Reductions in Cortico-Striatal Hyperconnectivity Accompany Successful Treatment of Obsessive-Compulsive Disorder with Dorsomedial Prefrontal rTMS

Katharine Dunlop1,2, Blake Woodside1,3,4,5, Marion Olmsted4,5, Patricia Colton3,4,5, Peter Giacobbe2,3,4 and Jonathan Downar*6,1,2,3,4,6

1Institute of Medical Sciences, University of Toronto, Toronto, ON, Canada; 2MRI-Guided rTMS Clinic, University Health Network, Toronto, ON, Canada; 3Department of Psychiatry, University Health Network, Toronto, ON, Canada; 4Department of Psychiatry, University of Toronto, Toronto, ON, Canada; 5Eating Disorders Program, University Health Network, Toronto, ON, Canada; 6Toronto Western Research Institute, University Health Network, Toronto, ON, Canada

Obsessive-compulsive disorder (OCD) is a disabling illness with high rates of nonresponse to conventional treatments. OCD pathophysiology is believed to involve abnormalities in cortico-striatal-thalamic-cortical circuits through regions such as dorsomedial prefrontal cortex (dmPFC) and ventral striatum. These regions may constitute therapeutic targets for neuromodulation treatments, such as repetitive transcranial magnetic stimulation (rTMS). However, the neurobiological predictors and correlates of successful rTMS treatment for OCD are unclear. Here, we used resting-state functional magnetic resonance imaging (fMRI) to identify neural predictors and correlates of response to 20–30 sessions of bilateral 10 Hz dmPFC-rTMS in 20 treatment-resistant OCD patients, with 40 healthy controls as baseline comparators. A region of interest in the dmPFC was used to generate whole-brain functional connectivity maps pre-treatment and post treatment. Ten of 20 patients met the response criteria (≥50% improvement on Yale-Brown Obsessive-Compulsive Scale, YBOCS); response to dmPFC-rTMS was sharply bimodal. dmPFC-rTMS responders had higher dmPFC-ventral striatal connectivity at baseline. The degree of reduction in this connectivity, from pre- to post-treatment, correlated to the degree of YBOCS symptomatic improvement. Baseline clinical and psychometric data did not predict treatment response. In summary, reductions in fronto-striatal hyperconnectivity were associated with treatment response to dmPFC-rTMS in OCD. This finding is consistent with previous fMRI studies of deep brain stimulation in OCD, but opposite to previous reports on mechanisms of dmPFC-rTMS in major depression. fMRI could prove useful in predicting the response to dmPFC-rTMS in OCD.

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INTRODUCTION

Obsessive-compulsive disorder (OCD) is a severely disabling psychiatric disorder with a lifetime prevalence of 1–3% (Abramowitz et al, 2009). OCD is characterized by intrusive, anxiety-provoking, ego-dystonic thoughts (obsessions), and associated repetitive behaviors (compulsions) (American Psychiatric Association, 1994). Of the OCD patients, 40–60% are refractory to conventional pharmacological and behavioral therapies (Pallanti et al, 2002). It is therefore crucial to develop novel therapies through a better understanding of the pathophysiology of OCD and the mechanisms of successful treatment.

Previous human and animal studies suggest that abnormalities in the cortico-striato-thalamic-cortical (CSTC) circuitry may be central to OCD pathophysiology (Menzies et al, 2008; Harrison et al, 2009; Ahmari et al, 2013; Admon et al, 2014). In healthy humans, specific CSTC loop circuits are important for self-regulation of affect, cognition, and behavior (Marsh et al, 2009; van den Heuvel et al, 2010; Lipsman et al, 2013). In OCD, these circuits display structural abnormalities relative to controls: volumetric gray matter reductions and reduced white matter integrity in the anterior cingulate cortex (Kühn et al, 2013), gray matter reductions in the orbitofrontal cortex (Rotge et al, 2010), and gray matter increases in thalamus and ventral striatum (Hou et al, 2013). On functional magnetic resonance imaging (fMRI), abnormal cortical-ventral striatal hyperconnectivity has been observed in OCD during a monetary incentive delay task (Beucke et al, 2012), during symptom provocation and during rest in many studies (Harrison et al, 2009; Cocchi et al, 2012; Figee et al, 2013; Anticevic et al, 2014). One study has shown the opposite, corticostrialtial hypoconnectivity, in a
group of unmedicated patients (Posner et al., 2014). In addition, altered anterior cingulate cortex metabolic activity has been observed in OCD on fluorodeoxyglucose–positron emission tomography and magnetic resonance spectroscopy (Perani et al., 1995; Ebert et al., 1997; Saxena et al., 2004, 2009; O’Neill et al., 2013). Taken together, these observations delineate a possible neuroanatomical substrate for OCD symptomatology.

