS studies target left dorsolateral prefrontal cortex (DLPFC). However, there is some evidence implicating right PFC in the pathophysiology of depression. In a double-blind placebo controlled clinical trial no significant differences emerged when comparing high-frequency left rTMS with low-frequency right rTMS. In addition, there is a study where abnormal
EEG sources (increase in current density) were observed in both hemispheres but with maximal inverse solution located mainly over right frontal lobe.27

Numerous authors26,28-35 have reported results of antidepressant effect for left high-frequency rTMS and few less for right low-frequency rTMS, but only one has compared these two stimulation strategies, finding no significant differences between left 10 Hz rTMS and right 1 Hz rTMS, both applied over DLPFC.26

Comparison of antidepressant efficacy of diverse stimulation frequencies is relevant since a main concern around rTMS is its potential to induce seizures;36 hence we consider that frequency of stimulation is an important aspect to be studied. In regards to that, a plausible rationale to target 5 Hz rTMS for antidepressant purposes, besides its promising effectiveness, is the fact that it has been poorly studied and also represents a lower risk for induced seizures besides of a more tolerable sensation for patients.

For this study we aimed to elucidate the clinical efficacy of rTMS comparing two groups of depressed patients stimulated over DLPFC, one over the left (at 5 Hz) and other over the right (at 1 Hz). We also meant to know if there were clinical and electroencephalographic differential long-term after-effects between those groups of treatment.

**MATERIALS AND METHOD**

**Participants**

Twenty right-handed patients (14 female, range 19-46 years, mean age=31.7, S.D.=7.38) with a DSM-IVR diagnosis of MDD37 were included in this study. Diagnosis was made by a psychiatrist using the Structured Clinical Interview for DSM-IVR (SCID-I). All patients scored higher than 18 points in the Hamilton Depression Rating Scale at baseline (mean score=28.3, S.D.=6.20).

Subjects with epilepsy, convulsive antecedents, drug abuse history, actual suicidal ideation or any axis I psychiatric disorder, excepting anxiety disorders were excluded. In order to reduce rTMS-induced seizure risk, the subjects showing epileptiform activity in the EEG recording were also excluded. In this sample, 15 patients were virgin to treatment. The remaining were medication-resistant patients; four of them were under a third-course and one under a second-course trial, having no response after at least eight weeks of treatment at maximum dose. Three of them were assigned to group 1 and two to group 2 (see below for assignment to group of treatment). A complete description of the study was given to every patient and afterwards an informed consent was signed by them on a form approved by the National Institute of Psychiatry (NIP) Research Ethics Committee. All subjects were recruited from the outpatient unit of the NIP in Mexico City.

**Clinical assessment**

A psychiatrist performed clinimetric assessment blinded to rTMS treatment by means of Hamilton Depression Rating Scale 21-item version (HDRS), Montgomery-Asberg Depression Rating Scale (MADRS), Beck Depression Inventory 21-item version (BDI) and Hamilton Anxiety Rating Scale (HARS).

Evaluation with those instruments was made at baseline and after sessions 5, 10 and 15. Response to treatment was considered as the reduction ≥50% in clinimetric scores.

**Treatment**

Repetitive Transcranial Magnetic Stimulation at 100% of motor threshold was administered using a Dantec MagPro rapid magnetic stimulator with a 50mm diameter figure-eight-shaped MC-B70 coil (Dantec; Skovlunde, Denmark). Motor threshold was determined at the beginning of every session using visual inspection method, assessing the motor response of the abductor of pollicis brevis muscle. The site of stimulation was defined as the region 5cm anterior to the point of maximum stimulation of the abductor pollicis brevis muscle.30,31,38,39 At every moment, the imaginary axis in the middle of the coil was held matching with the scalp parasagital line. Patients were randomly assigned into one of two groups of treatment. Group 1 received 5Hz rTMS over the left DLPFC (30 trains of 10 sec duration separated by 10sec; 22500 pulses per session). Group 2 received 1Hz rTMS over the right DLPFC (1 train of 15 min duration, 900 pulses per session). One daily session was administered from Monday to Friday until they all accomplished a total of 15.

