

Novel Tools and Methods

Persistent Enhancement of Hippocampal Network Connectivity by Parietal rTMS Is Reproducible

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Abstract

Wang et al. (2014) found that that five daily sessions of repetitive transcranial magnetic stimulation (rTMS) of the posterior parietal cortex (PPC) significantly increased functional connectivity (FC) in a network centered on the hippocampus, and caused a correlated increase in memory performance. However, this finding has not been reproduced independently and the requirement for five sessions has not been validated. We aimed to reproduce the imaging results of this experiment, focusing on hippocampal FC changes and using fewer days of rTMS. We measured resting state FC before and after three (N = 9) or four (N = 6) consecutive daily PPC rTMS sessions, using similar delivery parameter settings as Wang et al. (2014). Eight subjects received 3 d of rTMS delivered to the vertex as a control. We employed whole-brain and hypothesis-based statistical approaches to test for hippocampal FC changes. Additionally, we calculated FC in 17 brain networks to determine whether the topographic pattern of FC change was similar between studies. We did not include behavioral testing in this study. PPC, but not vertex, rTMS caused significant changes in hippocampal FC to the same regions as in the previous study. Brain-wide changes in hippocampal FC significantly exceeded changes in global connectedness, indicating that the effect of PPC rTMS was specific to the hippocampal network. Baseline hippocampal FC, measured before receiving stimulation, predicted the degree of rTMS-induced hippocampal FC as in the previous study. These findings reproduce the imaging findings of Wang et al. (2014) and show that FC enhancement can occur after only three to four sessions of PPC rTMS.

Key words: fMRI; functional connectivity; hippocampus; TMS

Significance Statement

One of the most striking recent findings in the area of neuromodulation is that of Wang et al. (2014), who reported that posterior parietal cortex (PPC) stimulation increased functional connectivity (FC) in a network related to declarative memory and centered on the hippocampus, a result with great potential experimental and clinical utility. We used a similar paradigm, with shorter treatment duration and reproduced the effects on connectivity, including specificity for the hippocampal network and dependence on the magnitude of baseline hippocampal connectivity. These results confirm and extend the initial finding and validate the technical approach.

Introduction

Enhancing memory in patients and healthy individuals is a potential application of repetitive transcranial mag-

netic stimulation (rTMS). Network connectivity modulation with non-invasive brain stimulation has been studied mostly in motor and procedural learning networks (Mu-

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ellbacher et al., 2002; Baraduc et al., 2004; Wilkinson et al., 2010, 2015; Hotermans et al., 2008; Rosenthal et al., 2009; lezzi et al., 2010; Teo et al., 2011) and the effects have not been shown to last longer than minutes or hours (Thut and Pascual-leone, 2010). The declarative memory system, on the other hand, has been less explored with rTMS, despite the fact that declarative memory deficits are among the most common and debilitating problems in neurology (Vakil, 2005; Nestor et al., 2006). Wang et al. (2014) increased declarative memory and resting hippocampal network functional connectivity (FC) by delivering multiple-session rTMS to individualized targets in the posterior parietal cortex (PPC), which is connected with the hippocampus via the retrosplenial and paraphippocampal cortices (Mesulam et al., 1977; Cavada and Goldman-Rakic, 1989). The FC increase and memory improvement persisted for 24 h after the final rTMS session and, with reduced strength, for up to approximately two weeks (Wang et al., 2014; Wang and Voss, 2015).

The Wang et al. (2014) findings are a dramatic demonstration of physiologic engagement of a specific brain target with correlated behavioral improvement and are, in this respect, unique in the noninvasive neuromodulation field. However, concern has grown over the rate of false positives in functional neuroimaging (Poldrack et al., 2017) and noninvasive neuromodulation (Nahas et al., 2008; Héroux et al., 2015), resulting in calls for reproduction of results. For example, Héroux et al. (2015) found that only between 45% and 60% of experienced researchers were able to reproduce a rTMS effect.

In this study, we used a similar paradigm to that of Wang et al. (2014), with identical targeting procedures and stimulation parameter values, but with fewer stimulation sessions and without memory testing. We also preprocessed the data somewhat differently and used vertex stimulation, instead of subthreshold or motor stimulation, as our control condition. Although the original researchers collaborated on this study and shared unpublished data and techniques with us, all data collection, implementation, and analysis were performed independently.

Materials and Methods

Subjects

Twenty-three healthy adults (nine females; age = 19-31 years), free of neurologic or psychiatric disorders or medications acting on the central nervous system,

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participated in the study. Fifteen received active rTMS delivered to the PPC and eight underwent a control procedure with identical stimulation applied to the vertex. All subjects reported being right-handed and passed screening for contraindications to TMS (Rossi et al., 2009) and MRI. Written informed consent was obtained and the study was approved by the local Institutional Review Board.

Procedures

All subjects underwent, in order, baseline scanning, three or four consecutive daily rTMS sessions, and a post-rTMS scan. Baseline scanning included an anatomical localizer, structural scan (for functional scan co-localization with anatomy, and neuro-navigation), a single resting state scan, and diffusion tensor imaging (not reported here). Nine subjects received three consecutive daily sessions of rTMS delivered to the PPC, six received four daily PPC sessions, and eight received three daily sessions of identical rTMS delivered to the vertex (see rTMS, below). The interval between rTMS sessions was ~24 h.

Twelve of our PPC subjects participated in a separate study to find the minimum number of days required to produce a conservative criterion change in hippocampal FC. We found no measurable difference in response between subjects receiving 3 and 4 d of rTMS (W=36, p=0.327, 95% CI [-0.074333, 0.210993]), so all were included here. The number of stimulation sessions in this study differed from Wang et al. (2014), who delivered stimulation on five consecutive days.

Unlike Wang et al. (2014), we used rTMS at the vertex, which produces auditory and somatosensory stimulation, but no significant changes in FC (Jung et al., 2016), as our control condition (see Discussion). Subjects underwent the first rTMS session within 36 h of baseline scanning. The second MRI session occurred on the day after the final rTMS session and within 3 h of the time of day of the first scanning session. Subjects were blind to the specific intent of the study and the stimulation condition.

fMRI acquisition and preprocessing

MRI was performed on a Siemen's Magnetom 3T scanner using a 16-channel head coil with foam padding to prevent head movement. Subjects were fitted with earplugs and supplied with headphones to protect hearing. During resting scans, subjects were instructed to lie still with their eyes open.

