Articles in PresS. J Neurophysiol (May 11, 2016). doi:10.1152/jn.00225.2016

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MOTOR LEARNING AND CROSS-LIMB TRANSFER

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3	Motor learning and cross-limb transfer rely upon distinct neural adaptation processes
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18	Running head: Motor learning and cross-limb transfer
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29	Abstract
30	Performance benefits conferred in the untrained limb after unilateral motor practice are
31	termed cross-limb transfer. Although the effect is robust, the neural mechanisms remain
32	incompletely understood. Here we use non-invasive brain stimulation to reveal that the
33	neural adaptations that mediate motor learning in the trained limb are distinct from those
34	that underlie cross-limb transfer to the opposite limb. Thirty-six participants practiced a
35	ballistic motor task with their right index finger (150 trials), followed by intermittent-theta
36	burst stimulation (iTBS) applied to the trained (contralateral) primary motor cortex (cM1
37	group), the untrained (ipsilateral) M1 (iM1 group), or the vertex (sham group). Following
38	stimulation, another 150 training trials were undertaken. Motor performance and
39	corticospinal excitability were assessed before motor training, pre- and post-iTBS, and
40	following the second training bout. For all groups, training significantly increased
41	performance and excitability of the trained hand, and performance, but not excitability, of
42	the untrained hand, indicating transfer at the level of task performance. The typical
43	faciltatory effect of iTBS on MEPs was reversed for cM1, suggesting homeostatic
44	metaplasticity, and prior performance gains in the trained hand were degraded, suggesting
45	that iTBS interfered with learning. In stark contrast, iM1 iTBS facilitated both performance
46	and excitability for the untrained hand. Importantly, the effects of cM1 and iM1 iTBS on
47	behaviour were exclusive to the hand contralateral to stimulation, suggesting that
48	adaptations within the untrained M1 contribute to cross-limb transfer. However, the neural
49	processes that mediate learning in the trained hemisphere versus transfer in the untrained
50	hemisphere appear distinct.
51	

52 **Keywords:** ballistic motor learning, inter-limb transfer, non-invasive brain stimulation,

53 corticospinal excitability, motor performance

54	New & Noteworthy
55	In the present study we observed that non-invasive brain stimulation interacted
56	differently with motor practice when applied to the motor cortex projecting to the trained
57	versus the untrained limb. This suggests that distinct neural processes underlie learning
58	obtained via direct motor practice, and learning conferred indirecty from practice with the
59	opposite limb (i.e. cross-limb transfer). The results provide a step forward in using non-
60	invasive brain stimulation methods to promote cross-limb transfer in motor rehabilitation.

61	Introduction
62	Generalization of learned actions is critical for flexible and adaptive human behavior;
63	it is clearly advantageous to be able to apply motor skill obtained in one context to
64	alternative spatial locations, movement directions and effectors. Cross-limb transfer
65	describes the behavioral benefit conferred in the untrained limb (i.e., inter-limb
66	generalization) following unilateral motor practice. Although this effect has been studied for
67	over a century (see Carroll et al. 2006; Farthing et al. 2009; Ruddy and Carson 2013 for
68	overviews) the neural mechanisms mediating performance gains in the untrained limb
69	remain incompletely understood.
70	While adaptations at the spinal level cannot be excluded, the available evidence
71	suggests that adaptations within cortical networks that project to the untrained limb are
72	likely to be primarily responsible for the phenomenon of cross-limb transfer (see Ruddy and
73	Carson 2013 for an overview). The data are consistent with Parlow and Kinsbourne's (1989)
74	cross-activation hypothesis, which suggests that during motor learning, task-relevant
75	information is simultaneously stored in both the trained and untrained hemispheres (also
76	Cramer et al. 1999; Dettmers et al. 1995). Transcranial magnetic stimulation (TMS) studies
77	also show that activation of one limb results in contraction intensity dependent excitability
78	changes of the pathways projecting to the opposite limb (e.g., Hess et al. 1986; Liepert et al.
79	2001); the stronger the contraction of one limb, the greater the change in excitability
80	observed in the projections to the opposite limb (Perez and Cohen 2008).
81	Motor learning paradigms utilizing simple ballistic movements, in which participants
82	aim to maximize the rate of force development or acceleration of the upper limb or hand
83	(e.g. Classen et al. 1998), represent an ideal model to study the mechanisms of adaptation
84	and transfer. Using a "virtual lesion" TMS approach in this paradigm, Lee et al. (2010)

