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# Proactive modulation of long-interval intracortical inhibition during response inhibition

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1	Proactive modulation of long-interval intracortical inhibition during response inhibition
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#### 23 Abstract

24 Daily activities often require sudden cancellation of pre-planned movement, termed response inhibition. When only a subcomponent of a whole response must be suppressed 25 26 (required herein on Partial trials), the ensuing component is markedly delayed. The neural 27 mechanisms underlying partial response inhibition remain unclear. We hypothesized that Partial trials would be associated with non-selective corticomotor suppression and that 28 GABA<sub>B</sub>-receptor mediated inhibition within primary motor cortex might be responsible for 29 30 the non-selective corticomotor suppression contributing to Partial trial response delays. 31 Sixteen right-handed participants performed a bimanual anticipatory response inhibition task 32 while single and paired-pulse transcranial magnetic stimulation was delivered to elicit motor 33 evoked potentials in the left first dorsal interosseous muscle. Lift times, amplitude of motor 34 evoked potentials and long-interval intracortical inhibition were examined across the different 35 trial types (Go, Stop-Left, Stop-Right, Stop-Both). Go trials produced a tight distribution of 36 lift times around the target, whereas those during Partial trials (Stop-Left and Stop-Right) 37 were substantially delayed. The modulation of motor evoked potential amplitude during Stop-38 Right trials reflected anticipation, suppression and subsequent re-initiation of movement. 39 Importantly, suppression was present across all Stop trial types, indicative of a "default" non-40 selective inhibitory process. Compared with blocks containing only Go trials, inhibition 41 increased when Stop trials were introduced but did not differ between trial types. The amount 42 of inhibition was positively correlated with lift times during Stop-Right trials. Tonic levels of 43 inhibition appear to be proactively modulated by task context and influence the speed at 44 which unimanual responses occur after a non-selective "brake" is applied.

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46

#### 47 New & Noteworthy

The ability to cancel a pre-planned movement, termed response inhibition, is essential for adaptable motor control. Participants performed a bimanual anticipatory response inhibition task while single and paired-pulse transcranial magnetic stimulation was delivered. The modulation of motor evoked potential amplitude during partial response trials reflected anticipation, suppression and subsequent re-initiation of movement. Importantly, suppression was present across all stop trial types, indicative of a "default" non-selective inhibitory process.

# 56 Introduction

57	The ability to cancel a pre-planned movement, termed response inhibition, is essential
58	for adaptable motor control. Response inhibition relies upon a cortico-subcortical network
59	(Aron and Poldrack 2006; Chambers et al. 2006; Coxon et al. 2009; Coxon et al. 2012;
60	Zandbelt et al. 2013) that inhibits corticospinal neurons (CSNs) within the primary motor
61	cortex (M1) in order to suppress descending motor output (Stinear et al., 2009). It is known
62	that gamma-aminobutyric acid (GABA) mediated interneurons within M1 exert powerful
63	inhibitory effects on CSNs (Jones 1993; Keller 1993). However, the role of GABA-ergic
64	inhibition during response inhibition is not fully understood.
65	Response inhibition can be proactive when stopping demands are anticipated, or
66	reactive when stop signals are presented unexpectedly (Aron and Verbruggen 2008).
67	Transcranial magnetic stimulation (TMS) has been used to assess the temporal modulation of
68	corticomotor excitability (CME) during both types of response inhibition. Proactive stopping
69	is suggested to recruit the indirect basal ganglia pathway to selectively decrease CME for
70	only the movement cued to stop (Aron and Verbruggen 2008; Cai et al. 2011; Majid et al.
71	2013). A topic of current debate is whether reactive stopping can be achieved selectively (Xu
72	et al. 2014), given that several lines of evidence indicate a transient process in which
73	stopping response preparation suppresses movement non-selectively. For example, when
74	successful stopping can be achieved by inhibiting all movement, CME is reduced in
75	response-irrelevant muscles (Badry et al. 2009; Cai et al. 2012; Coxon et al. 2006;
76	Greenhouse et al. 2012). When only a subcomponent of a prepared response is required to
77	stop (Partial trials), the remaining response is delayed (Coxon et al. 2007; 2009; Coxon et al.
78	2012; MacDonald et al. 2014; MacDonald et al. 2012). Interestingly there is also a
79	concomitant reduction in CME for the responding left hand on trials when only the right hand
80	is cued to stop (MacDonald et al. 2014), indicative of a non-selective inhibitory process that

cancels all prepared effectors. The subsequent delay may arise as a consequence of having to
initiate a new response. While these studies suggest reactive response inhibition is nonselective, the primary suppressive mechanism remains unclear.

Intracortical networks within M1 are the final cortical modulators of motor output. 84 Paired-pulse TMS can be used to investigate GABA-ergic inhibition within M1 and identify 85 the contribution of distinct intracortical networks to motor performance (Reis et al. 2008). 86 GABA<sub>A</sub>-receptor mediated short-interval intracortical inhibition (SICI), assessed using a 87 subthreshold conditioning stimulus followed 1-6 ms later by a suprathreshold test stimulus 88 89 (Kujirai et al. 1993), is involved with movement initiation and prevention. SICI is selectively reduced during movement initiation (Reynolds and Ashby 1999; Stinear and Byblow 2003; 90 91 Zoghi et al. 2003). While GABA<sub>A</sub>-mediated networks within M1 are essential for specific, 92 point-to-point control over motor representations, SICI is non-selectively increased following 93 stop signal presentation during unimanual response inhibition (Coxon et al. 2006). However, 94 MacDonald et al. (2014) could not temporally reconcile non-selective CME suppression with 95 an increase in SICI during partial response inhibition, making it an unlikely candidate 96 mechanism underlying the behavioral and neurophysiological findings observed when only a 97 subcomponent of a prepared response must be terminated.