Blood oxygen level-dependent (BOLD) data were recorded with a T2*-weighted gradient-echoplanar imaging sequence (EPI: TR = 2000 ms, TE = 27 ms, flip angle = 90°, 36 transversal contiguous interleaved slices per volume, 3.0 slice thickness, FOV 22 \times 22 cm, matrix size 64 \times 64, voxel size = 3.4 \times 3.4 \times 3.0 mm; scan length \sim 6.8 min). We acquired structural images with a magnetization-prepared rapid gradient echo sequence (MPRAGE; TR = 2530 ms, TE = 3.03 ms, 176 slices per volume, 1-mm thickness, FOV = 25.6 \times 25.6 cm², 256 \times 256 acquisition matrix, voxel size = 1.0 mm isotropic, 206 volumes, 6.83 min).



We processed the images with analysis of functional images (AFNI; Cox, 1996; RRID:SCR_005927) software. The first five volumes of 206 were removed to ensure that magnetization was stabilized. Preprocessing included motion correction, slice-timing correction to the first slice, functional/structural affine co-registration to Talairach space (TT_N27; Talairach and Tournoux, 1988), resampling to 2.0 mm isotropic voxel resolution, spatial smoothing using a 4-mm full-width half maximum (FWHM) Gaussian kernel, and linear detrending. We then scaled each voxel time series to a mean of 100, with a range of 0-200 and regressed head motion from each voxel time series using the mean and derivatives of six parameter estimates (pitch, roll, yaw, and rotation around each axis). Unlike Wang et al. (2014), we did not bandpass filter our data because test-retest reliability increases as the high pass cutoff is raised and even eliminated (Shirer et al., 2015). However, we achieved a high-pass filter via linear detrending using a 2nd or 3rd order polynomial, depending on the subject. We used spatial smoothing, which was omitted by Wang et al. (2014). Finally, frames which included movement displacement >0.3 mm were censored before statistical analysis to prevent inflated correlations (Power et al., 2012). We used a threshold of 0.3 mm of average head displacement across all frames, including censored ones, during any scan to exclude subjects (one subject).

We reprocessed and reanalyzed data from Wang et al. (2014), which were acquired on a Siemens 3T TIM Trio with a 32-channel head coil. Structural (MPRAGE T1-weighted scans, TR = 2400 ms, TE = 3.16 ms, voxel size = 1 mm³, FOV = 25.6 cm, flip angle = 8°, 176 sagittal slices) and functional whole-brain BOLD EPI (TR = 2500 ms, TE = 20 ms, voxel size = $1.72 \times 1.72 \times 3$ mm³, FOV = 22 cm, flip angle = 80° , 244 volumes, 10.2 min). We handled them identically to our own data, but resampled to 1.5-mm isotropic voxel resolution.

rTMS targeting

We based our targeting procedure on Wang et al. (2014) who chose the PPC subregion that was maximally connected to the hippocampus in each subject. They searched the anterior/middle hippocampus for the voxel with maximal FC to the PPC and chose the PPC location where this FC was strongest as the stimulation target. We applied a similar technique. For subjects receiving PPC stimulation, we guided rTMS to the PPC location with maximum FC to a seed location in the hippocampus. In each subject, the PPC target search volume was a sphere of 15-mm radius, cut to exclude non-brain voxels, around Talairach location x = -47, y = -68, z = +36, which included the supramarginal and angular gyri. The search for the hippocampal seed voxel involved two approaches, both employing automated scripts. For the first approach (12 subjects), we chose the maximally connected hippocampal voxel from six preselected locations along the longitudinal aspect of the hippocampus in Talairach-Tournoux space (seed 1: x = -26, y = -10, z = -17; seed 2: x = -22, y = -16, z = -13; seed 3: x = -30, y = -17, z = -18= -14; seed 4: x = -30, y = -22, z = -12; seed 5: x = -30,

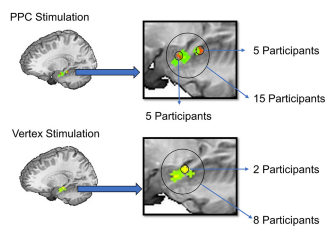


Figure 1. Seed locations from PPC (top; N=15) and vertex groups (bottom; N=8).

y = -27, z = -9; seed 6: x = -30, y - 32, z = -6). This deviated from the seeding procedure of Wang et al. (2014), who sampled only from the anterior/middle hippocampus. In the second approach (three subjects), we selected the maximally connected one of 97 preselected voxels in the anterior hippocampus. These included hippocampal voxels within 15 mm of the Talairach coordinates identified in Wang et al. (2014; x = -24, y = -18, z= -18). This approach was intended to provide wider sampling within the hippocampus. Figure 1 illustrates the seed locations for each subject. In both approaches, we created a 3-mm radius sphere around the coordinates of each voxel in the search and computed an average time series using the voxels in that sphere. We then searched the PPC sphere for the voxel with maximum correlation with the hippocampal seed, marked its location in standard space, and then back-transformed the location into subject space using the inverse matrix of the original affine transformation. Next, this location was transformed into a 3-mm radius sphere and overlaid on the subject's structural MRI for rTMS targeting with the Brainsight frameless stereotaxic system. For the PPC target, a stimulation trajectory was created in Brainsight, so that the plane of the coil was tangential to the scalp and the induced current field was oriented perpendicular to the long axis of the gyrus containing the stimulation target. For control stimulation, we located the vertex using the 10-20 International system (Steinmetz et al., 1989), and held the coil tangential to the scalp with the junction of the coil lobes in the sagittal axis.

rTMS

TMS was delivered with a MagStim Rapid² stimulator through a Double Airfilm coil. Wang et al. used a Nexstim eXimia NBS 4.3 air-cooled, MRI-guided system and a 70-mm figure eight coil. rTMS intensity was referenced to the individual motor evoked potential threshold, which was determined in the current experiment immediately before the first rTMS session using the TMS Motor Threshold Assessment Tool (MTAT 2.0; http://www.clinicalresearcher.org/software.htm). Stimulation parameter settings for PPC and vertex stimulation were identical to



those of Wang et al. (2014), i.e., 2-s trains at 20 Hz (40 pulses per train) with an intertrain interval of 28 s, at 100% of resting motor threshold (RMT). There were 40 trains, 1600 pulses, and a duration of 20 min per session.