**EEG acquisition**

We obtained two EEG studies, at enrollment and three days after the last rTMS session. EEG recordings were acquired with subjects awake with eyes closed, lying on a couch in a dimly lighted and not acoustically shielded room. The participants wore a polyester cap with surface electrodes distributed according to the 10-20 International System (Fp1, Fp2, F3, F4, C3, C4, P3, P4, O1, O2, F7, F8, T3, T4, T5, T6, Fz, Cz, Pz), using linked earlobes as reference. EOG was recorded from a supraorbital electrode and from an electrode on the external canthus of the left eye. Impedance levels were ≤5 KΩ. The amplifier bandwidth was set between 0.5 and 30 Hz and the EEG was sampled every 5 ms using Trackwalker v 2.0 and Medicid IV system from NeuronicTM. A photosimulator was used in order to reject subjects where photosensitive epileptiform activity was present. Total recording time was from 20 to 30 min.

**EEG analysis**

Two expert electroencephalographists selected 24 independent artifact-free EEG segments of 2.56 sec by visual
inspection. Segments were included only if both professionals agreed on the selection. EEG analysis was carried out off-line. Data sample spectral analysis was calculated by Fast Fourier Transform (FFT) and cross-segment averaging.

Cross-spectral matrices were calculated every 0.39 Hz, from 0.39 to 19.11 Hz, and Narrow Band Spectral Parameters (NBSP) were obtained (absolute power from 0.39 to 19.11 Hz). We also got Broad Band Spectral Parameters (BBSP: absolute power, relative power and total absolute power) for delta (1.5-19.0 Hz) (1.5-3.5 Hz), theta (3.5-7.5 Hz), and beta (12.5-19.0) bands.

In order to calculate Z values on both NBSP and BBSP, each individual EEG recording was compared to normal subject parameters data bases for each frequency and electrode position. All this analysis was done using Neuronic Quantitative and Tomographic EEG v 6.0 software.

Data analysis

Repeated measures ANOVA was performed to estimate decreases in HDRS, MSDRS, BDI and HARS scores over sessions. U-Mann Whitney tests were performed to investigate differences among the two groups in clinical and demographical variables. Post-hoc Bonferroni tests were only calculated where a significant effect was found in the analysis of variance. Significance was set at \( \alpha = 0.05 \). This statistical analysis was computed with PASW Statistics 18 software.

To explore differences after left or right rTMS, two factors (1dep/1indep) univariate ANOVA for Z values of BBSP and NBSP was performed. Post-hoc paired-t tests were only calculated when a significant effect was found in the analysis of variance. Significance was set at \( \alpha = 0.005 \). For this analysis we utilized Neuronic Statistics v4.0 software.

RESULTS

Clinical and demographic characteristics were compared by Mann-Whitney U (\( \alpha = 0.05 \)), no statistical differences at baseline among groups were found (table 1). We performed repeated measures ANOVA to estimate decreases in HDRS, MSDRS, BDI and HARS scores over sessions. Effects are presented in graphic 1. We observed response to treatment (reduction of \( \geq 50\% \) in clinical scores) for both groups of treatment. There was an overall effect group with significant differences between baseline and session 5 up to 15 assessed by HDRS (F=65.57; P= 0.001), MADRS (F=60.22; P=0.001)

Table 1. Demographic and baseline characteristics in 20 patients with major depression

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>1 Hz Group</th>
<th>5 Hz Group</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>34.9 (4.06)</td>
<td>28.7 (8.23)</td>
</tr>
<tr>
<td>Sex M/F No.</td>
<td>3/7</td>
<td>3/7</td>
</tr>
<tr>
<td>HAMD basal score</td>
<td>26.6 (4.88)</td>
<td>30.1 (7.12)</td>
</tr>
<tr>
<td>MADRS basal score</td>
<td>29.3 (6.18)</td>
<td>32.0 (4.80)</td>
</tr>
<tr>
<td>BDI basal score</td>
<td>33.5 (9.93)</td>
<td>33.1 (10.46)</td>
</tr>
<tr>
<td>HARS basal score</td>
<td>24.0 (5.31)</td>
<td>26.9 (9.10)</td>
</tr>
</tbody>
</table>

Groups were compared by Mann-Whitney U at \( \alpha = 0.05 \). No significant differences were observed (Data is presented as Mean [SD]), (n=10 right, n= left).

Graphic 1. Mean scores of HDRS, MADRS, BDI and HARS administered at baseline (1), session 5 (2), session 10 (2) and at the end of treatment (4) for each group (n=10 right, n= left). Significant changes with respect to basal scores are signalled with *
Clinical and electrophysiological effect of right and left rTMS

and HARS (F=58.79; P=0.001), and between baseline and session 10 and 15 assessed by BDI (F=46.84; P=0.001).