85	showed that adaptations within each hemisphere specifically mediate performance
86	improvements of the contralateral limb, irrespective of whether the performance gains are
87	due to direct practice or transfer. However, it remains unknown whether the synaptic
88	mechanisms of adaptation are similar in the two hemispheres.
89	Here we used a non-invasive brain stimulation (NBS) protocol that induces effects
90	that resemble long-term potentiation (LTP) in the resting brain (intermittent theta-burst
91	stimulation, iTBS; Huang et al. 2005), to study the synaptic mechanisms that underlie
92	performance improvements in the trained and untrained limbs. Specifically, following
93	unilateral ballistic motor learning, we administered iTBS to the trained (contralateral) or
94	untrained (ipsilateral) primary motor cortex. When applied following motor training, the
95	'expected' effects of NBS protocols that induce LTP-like effects at baseline can be occluded
96	or reversed (Rosenkranz et al. 2007; Stefan et al. 2006; Ziemann et al. 2004) according to
97	principles of homeostatic plasticity (i.e., Müller-Dahlhaus and Ziemann 2015 for a review),
98	which provides evidence that learning is driven by LTP-like plastic changes. Here, we tested
99	whether training-induced performance gains in the trained (direct learning) and untrained
100	hands (cross-limb transfer) are driven by similar, LTP-like, neural adaptations in the trained
101	and untrained motor cortices, respectively. If the synaptic mechanisms of learning and
102	transfer are similar in each hemisphere, then the LTP-like effects of iTBS should be reduced
103	or reversed in both the trained and untrained motor cortices (see Figure 1A). If however,
104	transfer represents a distinct neural process to learning, then iTBS applied to the untrained
105	hemisphere following training would be predicted to induce similar effects as when applied
106	in isolation (Figure 1B).

107

108

"Figure 1 about here"

109	Because it is of practical interest, for potential therapeutic applications, to
110	understand the impact of plasticity-inducing NBS on the capacity for subsequent
111	performance improvements via transfer, we also assessed performance changes due a
112	second block of unimanual training performed after iTBS. Prior induction of LTP-like plasticity
113	can enhance subsequent learning for the contralateral limb via non-homeostatic processes
114	(Teo et al. 2011), but the effects of synaptic plasticity induction upon subsequent transfer
115	have not been reported. If similar mechanisms apply to cross-limb transfer, we should see
116	the effects of iTBS to the untrained M1 reflected in subsequent performance gains according
117	to non-homeostatic processes (i.e. LTP-like effects should result in enhanced subsequent
118	performance gains, whereas LTD-like effects should impair subsequent performance gains).
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131 Task and procedure

132	The experiment was designed to use non-invasive brain stimulation to interact with
133	training-induced plasticity in the trained and untrained hemisphere following unilateral
134	motor training. We aimed to determine whether training-induced performance gains in the
135	trained (direct learning) and untrained hands (cross-limb transfer) are driven by similar, LTP-
136	like, neural adaptations in trained and untrained motor cortices, respectively. Figure 2
137	outlines the experimental procedure. Following Hinder et al. (2011, 2013) and Lee et al.
138	(2010), participants practiced a ballistic abduction of the right index finger (audio-paced at
139	0.5 Hz) where the performance goal was to maximise peak horizontal (abduction)
140	acceleration of each movement. This type of motor learning paradigm represents an ideal
141	model to study the mechanisms of adaptation and transfer for many reasons. Substantial
142	performance gains are exhibited within a single session, which simplifies the use of brain
143	stimulation methods such as transcranial magnetic stimulation (TMS) to assess the neural
144	underpinnings of adaptation (Carroll et al. 2008). Moreover, the neural responses to ballistic
145	motor training are similar to those observed after strength training (Selvanayagam et al.
146	2011). Accordingly the model provides a window into the mechanisms underlying an
147	important physical attribute that often limits function in old age and in patients with
148	neurological disorders. Triaxial accelerometers (Dytran Instruments, Chatsworth, CA;
149	Endevco, San Juan Capistrano, CA) were mounted to plastic splints and taped to the top of
150	the left and right index fingers such that one of the orthogonal axes of each accelerometer
151	was aligned to measure horizontal acceleration. A custom written Signal (CED) script (see
152	Hinder et al. 2011, 2013) allowed us to detect the first peak of the acceleration trace and
153	provide this information to participants as visual performance feedback according to the
154	feedback design (see below).

