Randomized Trial of Repetitive Transcranial Magnetic Stimulation for Apathy and Depression in Parkinson’s Disease

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Abstract

Objective: Repetitive transcranial magnetic stimulation (rTMS) has been reported to improve motor function and depression in Parkinson’s disease (PD) patients, but there has been only one randomized controlled trial for apathy. We evaluated the efficacy of rTMS for apathy and depression in Parkinson’s disease.

Methods: Fifteen PD patients received real rTMS (5 Hz, 500 pulses/day) and placebo stimulation over the supplementary motor area (SMA) for each 5 days with total amount of 2500 real pulses and 2500 placebo pulses, using a randomized real-first or placebo-first protocol. The modified apathy scale, the Zung Self-rating Depression Scale (SDS) and Unified Parkinson’s Disease Rating Scale (UPDRS) were used to assess apathy, depression and clinical status before and after each stimulative treatment.

Results: Real rTMS improved the apathy score by 3 points (P<0.05) compared to the baseline, while placebo stimulation produced no improvement, irrespective of the order of treatment. Real rTMS also improved the depression scale, SDS by 5 points (P<0.05) compared to the baseline, while placebo stimulation was ineffective. Combined analysis confirmed that real rTMS was significantly superior to placebo stimulation in apathy and depression (p<0.05). Real rTMS also improved UPDRS by 10 points (P=0.001), while placebo stimulation was ineffective. No side effects were observed in either real rTMS or placebo stimulation. Clinical factors including age, gender, disease duration, UPDRS score pre-rTMS, and daily dose of L-DOPA did not influence the improvement of UPDRS and apathy scores by real rTMS.

Conclusions: rTMS over the SMA appears to be effective for treatment of apathy and depression in PD patients in addition to UPDRS.

Keywords: Apathy; rTMS; Parkinson’s disease; Depression, SDS, UPDRS

Introduction

Alterations in mood, depression and apathy are the most frequent non-motor symptoms in Parkinson’s disease (PD) [1]. Repetitive transcranial magnetic stimulation (rTMS) has been used as a noninvasive stimulation for Parkinson’s disease, and has been modestly reported to improve motor function and bradykinesia compared with sham stimulation [2,3]. In addition, high-frequency rTMS over the left dorsolateral prefrontal cortex (DLPFC) was found to improve depression in PD patients with mood disorders, and low-frequency rTMS on the right DLPFC was also effective [4-9]. Apathy is considered a prevalent symptom with lack of motivation, and 17 to 70% of patients with PD also suffer from apathy [10]. However, there has been a few randomized controlled trial with well-defined inclusion criteria for the treatment of depression and apathy in PD patients [8,11,12]. Therefore, we examined whether rTMS improves apathy and depression in PD patients, using a randomized cross-over design of real and placebo stimulus sessions.

Patients and Methods

Participants included 15 subjects with Parkinson’s disease (PD) (mean age; 72.7, SD=8.4). The study protocol was approved by Shimane University Institutional Committee on Ethics (registered in the study number 231). Written informed consent was obtained from every participant before intervention. Their disease duration ranged from 33 to 218 months (mean 89.3 months, SD=48.9). All subjects underwent neurological examination and their total score on the unified Parkinson’s disease rating scale (UPDRS) was estimated. Apathy was evaluated according to the modified apathy scale (our revised version in Japanese [13]). Depression was also evaluated with the Zung Self-rating Depression Scale (SDS) [14]. Assessment of apathy, depression and UPDRS was performed before and after both real and placebo stimulations. Apathy scale, SDS and UPDRS were blind to stimulative allocation, real or placebo stimulation, and were obtained by blind raters. The patients continued to take their usual medications during the study. Medication for PD was only restricted to L-DOPA, because dopamine receptor agonist might influence on apathy and alter dopamine reward system [15-17].