FC calculations and voxel-wise analysis

For all hippocampal FC analyses, we conducted the following steps: Preprocessed data from the prestimulation and poststimulation resting state scans were seeded at the hippocampal location maximally connected with the PPC in the prestimulation scan, the area found for rTMS targeting. We created a 3-mm radius sphere around this location and averaged the BOLD time series of all voxels within it to derive a single hippocampal time series. Pearson's r values were then computed for the correlation between this time series and that from every voxel in the rest of the brain. Finally, all r values were r-to-r Fisher transformed to form a final connectivity metric $[z_{(r)}]$ across voxels for each scan.

Whole brain changes in hippocampal network FC and comparison to Wang et al. (2014)

To identify areas where PPC rTMS caused significant changes in hippocampal FC, and to see whether they were in the same areas reported by Wang et al. (2014), z_(r) values for each subject and time point, prestimulation and poststimulation, were fed into AFNI's 3dttest++ command for comparison. A group mask excluded ventricles and white matter. The results were false discovery rate (FDR) corrected at q = 0.05. We applied Bonferroni corrected post hoc tests to significant clusters in regions where Wang et al. (2014) reported significantly greater hippocampal FC increases with active compared to sham rTMS. These included the precuneus/retrosplenial, fusiform, lateral parietal, and superior parietal areas ($\alpha =$ 0.05/4 = 0.0125). Wilcox rank-sum testing was used for significance testing since these data were non-normally distributed.

Hypothesis-based comparison to the Wang et al. (2014) results

We performed this analysis to see whether PPC rTMS in the current study caused significant increases in hippocampal FC within a mask of regions showing significant hippocampal FC change in the reanalyzed data of Wang et al. (2014). To determine this region-of-interest, we searched for areas of the posterior left hemisphere that showed a significant increase in hippocampal FC after active rTMS, relative to sham, and calculated z_(r) values as described above. For each subject in the data set of Wang et al. (2014), the prestimulation correlation map was subtracted from poststimulation map, and the pre-sham map from the post-sham map. We then fed these subtractions into AFNI's 3dttest++ command for contrast. Like Wang et al. (2014), we applied a cluster size threshold of 290 voxels and identified a cluster encompassing the left precuneus and medial occipital lobe (left precuneus/occipital cortex; LPOC). We created a mask from these regions by applying the 3dclust command in AFNI and resampling the mask to the geometry of our own dataset (2-mm isotropic voxels). To account for variability across subjects, we dilated the mask by three voxels while restricting voxels to the left hemisphere. The pattern of results did not change based on the dilation of the mask. Finally, using the present data, we calculated prestimulation and poststimulation hippocampal FC in these regions and contrasted the resulting prestimulation and poststimulation $z_{(r)}$ values using a Wilcox rank-sum test, to look for a significant PPC rTMS-related change in FC between the hippocampus and the LPOC region, like that reported by Wang et al. (2014).

We also calculated the change in hippocampus-LPOC FC with vertex rTMS. Here, we used the hippocampal seed that was maximally connected with the PPC target at baseline, and the same automated script applied to the PPC subjects, to avoid potential bias in the selection of seeds. To determine whether changes in hippocampal FC with the LPOC mask were specific to PPC stimulation, we compared the rTMS-related change in hippocampus-LPOC FC between groups with the Mann-Whitney test. Additionally, to determine whether our results were affected by differences in sample size between groups, we performed a permutation test using matched sample sizes. This was performed by subtracting the mean FC change of the vertex group from the mean FC change in eight subjects randomly selected from the PPC group. This was performed 1000 times to form a distribution of possible outcomes, which we then compared to the observed mean difference.

Specificity analysis

To gauge the specificity of the Wang et al. (2014) effect on FC, we compared the changes in hippocampal FC and global connectedness (GC) occurring in the LPOC mask (LPOC-GC) with PPC rTMS. To calculate LPOC-GC, we found Pearson's r values for each voxel in the brain for the correlation of its time series with those of every other voxel. Next, we calculated the mean of all of the r values for each voxel within the LPOC mask (Gotts et al., 2012). The mean r values were then r-to-z Fisher transformed to create a GC value for each voxel. Finally, all voxel GC values in the LPOC mask were averaged.

As an additional control, we calculated the change in FC between the left dorsolateral prefrontal cortex (DLPFC) and the LPOC mask with the expectation that PPC stimulation would not significantly enhance FC between these regions. We created the DLPFC seed by forming a 3-mm radius sphere around Talairach and Tournoux location x = -41, y = 44, z = 5, a peak area of activation found during procedural learning (Poldrack et al., 2001). The mean time series from this sphere was then compared with that from every voxel in the LPOC mask. Finally, we took the mean of all r-to-z transformed values in the LPOC mask. Wilcox rank-sum tests were performed to determine whether the hippocampal-LPOC FC change differed significantly from the DLPFC-LPOC FC and LPOC-GC changes.

Comparison of topographic changes

We assessed the topographic pattern of hippocampal FC changes from PPC stimulation by calculating the change in hippocampal FC with 17 segregated networks (Yeo et al., 2011) using AFNI's 3dBrickStat command. We



also calculated within-network GC for this analysis using the time series of all voxels in each of the 17 networks. GC for each network was calculated as the mean $z_{(r)}$ value across all voxels in that network. We then compared the hippocampal FC and GC changes. The same steps were performed using the pre- to post-active stimulation data from Wang et al. (2014). We performed hippocampal-FC to GC comparisons for each study with one-sample, two-tailed, t tests, since these data were normally distributed.

Finally, to test the hypothesis that the magnitude of hippocampal FC changes across networks were correlated across studies, we conducted a simple correlation analysis to test this hypothesis ($\alpha=0.05$).

