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Psychophysical measurement of the effects and non-effects of TMS on contrast perception

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ACCEPTED MANUSCRIPT	
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1	Title: Psychophysical measurement of the effects and
2	non-effects of TMS on contrast perception
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18 Highlights

- 19
- Compared the effects of four TMS protocols on neural signals and
 noise.
- Single pulse TMS suppressed neural signals and repetitive TMS
 increased neural noise.
- Theta burst stimulation did not affect perceptual task performance.
- Participants differed in TMS susceptibility, determined by phosphene
 perception.
- Findings suggest systematic inter-protocol and inter-participant
- 28 differences in TMS effects.
- 29
- 30

31 Key words

- 32 Sensory processing, neural effects, theta burst, online stimulation,
- 33 psychophysics

34 To the editor:

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Transcranial magnetic stimulation (TMS) is widely used to establish causal 36 37 relationships between brain areas and behavior, but its effects on task performance are not fully understood and have rarely been directly compared 38 39 between protocols. Decreases in performance on psychophysical tasks, such as those observed when applying TMS, can be attributed to either 40 41 suppression of stimulus-related neural signals, increased random activity (i.e. neural noise), or a combination of both [1,2]. Indeed, evidence for all three 42 43 hypotheses has been found when using differing methodologies and online 44 stimulation protocols [3–5]. Similarly, theta burst stimulation (TBS) has been 45 shown to have variable or bimodal effects between participants and between exact stimulation protocols [6,7]. Despite different TMS protocols (e.g. online, 46 offline, repetitive, single pulse) potentially having vastly different effects, they 47 48 are often used interchangeably in sensory and cognitive research.

49

We directly compared the neural effects of four commonly used TMS protocols: online single pulse (spTMS), online 3-pulse repetitive (rTMS; 50ms between pulses), offline continuous theta burst (cTBS) and offline intermittent theta burst (iTBS), during a well-understood neural computation – contrast transduction. As a secondary objective, we investigated natural TMSsusceptibility by comparing participants who could and could not perceive phosphenes to address inter-participant variability in TMS effectiveness.

58 We tested all stimulation protocols using the same area (occipital cortex, 59 Supp. 2A) and a highly sensitive double-pass paradigm [8] to dissociate TMS 60 induced changes in stimulus-related neural signal strength (i.e. suppression) 61 and neural noise. On each trial (200 total per TMS condition) two luminancemodulated stimuli (3 deg. vis. ang.) of randomly-selected contrast were 62 63 presented peripherally. Half of the trials contained a 4% contrast increment in one of the intervals (see Supp. 1A,B for examples). The exact same trials 64 65 were then repeated with randomized interval order. Full details of stimuli and 66 the double-pass paradigm be found in [9]. Using standard protocols with a Magstim Super Rapid² 'figure of 8' coil spTMS and rTMS (Supp. 2C, 70%) 67 68 stimulator output) were applied 50ms after stimulus onset in each interval, and 69 offline TBS (Supp. 2D, 30% stimulator output) was applied before the start of 70 the task. Consistency between the first and second presentation of the trials was calculated as a direct index of neural noise. Accuracy on the task was 71 72 calculated as a measure of stimulus-related signal strength.

73

74 During phosphene localization pre-screening, six participants (4 females, age 75 22-34) consistently perceived phosphenes and completed the main experiment (a further 19 participants were screened but did not report seeing 76 phosphenes). Study was approved by YNiC ethics committee. All TMS 77 78 protocols were tested on different days (rTMS was tested over four days due 79 to high numbers of pulses). Phosphene localization was performed before each testing session and the location of the phosphenes (as indicated with a 80 81 computer interface, Fig. 1A) was used to subsequently present stimuli.

82 Control trials (no TMS) were performed before stimulation for each TMS83 protocol separately.

84

We simulated predictions using a linear amplifier model (LAM). Simulations showed that if TMS reduced neural signal strength (lowered sensitivity), we would observe a steep drop in task accuracy but no change in double-pass consistency. Alternatively, if TMS increased neural noise, we would see a small reduction in accuracy and a larger drop in consistency. Finally, if TMS both reduced stimulus-related signals and increased noise, we would observe a large reduction in both measures (Supp. 1C-E).

92

93 We found a significant drop in accuracy (t(5)=2.83, p=0.037, Bayes factor)94 (BF)=2.83) when applying spTMS compared to the no TMS condition, but no 95 change in consistency (p=0.601, BF=0.29, Fig. 1B). This closely resembles 96 our LAM model predictions for an increase in neural suppression and suggests that spTMS suppresses neural signals. Conversely, applying rTMS 97 98 showed a small non-significant change in accuracy p=0.848, BF=0.33) 99 compared to the no-TMS condition, and a significant decrease in consistency 100 (t(5)=2.74, p=0.041, BF=2.38, Fig. 1C) – consistent with model predictions for 101 an increase in neural noise. Neither protocol produced data consistent with 102 change in both suppression and noise. This comparison between spTMS and 103 rTMS is consistent with previous research that tested these protocols 104 separately [4,3] and suggests suppressive and noise-inducing effects are 105 protocol-specific.