"Figure 2 about here"

158	Participants undertook a total of 300 practice trials within two training blocks, each
159	consisting of 150 movements (cf. Hinder et al. 2011, 2013). 30s rest breaks were provided
160	every 15 movements (i.e., ten 15-movement sub-blocks per block) to avoid fatigue. Visual
161	feedback of the movement outcome was provided on 50% of the movements (i.e., odd-
162	numbered sub-blocks) to assist in promoting performance gain (Winstein and Schmidt,
163	1990).
164	In order to specifically interact with the neural adaptations mediating performance
165	gains in the trained hand (i.e., direct motor learning gains) and the untrained hand (i.e.,
166	cross-limb transfer), we applied intermittent theta burst stimulation (iTBS) to the trained or
167	untrained M1, or to the vertex as a 'sham' condition, after the first training block. iTBS has
168	been shown to increase motor evoked potential (MEP) amplitude in a manner consistent
169	with LTP-like plasticity (Huang et al. 2005). Consistent with principles of homeostatic
170	plasticity (Müller-Dahlhaus and Ziemann 2015, as well as Karabanov et al. 2015 for an
171	overview), we postulated that the LTP-like effects of iTBS on MEPs would be reduced or
172	reversed in both the trained or untrained motor cortices, if both learning and transfer are
173	driven by LTP-like plastic changes. If untrained hand performance gains following unilateral
174	ballistic practice (cross-limb transfer) are not driven by LTP-like plastic changes in the
175	untrained motor cortex, iTBS should be able to act in the 'expected' direction (Huang et al.
176	2005) and facilitate MEPs within the cortical network that projects to the untrained limb.
177	iTBS (600 pulses, 190s stimulation; cf. Huang et al. 2005) was administered (Magstim Super
178	Rapid ² stimulator and 70mm figure-of-eight-coil) at an intensity of 80% of active motor
179	threshold (AMT) over the motor hotspot (coil handle 45° to the midline) of the trained first

180	dorsal interosseus (FDI) muscle (cM1 group), the untrained FDI muscle (iM1 group), or over
181	the vertex (handle backwards) with the coil tilted by 90° (coil surface orthogonal to the scalp
182	surface) with one side of the coil remaining contact with the head (sham group; Mistry et al.
183	2012). The active motor threshold was defined as the minimum stimulator intensity required
184	to evoke MEPs of \geq 200 μV (in three out of five trials) (Huang et al. 2005) during a light
185	isometric contraction of the corresponding FDI muscle at about 10 % of maximum force.
186	Motor performance (i.e., peak acceleration in 10 test movements per hand) and
187	neurophysiological measures (i.e., cortical excitability and intracortical inhibition as assessed
188	with TMS) were obtained for both hands/ motor cortices before motor training commenced
189	(pre-test), after the first motor training block but before iTBS administration (pre-iTBS),
190	immediately following iTBS (post-iTBS), and following completion of the second training
191	block (post-test). TMS testing always preceded motor performance testing at each of the
192	time points such that changes in neurophysiological measures could be attributed to the
193	unilateral training block rather than the test phases conducted with both hands; the hand-
194	order during motor performance and TMS testing was counterbalanced across participants
195	within each group.