Neuromodulation treatments offer a novel, anatomically targeted approach to refractory psychiatric conditions. Deep brain stimulation (DBS) has shown promising effects for OCD in recent studies (Greenberg et al., 2010); several CSTC targets have been explored, including the anterior limb of the internal capsule (Abelson et al., 2005), subthalamic nucleus (Mallet et al., 2008), and ventral striatum/nucleus accumbens (NAcc) (Denys et al., 2010; Figue et al., 2013). Regarding therapeutic mechanisms, a recent MRI study found that NAcc-DBS normalized excessive functional connectivity between NAcc and dorsomedial and dorsolateral prefrontal cortex (dPFC) in OCD patients; the degree of reduction correlated to the degree of symptomatic improvement (Figue et al., 2013).

Although DBS has shown promising effects in severe, refractory OCD cases, noninvasive forms of neuromodulation could be offered to a much wider range of patients. Repetitive transcranial magnetic stimulation (rTMS) could present a noninvasive alternative to DBS in OCD, if directed at a suitable stimulation target. rTMS to the dPFC, although successful in major depression (O’Reardon et al., 2007; Berlim et al., 2014), has shown minimal clinical benefit over sham in double-blind trials for OCD (Alonso et al., 2001; Sachdev et al., 2007). However, medial prefrontal targets appear more promising: 1 Hz rTMS of the supplementary motor area (SMA) and pre-SMA has achieved substantial symptom improvement in case reports and randomized controlled trials (Mantovani et al., 2006, 2010a, b). Likewise, with transcranial direct current stimulation, catholic but not anodal stimulation of the SMA has been reported to improve OCD symptoms (D’Urso et al., 2015).

A neighboring potential target is the dorsomedial prefrontal cortex (dmPFC), just anterior to the pre-SMA. Abnormally high resting state functional connectivity (Stern et al., 2012) and gray matter volume reductions in dmPFC (Radua et al., 2010) have been observed in OCD patients relative to controls. With NAcc-DBS (Figue et al., 2013), therapeutic efficacy correlated to reduction in excessive frontostriatal connectivity through the dmPFC regions. dmPFC-rTMS has not yet been studied in OCD. However, in major depressive disorder (MDD), a recent open-label study of 10 Hz dmPFC-rTMS for treatment-resistant MDD achieved $\geq 50\%$ symptom improvement in approximately half of the patients (Downar et al., 2014). Notably, therapeutic effects correlated to change in CSTC connectivity through the dmPFC on fMRI, as with NAcc-DBS in MDD (Salomons et al., 2014). These observations raise the possibility that dmPFC-rTMS might be able to engage a similar therapeutic mechanism through noninvasive means in OCD.

Here, we used resting-state fMRI to identify neural predictors and correlates of treatment response to 20–30 sessions of open-label, 10 Hz dmPFC-rTMS for refractory OCD. On the basis of our previous findings in MDD (Bakker et al., 2014; Downar et al., 2014; Salomons et al., 2014) and the aforementioned findings with NAcc-DBS in OCD (Figue et al., 2013), we hypothesized that pre-treatment resting-state dmPFC functional connectivity would predict response to dmPFC-rTMS in OCD, and that changes in resting-state dmPFC functional connectivity from pre- to post-treatment would correlate with the degree of symptomatic improvement.