The rTMS was performed under two conditions (real rTMS and placebo stimulation). For real rTMS, a figure-of-eight-shaped coil connected to a magnetic stimulator (Magstim Rapid; The Magstim...
Co.UK) was attached on the cranial surface over the SMA vertically in relation to the parasagittal plane. We selected the supplementary motor area (SMA) as the stimulus site, because SMA stimulation has been reported to improve motor function in PD patients. The stimulus site was set 3 cm anterior to the site of the anterior tibial muscle. Stimulus intensity, expressed as a percentage of the maximum stimulator output, was set at 110% resting motor threshold (MT) for the right abductor of pollicis brevis muscle [18]. One TMS session consisted of 10 trains of 10 s duration with 5 Hz frequency; 5 trains were applied over each hemisphere. Placebo stimulation was delivered with the coil angled at 90°, so that only the edge of the coil rested on the scalp. The stimulus parameters were the same as in real rTMS. The patients were randomized into real-first (n = 8) and placebo-first (n = 7) groups. Each of real rTMS and placebo stimulation was delivered for five days, and wash out period was set for two days between real and placebo stimulation (Figure 1).

**Results**

Baseline characteristics of 2 groups (procedure of stimulation), real placebo group and placebo real group. Mean age, gender, length of illness, daily dose of L-DOPA, total UPDRS on pre-stimulation, MMSE, apathy, depression and motor threshold intensity were all not different between two groups. Stimulation intensity at 110% resting motor threshold ranged from 48 to 93 (mean value is 73 intensity, SD=14) (Table 1). Total UPDRS score ranged from 20 to 88 (mean 46, SD=17), and all participants were usually on state with no antipsychotic drugs.

<table>
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<th>1st Real &amp; 2nd placebo group (n=8)</th>
<th>1st Placebo &amp; 2nd Real group (n=7)</th>
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<tr>
<td>Mean age (years)</td>
<td>70.0 ± 8.4</td>
<td>76.0 ± 7.7</td>
<td>0.163*</td>
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<td>Male (%)</td>
<td>2 (25.0)</td>
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Figure 1: The rTMS was performed under two conditions (real rTMS and placebo stimulation) by randomized and cross over designed study. The patients were randomized into real-first (n = 8) and placebo-first (n = 7) groups. SDS: Zung Self-rating Depression Scale, UPDRS: the unified Parkinson’s disease rating scale
Apathy scale before rTMS ranged from 3 to 29 (mean 16, SD=8), and 6 participants (40%) with being more than 16 were diagnosed by apathy. Real rTMS improved the apathy scale by 3 points compared to the baseline (pre=18, post=15, p<0.05), whereas placebo stimulation worsened the score by 3 points (pre=15, post=18, p<0.05). When the changes of apathy scale between the two sessions (real rTMS and placebo stimulation) were evaluated with repeated-measures ANOVA, real rTMS produced a significant improvement in apathy score compared with placebo stimulation (p<0.05) (Figure 2). Stimulative procedure of real and placebo have no significant interaction (stimulation x timing: F=1.826, p=0.1818). SDS before rTMS ranged from 25 to 60 (mean 46, SD=9), and 6 participants (40%) with being more than 50 were diagnosed by depression. SDS statistically changed after real TMS (pre=47, post=42, p<0.05) and placebo stimulation (pre=44, post=46, p<0.05). When the changes of SDS between the two sessions (real rTMS and placebo stimulation) were evaluated with repeated-measures ANOVA, real rTMS produced a significant improvement in SDS compared with placebo stimulation (p<0.05) (Figure 3). Stimulative procedure of real and placebo have no significant interaction (stimulation x timing: F=2.963, p=0.079).
Real rTMS improved UPDRS by 10 points (pre = 46, post = 36, P=0.001), but UPDRS did not change after placebo stimulation (pre = 43, post = 43). When the changes of UPDRS between the two sessions were compared with repeated-measures ANOVA, real rTMS produced a significant improvement of UPDRS compared with placebo stimulation (p<0.01) (Figure 4). Stimulative procedure of 1st real and 2nd placebo was significantly more effective than 1st placebo and 2nd real with interaction (stimulation x timing: F=7.306, p=0.001). No side effects were observed in either real or placebo stimulation. Clinical factors (i.e., age, gender, disease duration (months), UPDRS score pre-rTMS, and daily dose of L-DOPA) did not influence the improvement of UPDRS and apathy scores by real rTMS. Between the two sessions there was no correlation among the changes in apathy scale, SDS and UPDRS scores.