Correlation between baseline hippocampal FC and rTMS-induced changes in FC among hippocampal network nodes

The purpose of this analysis was to determine whether we could reproduce the finding of Wang et al. (2014), that baseline hippocampal FC predicted the degree of PPC rTMS-induced change in hippocampal FC among brain areas. We first found clusters of voxels in our data where rTMS produced a significant increase in hippocampal FC at a threshold of p < 0.01, with no spatial extent threshold. Like Wang et al. (2014), we applied a liberal threshold to include a range of change values. This resulted in 183 significant clusters, which we then divided into automated anatomic labeling (AAL; Tzourio-Mazoyer et al., 2002)-defined anatomic regions and all regions with >15 voxels were included in the analysis. The 15-voxel threshold was applied to ensure that each cluster contained enough voxels to calculate a reliable mean time series. This resulted in 95 clusters. We then formed a correlation matrix for each subject and time point by comparing the mean time series of each cluster with that of each other cluster (3dNetCorr). Next, we averaged the correlation matrices within each time point across subjects and subtracted the prestimulation correlation matrix from the poststimulation matrix. This resulted in a single matrix, which we sorted by baseline hippocampal FC. Then, to determine whether baseline hippocampal FC predicted the rTMS-induced change in FC, we plotted the baseline hippocampal FC of each cluster against the mean change in FC between that cluster and every other cluster. Finally, to determine whether these changes were specific to FC with the hippocampus, we performed the same analyses, but replaced hippocampal FC with GC for each cluster. Additionally, we re-sorted these matrices by region to reveal, qualitatively, areas where hippocampal nodes and nodes that increased in GC, showed the highest change in FC.

Statistical analyses

All analyses were conducted using R software. Shapiro–Wilks tests of normality were conducted before each analysis. Table 1 lists the specifications of each test, including critical values, the data used in each test, and confidence intervals. In the Results, an alphabetic code is listed with each test linking it to additional details in Table 1.

Results

The interval between rTMS sessions was 23.9 \pm 3.0 h for the PPC group and 24.3 \pm 2.7 h for the vertex group (non-significant; Table 1, a, W = 36, p = 0.327, 95% CI [-0.0743, 0.2110]). Head motion, calculated as average head frame displacement in six directions, did not significantly differ between scans (prestimulation vs poststimulation; Table 1, b, V = 167, p = 0.194, 95% CI [-0.0743, 0.2110]) or groups (parietal vs vertex; Table 1, c, W =199.5, p = 0.350, 95% CI [-0.0225, 0.0096]). The same was true for the number of censored TRs during denoising (prestimulation vs poststimulation; Table 1, d, V = 118.5, p = 0.155, 95% CI [-0.9999, 6.5000]; parietal vs vertex; Table 1, e, W = 218.5, p = 0.604, 95% CI [-3.0000, 0.00004]). Average head displacement was 0.089 \pm 0.005 mm per frame. The average number of censored TRs per scan was 5.348 ± 1.561.

Figure 2 shows regions that changed in FC with the hippocampus (FDR corrected, q=0.05) in the current sample. These changes were all increases. PPC rTMS produced significant increases in hippocampal FC in all of the areas reported by Wang et al. (2014), including left retrosplenial cortex (Table 1, f, V=7, $p=1.16\times10^{-3}$, 95% CI [0.0654, 0.2590]), left fusiform gyrus (Table 1, g, V=4, $p=4.27\times10^{-4}$, 95% CI [0.0951, 0.2132]), left lateral PC (Table 1, h, V=1, $p=1.22\times10^{-4}$, 95% CI [0.0777, 0.2034]), left superior PC (Table 1, i, V=2, $p=1.83\times10^{-4}$, 95% CI [0.0815, 0.2294]; all results Bonferroni corrected).

In our reanalysis of the Wang et al. (2014) data, the LPOC region of interest showed significantly increased FC with the hippocampus after active rTMS, relative to sham. In the current sample, we also found that PPC rTMS caused significant increases there (Table 1, j, V = 95, p = 0.048, 95% CI [0.0013, 0.2053]; Fig. 3A). This increase [$z_{(r)} = 0.20 \pm 0.04$; mean_(SEM)] was larger than, and opposite in direction to, the mean change after vertex rTMS ($z_{(r)} = -0.08 \pm 0.06$; Fig. 3A). The changes in the PPC rTMS group were significantly greater than the changes in the vertex group (Table 1, k, W = 93, p =0.034, 95% CI [0.0195, 0.3257]). Vertex stimulation did not cause changes in hippocampal-LPOC FC (Table 1, I, V = 7, p = 0.148, 95% CI [-0.055, 0.2357]). Resampling the group differences in hippocampal-LPOC FC in 1000 matched groups of eight subjects showed no instances where changes were greater in the vertex group, including those bounded by 95% of the distribution (Table 1, m, observed mean = 0.1795, 95% of distribution [0.02513, 0.03392]; Fig. 4). Thus, it is unlikely that our results were driven by differences in sample size between groups. Whole-brain analyses of hippocampal FC changes in the vertex group did not reveal any significant clusters (all p >0.05). The same was true when measuring FC from the vertex stimulation site.

In the current data, DLPFC-LPOC FC did not increase significantly after PPC rTMS (Table 1, n, V=81, p=0.2524, 95% CI [-0.0344, 0.1270]; Fig. 3A), but the DLPFC-LPOC FC change did not differ significantly from the hippocampal-LPOC FC change (Table 1, o, V=75, p=0.4212, 95% CI [-0.0636, 0.1593]), nor did LPOC-GC