**MATERIALS AND METHODS**

**Subjects**

Twenty OCD patients (4 male, mean age $= 37.3 \pm 15.5$, range $= 21–63$) participated in the study. OCD and comorbid Axis I and II disorders were diagnosed by a board-certified psychiatrist (authors JD, PG, PC, or BW) using DSM-IV criteria during a semi-structured clinical psychiatric evaluation incorporating the Mini International Neuropsychiatric Interview (MINI 6.0). Controls were screened using the same instrument. All patients reported at least one previous failed medication trial on clinical interview (ie not tolerated or clinically nonresponsive) (mean $= 5.7 \pm 4.1$ trials) and 19 had failed at least one attempt at cognitive/behavioral intervention (ie did not complete the intervention or clinical nonresponsive). Mean illness duration was $24.2 \pm 15.2$ years (range $= 8–54$). No patients had hoarding symptomatology. Comorbidities included MDD ($n=16$), bipolar disorder ($n=2$), anorexia nervosa ($n=5$), bulimia nervosa ($n=4$), post-traumatic stress disorder ($n=4$), and Tourette’s syndrome ($n=1$). No patients had a history of tics. Current medications included neuroleptic agents ($n=12$), selective serotonin reuptake inhibitors ($n=8$), serotonin-norepinephrine reuptake inhibitors ($n=3$), trazodone ($n=2$), lithium ($n=1$), and benzodiazepines ($n=10$); maximum daily benzodiazepine doses were 2 mg clonazepam, 4.5 mg bromazepam, and 0.75 mg alprazolam. Subjects were required to maintain a stable medication regimen for at least 4 weeks prior to and throughout rTMS treatment. All patients provided informed consent, and the study was approved by the University Health Network Research Ethics Board.

Forty healthy controls (17 male, mean age $= 34.88 \pm 11.76$, range $= 18–66$) were recruited to provide a comparator group for resting-state connectivity post hoc analyses. There was no difference in age and sex between the OCD and healthy control group (age $t_{48} = 1.85$, $p = n.s.$; sex $\chi^2 = 2.97$, $p = n.s.$). Healthy participants had no previous psychiatric diagnoses, no current psychiatric medication, and no current substance abuse or dependence, as verified by a screening interview by trained research staff. Healthy participants underwent MRI, but did not undergo a course of rTMS.

**Clinical Measures**

Clinical measures were collected at baseline and 2 weeks post treatment. The primary outcome measure was the Yale-Brown Obsessive Compulsive Scale (YBOCS) (Goodman et al., 1989); in this study, treatment response was defined as $\geq 50\%$ improvement on the YBOCS. Secondary clinical measures included the 17-item Hamilton Rating Scale for Depression (HamD17), Beck Depression...
Index II (BDI-II), and Beck Anxiety Index (BAI). Additional clinical variables comprised the duration of illness, number of previous hospitalizations and outpatient treatment programs, number of previous medication trials, and current medication type and dose. Kernel density estimation of the distribution of clinical responses was performed in Stata13 (College Station, TX, USA).

**Intervention**

dmPFC-rTMS was performed according a protocol we have previously reported for major depression (Downar et al, 2014), described in detail in the Supplementary Material. In summary, rTMS was delivered using the MagPro R30 system equipped with a Cool-DB80 coil (MagVenture, Farum, Denmark) and a Visor 2.0 neuronavigation system (Advanced NeuroTechnologies, Enschede, the Netherlands). Neuronavigation was performed for anatomical landmarking and co-registration of the brain to Talairach stereotaxic space, with the co-registered coil vertex coordinate (x0 y+60 z+60 in Talairach stereotaxic space), as performed in previously published work on dmPFC-rTMS (Bakker et al, 2014; Downar et al, 2014; Salomons et al, 2014). This location corresponds to approximately 25% of the distance from nasion to inion, slightly anterior to the location specified by previous authors targeting the pre-SMA for rTMS in OCD, which was at 35% of the nasion–inion distance (Mantovani et al, 2010a). Lateral coil orientation for preferential stimulation (Harmer et al, 2001; Terao et al, 2001) of the left then right dmPFC at 10 Hz, at 120% of the extensor hallucis longus muscle resting motor threshold, with a duty cycle of 5 s on, 10 s off, for 60 trains (3000 pulses per hemisphere per session) for 20 daily sessions on weekdays, with non-remitters offered extension to 30 sessions.