98 Within M1, GABA<sub>B</sub>-receptor mediated inhibition can be assessed as long-interval intracortical inhibition (LICI) using suprathreshold conditioning and test stimuli separated by 99 100 50-200 ms (Valls-Solé et al. 1992; Wassermann et al. 1996). LICI engages pre-synaptic 101 receptors (Bettler et al. 2004), typically associated with tonic inhibitory effects. Interestingly, 102 one study reported decreased LICI in a Go/No-Go task (Sohn et al. 2002) but did not assess 103 LICI using an optimal interstimulus interval (Sanger et al. 2001) or use conditioning stimulus 104 intensities that allowed increases in LICI to be observed. Therefore, the role of LICI in 105 reactive response inhibition is yet to be established.

106	There were three aims of the present study. First, we wanted to confirm and extend on
107	the finding that Partial trials are associated with non-selective CME suppression as
108	demonstrated by MacDonald et al. (2014). We hypothesized that CME, as evident by MEP
109	amplitude, would be reduced during Partial trials, supporting a model of non-selective
110	suppression (MacDonald et al. 2014), and that CME suppression would occur at equivalent
111	time points whether one or both sides were cued to stop. Second, we hypothesized that LICI
112	would be up-regulated during response inhibition trials compared with Go trials irrespective
113	of Stop trial type i.e., indicative of a non-selective inhibitory mechanism. Finally, we wanted
114	to examine CME and LICI at the onset of trials to determine if either could explain why lift
115	times are influenced by previous trial type (Coxon et al. 2007; MacDonald et al. 2012).
116	Methods
117	Participants
118	Sixteen participants without neurological impairment participated in the experiment (mean
119	age 24 y, range 18-49 y, 14 male). Thirteen of these completed the LICI protocol (mean age
120	25 y, range 18-49 y, 11 male). All were right handed (laterality quotient mean 0.92, range
121	0.71-1) as determined using a four-item version of the Edinburgh handedness inventory
122	(Veale 2014). Written consent was obtained before participation and the study was approved
123	by the University of Auckland Human Participants Ethics Committee.
124	Response inhibition task
125	The experiment utilized a bimanual anticipatory response inhibition task (Coxon et al.
126	2007; 2009; MacDonald et al. 2014; MacDonald et al. 2012). Participants were seated with
127	forearms resting on a table positioned midway between pronation and supination. This
128	allowed the distal and medial aspect of each index finger to occupy a mechanical switch
129	positioned 55 cm away from the computer monitor. The display consisted of two indicators

(vertical rectangles) each 16 cm in length and 1.5 cm in width (Figure 1A). Control of the left
or right indicator was via the corresponding left or right switch. Switch "up/down" state was
precisely recorded through an Arduino and synchronized to the display through an analogdigital USB interface (NI-DAQmx 9.7; National Instruments). Customized software written
in MATLAB (R2011a, version 7.12; The MathWorks) generated the trial order, recorded trial
data and controlled the visual output during the task.

136 Participants were instructed to let the weight of their fingers passively depress the 137 switches. Switch height was adjusted to eliminate any positional related muscle activity. 138 Depression of both switches initiated the trial after a 400-900 ms variable delay. If depression 139 continued, both indicators would ascend vertically at a constant velocity reaching a horizontal 140 line after 800 ms and the top of the display after 1000 ms. Participants were informed that 141 releasing the switch (index finger abduction) would stop the corresponding indicator. Go 142 trials (Go-Left Go-Right; GG) required participants to stop both indicators at the target by 143 releasing both switches (Figure 1B). Stop trials were cued by one or both indicators stopping 144 before the target, requiring participants to inhibit the response of one or both hands (Figures 145 1C and D). In the protocols a 2:1 ratio of Go to Stop trials existed, establishing Go trials as 146 the default response. When both hands were required to stop (Stop-Both i.e. Stop-Left Stop-147 Right; SS), both indicators stopped 600 ms into the trial. On Partial trials only one hand was 148 required to stop (Go-Left Stop-Right; GS, or Stop-Left Go-Right; SG). On Partial trials a 149 single indicator stopped 550 ms into the trial while the other continued to ascend. The time 150 the indicator stopped on Stop trials (550 & 600 ms) did not change. This enabled constant 151 TMS times relative to both the stopping of the indicator and the start of the trial. Feedback 152 was visually displayed during each inter-trial interval.

#### 153 *Electromyography*

154 MEP amplitude was recorded using electromyography (EMG) over the left first dorsal interosseous (FDI) muscle as the non-dominant hand is more strongly affected than the 155 156 dominant hand by the processes required to successfully cancel a subset of a movement 157 (MacDonald et al. 2012). A belly-tendon electrode montage was used with a ground electrode 158 on the posterior hand surface. EMG activity was recorded using a National Instruments A/D 159 acquisition system, displayed using custom LabVIEW software, and stored to disk for offline analysis. Electrical activity was amplified (Grass P511AC), band-pass filtered (10-1000 Hz) 160 and sampled at 2 kHz. 161 162 Transcranial magnetic stimulation 163 To examine CME during successful Stop and Go trials single-pulse TMS was delivered 164 using a single Magstim 200 stimulator (Magstim, Dyfed, United Kingdom). A figure-of-eight 165 coil (70 mm diameter) was held tangentially over the right M1 of the participant. The optimal coil position for eliciting MEPs in the left FDI was determined using a slightly suprathreshold 166 167 intensity and marked on the scalp. The handle was posteriorly positioned and the coil 168 orientated at a 45° angle to the midline, inducing a posterior to anterior directed current 169 (Brasil-Neto et al. 1992). To examine LICI we delivered paired-pulse TMS from two 170 Magstim 200 units connected via a Bistim unit (Magstim, Dyfed, United Kingdom). The 171 CME protocol always preceded the LICI protocol.