Table 1. Statistics table indicating the results of all analyses

Manuscript	Figure	Sample	Data type	Data structure	Type of test	Multiple comparison correction		Statistics	p values	Confidence intervals
a		Current	Spacing between stimulation sessions	Non-normal distribution	Mann–Whitney (between groups; PPC group: participants receiving 3 vs 4 d of		R	W = 36	p = 0.327	Mean = 0.1092, 95% CI [-0.0743 0.2110]
b		Current	Average motion displacement	Non-normal distribution	stimulation) Wilcox rank-sum (all subjects; post vs pre)		R	V = 167	p = 0.194	Mean = 0.0074 959 CI [-0.0032, 0.0204]
С		Current	Average motion displacement	Non-normal distribution	Mann-Whitney (parietal vs vertex)		R	W = 199.5	p = 0.350	Mean = -0.0099 95% CI [-0.0225 0.0096]
d		Current	Number of censored trials	Non-normal distribution	Wilcox rank-sum (all subjects; post vs pre)		R	V = 118.5	p = 0.155	Mean = 2.5000 959 CI [-0.9999, 6.5000]
Э		Current	Number of censored trials	Non-normal distribution	Mann-Whitney (parietal vs vertex)		R	W = 218.5	p = 0.604	Mean = -2.1121 95% CI [-3.0000, 0.00004]
f	2	Current	Whole-brain FC analysis, retrosplenial cortex	Non-normal distribution	Wilcox rank-sum (within- groups; PPC group, post hoc test, post vs pre active stimulation)	Bonferroni	R	<i>V</i> = 7	$p = 1.16 \times 10^{-3}$	
g	2	Current	Whole-brain FC analysis, fusiform gyrus	Non-normal distribution	Wilcox rank-sum (within- groups; PPC group, post hoc test, post vs pre active stimulation)	Bonferroni	R	<i>V</i> = 4	$p = 4.27 \times 10^{-4}$	Mean = 0.1475, 95% CI [0.0951, 0.2132]
h	2	Current	Whole-brain FC analysis, lateral PC	Non-normal distribution	Wilcox rank-sum (within- groups; PPC group, post hoc test, post vs pre active stimulation)		R	<i>V</i> = 1	$p = 1.22 \times 10^{-4}$	Mean = 0.1331, 95% CI [0.0777, 0.2034]
i	2	Current	Whole-brain FC analysis, superior PC	Non-normal distribution	Wilcox rank-sum (within- groups; PPC group, post hoc test, post vs pre active stimulation)	Bonferroni	R	<i>V</i> = 2	$\rho = 1.83 \times 10^{-4}$	Mean = 0.1682, 95% CI [0.0815, 0.2294]
İ	3.4	Current	Hippocampal-LPOC FC changes (a priori)	Non-normal distribution	Wilcox rank-sum (within- groups; PPC group, post vs pre active stimulation)		R	<i>V</i> = 95	p = 0.048	Mean = 0.0867, 95% CI [0.0013, 0.2053]
k	3 <i>A</i>	Current	Hippocampal-LPOC FC changes (a priori)	Non-normal distribution	Mann-Whitney (between groups; PPC group vs vertex group)		R	W = 93	p = 0.034	Mean = 0.1367, 95% CI [0.0195, 0.3257]
I	4	Current	Hippocampal-LPOC FC changes	Non-normal distribution	Permutation test (between groups; PPC group vs vertex group)		R	Observed mean difference = 0.1795		95% of distribution [0.1098, 0.2536]
m	3 <i>A</i>	Current	Hippocampal-LPOC FC changes (a priori)	Non-normal distribution	Wilcox rank-sum (within groups; vertex group, post vs pre active stimulation)		R	<i>V</i> = 7	p = 0.148	Mean = -0.0477, 95% CI [-0.0554, 0.2357]
n	3 <i>A</i>	Current	DLPFC-LPOC FC changes (a priori)	Non-normal distribution	Wilcox rank-sum (within groups; PPC group, post vs pre active stimulation)		R	V = 81	p = 0.252	Mean = 0.0444, 95% CI [-0.0344, 0.1270]
0	3 <i>A</i>	Current	DLPFC and Hippocampal- LPOC changes (a priori)	Non-normal distribution	Wilcox rank-sum (within groups; PPC group, DLPFC-LPOC vs hippocampal-LPOC FC)		R	<i>V</i> = 75	p = 0.421	Mean = 0.0344, 95% CI [-0.0636, 0.1593]
р		Current	GC-LPOC changes (a priori)	Non-normal distribution	Wilcox rank-sum (within groups; PPC group, post vs pre active stimulation)		R	V = 94	p = 0.055	Mean = 0.0179, 95% CI [-0.0006, 0.0434]
q		Current	GC and Hippocampal-FC changes (a priori)	Non-normal distribution	Wilcox rank-sum (within groups; PPC group, post vs pre active stimulation) (Continued)		R	<i>V</i> = 92	p = 0.073	Mean = 0.0641, 95% CI [-0.0066, 0.1547]



Table 1. Continued

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Manuscript	Figure	Sample	Data type	Data structure	Type of test	correction	Program	Statistics	p values	Confidence intervals
r		Current	DLPFC and hippocampal target changes (control analysis)	Normally distributed	Paired t test (within-groups; PPC group, post vs per active stimulation)		R	$t_{(14)} = 0.949$	p = 0.359	Mean = 0.048 95% CI [-0.061, 0.157]
S		Current	DLPFC and stimulus location changes (control analysis)	Non-normal distribution	Wilcox rank-sum (within- groups; PPC group, post vs per active stimulation)		R	V = 68	<i>ρ</i> = 0.679	Mean = 0.026 95% CI [-0.103, 0.121]
t		Current	Changes in PPC GC	Non-normal distribution	Wilcox rank-sum (within- groups; PPC group, post vs per active stimulation)		R	V = 92	<i>ρ</i> = 0.073	Mean = 0.022 95% CI [-0.005, 0.050]
u	3 <i>B</i>	Current	Hippocampal FC changes within Yeo Networks	Normally distributed	Paired <i>t</i> test (within groups; PPC group, post vs pre active stimulation)		R	$t_{(16)} = 10.96$	$\rho = 7.6 \times 10^{-9}$	Mean = 0.0900, 95% CI [0.0725, 0.1073]
V	3 <i>B</i>	Wang et al. (2014)	Hippocampal FC changes within Yeo networks	Normally distributed	Paired t test (within groups; PPC group, post vs pre active stimulation)		R	$t_{(16)} = 11.27$	$\rho = 5.10 \times 10^{-9}$	Mean = 0.0169, 95% CI [0.0138, 0.0201]
W	3 <i>B</i>	Both samples	Hippocampal FC changes within Yeo networks	Normally distributed	Paired <i>t</i> test (between groups; current vs Wang, active stimulation)		R	$t_{(32)} = 8.75$	$\rho = 5.42 \times 10^{-10}$	95% CI [0.0560, 0.0900]
x	6	Both samples	Hippocampal FC changes within Yeo networks	Non-Normally distributed	Spearman correlation across samples (current and Wang)		R	r = 0.51	p = 0.037	95% CI [0.0389, 0.7956]
у		Both samples	GC changes within Yeo networks	Non-Normally distributed	Spearman correlation across samples (current and Wang)		R	<i>r</i> = 0.16	p = 0.536	95% CI [-0.3419, 0.5989]
Z	8	Current	Hippocampal FC changes in significant regions (p < 0.01)	Non-normal distribution	Spearman correlation (within groups; baseline hippocampal FC and hippocampal- FC changes)		R	<i>r</i> = 0.39	$p = 1.0 \times 10^{-4}$	95% CI [0.2002, 0.5453]
aa		Current	Hippocampal FC changes in significant regions (p < 0.01), outlier removed	Normally distributed	Pearson correlation (within groups; baseline hippocampal FC and hippocampal- FC changes)		R	$r_{(92)} = 0.47$	$p = 1.14 \times 10^{-6}$	95% CI [0.2955, 0.6141]
bb	8	Current	GC changes in significant regions ($p < 0.01$)	Non-normal distribution	Spearman correlation (within groups; baseline GC and GC changes)		R	r = -0.08	ρ = 0.39	95% CI [-0.2593, 0.1046]