196

197 *Recording of muscle activity*

In order to quantify muscle activity (during the execution of the motor task and in
response to suprathreshold pulses of TMS) we recorded EMG activity with Ag/AgCl
electrodes (Meditrace 130, Tyco Healthcare, Mansfield, MA) from the FDI in both hands in a
belly-tendon montage (as per Hinder et al. 2011, 2013). EMG signals were fed into a CED
1401 amplifier (Cambridge, UK), where a notch filter (50 Hz) was applied before
amplification (gain 300–1,000), and stored for off-line analysis. Participants' EMG activity

- was constantly monitored by the experimenter to guarantee strong movement-related FDI
 bursts in the activated hand and a relaxation of the muscle between trials.
- 206
- 207 Transcranial magnetic stimulation

208 Transcranial magnetic stimulation (TMS) was delivered using two Magstim 200² units (Magstim Company, Dyfed, UK) connected via a Bistim² unit and a single figure-of-eight coil 209 210 (70 mm external diameter). Motor 'hotspots' for the left and the right FDI (with posterior- to anterior-induced current in the cortex) were determined and resting motor thresholds 211 (RMT) were established as the minimum intensities required to elicit MEPs > 50 μ V in the 212 213 right and left FDI muscles in three out of five consecutive trials when stimulating at the hotspots (Carroll et al. 2001; Hinder et al. 2010). Participants were instructed to relax their 214 limbs during RMT determination and visual feedback of muscle activity helped to keep 215 216 muscle activity to a minimum.

217 During TMS test blocks, 30 stimulations (with an interstimulus interval of 4-6 s) were administered to the right (untrained) or left (trained) motor hotspots, respectively. Half of 218 the stimulations involved a single 'test' pulse (130 % RMT) to assess the net excitability of 219 220 the corticospinal projections to the trained/untrained hand, while half of the trials involved 221 paired-pulse stimulation (Kujirai et al. 1993) in which a subthreshold conditioning pulse (70 222 % RMT) preceded the same test pulse. The ratio of the average MEP evoked following 223 paired-pulse trials (within one TMS test block) to the average MEP amplitude evoked in the 224 single-pulse trials (within the same TMS test block) is referred to as the short-interval 225 intracortical inhibition ratio, SICI (Kujirai et al. 1993), and reflects activity of intracortical inhibitory circuits. The order of single- and paired-pulse stimulations was randomised within 226 227 each TMS block.

228

229 Control Experiment

Because the results of the main experiment showed that both performance and 230 231 MEPs increased in the untrained hand after iTBS to the untrained M1, it was important to 232 determine whether the performance gain reflected a general improvement in motor 233 function due to enhanced excitability, or was due to an interaction with the recently 234 transferred (improved) motor skill. We therefore conducted a control experiment, for which 235 another cohort of healthy, right-handed young adults (n = 12, 3 males, average age = 25.9 years, SD = 7.3) was recruited. Here we examined the effects of iTBS delivered to the right 236 237 M1 (N.B. to correspond with the untrained hemisphere in the main experiment) without prior motor training. MEP amplitudes and motor performance were assessed for right and 238 left hands before and after iTBS, in the absence of a preceding training block. Thus, the 239 240 results of the control experiment allowed us to isolate the effects of iTBS, applied over the right M1, on MEP amplitudes and motor performance without being influenced by prior 241 242 motor training.

243

244 Data analysis

Acceleration data were low-pass filtered at 20 Hz prior to analysis. As per Hinder et al. (2011, 2013) peak acceleration of the ballistic abduction was determined as the first peak in the horizontal acceleration for each movement trial (referred to as ACC). Performance of right and left hand movements at each test phase was calculated as the average peak acceleration across the 10 trials in each test for the respective hand. Performance at preiTBS and post-iTBS was subsequently normalized to pre-test values (referred to as nACC [nACC > 1 indicating increased performance and nACC < 1 decreased performance relative to

252 performance at pre-test]) to explore the effect of iTBS on prior motor training-induced 253 changes in the trained and untrained hands. Performance data at post-test were normalized to values obtained at post-iTBS to examine the influence of iTBS on changes in performance 254 255 in both hands following a second block of motor training. Performance of right hand 256 movements during training was expressed as the average peak acceleration across the 15 trials in each sub-block. Average performance of the trained, right hand in the penultimate, 257 258 ninth sub-block was then normalized to the average performance obtained during the first sub-block of training for the right hand as a measure of training-related changes in trained 259 260 hand performance (referred to as nACC_{training}). The penultimate block was chosen such that 261 we compared sub-blocks in which visual feedback was consistent (i.e., visual feedback of performance was provided in both the first and ninth sub-block, but not the tenth sub-262 block). 263 264 Responses to TMS were sampled at 10 kHz from 3 s before to 2 s after the test pulse.