172 *Protocol* 

To examine CME, Task Motor Threshold (TMT) was determined while the participant rested their index fingers on the switches. TMT was determined as the minimum stimulus intensity required to elicit a MEP of at least 50  $\mu$ V in the FDI in 4 out of 8 consecutive trials. Stimulus intensity was adjusted in 1-2 % increments from TMT to produce an average MEP amplitude of 0.1-0.2 mV at 200 ms before the target while not disrupting task performance.

This intensity was then used for all remaining stimulated trials. Participants completed anunstimulated practice block of 36 trials containing pseudo-randomized Stop trials.

180	The task in the CME protocol consisted of 432 trials split into 12 pseudo-randomized
181	blocks of 36 with 288 Go and 144 Stop trials. There were 98 Go trials where TMS was
182	delivered at either 250, 225, 200, 175, 150, 125 or 100 ms before the target to obtain 14
183	stimuli at each time point. All Stop trials were stimulated and distributed as 84 GS, 30 SG
184	and 30 SS because GS trials were of primary interest. For GS trials, 14 stimuli were delivered
185	150, 125, 100, 75, 50 and 25 ms before the target. Timing was delayed by 100 ms relative to
186	Go trials because of the expected $\sim$ 75 - 100 ms delay in the responding hand (Coxon et al.
187	2007; 2009; Coxon et al. 2012; MacDonald et al. 2014; MacDonald et al. 2012). For SG and
188	SS trials, 15 stimuli were delivered 175 and 200 ms after the stop signal.
189	To examine LICI, paired-pulse TMS was delivered at an ISI of 100 ms (Sanger et al.
190	2001). Stimulation intensity was adjusted to elicit a cortical silent period (CSP) duration of
191	175 ms while left FDI was activated at ~10% of maximum voluntary contraction. This
192	intensity was used in LICI for both the conditioning stimulus (CS) and the test stimulus (TS).
193	CS and TS were then adjusted to produce approximately 50-85% inhibition of the MEP
194	amplitude (%INH). This intensity was used for all following trials.
195	During the LICI protocol, participants performed a Go Only block consisting of 30
196	Go trials. %INH was measured at the start of each trial (0 ms). Each participant then
197	performed the task, which consisted of 360 trials split into 10 blocks of 36 trials. Of these,
198	240 (67%) were Go trials and 120 (33%) were Stop trials. Stop trial types (GS, SG and SS)
199	were equally represented, each condition consisting of 40 trials. The %INH obtained at 0 ms
200	provided information for the previous and upcoming trial. All trials following Stop trials and
201	185 trials following Go trials were stimulated.

Peak-to-peak MEP amplitude was calculated from EMG 10 to 50 ms after the 203 204 stimulus. MEPs were excluded when root mean square (rms) EMG was  $> 10 \ \mu$ V in the 50 ms 205 preceding stimulation. Also, EMG traces were excluded if any activity was present between 206 the stimulus and MEP evident from visual inspection. Average MEP amplitude was calculated following trimming of the upper and lower 10 % (if > 8 MEPs were present for 207 208 that time point) or  $\pm 1.5$  standard deviations (if 4 - 8 MEPs were present for that time point). 209 To reduce inter-subject variability MEP amplitude was normalised across Trial Types and 210 Stimulations Times such that the condition with the largest mean MEP amplitude was 211 reassigned the value 1, and all other conditions scaled accordingly for the participant. For the 212 LICI protocol, mean stimulated and unstimulated left hand LTs were calculated from Go 213 trials after Go trials. MEP amplitude was calculated for each stimulated trial in the left hand 214 for both CS and TS. The primary dependent measure was %INH, which was calculated as 215 %INH = 100 – (TS MEP amplitude/ CS MEP amplitude)  $\times$ 100, where TS and CS MEP 216 amplitude are the mean values for each condition from each participant.  $\Delta$ %INH and  $\Delta$ CS 217 MEP amplitude was calculated for Partial trials followed and preceded by Go trials.  $\Delta =$ 218 [(subsequent Go trial – Partial trial)/ Partial trial]  $\times$  100.

219 To assess behaviour, lift times (LTs) were recorded and are reported relative to the 220 target. Mean LTs from Go and successful Partial trials were calculated after the removal of 221 outliers ( $\pm$  3 SD; 1% removed for Go and Partial trials in CME protocol, 2% removed for Go 222 and Partial trials in LICI protocol). In the LICI protocol only, lift time asynchronies (LTAs) 223 were calculated from LTs in Go trials following Go and successful Stop trials (LTA = [Left hand LT] - [Right hand LT]). For Stop trials, stop signal reaction time (SSRT) and 224 225 percentage of successful trials were determined. SSRT was calculated using the integration 226 method (Logan and Cowan 1984; Verbruggen et al. 2013).