No statistical differences in response to treatment were observed when weekly comparisons of clinimetric scores among groups were performed using Mann-Whitney U test (α=0.05) (table 2).

After a two factor univariate ANOVA analysis carried out on BBSP Z values we found no significant results (α=0.05).

Regarding NBSP Z values, we got significances for interaction (α=0.005; DF=1.18; F=10.218). Subsequent paired-t tests showed a significant effect on pre-post differences for both groups of treatment. The major effect was observed in the right 1 Hz rTMS group where changes were elicited mainly over frontal, central and temporal regions on alpha and beta frequency bands, particularly on the last one; interestingly all t-scores for this group were positive (α=0.025) (figure 1). For the left 5 Hz rTMS group, significant pre-post changes were spread across all frequency bands and topographies although the main effect was observed on anterior regions for fast frequency bands. Remarkably and opposite to the right 1 Hz rTMS group, the majority of t-scores (17/21) were negative (α=0.025) (figure 2).

As side-effects, three subjects reported headaches along the first three days of treatment. Pain responded to treatment

<table>
<thead>
<tr>
<th>Time and Group</th>
<th>HDRS Scores</th>
<th>MADRS Scores</th>
<th>BDI Scores</th>
<th>HARS Scores</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean ± S.D.</td>
<td>Mean ± S.D.</td>
<td>Mean ± S.D.</td>
<td>Mean ± S.D.</td>
</tr>
<tr>
<td>Baseline</td>
<td>1 Hz 26.6 ± 4.88</td>
<td>29.3 ± 6.18</td>
<td>33.5 ± 9.93</td>
<td>24.0 ± 5.31</td>
</tr>
<tr>
<td></td>
<td>5 Hz 30.1 ± 7.12</td>
<td>32.0 ± 4.80</td>
<td>33.1 ± 10.46</td>
<td>26.9 ± 9.10</td>
</tr>
<tr>
<td>Week 1</td>
<td>1 Hz 16.8 ± 8.25</td>
<td>18.9 ± 10.12</td>
<td>23.1 ± 17.19</td>
<td>15.1 ± 8.84</td>
</tr>
<tr>
<td></td>
<td>5 Hz 12.9 ± 4.77</td>
<td>16.2 ± 8.16</td>
<td>14.8 ± 8.01</td>
<td>13.1 ± 7.65</td>
</tr>
<tr>
<td>Week 2</td>
<td>1 Hz 10.5 ± 6.72</td>
<td>10.0 ± 5.86</td>
<td>13.9 ± 10.04</td>
<td>9.5 ± 6.31</td>
</tr>
<tr>
<td></td>
<td>5 Hz 10.7 ± 5.45</td>
<td>10.5 ± 5.19</td>
<td>11.7 ± 7.57</td>
<td>9.4 ± 6.39</td>
</tr>
<tr>
<td>Final</td>
<td>1 Hz 9.1 ± 5.08</td>
<td>7.4 ± 3.97</td>
<td>13.9 ± 8.96</td>
<td>8.0 ± 4.54</td>
</tr>
<tr>
<td></td>
<td>5 Hz 7.6 ± 5.94</td>
<td>7.9 ± 8.98</td>
<td>9.0 ± 7.8</td>
<td>7.0 ± 5.33</td>
</tr>
</tbody>
</table>

Groups were compared by Mann-Whitney U at α=0.05. No significant differences were observed (n=10 right, n= left).