265 Trials in which background root mean square EMG exceeded 25 μ V in a 40 ms time window immediately prior to TMS stimulation were excluded from further analysis. The peak-to-peak 266 amplitudes of the motor evoked response (MEP) were measured in a window 15–50 ms 267 after stimulation in the limb contralateral to the stimulated cortex. For both 268 269 neurophysiological measures (MEP, SICI), data at pre-iTBS and post-iTBS were normalized to 270 those values obtained at pre-test (referred to as nMEP [nMEP > 1 indicates a facilitatory change while nMEP < 1 indicates suppression of evoked responses, relative to pre-test 271 272 responses] and nSICI [nSICI > 1 indicates a release of inhibition and nSICI < 1 indicates 273 increased inhibition relative to pre-test]) to explore the effect of iTBS on prior motor 274 training-induced changes in excitability and inhibition in both motor cortices. Post-test

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275	values were normalized to those values obtained at post-iTBS to examine the influence of
276	iTBS on changes in excitability/inhibition following a second block of motor training.
277	As potential predictors of cross-limb transfer, we quantified the training-induced
278	change in FDI activity of the trained (right) and untrained (left) hands, as well as a measure
279	of relative mirror muscle activity of the left hand during right hand movements. As per
280	Hinder et al. (2011, 2013) individual EMG signals of trained and untrained FDIs assessed
281	during task execution were processed to only represent movement-related muscle activity
282	during the ballistic action. That is, movement-related EMG data during training trials was
283	rectified and low-pass filtered (20 Hz) before determining the peak EMG amplitude in the
284	active FDI (i.e., of the hand performing the ballistic abduction). Movement onset was
285	defined as the time when FDI activity in the active hand first exceeded 4 times background
286	EMG determined 50–100 ms before the 'go' tone. Movement offset was defined as the time
287	when FDI activity in the hand performing the task first dropped below 0.2 times the peak
288	amplitude (Carroll et al. 2008; Hinder et al. 2011). In this time window, the average burst-
289	related EMG of the FDI in the hand performing the task was calculated minus the average
290	value of background EMG. During the same time window the average FDI EMG in the
291	contralateral hand (i.e., mirror activity) was determined. EMG values were then averaged
292	over the 15 trials of each sub-block of the training. The average values for both the trained
293	and untrained FDI of the penultimate, ninth sub-block were normalized to the average EMG
294	values obtained during the first sub-block of training for the respective hand as a measure of
295	training-related changes in FDI activity in the trained and untrained hands (referred to as
296	nEMG [nEMG > 1 indicating increased FDI activity and nEMG < 1 decreased FDI activity
297	relative to the first sub-block). Additionally, FDI activity of the untrained hand averaged
298	across the 150 training trials was normalized to FDI activity of the trained hand averaged

across these 150 trials as a measure of relative mirror muscle activity during training
(EMG_{mirror}).

301

302 Statistical analysis

303 To ascertain that 1) pre-test values (relating to both behaviour and cortical 304 excitability/inhibition) and training-induced changes from pre-test to pre-iTBS in these 305 parameters were similar across groups and 2) to ensure significant learning and transfer 306 effects following the first motor training block were apparent, we separately submitted raw 307 (non-normalized) peak acceleration, MEP and SICI values to time (pre-test, pre-iTBS) x hand 308 (trained, untrained) x group (cM1, iM1, sham) ANOVAs. Subsequently, normalized (relative 309 to pre-test) performance (nACC) and TMS measures (nMEP, nSICI) were subjected to time (pre-iTBS, post-iTBS) x hand (trained, untrained) x group (cM1, iM1, sham) ANOVAs (for each 310 311 dependent variable separately) to examine the effect of iTBS on prior motor learning gains 312 of the trained and untrained hands, and associated changes in excitability/inhibition of the 313 corresponding motor cortices. Additionally, separate hand (trained, untrained) x group (cM1, 314 iM1, sham) ANOVAs were performed on the (normalized) post-test values for performance and TMS measures to examine the impact of iTBS on subsequent motor training gains (i.e., 315 316 gains in block 2 normalized to post-iTBS values). Significant main or interaction effects were 317 further explored using post hoc pairwise comparisons (using the Sidak adjustment). Main 318 inferential analyses (ANOVA) were complemented by correlation statistics (with Benjamini-319 Hochberg procedure applied to correct for multiple comparisons) where appropriate (e.g., to 320 explore the nature of the interaction between use-dependent and iTBS-induced changes in 321 performance and corticospinal excitability).