107 No effects on the accuracy (p=0.790, BF=0.30) or consistency (p=0.132, 108 BF=0.93) were observed when applying cTBS (Fig. 1D). Similarly, no 109 changes in accuracy (p=0.773, BF=0.30) or consistency (p=0.244, BF=0.58) were observed when applying iTBS (Fig. 1E), indicating that neither protocol 110 changed the levels of neural noise or sensory signals. This may seem to 111 oppose the large number of successful TBS studies, particularly in the motor 112 cortex. However, most previous research into TBS effects measured motor 113 evoked potentials, which reflect an overall increase or decrease in neural 114 activity (e.g. [10]). It may be that TBS changes overall neural activation but 115 116 does not have particular effects on perceptually-relevant signals that would 117 affect sensory task performance. Alternatively, the effectiveness of TBS may be overstated in the literature, as indicated by a recent large scale meta-118 analysis [11] which found a large positive publication bias in the TBS 119 literature. 120

121

To investigate the effects of TMS susceptibility on task-relevant effects, a 122 further six participants (3 females, age 23-55) who did not report seeing 123 124 phosphenes also completed the experiment. For these participants, stimuli were presented at the mean location of phosphenes experienced by the other 125 group. None of the four TMS protocols had any significant effect on accuracy 126 127 or consistency scores in these individuals, indicating that the participants who did not perceive phosphenes during phosphene localization were not affected 128 129 by TMS during the task. Anatomical differences in cortical folding and skull 130 thickness may explain these individual differences in TMS susceptibility.

132	The inter-participant and inter-protocol differences in TMS effects found here		
133	shed light on the interpretation of findings in the existing TMS literature and		
134	inform future methodological choices. The individual differences in		
135	susceptibility and the use of different stimulation protocols in the literature		
136	may be some of the major factors in the TMS 'replication crisis' [12]. The		
137	effects of TMS are subtle and can often only be detected in reaction time data		
138	rather than task performance [13]. In this respect, the sensitivity and precision		
139	of the double-pass paradigm is a valuable tool for further investigating TMS		
140	inter-protocol and inter-participant variability in other brain areas and with		
141	large	r samples.	
142			
143	Conflict of Interest		
144	There is no conflict of interest relating to this manuscript.		
145			
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191 Figures

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Figure 1. Phosphene locations, mean accuracy and consistency scores for the individuals seeing phosphenes. Phosphene locations were similar for all six participants, centered around the midline of the left visual field (A), within 15 degrees of the fixation cross. Phosphene locations were consistent across the four experiments using different stimulation protocols: spTMS (blue), rTMS (yellow,

199 averaged over four sessions), cTBS (green) and iTBS (purple), as indicated by filled 200 ovals. In Exp 1, single pulse TMS (B) significantly reduced the mean accuracy scores 201 (dark bars) compared to the no-TMS condition (light bars) but not consistency scores 202 which indicates increased suppression resulting from TMS stimulation. Repetitive 203 TMS (C) significantly reduced task consistency but not task accuracy, indicating a 204 TMS-induced increase in neural noise. Neither cTBS (D) nor iTBS (E) produced any 205 significant change in task performance. Error bars indicate bootstrapped 95% 206 confidence intervals.



210 Supplementary figure 1. Each interval during a trial was drawn from the target 211 (blue) and non-target (vellow) stimulus distributions (A). Participants were asked to 212 choose the interval with the more positive contrast (B; example correct intervals are 213 shown with a blue circle). Stochastic simulations were used to generate model 214 predictions of double-pass data (C-E). Light bars in all panels indicate a system with 215 low neural noise and low suppression (high sensitivity) in the system. Dark bars 216 model an increase in either suppression, noise, or both. If TMS suppresses neural 217 signals (lowers sensitivity) then we should expect double-pass data to be similar to 218 the prediction in panel C. On the other hand, if TMS increases neural noise the data 219 should resemble panel D. If both suppression and neural noise are increased we 220 would expect data to be similar to panel E.



Supplementary figure 2. The TMS coil was positioned (red dot) approximately 2cm above and 1cm to the right of the inion (blue line intersection) to induce phosphenes (A). Before phosphene localization participants were trained to indicate the location and shape of a simulated phosphene on the screen (B; see section 2.3). During spTMS and rTMS protocols either one or three pulses (50ms apart) were delivered 50ms after stimulus onset (C). Pulses during offline cTBS and iTBS were delivered as shown in D.