EEG regions with significant changes in Z-values after 1 Hz right rTMS treatment

<table>
<thead>
<tr>
<th>Frequency (Hz)</th>
<th>Electrode position (t scores, ±2.26)</th>
</tr>
</thead>
<tbody>
<tr>
<td>7.42</td>
<td>O2 (2.17)</td>
</tr>
<tr>
<td>9.37</td>
<td>F4 (2.28), FZ (2.34)</td>
</tr>
<tr>
<td>9.77</td>
<td>C4 (2.44), CZ (2.60)</td>
</tr>
<tr>
<td>10.55</td>
<td>C3 (2.52)</td>
</tr>
<tr>
<td>10.94</td>
<td>C4 (2.81), T3 (2.32)</td>
</tr>
<tr>
<td>11.33</td>
<td>F1 (2.70), F7 (2.50), F8 (2.40), T3 (2.67)</td>
</tr>
<tr>
<td>13.67</td>
<td>T3 (2.70)</td>
</tr>
<tr>
<td>14.06</td>
<td>T3 (3.02), T4 (2.36)</td>
</tr>
<tr>
<td>14.45</td>
<td>F2 (2.51), T3 (2.56), P3 (2.65)</td>
</tr>
<tr>
<td>14.84</td>
<td>T3 (2.36)</td>
</tr>
<tr>
<td>15.23</td>
<td>F4 (2.34), F8 (2.66), FZ (2.65), P3 (2.78)</td>
</tr>
<tr>
<td>15.62</td>
<td>F1 (4.10), F2 (2.31)</td>
</tr>
<tr>
<td>16.02</td>
<td>F1 (2.43), F2 (2.37), F4 (2.30), T3 (2.41), P3 (2.7, 5)</td>
</tr>
<tr>
<td>16.41</td>
<td>F1 (2.42), F2 (2.54), F3 (2.61), F7 (2.36), FZ (2.97), T6 (3.14), P3 (2.30), F4 (2.39), O2 (2.34)</td>
</tr>
<tr>
<td>16.80</td>
<td>F1 (2.56), F2 (2.28), F3 (2.80), C4 (2.62), CZ (2.41), T3 (3.21), T4 (2.68), O2 (2.30)</td>
</tr>
<tr>
<td>17.19</td>
<td>F1 (2.58), F2 (2.36), C3 (2.31)</td>
</tr>
<tr>
<td>17.58</td>
<td>F3 (2.37), F7 (2.46), CZ (3.38), O2 (2.28)</td>
</tr>
<tr>
<td>17.97</td>
<td>T3 (2.56)</td>
</tr>
<tr>
<td>18.36</td>
<td>CZ (2.93), T3 (2.92), T4 (2.58), P3 (2.65)</td>
</tr>
<tr>
<td>18.75</td>
<td>F1 (2.31), T3 (2.32), T3 (2.63)</td>
</tr>
<tr>
<td>19.14</td>
<td>F1 (2.33), F4 (2.43), F8 (2.56), CZ (2.79), T3 (2.28)</td>
</tr>
</tbody>
</table>

Figure 1. T scores ±2.26 for each electrode position were 1 Hz right rTMS induced change after treatment (n=10, p=0.025). Concentric circles represents the number of NBSP frequencies with changes at each derivation (the signal inside correspond to 1 signal).
with aspirin. One of those patients also experienced facial paresthesia that spontaneously disappeared after the third session of treatment.

Since all subjects achieved response to treatment after 15 rTMS sessions, they all were invited to receive maintenance treatment with one session per week for eight weeks (data to be published later). After the follow up, all patients were referred to the NIP outpatient service for their further management. At the moment patients were referred, pharmacological treatment was left unchanged in subjects who formerly had it and no pharmacological adding was made on the remaining ones.

**DISCUSSION**

After slightly more than two decades of studies focused on rTMS therapeutic usages, an important amount of evidence points out the antidepressant effect of this technique.8-11,32,42 According to those findings, significant decreases on weekly assessed scores in HDRS, MADRS, BDI and HARS observed in our study show that both prefrontal left and right rTMS are able to reach response to treatment after 15 sessions.

Joint together with the study of Fitzgerald et al.,26 where right (1 Hz) and left (10 Hz) rTMS resulted in the same antidepressant effect, our results support that rTMS applied in either one or other PFC is equivalent in terms of efficacy. Unlike Fitzgerald’s study, we administered 5 Hz rTMS for left PFC, which raises an important clinical consideration. Since the unpleasant feeling and induced seizure risk increases at higher stimulation frequencies,16 the relevance of our results lies in the need to highlight that lower frequencies applied into rTMS protocols could bring benefits to patients in terms of safety and comfort.

The study of rTMS by means of EEG or neuroimaging techniques has brought traces of its effect on brain connectivity. The number of reports combining QEEG and rTMS is not extensive35,43-48 and they all differ in terms of hypothesis, aims and stimulation parameters. Among them all, only three assessed the long-term effect of rTMS (considered as more than 1 session)35,45,49 and just two of them included depressed subjects.35,49

Griskova et al.45 administered, on separate days, one real and one sham rTMS session at 10 Hz over left DLPFC in healthy subjects. They found a significant increase in delta power over frontal, central and parietal regions after real stimulation.