**Discussion**

Apathy is defined as lack of motivation, manifested as diminished goal-directed behavior and cognition and decreased emotional engagement. It reduces the quality of life of PD patients with increasing functional dependency, as well as caregivers [19,20]. Dysfunction of the frontal-subcortical circuits is thought to be a cardinal pathophysiological feature of apathy in PD patients, because a reduction in dopaminergic afferents to the striatum disturbs connections between frontal lobe, caudate, anterior cingulate circuits and the basal ganglia [21,22]. This is also consistent with our report that a dopamine D2/3 receptor agonist was effective for the treatment of apathy [23]. Further, we have shown that cerebral blood flow is reduced in the dorsolateral frontal lobe and basal ganglia of apathetic stroke patients [13,24]. Hypoactivity in these regions appears to contribute to the emergence of apathy symptoms. Mottaghy et al. reported that rTMS at the DLPFC caused a local increase of regional cerebral blood flow in several neocortical areas [25]. Boggio et al. reported that rTMS stimulation improves the score in the Stroop test, which reflects attention and executive function associated with the frontal lobe [9]. Further, the score in the Stroop test was significantly correlated with the apathy scale score in PD and depressed patients [26,27]. These reports also suggest that rTMS may be effective to improve frontal lobe functions.

We adopted the method described in the University Tokyo research group’s reports, in which the SMA was stimulated for the treatment of PD [2,3,12]. The SMA is functionally impaired in PD patients, because of decreased positive efferent feedback involving the motor circuit. High-frequency rTMS activates neurons of the SMA and ameliorates motor circuit dysfunction [2,3]. The majority of previous studies have employed a rTMS intensity of 110% of MT for the treatment of depression, as we did here [6]. Our findings are in agreement with the University Tokyo research group’s reports as regards UPDRS improvement in response to rTMS over the SMA [2,12]. Although four reports showed rTMS on let DLPFC in PD are effective on depressed PD [6-9], our rTMS on SMA had been also effective on depression. Our stimulus intensity 5Hz is lower than these reports
(10–15Hz) and our stimulus position SMA is different from left DLPFC in four reports. Although we could not clearly explain why placebo stimulation worsened depression scale, only placebo stimulation without any drug control might affect depressive state.

There is some uncertainty as to whether stimulation of the SMA can cause activation of the frontal-subcortical network, which is compromised in patients with apathy. SMA has been reported to be activated in association with DLPFC activation in several cognitive activities, such as the Stroop task [28] and retrieval of motor memory following interleaved practice and skill learning [29], suggesting that there is functional connectivity between the SMA and DLFPC. We speculate that SMA stimulation by rTMS indirectly up-regulates neuronal activity in the frontal-subcortical circuit and contributes to improvement of apathy and depression in PD patients. The degree of improvement in apathy and depression was not correlated with UPDRS score in our study, so it is unlikely that alleviation of apathy and depression was caused by improved motor performance [7].

One randomized controlled trial for the treatment of 1Hz or 10 Hz rTMS of apathy in 70 PD patients showed no efficacy [12]. This research design was a weekly intervention with 8 times stimulation with daily 1000 pulses (total amount of 8000 pulses for 8 weeks). It was different from our rTMS stimulative frequency 5 Hz with daily 500 pulses and total amount of 2500 pulses for 5 days. So our real rTMS design with every 5 days (2500 pulses per a week) might be slightly stronger than theirs (1000 pulses per a week). In rTMS intervention for PD with apathy, our report is second randomized trial with original stimulative method on every day compared to Shirota’s first report with weekly rTMS intervention [12].

Our rTMS session is consists of twelve days in admission and we set wash out period for two days between real and placebo stimulation. The lasting time of rTMS efficacy on motor cortical excitability varied from one hour to one month with various stimulative intensities. It can last one hour by frequency 1 Hz, 10 days by 5 Hz, and one month by 25Hz [30,31]. However effective time could be modified with various protocol (rTMS frequency, intensity, position and total pulses), definite protocol would be required to determine.

The main limitation of this study is the small sample size. However, the finding that the effect of rTMS was statistically significant even in as few as 15 patients indicates that the effect is robust. Another possible limitation is that rTMS was conducted at a location known to produce an improvement of motor dysfunction as well as apathy. We think a further study that includes a specific protocol to identify optimal brain regions for treatment to improve apathy would be worthwhile. And we need to investigate hippocampus as a treated target which is a central area in modulation of mood and apathy. rTMS with double cone coil might be useful for stimulating deep brain region.

References