**MRI acquisition and Analysis**

MRI acquisition parameters follow a protocol we have previously described in detail (Salomons et al, 2014), presented in full in the Supplementary Material. In summary, patients underwent MRI 1 week before and after rTMS treatment, to acquire a T1-weighted (0.94 × 0.94 × 1.5 mm) followed by a 10-min resting-state, eyes-closed T2* series (3.4 × 3.4 × 5 mm, TR = 2 s). Healthy controls underwent a single session using identical parameters. Data preprocessing and analysis were performed in FSL (Jenkinson et al, 2012). Briefly, preprocessing included motion correction, slice-timing correction, spatial smoothing (6 mm FWHM Gaussian kernel), nuisance regression using 6 motion parameters and extracted white matter and cerebrospinal fluid mean time series, bandpass filter (0.009–0.09 Hz), and co-registration to the MNI-152 standard brain. Global signal regression was omitted from the pipeline to avoid the risk of introducing spurious anticorrelations in the results (Chai et al, 2012). Simple white matter and cerebrospinal fluid time series regression was performed as opposed to more complex techniques (eg, aCompCor) to maintain an identical pipeline to the atlas of resting-state functional connectivity used to obtain our seed as described below. FSL was then used to generate whole-brain maps of pre-treatment functional connectivity and of change in functional connectivity from pre-to post-treatment, to the dmPFC region-of-interest. The dmPFC region-of-interest (center of mass = MNI X 0, Y+38, Z+24) was defined from a resting-state connectivity-based atlas (Craddock et al, 2012), as previously for dmPFC-rTMS in MDD (Salomons et al, 2014). Other subcortical targets (the medial dorsal thalamus, three sites within ventral striatum, and subthalamic nucleus) were selected as exploratory seeds on the basis of their use as targets for DBS elsewhere in the literature, as reviewed in the Introduction; these seeds and the corresponding results are presented in the Supplementary Material.

The seed region-of-interest was co-registered to each subject, and its mean time series used as a regressor in a first-level analysis. To localize regions where pre-treatment functional connectivity correlated to treatment response, FSL’s FLAME mixed effects model (Beckmann et al, 2003) was then applied for the group-level analysis using the responder/nonresponder status of each subject as a categorical, group-level regressor.

To localize regions where the changes in functional connectivity from pre- to post-treatment correlated to the degree of treatment response, a within-subjects, fixed-effects general linear model analysis was performed for each subject and the dmPFC seed region. For group-level analysis, these individual-subject beta-weighted change maps (ie, post- minus pre-rTMS) were then entered as first-level statistical maps into a between-subjects, mixed-effects linear regression analysis, using the responder/nonresponder status of each subject as a categorical, group-level independent variable.

Corrections for multiple comparisons were performed using Gaussian random field theory (Z > 1.96, cluster significance p < 0.05 corrected).

Parameter estimates for individual subjects’ functional connectivity values (mean z-score) were then extracted for post hoc analysis. An 8-mm sphere centered at the peak cluster voxel (masked by the relevant cluster to ensure specificity) was registered from standard space to each individual subject using the transformation matrix from the original registration, extracting the mean z-score values from relevant connectivity maps. Healthy control parameter estimates were extracted in the same way.

**RESULTS**

**Clinical Outcomes**

Subjects completed a mean 21.3 ± 4.1 sessions of 10 Hz dmPFC-rTMS (range = 14–30). Treatment was well tolerated, with no serious or treatment-limiting adverse effects occurring. One subject discontinued treatment after 14 sessions owing to nonresponse, and was analyzed as a nonresponder using baseline measures.