228 Repeated measures Analysis of Variance (RM ANOVA) were used to test our 229 hypotheses. To test the first hypothesis, normalized MEP amplitude was first compared in an 230 omnibus RM ANOVA with a 2 Trial Type (GG, GS) × 6 Stimulation Time (-225, -200, -175, 231 -150, -125 and -100 ms relative to expected LT i.e. 0 ms on GG, 75 ms on GS) design. To 232 determine if the hypothesised pattern of facilitation, suppression and facilitation on Partial 233 Trials was present, one-way RM ANOVAs with the factor Stimulation Time (6 levels) were 234 run separately on GS and GG trials. Additionally, MEP amplitude for Stop trials was 235 analyzed with a 3 Trial Type (GS, SG, SS) x 2 Stimulation Time (175, 200 ms post stop 236 signal) RM ANOVA. Participants with fewer than 4 MEPs for more than one stimulation 237 time across trial types were excluded from analysis. For all other analyses in the CME 238 protocol, missing data points were replaced with the average of the row and column mean. To 239 determine task compliance behavioral data were analyzed as follows. LTs on Go and Partial 240 trials were analyzed with a two-way RM ANOVA with factors Trial Type (Go, Partial) and 241 Hand (Left, Right). LTAs from Go trials in the LICI protocol were analyzed using a one-way 242 RM ANOVA with Preceding Trial Type (GG, GS, SG, SS). LTs that contributed to LTAs 243 were analyzed in a two-way RM ANOVA with factors Preceding Trial Type (GG, GS, SG, 244 SS) and Hand. A one-way RM ANOVA with factor Stop Trial Type (GS, SG, SS) tested for differences in SSRT. 245

The second hypothesis was examined in the LICI protocol. A one-way RM ANOVA with 5 Preceding Trial Type (Go Only, GG, GS, SG and SS) was used to analyse CS MEP amplitude and %INH. Linear regression analyses were performed to examine correlations between each of %INH and CS MEP amplitude with LTs of the same trial. To assess the relationship between partial trial CME suppression with both inhibition and response delays, linear regression analyses were performed investigating correlations between GS trial MEP

amplitude 175 ms after the stop cue and each of average LICI across trials and GS lift times. To explore the effects of CME and LICI on lift times linear regression analyses were performed to assess the correlations of  $\Delta$ %INH with LTAs and  $\Delta$ CS MEP amplitude with LTAs. A paired t-test was used to examine the difference between left hand LTs in stimulated (following GG trials) and unstimulated GG trials.

257 Statistical significance was determined by  $\alpha = 0.05$ . *Post hoc* comparisons were

258 performed using t-tests. Normality was assessed prior to ANOVA using the Shapiro-Wilk

test. Non-spherical data were reported using Greenhouse-Geisser corrected *P* values. Values

are reported as mean  $\pm$  standard error.

261 **Results** 

#### 262 *MEP amplitudes and LICI*

263 For normalized MEP amplitudes of GG and GS trials in the CME protocol (Figure 264 2A) there was a main effect of Time ( $F_{5,75} = 23.8$ , P < 0.001), but not Trial Type ( $F_{1,15} = 0.79$ , P = 0.387) or Trial Type × Time interaction ( $F_{5,75} = 1.59$ , P = 0.211). During GG trials, there 265 266 was a main effect of Time ( $F_{6,90} = 26.6$ , P < 0.001) where MEP amplitudes at -175, -150, -125 and -100 ms were greater than baseline at -250 ms (all P < 0.008). During GS trials, there 267 was a main effect of Time ( $F_{5,75} = 8.7$ , P < 0.001) where MEP amplitudes at -100, ( $t_{15} = 3.9$ , 268 P = 0.001) and -25 ms ( $t_{15} = 4.2$ , P < 0.001) were greater than baseline at -150 ms, whereas -269 75 ms ( $t_{15} = 1.1$ , P = 0.289) and -50 ms ( $t_{15} = 1.7$ , P = 0.096) were not. Additionally, MEP 270 amplitude at -75 ms was less than -100 ms ( $t_{15} = 2.3$ , P = 0.035), indicative of non-selective 271 272 braking 175 ms after the stop signal. For Stop trials (Figure 2B; n = 9), there was no main 273 effect of Trial Type ( $F_{2,16} = 0.78$ , P = 0.426) or Time ( $F_{1,8} = 32.5$ , P = 0.152) and no Trial 274 Type × Time interaction ( $F_{2,16} = 0.61$ , P = 0.555). In summary, left FDI MEP amplitudes

demonstrated suppression 175 ms after a stop signal even when the left side was not cued tostop (GS trials).

Figure 3 shows averaged left FDI EMG and MEP amplitudes from an individual participant in the LICI protocol during the Go Only block (Figure 3A) and GS trials in the Stop task (Figure 3B). For LICI, there was a main effect of Trial Type ( $F_{4,48} = 6.5$ , P =0.018). *Post hoc* comparisons revealed that %INH was less during blocks containing only Go trials (53 ± 6 %) compared with all trial types once Stop trials were introduced (all > 70 %; P< 0.025) (Figure 4A).

There was a positive correlation between %INH at the start of GS trials and the resulting LT (r = 0.660, P = 0.014; Figure 4B) such that greater %INH was associated with longer LTs of the left hand during GS trials. There was no correlation for GG trials (r =0.032, P = 0.917). There was no correlation between  $\Delta$ %INH on a GS trial and the LTA on the subsequent GG trial (r = 0.081, P = 0.792).