Each analysis includes a letter indicator ("manuscript" column) linking the test in the table to the analysis in the text. The link to the corresponding figure, if any, and the sample used for the test are indicated in the "figure," and "sample," columns, respectively. The "current" sample includes tests using data from the current work, and the previous study is indicated as Wang et al. (2014), the dependent variables for each test are listed as "data type," and the "data structure" column indicates whether the data are normally distributed. The type of test, contrast, and the groups used for the analysis are listed in the "type of test" column. The multiple correction method is listed under "multiple comparisons correction." the program used to perform the analysis is included under "program." The critical value and degrees of freedom are listed for each test under "statistics." Finally, the p value and confidence intervals are listed in the final two columns. DLPFC, dorsolateral prefrontal cortex; GC, global connectedness; LPOC, left precuneus and medial occipital cortex.

(Table 1, p, V=94, p=0.055, 95% CI [-0.0006, 0.04139]). However, there was a trend-level difference between the GC and hippocampal FC changes in the LPOC region (Table 1, q, V=92, p=0.073, 95% CI [-0.0066, 0.1547]). We conducted additional control analyses to determine whether stimulation caused significant increases in FC between the DLPFC and the hippocampus, but it did not (Table 1, r, $t_{(14)}=0.949$, p=0.359, 95% CI [-0.061, 0.157]), nor were there changes in FC between the DLPFC and the stimulus location in the PPC (Table 1, s, V=68, p=0.679, 95% CI [-0.103, 0.121]), nor did PPC-GC increase (Table 1, t, V=92, P=0.073, 95% CI [-0.005, 0.050]).

In the current sample, there was an increase in hippocampal FC with the 17 networks identified by Yeo et al.

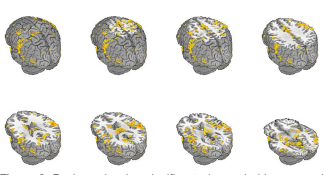


Figure 2. Regions showing significant change in hippocampal FC following PPC rTMS from the current study (FDR corrected, q=0.05).



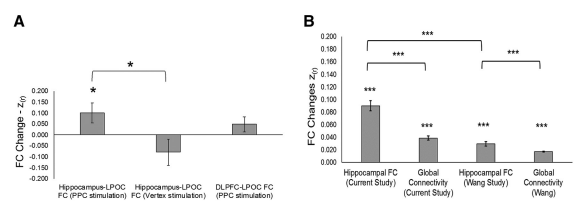


Figure 3. A, Average change in hippocampal-LPOC FC for subjects receiving PPC stimulation (left bar) and vertex stimulation (middle bar). Average DLPFC-LPOC FC changes for subjects receiving PPC stimulation is represented by the right bar. B, Mean changes in hippocampal FC within 17 segregated networks from Yeo et al. (2011) after PPC rTMS in this study and Wang et al., and change in GC within networks from both studies. Error bars represent the standard error of the mean; *p < 0.05, ***p < 0.0001.

(2011), which was significantly stronger than the GC changes in these networks (Table 1, u, $t_{(16)} = 10.96$, 7.6×10^{-9} , 95% CI [0.0725, 0.1073]). We found the same effect in the data from Wang et al. (Table 1, v, $t_{(16)} = 11.27$, $p = 5.10 \times 10^{-9}$, 95% CI [0.0138, 0.0201]; Fig. 3*B*). Comparing the hippocampal FC changes between studies, we found that they were larger in the current study, despite using fewer stimulation sessions (Table 1, w, $t_{(32)} = 8.75$, $p = 5.42 \times 10^{-10}$, 95% CI [0.0560, 0.09]), although GC changes were also larger in the current results than in the Wang et al. data. This suggests overall differences in the magnitude of FC changes across experiments, which could reflect factors such as different scan variables be-

tween studies. After PPC stimulation in both studies, increases in hippocampal FC were maximal in networks that included the cuneus and retrosplenial, somatosensory, and superior temporal, areas (Fig. 5).