Across all subjects, baseline YBOCS scores significantly decreased from 30.5 ± 4.3 to 18.4 ± 10.8 (Wilcoxon rank-sum test, W18 = 3.41, p = 0.007). However, kernel density estimation revealed a sharply bimodal response distribution, with distinct responder and nonresponder subpopulations (Figure 1); these subpopulations are therefore considered separately hereafter. Ten of 20 subjects met the response criterion of ≥ 50% improvement. Among responders,
YBOCS scores decreased 67.2%, from 29.3 ± 4.6 to 9.6 ± 3.9 (W₁₀ = 2.81, p = 0.005); among nonresponders, YBOCS scores decreased nonsignificantly by 11.4%, from 31.7 ± 4.1 to 28.1 ± 7.8 (W₁₀ = 1.45, p = 0.15). Of note, responders and nonresponders did not differ in YBOCS severity at baseline (29.3 ± 4.6 vs 31.7 ± 4.1; Mann–Whitney U₁₀ = 1.21, p = 0.22).

Regarding secondary measures, across all subjects, depression severity significantly improved on HamD₁₇ from 17.7 ± 7.7 to 9.9 ± 7.3 (t₁₀ = 3.08, p = 0.008) and on BDI-II from 29.9 ± 15.8 to 19.4 ± 15.6 (t₁₀ = 3.27, p = 0.005). Again, outcomes were sharply dichotomous; YBOCS responders improved significantly on HamD₁₇ from 15.8 ± 5.8 to 5.8 ± 4.0 (t₉ = 3.64, p = 0.01) and on BDI-II from 26.9 ± 18.4 to 8.7 ± 9.6 (t₉ = 5.51, p = 0.0006). YBOCS nonresponders showed no significant improvement on HamD₁₇, from 18.0 ± 6.8 to 13.7 ± 7.6 (t₉ = 1.24, p = 0.25), or on BDI-II, from 30.6 ± 16.3 to 29.7 ± 13.7 (t₉ = 0.37, p = 0.72). Anxiety symptomatology significantly improved on BAI from 28.7 ± 15.0 to 15.9 ± 12.4 (t₁₀ = 4.01, p = 0.0009) across all subjects, and from 29.8 ± 18.3 to 10.4 ± 10.9 (t₉ = 5.11, p = 0.0009) in YBOCS responders. YBOCS nonresponders showed nonsignificant improvement on BAI from 25.5 ± 13.1 to 20.2 ± 13.4 (t₉ = 1.47, p = 0.18). There was no difference in baseline severity for YBOCS responders vs YBOCS nonresponders on HamD₁₇ (15.8 ± 5.8 vs 18.0 ± 6.8, t₁₀ = 0.696, p = 0.49), BDI-II (26.9 ± 18.4 vs 30.6 ± 16.3, t₁₀ = 0.476, p = 0.64), or BAI (29.8 ± 18.3 vs 25.5 ± 13.1, t₁₀ = 0.602, p = 0.55).

A Spearman’s rank correlation coefficient was performed on baseline psychometric and clinical measures to determine whether any clinical factors predicted treatment outcome. No baseline factors (number of rTMS sessions, age, baseline severity, duration of illness, number of failed medications/treatments, comorbidities or current medication (antipsychotics, SSRI, or benzodiazepines), or baseline HamD₁₇, BDI-II, or BAI) showed a significant correlation to treatment outcome, either before or after Bonferroni correction for multiple comparisons.

### MRI Predictors of Treatment Response

Analysis of pre-treatment resting-state fMRI data revealed significant differences in baseline functional connectivity between responders and nonresponders for several exploratory seed regions-of-interest, including ventral rostral putamen to the dmPFC region (p = 0.05) (see Supplementary Materials). The contrast of baseline functional connectivity between responders and nonresponders using the a priori dmPFC region showed trend toward higher connectivity among responders; however, the difference did not reach significance (t₁₀ = 2.01, p = 0.06).