For CS MEP amplitude there was a main effect of Trial Type ( $F_{4,48} = 3.9, P = 0.034$ ) 288 (Figure 5). Post hoc comparisons revealed that CS MEP amplitudes were greater following 289 Go trials in the Stop task ( $2.3 \pm 0.3 \text{ mV}$ ) than during the Go Only block ( $1.9 \pm 0.3 \text{ mV}$ ;  $t_{12} =$ 290 2.4, P = 0.034). Therefore, responding in the context of the Stop task increased CME. 291 292 Furthermore, CS MEP amplitudes after GS trials  $(2.4 \pm 0.4 \text{ mV})$  were larger than after SG, 293 SS and Go Only trials (all  $\leq 2.2 \pm 0.3$  mV;  $P \leq 0.047$ ). This indicates that the re-initiation of movement on a Partial trial increased contralateral M1 excitability which persisted to the start 294 295 of the subsequent trial.

For GS trials, there was an association between MEP amplitude 175 ms post stop cue and LTs (r = -0.544) as well as with LICI (r = -0.504). However, both correlations failed to reach statistical significance (LTs, P = 0.054; LICI, P = 0.079). There was no correlation

between CS MEP amplitude at the start of a GG or GS trial and the resulting LT (both r < r

300 0.178, P > 0.560). Likewise, there was no correlation between the  $\Delta$ CS MEP amplitude on a

GS trial and the LTA on the subsequent Go trial (r = 0.019, P = 0.951).

In the CME protocol, TMT =  $38 \pm 2\%$  MSO and stimulation intensity =  $39 \pm 2\%$ 

MSO (104  $\pm$  2% TMT). In the LICI protocol, TMT = 43  $\pm$  2% MSO, CS and TS = 56  $\pm$  3%

MSO ( $129 \pm 3 \%$  TMT). The number of trials excluded for rmsEMG > 10  $\mu$ V was 28  $\pm 4 \%$ 

in the CME and  $9 \pm 2\%$  in the LICI protocol. In the CME protocol under the GS condition, 7

306 out of 96 values for MEP amplitude were missing due to pre-trigger EMG and replaced

307 according to the method described.

#### 308 *Lift times and asynchronies*

The task was performed successfully as evident in LTs that were close to the target 309 310 (Table 1), and as noted previously for this task (Coxon et al. 2007; 2009; Coxon et al. 2012; 311 MacDonald et al. 2014; MacDonald et al. 2012). For LTs in the CME protocol, there was a main effect of Trial Type ( $F_{1,15} = 100.0, P < 0.001$ ). No main effect of Hand ( $F_{1,15} = 0.7, P =$ 312 0.409) or Trial Type × Hand interaction ( $F_{1,15} = 0.0, P = 0.887$ ) existed. For the LICI 313 protocol, there was a main effect of Trial Type ( $F_{1,12} = 111.6$ , P < 0.001). There was a main 314 effect of Hand ( $F_{1,12} = 31.9$ , P < 0.001) with right LTs ( $35 \pm 4$  ms) faster than left ( $47 \pm 4$ 315 ms). There was also a Trial Type × Hand interaction ( $F_{1,12} = 4.9$ , P = 0.048), which likely 316 317 arose from a trend for longer left hand delays ( $61 \pm 5$  ms) than right hand delays ( $55 \pm 6$ ms;  $t_{12} = 2.1$ , P = 0.054) between Partial and Go trials. There was no difference in left LTs 318 between stimulated (19  $\pm$  3 ms) and unstimulated (21  $\pm$  3 ms;  $t_{12}$  = 1.1, P = 0.294) Go trials. 319 Lift time asynchronies (LTAs) were analysed from GG trials in the Stop task of the 320 321 LICI protocol (Table 1). For the Stop task of the LICI protocol, there was a main effect of

Preceding Trial Type ( $F_{3,36} = 16.1$ , P < 0.001) such that LTAs decreased after Partial GS

compared with after GG trials ( $t_{12} = 3.9$ , P = 0.002). In contrast, LTAs increased after Partial 323 SG compared with after GG trials ( $t_{12} = 4.9$ , P < 0.001). LTAs were not different after SS 324 trials compared with after GG trials ( $t_{12} = 0.3$ , P = 0.769). Figure 6 shows the LTs of GG 325 326 trials in the Stop task used for LTAs in the LICI protocol. Interestingly, LT on either side was 327 faster on a subsequent GG trial if that side had previously responded on a Partial trial. There was a main effect of Preceding Trial Type ( $F_{3,36} = 5.6$ , P = 0.003) and Hand ( $F_{1,12} = 35.6$ , P < 0.003) 328 329 0.001). There was also a Preceding Trial Type × Hand interaction ( $F_{3,36} = 15.4, P < 0.001$ ). Left LTs were faster after GS trials compared with after GG ( $t_{12} = 4.6$ , P = 0.001) and SS 330 trials ( $t_{12} = 2.9$ , P = 0.014) which did not differ from each other ( $t_{12} = 1.0$ , P = 0.329). 331 332 Similarly, right LTs after SG trials were faster compared with after GG ( $t_{12} = 5.3$ , P < 0.001) and SS trials ( $t_{12} = 3.1$ , P = 0.009) which did not differ from each other ( $t_{12} = 0.7$ , P = 0.500). 333 334 For the left hand, LTs were faster if the hand had previously stopped on a Partial trial, although to a lesser extent than if it had responded ( $t_{12} = 4.6$ , P = 0.001). For the right hand, 335 LTs were faster after responding versus stopping on a Partial trial ( $t_{12} = 3.4$ , P = 0.005). 336