322	To benefit from cross-limb transfer effects (e.g., in rehabilitation settings) it is critical
323	to know which factors (e.g., motor learning itself, mirror muscle activity, corticospinal
324	excitability) predict and mediate performance gains in an untrained hand. It is also important
325	to know whether performance gains in an untrained limb after unilateral practice are driven
326	by adaptations in the untrained hand/ motor cortex during training (i.e., via cross activation)
327	or in the trained hand/ motor cortex upon retrieval (i.e., via callosal access). To this end, a
328	multiple regression analysis was employed to identify the main predictors of cross-limb
329	transfer (i.e., normalized performance gains of the untrained hand relative to pre-test
330	performance of that hand) following an initial unilateral practice period (i.e., at pre-iTBS),
331	and to study their relative predictive strength (when controlling for other predictor
332	variables). Two regression models were tested. The first one of these models included three
333	variables derived from the trained (active hand). These were: the normalized performance
334	change of the trained hand from the first to the penultimate, ninth training sub-block
335	(nACC _{training}), the change in burst-related FDI activity of the active (trained) hand from the
336	first to the ninth training block (nEMG _{trained}), and the training-induced change in corticospinal
337	excitability of the trained M1 from pre-test to pre-iTBS (nMEP _{trained}). The second model was
338	complemented by the inclusion of three additional variables related to the untrained hand.
339	Specifically, we considered the change in FDI activity of the untrained hand (as defined
340	above, nEMG _{untrained}), relative mirror muscle activity during training (EMG _{mirror}) and the
341	training-induced change in untrained M1 excitablity (see above, nMEP _{untrained}).

All data are reported as normalized values: to assess the effects of the first training block and subsequent iTBS, behavioural and neurophysiological parameters are expressed relative to the corresponding pre-test value; to assess the affect of the second training block, post-training values are expressed relative to post-iTBS values. Corresponding 95 %

- 346 confidence intervals (CI) provide a measure of variability, while partial eta-squared (η_p^2) and 347 Cohen's *d* are reported as measures of effect size.
- 348
- 349

Results

350 *Motor performance*

351 Training-induced effects on motor performance

Average peak accelerations at pre-test were 20.5 ± 9.4 ms⁻², 20.9 ± 4.6 ms⁻² and 14.8352 \pm 2.4 ms⁻² for the right hand and were 18.9 \pm 5.9 ms⁻², 20.1 \pm 3.7 ms⁻² and 14.4 \pm 2.7 ms⁻² for 353 the left hand for the cM1, iM1 and sham groups, respectively. Upon completion of the first 354 355 training block (i.e., at pre-iTBS), peak acceleration of the index finger had increased to $36.5 \pm$ 10.0 ms⁻², 40.1 \pm 6.7 ms⁻² and 28.5 \pm 8.4 ms⁻² in the trained hand, and to 27.8 \pm 7.6 ms⁻², 31.3 356 \pm 7.1 ms⁻² and 21.7 \pm 4.3 ms⁻² in the untrained hand in the cM1, iM1 and sham groups, 357 358 respectively. ANOVA revealed a significant time x hand interaction, F(1,33) = 25.62, p < .001, η_{ρ}^{2} = .44, with posthoc pairwise comparisons revealing a significant increase in performance 359 from pre-test (trained: $18.8 \pm 3.7 \text{ ms}^{-2}$, untrained: $17.8 \pm 2.6 \text{ ms}^{-2}$) to pre-iTBS (trained: 35.0 ± 10^{-2}) 360 5.2 ms⁻², untrained: 26.9 ± 4.0 ms⁻²) for both, trained and untrained, hands (both p < .001, d 361 362 > .89) averaged across all three groups. Peak acceleration did not differ significantly between hands at pre-test (p = .39, d = .09; pairwise Sidak adjusted post-hoc tests confirmed that 363 baseline performance was not significantly different between any of the groups, all p > 0.27), 364 but was greater for the trained as compared to the untrained hand after motor training at 365 366 pre-iTBS (p < .001, d = .58). ANOVA revealed no significant main or interaction effects including the factor group. 367