### MRI Correlates of Treatment Response

Comparisons of pre-treatment and post-treatment resting-state fMRI data also revealed significant differences between responders and nonresponders in how functional connectivity to the seed regions changed over the course of treatment (Table 1). For the dmPFC seed, successful treatment response was associated with increased connectivity to the bilateral pre- and post-central gyrus and left precuneus, and decreased connectivity to the bilateral caudate nucleus, midbrain, thalamus, superior frontal gyrus, and right hippocampus (Table 1, Figure 2a). Responders showed significant decreases in dmPFC functional connectivity to the bilateral caudate (pre-treatment z = 4.81 ± 1.27, post-treatment z = 1.65 ± 1.16, t₈ = 2.61, p = 0.03), and thalamus (pre-treatment z = 0.64 ± 1.24, post-treatment z = −3.37 ± 1.01, t₈ = 2.57, p = 0.03). Conversely, nonresponders showed significant increases in functional connectivity from dmPFC to the hippocampus (pre-treatment z = −3.19 ± 0.83, post-treatment z = −0.86 ± 0.88, t₈ = 3.24, p = 0.01) and midbrain (pre-treatment z = −3.71 ± 1.00, post-treatment z = −1.20 ± 0.86, t₈ = 2.50, p = 0.04). Again, across all subjects, the percent YBOCS improvement correlated significantly to the degree of reduction in dmPFC functional connectivity to the caudate (r₁₀ = −0.56, p = 0.02) and hippocampus (r₁₀ = −0.58, p = 0.02).

Compared with healthy controls, responders’ dmPFC-caudate connectivity was significantly higher at baseline.
Table 1 Brain Regions Where the Pre-to-Post Treatment Change in Functional Connectivity to the dmPFC Differed Significantly Between rTMS Responders and Nonresponders

| Seed       | Region              | Brodmann area | X   | Y   | Z   | Peak Z score |
|------------|---------------------|---------------|-----|-----|-----|--------------|
| dmPFC      |                     |               |     |     |     |              |
| FC increase in Resp > Nonresp |                     |               |     |     |     |              |
| L          | Precuneus           | 7             | −6  | −48 | 62  | 3.93         |
| L          | Postcentral gyrus   | 7             | −16 | −30 | 54  | 3.83         |
| R          | Precentral gyrus    | 6             | 4   | −20 | 60  | 2.87         |
| R          | Postcentral gyrus   | 7             | 14  | −36 | 48  | 2.74         |
| FC reduction in Resp > Nonresp |                     |               |     |     |     |              |
| L          | Superior frontal gyrus | 6       | −8  | 14  | 6   | 2.72         |
| R          | Superior frontal gyrus | 6       | 12  | 26  | 58  | 3.33         |
| L          | Caudate nucleus     | 6             | −6  | −14 | 2   | 3.02         |
| R          | Thalamus            | 22            | −4  | 8   | 3.07 |
| L          | Thalamus/Putamen    | 30            | −22 | −8  | 3.39 |
| R          | Hippocampus         | −6            | −28 | −10 | 2.57 |
| B          | Dorsal midbrain     | −2            | −16 | 16  | 2.61 |

Abbreviations: dmPFC, dorsomedial prefrontal cortex; FC, functional connectivity; MNI, Montreal Neurological Institute; Nonresp, nonresponder; Resp, responder. All activations are Gaussian random field theory corrected for multiple comparisons at a cluster threshold $p < 0.05$.

DISCUSSION

To our knowledge, this is the first case series using fMRI to identify neural predictors and correlates of response to any form of noninvasive brain stimulation in OCD. Previous studies of rTMS in OCD have encountered little therapeutic...
benefit with stimulation of lateral prefrontal targets such as the dlPFC (Alonso et al, 2001; Prasko et al, 2006; Sachdev et al, 2007; Sarkhel et al, 2010), but somewhat more success with medial prefrontal targets such as the SMA or pre-SMA (Mantovani et al, 2006, 2010a,b; Kumar and Chadda, 2011). The results of the present study are congruent with this latter literature in suggesting that rTMS of a slightly more anterior medial prefrontal target, the dmPFC, can also yield substantial symptom reduction in a proportion of OCD cases, even when multiple previous medication trials have failed. However, as previously observed in patients with depression (Downar et al, 2014), dmPFC-rTMS outcomes appear sharply bimodal in OCD.