#### 337 Stop signal reaction times and stopping success

338	Average success on Stop trials was as follows: CME protocol: GS: $76 \pm 3\%$ , SG: $61 \pm 3\%$
339	6%, SS: $58 \pm 4\%$ . LICI protocol: GS: $83 \pm 3\%$ , SG: $82 \pm 3\%$ , SS: $65 \pm 3\%$ . In both protocols,
340	average SSRT showed a main effect of Stop Trial Type (CME: $F_{2,30} = 57.3$ , $P < 0.001$ , LICI:
341	$F_{2,24} = 55.5, P < 0.001$ ). SSRT was shorter for SS (CME: 209 ± 3 ms, LICI: 201 ± 2 ms) than
342	GS (CME: 245 ± 4 ms; $t_{15}$ = 11.5, $P < 0.001$ , LICI: 236 ± 4 ms; $t_{12}$ = 9.3, $P < 0.001$ ) and SG
343	trials (CME: $256 \pm 4$ ms; $t_{15} = 8.6$ , $P < 0.001$ , LICI: $236 \pm 4$ ms; $t_{12} = 8.1$ , $P < 0.001$ ). Partial
344	trial types did not differ from each other in the LICI protocol ( $t_{12} = 0.1$ , $P = 0.956$ ), whereas
345	GS SSRT was shorter than SG in the CME protocol ( $t_{15} = 2.3, P = 0.038$ ).

346 **Discussion** 

This study provides novel insights into the non-selective suppression of motor output 347 348 in the context of reactive response inhibition. As expected, responses were delayed on Partial 349 trials and temporal modulation of CME for partial response cancellation was consistent with 350 the anticipation, suppression, and subsequent re-initiation of movement. Furthermore, CME 351 suppression was evident when one or both hands were required to stop. CME at trial onset 352 reflected the sum of inhibitory and facilitatory processes required to successfully perform the 353 previous trial. Changes to LTAs after Partial trials were driven by speeding up of LTs of the 354 hand that had previously responded. The magnitude of LICI at trial onset was positively 355 associated with the extent of the delay during GS trials. Interestingly, LICI increased when 356 Stop trials were introduced compared with a block of trials in which stopping was not a 357 possibility. These results may indicate that LICI is a proactive mechanism capable of 358 influencing the interference effect during partial cancellation performed in a reactive context. Response delays and CME modulation during Stop trials indicated non-selective 359 360 suppression. As observed previously, LTs in Partial trials were delayed relative to Go trials 361 (Coxon et al. 2007; 2009; Coxon et al. 2012; MacDonald et al. 2014; MacDonald et al. 2012). These substantial delays were observed despite participants achieving relatively high success 362 363 rates on Partial trials, especially in the LICI protocol (> 80%). It is important to note that 364 response delays were not eliminated, or even reduced, when success rate increased as a result 365 of familiarity with Partial trials; c.f. (Xu et al. 2014). Modulation of MEP amplitude in the 366 CME protocol supported a model of non-selective suppression during Partial trials. Go trials 367 displayed a sustained increase in CME from 200 ms before the target. The delay on Partial 368 trials was a result of an initial facilitation, a dip back to baseline, followed by a secondary 369 increase in CME. This pattern of CME replicates those of MacDonald et al. (2014). At 370 equivalent times relative to the stop signal, MEP amplitude did not differ between the three

371 types of Stop trials. This finding is consistent with functional magnetic resonance imaging 372 studies showing a similar pattern of neural activation between the three Stop trial types 373 (Coxon et al. 2016; Coxon et al. 2009; Coxon et al. 2012). Conversely, functional imaging 374 results from Majid et al. (2013) suggest a distinct role of the selective indirect basal ganglia 375 pathway during partial stopping. Activation of the indirect pathway should have no effect on 376 MEP amplitude in the responding finger during Partial trials. However, all trials in their study 377 were preceded by a warning cue about stopping requirements. The neural activation in 378 response may differ if the cue is unexpected. The present study adds weight to the model of 379 non-selective response inhibition following an unexpected stop cue.

380 The present study also provides insight into the modulation of intracortical inhibition 381 during response inhibition. Compared with Go Only blocks, LICI increased when Stop trials 382 were introduced. The amount of LICI was comparable between Go and Stop trials suggesting 383 that LICI is not specifically associated with stopping. Previous results using paired-pulse 384 TMS indicated that increased SICI did not coincide with CME suppression in the responding 385 effector during partial response inhibition conditions (MacDonald et al. 2014). Together, 386 these findings indicate that GABAergic circuits within M1 are not primarily responsible for 387 non-selective suppression during reactive response inhibition. Why then, did LICI increase 388 during response inhibition trials? It is likely that increased LICI reflects the proactive 389 modulation of tonic inhibitory circuits as a result of expecting to occasionally stop one or 390 both hands. Studies have demonstrated that CME is modulated as a result of foreknowledge 391 about an ensuing response. When response *execution* is forewarned, SICI and LICI are 392 decreased in the foreperiod for the muscles cued to respond (Sinclair and Hammond 2008) 393 while CME is simultaneously reduced, most likely to prevent a premature response 394 (Davranche et al. 2007; Duque and Ivry 2009). When response prevention is forewarned, 395 CME is similarly reduced during the foreperiod (Cai et al. 2011), acting as a mark of

advanced inhibitory control. Prior to this study, there had been no examination of ICI

modulation as a result of the knowledge that a prepared response may need to be prevented.