368

369 *iTBS-induced effects on motor performance*

370	To assess the impact of the different iTBS protocols on trained and untrained hand
371	performance, we compared normalized peak acceleration values (relative to pre-test) of
372	cM1, iM1 and sham group participants before and after the application of iTBS. As seen in
373	Figure 3, iTBS resulted in a <i>reduction</i> of normalized performance (relative to pre-test) of the
374	trained hand from 2.14 (\pm 0.47) to 1.77 (\pm 0.47) for the cM1 group, whereas normalized
375	performance of the trained hand in the iM1 group (pre-iTBS: 2.01 \pm 0.25; post-iTBS: 1.88 \pm
376	0.40) and sham group (pre-iTBS: 2.02 \pm 0.53; post-iTBS: 2.05 \pm 0.62) was much less affected
377	by iTBS. In contrast, iTBS increased normalized performance in the untrained hand from 1.62
378	(\pm 0.26) at pre-iTBS to 2.03 (\pm 0.54) at post-iTBS in the iM1 group, whereas normalized
379	performance of the untrained hand appeared to be unaffected by iTBS in the cM1 (pre-iTBS:
380	1.57 \pm 0.21; post-iTBS: 1.49 \pm 0.25) and sham groups (pre-iTBS: 1.51 \pm 0.22; post-iTBS: 1.66 \pm
381	0.37). ANOVA conducted on normalized performance revealed a significant time x hand x
382	group interaction, $F(2,33) = 3.75$, $p = .03$, $\eta_p^2 = .19$, confirming the changes described above;
383	i.e., performance changes in response to iTBS were hand- and iTBS location (group)-specific.
384	Indeed, posthoc pairwise comparisons revealed that the performance decrease of the
385	trained hand in the cM1 group ($p = .004$, $d = .43$) and the performance <i>increase</i> of the
386	untrained hand in the iM1 group ($p = .002$, $d = .52$) following iTBS were statistically
387	significant; all other pairwise comparisons were not statistically significant (all $p > .23$, $d <$
388	.26).

Correlation analyses revealed significant (positive) relationships between trained and untrained hand performance gains following motor training at pre-iTBS (r = .68, p < .001) across all participants. That is, the greater trained hand performance improvements, the greater improvements in the untrained hand. Moreover, for the iM1 group, analyses revealed that the greater the increase in untrained hand performance following the first

394	training block, the greater the subsequent iTBS-induced improvements in that hand. That is,
395	we observed a positive relationship between the extent of iTBS-induced change in
396	performance in the untrained hand and the extent of the previous performance gains in the
397	untrained hand (i.e., as a result of cross-limb transfer) following the first training block (r =
398	0.72, $p = .004$). In contrast, for the trained hand of the cM1 group, there was a negative
399	correlation between iTBS-induced performance changes and the previous use-dependent
400	performance gains as a result of the first training block ($r = -0.55$, $p = .03$). This illustrates
401	that the greater the use-dependent performance increase in the trained hand following the
402	first training block, the more performance of that hand is reduced following the application
403	of iTBS.
404	
405	"Figure 3 about here"
406	
407	iTBS-induced effects on subsequent motor training
408	To test for the influence of iTBS on subsequent learning and transfer, post-test data
409	following the second training block were analysed relative to post-iTBS values. Upon
410	completion of the second training block, the normalized performance of participants in the
411	cM1, iM1 and sham groups increased (relative to post-iTBS) to 1.64 (\pm 0.26), 1.34 (\pm 0.24)
412	and 1.18 (± 0.13) in the trained hand, and to 1.29 (± 0.21), 1.15 (± 0.25) and 1.11 (± 0.06) in
413	the untrained hand, respectively. ANOVA revealed the greater performance gains of the cM1
414	group as compared to iM1 and sham groups to be significant averaged across hands (main
415	effect for group: $F(2,33) = 4.22$, $p = .023$, $\eta_p^2 = .20$). Interestingly, across all participants, and
416	for both the trained and untrained hands, there were negative correlations between the
417	performance gains as a result of the second training block, and the extent of previous iTBS-

induced change in performance (trained hand: r = -0.57, p < .001; untrained hand: r = -0.49,