In our test for whether the whole-brain topographic patterns of rTMS-induced hippocampal FC were similar between studies, we found that FC changes were correlated between studies (Table 1, x, r=0.51, n=17, p=0.037, 95% CI [0.0389, 0.7956]; Fig. 6). There was no significant correlation between the GC changes in the current study with the hippocampal FC changes of Wang et al. (Table 1, y, r=0.16, n=17, p=0.536, 95% CI [-0.3419, 0.5989]). These results indicate that the magni-

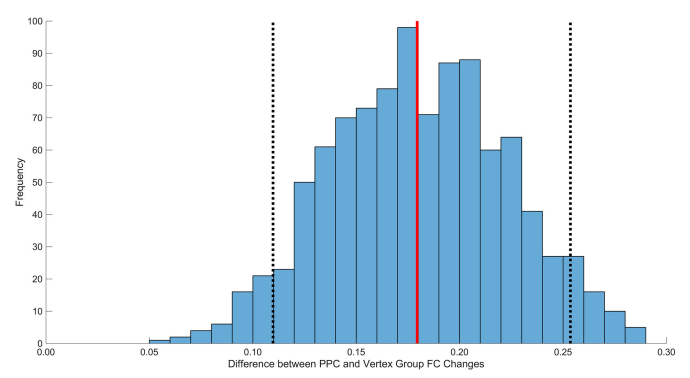


Figure 4. Histogram representing the result of 1000 group mean differences using eight subjects from each group, where the eight PPC subjects are randomly sampled each time. The black dotted lines represent the upper (0.2536) and lower (0.1098) limit of 95% of the distribution. The observed mean difference between the PPC and vertex group is shown by the red line (0.1795).



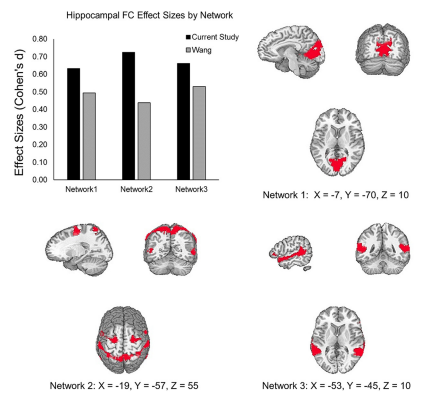


Figure 5. Effect size of increases in hippocampal FC within three representative networks from Yeo et al. (2011) after PPC rTMS in this study and Wang et al. Network 1 includes cuneus and retrosplenial cortex. Network 2 includes somatosensory areas. Network 3 includes superior temporal areas.

tude of hippocampal FC changes across networks was similar between studies and that their topographic distribution was reproducible.

Finally, we reproduced the finding that, among areas showing significant increases in hippocampal FC after PPC rTMS, prestimulation hippocampal FC predicted the magnitude of the increase (Fig. 7*A*). This was confirmed by the relationship between the baseline and mean change in hippocampal FC across areas (Table 1, z, r = 0.39, n = 95, $p = 1.0 \times 10^{-4}$, 95% CI [0.2002, 0.5453]; Fig. 8*A*). Removing the single outlier did not change the

significance of the correlation (Table 1, aa, $r_{(92)} = 0.47$, $p = 1.14 \times 10^{-6}$, 95% CI [0.2955, 0.6141]). We did not observe the same pattern of results when performing the same analyses using GC as the dependent variable (Table 1, bb, r = -0.08, n = 115, p = 0.39, 95% CI [-0.2593, 0.1046]; Figs. 7B, 8B). These findings indicate a specific effect of PPC rTMS on the hippocampus and rule out non-specific enhancement of FC across the brain. Re-sorting the matrices in Figure 7 revealed no regional differences in the change in GC (Fig. 7B), but did show that, among regions connected to the hippocampus at

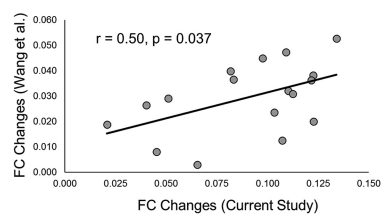


Figure 6. Scatterplot of PPC rTMS-induced hippocampal FC $[z_{(r)}]$ changes across networks from Yeo et al. (2011). Each dot represents the rTMS-induced hippocampal FC change from the current study (x-axis) and Wang et al. (y-axis) within one of the 17 networks from Yeo et al. (2011). The black line represents the regression line across individual data points.



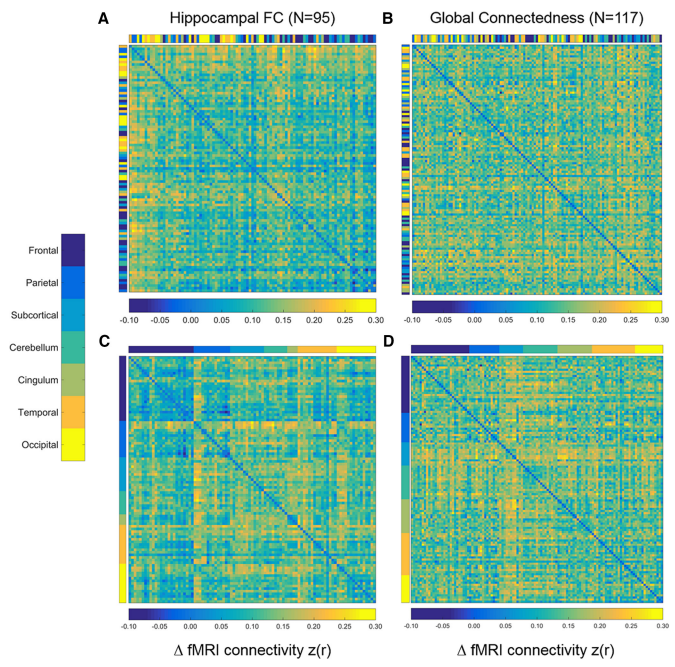


Figure 7. Correlation matrices of regions demonstrating significant (p < 0.01) changes in hippocampal (**A**) and global (**B**) FC. Matrices are sorted by baseline FC with the highest values represented at the top of the matrices on the *y*-axis and to the left on the *x*-axis. Color bars aligned with each axis represent AAL-defined regions. **C**, **D**, Identical to **A**, **B** but are sorted by region.

baseline, frontal regions showed qualitatively less change than more posterior regions, such as the PPC, similar to the results of Wang et al. (2014) and consistent with the interpretation that areas with higher baseline FC with the hippocampus change most with PPC rTMS.