LICI increased during the foreperiod (trial onset) with the foreknowledge that stopping was a
possibility. Therefore, proactive inhibitory control is at least in part, mediated by changes in
LICI.

401 The implications of proactively increasing LICI for reactive response inhibition can 402 be understood within the framework of an activation threshold model (e.g., MacDonald et al. 403 2014). Tonic levels of inhibition mediate premature response prevention (Duque et al. 2010; 404 Jaffard et al. 2008) requiring a facilitatory process in order to initiate movement. In the 405 present study, a concurrent increase in LICI and CME observed on Go trials when Stop trials 406 were introduced provide candidate mechanisms. The increased CME (facilitation) counteracts 407 the rise in tonic inhibition and Go trial LTs are thereby maintained on target. However on GS 408 trials, LTs were markedly delayed. In response to the stop signal, a reactive inhibitory input is 409 superimposed onto the tonic resting level, raising the threshold for responding (activation 410 threshold) and effectively cancelling all movement. The trend in the association between 411 greater CME suppression and greater non-selective inhibition (LICI) on GS trials in the 412 current study supports such a model. The initial facilitatory process is inadequate to surpass 413 the activation threshold and a second phase of facilitation must be added, resulting in the 414 response delay (MacDonald et al. 2014). It is worth noting that longer GS response delays 415 were associated with higher levels of LICI, supporting the idea that a second phase of 416 facilitation is required to overcome the tonic resting level. The trend between longer response 417 delays and CME suppression for GS trials is in agreement with such a second phase of 418 facilitation. It is interesting that the association between MEP amplitude and response delay 419 was not stronger for CME evaluated closer in time to the response than LICI measured at trial 420 onset. A likely explanation is that MEP amplitude reflects the net excitatory and inhibitory

inputs whereas LICI provides a measure of inhibition only. How LICI is modulated within
the proactive response inhibition network remains unclear (Majid et al. 2013; Van Belle et al.
2014). The present study supports the idea that proactive and reactive control mechanisms are
not independent but rather, reactive stopping depends on ongoing proactive control (Dunovan
et al. 2015).

426 The within trial processes outlined above have neurophysiological and behavioural 427 consequences for the subsequent trial. Left hand LT on a Go trial was quicker if preceded by 428 a GS compared with a Go trial. At the same time, CS MEP amplitude was increased after GS 429 trials compared with other Stop trial types. Therefore, it may be that the second phase of 430 facilitation required to respond on GS trials has a carry-on effect which is evident on the next 431 trial. The second phase of facilitation on SG trials also explains the speeding up of the right 432 hand on a subsequent Go trial. Interestingly, the hand that stops on a Partial trial also speeds 433 up to some degree on the subsequent Go trial. We suspect that Partial trials require 434 "uncoupling" of the two effectors involved in the default Go response, in order to selectively 435 initiate a unimanual response (MacDonald et al. 2012). Some residual effect of uncoupling is 436 still present on the subsequent Go trial as is evident by the heightened LTAs after Partial 437 trials. The presence of (weaker) coupling suggests that the required second phase of 438 facilitation for the unimanual response on the Partial trial will affect the entire bimanual 439 response on the following trial. In other words, the hand that stops on a Partial trial to some 440 extent "comes along for the ride" on a subsequent Go trial. The fact that this dependence is 441 seen more strongly in left hand LTs aligns with the idea that the nondominant hand is more 442 stringently coupled to the dominant than vice versa (Byblow et al. 2000; Carson 1993). 443 Therefore, the after-effects of Partial trials on corticomotor excitability and performance 444 likely result from the second phase of facilitation, rather than any lingering effects of 445 inhibition.

446 A potential limitation of the present study was the timing of the LICI measurement. 447 At a cellular level, postsynaptic hyperpolarization mediated by GABA<sub>B</sub> receptors has been 448 observed up to 500 ms (Lacaille 1991; Otis et al. 1993). However, it is not feasible to record 449 LICI with the required stimulus intensities within the time window between CME 450 suppression and LT without disrupting task performance. Furthermore, it is also difficult to 451 maintain a comparable level of CME or interpret LICI with a constant test stimulus during or 452 immediately following trials in a task where there is dynamic modulation of CME. However, 453 if LICI is responsible for non-selective suppression on Stop trials and the resulting LTAs on 454 the subsequent trials, one would expect LICI to still be modulated at the time we applied 455 TMS (i.e. at the onset of trials where LTAs are present). Stimulation at this time is unlikely to 456 affect task performance (Leocani et al. 2000) with CME variability reduced compared with 457 times closer to the target (Coxon et al. 2006). However CS MEP amplitude varied following 458 different trial types. The strength of the CS influences the amount of LICI, with a higher CS 459 intensity resulting in reduced LICI (Sanger et al. 2001). This has the potential to complicate 460 the comparison of LICI across and between trials. Nonetheless, it is important to note that the 461 mean CS MEP amplitude was between 1.9 and 2.4 mV (~130 % of TMT), where similar 462 amounts of LICI are reported (Opie and Semmler 2014). Furthermore, CS MEP amplitude 463 did not correlate with GS trial LTs. Thus it is unlikely that CS MEP amplitude could solely 464 account for the observed pattern of LICI. It is possible that reduced LICI in Go Only blocks 465 may reflect, in part, an order effect. Go Only trials were always presented first in the LICI 466 protocol. However, an order effect seems unlikely given that all participants had completed 467 the CME protocol prior to the LICI protocol, and stop trials were presented throughout the 468 entire CME protocol.