Spronk et al.35 applied a scheme of sessions varying from 15 to 25 over the left DLPFC at 10 Hz, depending on the clinical course of each patient. Besides a highly significant clinical improvement evidenced by a decrease in BDI scores from baseline to end of treatment (p<0.001), they found a trend to increase in power of delta and alpha-2 bands, as well as a decrease in theta 2 power after treatment. In a preliminary analysis of 10 Hz rTMS effect, Funk et al.49 calculated a hemispheric ratio for each frequency band considering values pre-rTMS and post-rTMS every treatment session. For all bands, EEG power was stronger in the right hemisphere at baseline and throughout the course of treatment it tended to become stronger in left hemisphere until reaching a reversal of pre and post values, except for theta were the pre-post ratio reached equilibrium.

In contrast with the aforementioned studies, we did not find significant results in BBSP data when intra- and intergroup rTMS effect was compared. Nevertheless, NBSP analysis showed significant changes after rTMS treatment for both grups, mainly on frontal, central and temporal...
regions for alpha and beta frequency bands. Our results can hardly be compared to studies mentioned above mostly for three reasons: 1. they do not comprise the whole spectrum of every frequency band, 2. we explored different stimulation parameters and laterality and 3. none of the studies previously reported assessed rTMS long-term EEG after effects.

However, we observed an interesting pattern in relation to the effect of the frequency of stimulation used on each hemisphere. Remarkably, the direction of change of Z values was the same for all t scores for each group of treatment. With regard to right 1 Hz rTMS group, all t scores were positive for electrodes where significances were observed, which means an after treatment decrease in spectral power on alpha and beta frequency bands. As for the left 5 Hz rTMS group, except for FZ and F8 (on delta frequency band) and T5 and T6 (on beta band), all t scores were negative, meaning an increase on spectral power for theta, alpha and partially beta frequency bands after treatment.

In consequence, those decreases and increases on spectral power could be explained according to this major hypothesis concerning the underlying neural mechanisms of rTMS that points at the excitatory effect of high stimulation frequencies (≥1 Hz), as well as the inhibitory effect of low stimulation frequencies (≤1 Hz).

Additional supporting evidence for this rationale may stand on studies where rTMS at high and low frequency has been administered over either left or right prefrontal and primary motor cortex has induced either increased or decreased regional cerebral blood flow (rCBF) on ipsilateral and contralateral functional-related areas to the site of stimulation.

Based on all these data, it is feasible to expect an after-rTMS resultant cascade of effects throughout the brain which can also be measured at more locations, suggesting an underlying interhemispheric modulatory effect that probably entails the reorganization of neural circuits.

If such interhemispheric modulatory effect is so, a plausible explanation to the underlying neural mechanism related to changes observed in our study could be partially found in a way analogous to single-cell long-term depression and long-term potentiation acting over the stimulated hemisphere and perhaps giving place to those after-rTMS changes on spectral power.

CONCLUSION

To our knowledge, this is the first study to examine rTMS effect on EEG by NBSP, to explore the effect of 5 Hz rTMS on EEG and also to investigate long-term after effects of laterality of rTMS.

Clinically, our results demonstrate that administration of 15 sessions on either left (5 Hz) or right (1 Hz) rTMS over DLPFC is sufficient to reach response to treatment, assessed by HDRS, MADRS, BDI and HARS in subjects with MDD. Moreover, in both cases rTMS was able to induce an equivalent antidepressant effect.

Electrophysiological, right 1 Hz rTMS was able to elicit more changes than left 5 Hz rTMS. Those changes were observed mainly on beta frequency band for right 1 Hz group and alpha frequency band for left 5 Hz group. Additionally, NBSP were more useful to reflect EEG changes induced by rTMS. These findings suggest a reorganization of neural circuits secondary to long-term stimulation.

Despite that the main limitation of this study was the small number of patients included, its strength laid in our contribution to clinical and electrophysiological characterization of long-term effects of right (1 Hz) and left (5 Hz) rTMS, as currently there is a lack of studies comparing laterality of stimulation, especially with lower frequencies, and the neurophysiological mechanisms underlying to this promising therapeutic tool.

Finally, it is pertinent to continue exploring the therapeutic potential of lower stimulation frequencies, for what further research including larger samples is still necessary to confirm these trends.

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