Discussion

We independently reproduced the highly specific increase in hippocampal FC, reported by Wang et al. (2014), resulting from high-frequency rTMS of PPC, using a partial replication of their technique and adding additional new controls. We applied a whole-brain analysis as well

as a hypothesis-based approach, predicated on the anatomic distribution of changes reported by Wang et al. (2014). We also looked for changes in hippocampal FC within 17 additional segregated brain networks (Yeo et al., 2011). The whole-brain comparison to Wang et al. (2014) revealed that PPC rTMS caused significant hippocampal FC changes in all of the regions reported by Wang et al., as well as several others. The hypothesis-based approach revealed significant increases in hippocampal FC with the LPOC, a region derived from our re-analysis of the Wang et al. (2014) data. These changes were specific to FC with



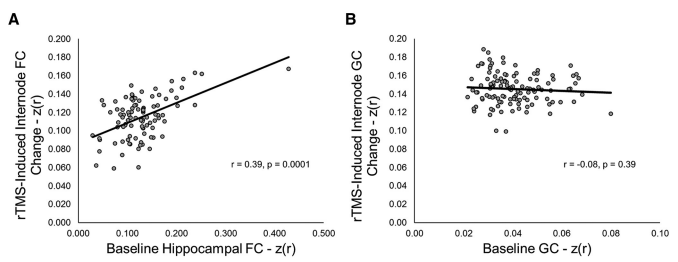


Figure 8. A, Scatterplot of baseline hippocampal FC for regions demonstrating significant (p < 0.01) changes in hippocampal FC and average rTMS-induced FC change in those regions. **B**, Scatterplot of baseline GC for regions demonstrating significant (p < 0.01) changes in GC and average rTMS-induced internode GC change in those regions.

the hippocampus: PPC rTMS did not significantly increase FC between the DLPFC, an area active in many cognitive processes including learning, and the LPOC. We also ruled out the possibility that the findings reflected a general increase in brain connectivity: Hippocampal FC was significantly greater than GC across all networks examined in both the present and the Wang et al. data. Although our vertex control sample was small, we found no significant FC changes in this group and hippocampus-LPOC FC was significantly greater for the PPC rTMS group than the vertex group.

As in the data of Wang et al. (2014), baseline hippocampal FC predicted PPC rTMS-induced FC changes and we demonstrated the specificity of this relationship by showing that baseline GC did not predict GC increases after rTMS. Finally, the spatial pattern of rTMS-induced FC change was similar and correlated between studies. Taken together, this is strong evidence that the effect of 20-Hz rTMS on the PPC on hippocampal FC is robust, reproducible, and highly specific in anatomic terms.

Notably, we were able to reproduce, and possibly to exceed, the results of Wang et al. (2014) with fewer stimulation sessions. Multiple consecutive rTMS sessions are burdensome to subjects and investigators alike and reducing the requirement increases the attractiveness of the PPC rTMS paradigm.

Our vertex rTMS group showed decreased hippocampal FC with almost every network, including the LPOC. This unexplained time-related drift could be due to a physiologic effect and might represent a potential confound. However, as noted above, others (Jung et al., 2016) have found no evidence of FC changes from vertex rTMS. Additionally, the average change in hippocampal FC across the networks from Yeo et al. (2011) in the Wang et al. (2014) sham data did not differ significantly from zero.

There were several procedural differences between the current work and that of Wang et al. (2014), the most obvious of which was the absence of behavioral testing.

Therefore, we do not know whether the changes in hippocampal FC were associated with an improvement in declarative memory. Additionally, there were differences in how we preprocessed our resting-state data. We did not bandpass filter our data and, unlike Wang et al. (2014), we included spatial smoothing to reduce the influence of spatial noise and increase signal-to-noise ratio (SNR).

Another difference between studies was our use of a vertex stimulation control. Wang et al. (2014) used subthreshold stimulation (10% of RMT) of the PPC as their within-subjects control, which may have caused weak local brain effects without reproducing the somatosensory effect of full-intensity rTMS. They also used fullintensity stimulation of the motor cortex in an independent group as a secondary control, and this was all but certain to produce widespread changes in FC. We chose activeintensity stimulation at the vertex as our control because others (Jung et al., 2016) found no effect on FC from stimulation there. This site lies over the sagittal sinus and the interhemispheric fissure, where the cortex is relatively distant from the coil and the nearest regions are out of the plane of the stimulating current. Neither in this, nor the study of Wang et al. (2014), did control stimulation produce any measurable increase in hippocampal FC.

Finally, for reasons described above, we used three and, in some cases, 4 d of rTMS, while Wang et al. (2014) used five. This study contains no basis for a quantitative comparison of the strength or duration of the connectivity or behavioral changes produced by various treatment durations, but 3–4 d appears adequate to produce FC changes similar to those of Wang et al. (2014) at a 24-h delay. These procedural differences do not allow us to claim a strict replication of the paradigm, but they do not detract from the substantial reproduction of the result and could not have caused it by themselves.

Both we and Wang et al. (2014) were able to produce dramatic increases in hippocampal network FC with a few sessions of PPC rTMS, making this one of the strongest and most reliable effects in noninvasive neuromodulation.



The differences in the treatment paradigms and image processing procedures decrease the likelihood that both studies arrived at a similar result due to an artifact or systematic noise. Others (van der Werf et al., 2010; Vercammen et al., 2010; Gratton et al., 2013; Rahnev et al., 2013; Steel et al., 2016; Rastogi et al., 2017) have also used FC to study how rTMS affects brain function at the network level. FC in the default mode network appears to be particularly sensitive to modulation with rTMS (Eldaief et al., 2011; Halko et al., 2014; Wang et al., 2014) and can be modulated by stimulating the PPC (Eldaief et al., 2011; Wang et al., 2014) and the cerebellum (Halko et al., 2014). The latter study also used individual FC to choose the stimulation target.

Future studies may consider examining whether the stimulation regimen itself, largely inspired by conventional rTMS treatment for depression (George et al., 1997), where it was adopted without systematic exploration of the parameter space, is optimal, and whether even more dramatic or faster responses are attainable using optimized stimulation parameter settings.

Conclusion

The hippocampal network FC changes reported by Wang et al. (2014) after PPC rTMS are reproducible in magnitude, specificity, and topographic distribution. Our additional analyses, ruling out changes in global correlation, further strengthen the evidence for the selectivity of this approach for the hippocampal network. Moreover, our findings suggest that these effects are achievable with fewer than five stimulation sessions. This provides encouraging support for PPC rTMS as a means of enhancing memory network FC and for rTMS in general as a technique for producing targeted changes in brain network connectivity.

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