- In summary, this study provides novel insight into the role of LICI during movement
- 470 cancellation. LICI is a proactive mechanism capable of influencing the interference effect
- 471 during partial cancellation performed in a reactive context.

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# 477 Disclosures

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600

#### 602 Figure Captions

603 Figure 1. Bimanual anticipatory response inhibition task. A. Each trial began with the target 604 line (horizontal black bar) displayed and trial type ambiguous. Go trials (GG) were deemed 605 successful if both ascending indicators were stopped within 30 ms of the target line (800 ms 606 into the trial) by lifting the left and right index fingers from switches. To indicate "Success" 607 the target line turned green. **B.** Partial trial (GS = Go-Left Stop-Right): the right indicator 608 automatically stops 550 ms into the trial. This trial type requires the right switch to remain 609 depressed while the left switch must be released as the rising indicator reaches the target line. 610 The target line turned red if the indicator stopped greater than 30 ms of the target line or the 611 hand response was not correctly inhibited ("Miss"). C. Stop-Both trial (SS): Both indicators 612 stop automatically 600 ms into the trial and a successful trial is achieved when both switches 613 remain depressed. **D.** Transcranial magnetic stimulation over the right motor cortex. Motor 614 evoked potentials (MEPs) were recorded from the first dorsal interosseous (FDI) muscle during task performance. 615

616

617 Figure 2. Modulation of left first dorsal interosseous normalized motor evoked potential 618 (MEP) amplitude during the corticomotor excitability protocol. A. MEP amplitudes before 619 the target (0 ms) during GG (unfilled circles) and GS (shaded squares) trials (n = 16). Filled 620 symbols represent values significantly different ( $P \le 0.05$ ) to baseline (GG: -250 ms, GS: -621 150 ms). Note the dip in corticomotor excitability on GS trials at -75 ms. B. MEP amplitudes 622 after the stop signal. Bars indicate group means (n = 9). Data correspond to -75 and -50 ms 623 relative to lift time from Panel A. Note that MEP amplitudes were suppressed 175 ms after the stop signal regardless of Stop trial type. Error bars indicate 1 SE. \* P < 0.05. GG = Go 624 trial, GS = Go-Left Stop-Right, SG = Stop-Left Go-Right, SS = Stop-Both. 625

**Figure 3.** Representative left first dorsal interosseous electromyography from a single

627 participant in the long-interval intracortical inhibition protocol. Vertical dashed line

represents target (800 ms into the trial) and arrows represent average lift time (LT). A. Go

- Only block. % inhibition = 48 %, LT = 786 ms, CS motor evoked potential (MEP) amplitude
- = 2.2 mV. **B.** Successful GS trials. % inhibition = 72 %, LT = 868 ms, CS MEP amplitude =
- 631 2.2 mV. CS = conditioning stimulus, TS = test stimulus.
- **Figure 4.** Group averages (n = 13) for measures of long-interval intracortical inhibition

633 (LICI) at trial onset. A. Percent inhibition relative to the previous trial. B. Linear regression

between % LICI and lift times within GG (unfilled circles) and successful GS trials (filled

squares). Error bars indicate 1 SE. \* P < 0.05. Go Only = block with only Go trials possible,

GG = Go trial in the context of the response inhibition task, GS = Go-Left Stop-Right, SG = Go

637 Stop-Left Go-Right, SS = Stop-Both.

**Figure 5.** Corticomotor excitability at trial onset. Bars are group averages (n = 13) of the

639 conditioning stimulus (CS) motor evoked potential (MEP) amplitude at trial onset relative to

640 previous trial type. Only data following successful Stop trials were included in the analysis.

Error bars indicate 1 SE. \* P < 0.05. Go Only = block with only Go trials possible, GG = Go

trial in the context of the response inhibition task, GS = Go-Left Stop-Right, SG = Stop-Left

 $643 \qquad \text{Go-Right, SS} = \text{Stop-Both.}$ 

**Figure 6.** Lift times on Go trials following different trial types. Bars indicate group means (n

645 = 13). Black horizontal lines denote when the hand had previously stopped on a Partial trial.

Note that for both hands lift times were faster on a subsequent Go trial if the hand had

previously responded on a Partial trial. Error bars indicate 1 SE. \* P < 0.05, \*\* P < 0.001.

648

# 650 Tables

	LTs						
	Go (L)	Go (R)	Go Avg	Partial GS	Partial SG	Partial Avg	
CME protocol (ms)	16±3	13±2	14±2	70±8	66±7	68±6*	
LICI protocol (ms)	17±2	7±2	12±2	78±6	62±6	70±6*	
LICI protocol		Preceding trial type					
		Go	Partial (	GS Part	ial SG	Stop Both	
LTA (ms)		10±2	3±2†	14±	2†	9±1	

**Table 1.** Summary of behavioural data from both protocols.

Values are mean±SE lift times (LTs) displayed relative to the target (800 ms). Lift time asynchronies (LTAs) are determined as left LT – right

LT in Go trials. GS = Go-Left Stop-Right, SG = Stop-Left Go-Right. \* P < 0.001 compared with Go Avg, † P < 0.01 compared with Go trials.

654





Stimulation Time Relative To Stop Signal (ms)