In keeping with our first hypothesis, responders showed significant differences from nonresponders in baseline dmPFC-striatal resting-state functional connectivity. Furthermore, in keeping with our second hypothesis, the amount of symptomatic improvement correlated to the degree of reduction in functional connectivity through a very similar frontal-striatal-thalamic-subthalamic circuit connecting the dmPFC to VS, caudate nucleus, thalamus, and midbrain. In relation to healthy controls, responders initially showed hyperconnectivity from dmPFC to caudate before treatment, and this hyperconnectivity normalized following treatment; no such pattern was evident in nonresponders.

These observations are remarkably consistent with the recently reported results of an fMRI study of therapeutic mechanisms in OCD, using ventral striatum-DBS rather than rTMS of the dmPFC (Figee et al, 2013). In that study, OCD patients also showed baseline functional hyperconnectivity from ventral striatum to a dmPFC region just immediately anterior to the region identified in Figure 2 of the present study. DBS reduced functional connectivity from ventral striatum to dmPFC, and the degree of reduction correlated well with the amount of symptomatic improvement. The results of the present study suggest that similar therapeutic effects in OCD might also be achieved via noninvasive stimulation of the more superficial dmPFC, via a similar mechanism. Notably, the circuit identified in the present study also includes nodes in lateral orbitofrontal cortex, ventral striatum, medial thalamus, and subthalamic nucleus, each of which has shown promise as a therapeutic target for either DBS (Abelson et al, 2005; Mallet et al, 2008; Denys et al, 2010; Lipsman et al, 2013), or in the case of lateral orbitofrontal cortex, rTMS (Ruffini et al, 2009).

More generally, the observations of the present study support the growing body of evidence that OCD pathophysiology may arise from functional hyperconnectivity through specific cortico-striatal loop circuits projecting from ventral striatum to the medial PFC (Harrison et al, 2009; Sakai et al, 2011; Beucke et al, 2013). Recent studies, in larger patient samples, have even suggested that different OCD symptom dimensions may map on to anatomically distinct cortico-striatal pathways (Harrison et al, 2013). Animal studies also suggest that the dynamics of the hyperactivation may be important. For example, a recent study showed that repeated exposures to brief optogenetic hyperactivation of an orbitofrontal cortex-ventral striatal circuit in mice led to progressively increased compulsive grooming behaviors that were reversible by fluoxetine, a standard pharmacotherapy for OCD (Ahmari et al, 2013). These more nuanced techniques could lead to a better understanding of the variability of response to current neuromodulation therapies in OCD.

We also note that the fMRI predictors and correlates of successful 10 Hz dmPFC-rTMS for OCD appear opposite to those we have previously reported for the same intervention in major depression (Salomons et al, 2014). In the MDD patients, low baseline connectivity from dmPFC to the putamen and thalamus predicted better response to treatment, and the degree of increase in frontal-striatal-thalamic connectivity correlated to symptomatic improvement. The results of the present study suggest that dmPFC-rTMS may exert therapeutic effects via a similar CSTC pathway in both MDD and OCD, but via opposite mechanisms (i.e., via reduction of a pathologically high baseline CSTC connectivity in OCD, rather than via strengthening of a pathologically low baseline CSTC connectivity in MDD). These seemingly contrary findings could potentially be reconciled if dmPFC-rTMS exerts its therapeutic effects on OCD and MDD not directly but indirectly, by relieving intrusive thoughts, as suggested by a recent study (Carew et al, 2013). However, this issue will require further study.

One important difference in technique between the present study and previous studies of rTMS in OCD targeting the medial wall is that the previous studies used 1 Hz rather than 10 Hz stimulation. Classically, 1 Hz stimulation is considered inhibitory, and 10 Hz stimulation excitatory (Hallett, 2007); suppression of overactive regions in SMA and pre-SMA provided a rationale for using 1 Hz stimulation in these previous studies (Mantovani et al, 2006, 2010a,b). Yet it is increasingly recognized that the effects of many rTMS protocols, including both 1 and 10 Hz stimulation, can be quite heterogeneous: a substantial proportion of individuals show ‘paradoxical’ excitatory responses to 1 Hz stimulation or inhibitory responses to high-frequency stimulation, both on motor evoked potentials (Maeda et al, 2000; Hamada et al, 2013) and on resting-state fMRI (Eldaief et al, 2011). In the present study, as well as in our previous study of dmPFC-rTMS in MDD, the effects of 10 Hz stimulation on cortical-striatal-thalamic activity were in fact quite variable across individuals, with ‘paradoxical’ reductions in functional connectivity appearing in up to 40% of MDD patients in the previous study and in more than half of the OCD patients in the present study (Figure 3). Thus, it is possible that 1 Hz SMA-rTMS and 10 Hz dmPFC-rTMS are achieving successful OCD treatment outcomes via similar mechanisms, notwithstanding the differences in technique. Alternatively, individual patients may require different sites or patterns of stimulation to achieve a therapeutic effect, as has been reported for dPFC-rTMS in depression (Speer et al, 2009). The question of whether 1 Hz SMA-rTMS and 10 Hz dmPFC-rTMS treat similar or different subpopulations of OCD patients will be an important topic for future study. In either case, resting-state fMRI is likely to have an important role both in stratifying patients and in clarifying therapeutic mechanisms.

One limitation of the present study is use of an open-label design, leaving open the possibility that the observed symptomatic improvements were due to non-specific or placebo effects. However, it should be noted that in the setting of refractory OCD, previously reported effects of sham rTMS are relatively minimal, ranging from 1% to ~20% improvement in YBOCS scores; indeed, even active
stimulation of lateral targets such as dIPFC has consistently achieved less than 25% YBOCS improvement across several independent studies (Alonso et al, 2001; Sachdev et al, 2007; Mantovani et al, 2010a; Sarkhel et al, 2010). Thus, it is unlikely that placebo effects can fully account for the present observations of a ~ 40% overall improvement in the present study, a bimodal outcome distribution, distinct patterns of functional connectivity through the dmPFC-ventral striatal target circuit in responders vs nonresponders (who were not otherwise distinct on clinical measures), distinct patterns of change in this circuit in responders and nonresponders, and the concordance of these results with the previous findings of an independent study using DBS rather than rTMS. None-theless, replication of the present findings under sham-controlled conditions would be an important next step in this line of research.

Another potential criticism of the current study relates to the small sample size. Although the sample size here is comparable with or larger than that in other dmPFC rTMS-MRI and DBS-MRI studies (Figee et al, 2013; Salomons et al, 2014), open-label and randomized control rTMS treatment studies for OCD (Mantovani et al, 2006, 2010a,b) and neuroimaging studies for OCD (Admon et al, 2012), it is nonetheless underpowered to capture the full heterogeneity of OCD symptomatology, comorbidity, and treatment types. Hence, the present observations are as yet insufficient to properly address the question as to whether dmPFC-rTMS response could also be predicted on the basis of clinical features such as OCD subtype (eg, hoarding vs checking), presence or absence of comorbid symptoms (eg, major depression, tics), or adjunctive treatment types (eg, use of SSRIs or neuroleptics). They also cannot yet address the question of whether dmPFC-rTMS might selectively improve some OCD symptom dimensions but not others, as might be suspected if different symptom clusters map reliably on to distinct neural substrates. Resolution of such issues must await a larger sample of patients.

In summary, 10 Hz dmPFC-rTMS may offer a promising, noninvasive therapeutic option for medication-refractory OCD, achieving ≥ 50% reductions in YBOCS scores in 50% of the patients in the sample. In agreement with recent findings with ventral striatal DBS for OCD, rTMS may be most effective in patients with greater hyperconnectivity between dmPFC and ventral striatum on resting-state fMRI. Also in agreement with DBS findings, therapeutic effects of dmPFC-rTMS correlated with reductions in dmPFC-ventral striatum functional connectivity. A randomized controlled trial incorporating a sham rTMS arm would be a logical next step in evaluating dmPFC-rTMS as a noninvasive alternative to DBS in medication-refractory OCD. Future studies of rTMS in OCD may also benefit from using fMRI to characterize cortico-striatal connectivity prior to treatment, in order to predict response and/or tailor the stimulation target and parameters in individual patients (Fox et al, 2013). Properly optimized, rTMS could evolve into a potent, novel treatment option for patients faced with this challenging and crippling illness.

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