419 p = .001). That is, the greater the iTBS-induced performance decrement in the trained hand,

420 the greater the subsequent learning in that hand in the second block. For the untrained

421 hand, the greater the iTBS-induced gain, the lower the subsequent performance gain in that

- 422 hand resulting from the second training block.
- 423

424 *Neurophysiological measures*

425 Corticospinal excitability

426 Training-induced effects on corticospinal excitability

427 RMTs (as a % of maximum stimulator output, \pm 95% CI) for the right hand were 42.5 \pm 4.2%, 40.7% \pm 3.5% and 43.1 \pm 3.5%; and were 42.1 \pm 4.1%, 40.8 \pm 4.0% and 44.1 \pm 2.4% for 428 the left hand for cM1, iM1 and sham group participants, respectively. There were no 429 430 significant differences between groups, F(2,33) = 0.60, p = .56, $\eta_p^2 = .04$, or hands, F(1,33) =0.11, p = .75, $\eta_p^2 = .003$, and no interaction between hand and group, F(1,33) = 0.30, p = .75, 431 η_p^2 = .02. AMTs were 48.0 ± 3.4%, 47.2 ± 2.5% and 49.4 ± 3.6% of maximum stimulator 432 output for cM1, iM1 and sham group participants, respectively, and did not differ between 433 groups, F(2,33) = 0.45, p = .65, $\eta_p^2 = .03$. (NB: AMT appears higher than RMT because it was 434 determined on the less powerful Magstim Super Rapid² stimulator which was used to 435 subsequently administer iTBS, whereas RMT and single/paired pulse TMS was administered 436 using two Magtim 200² units connected with a BiStim module). Average MEP amplitudes at 437 438 pre-test were 1.49 \pm 0.51 mV, 1.34 \pm 0.56 mV and 0.90 \pm 0.17 mV for the right FDI and were 1.37 ± 0.63 mV, 1.47 ± 0.52 mV and 1.40 ± 0.32 mV for the left FDI for cM1, iM1 and sham 439 440 group participants respectively. Upon completion of the first training block (i.e., at pre-iTBS), 441 cM1, iM1 and sham group participants' excitability increased to 2.18 ± 0.70 mV, 1.94 ± 0.74

442	mV and 1.25 ± 0.35 mV in the trained hand, respectively. However, excitability of the
443	untrained hand was relatively unaffected by training in all three groups (cM1: 1.41 \pm 0.80
444	mV, iM1: 1.42 \pm 0.61 mV, sham: 1.36 \pm 0.33 mV). ANOVA revealed a significant main effect
445	for time, $F(1,33) = 13.11$, $p = .001$, $\eta_p^2 = .28$; averaged across both hands excitability
446	increased from pre-test (1.33 \pm 0.28 mV) to pre-iTBS (1.59 \pm 0.37 mV); however the
447	significant time x hand interaction, $F(1,33) = 18.05$, $p < .001$, $\eta_p^2 = .35$ indicates that this
448	effect was driven by changes in excitability in the trained hand. Indeed, posthoc pairwise
449	comparisons revealed that (averaged across all groups) a significant increase in corticospinal
450	excitability occurred from pre-test to pre-iTBS for the trained hand ($p < .001$, $d = .54$), but
451	not for the untrained hand ($p = .87$, $d = .02$). There were no significant differences between
452	the excitability of trained and untrained hands at pre-test ($p = .25$, $d = .19$); however, the
453	trained hand exhibited greater excitability at pre-iTBS than the untrained hand ($p = .03$, $d =$
454	.35). All other main or interaction effects were not significant (all <i>F</i> < 2.28, <i>p</i> > .12, η_p^2 < .12).

455

456 *iTBS-induced effects on corticospinal excitability*

To assess the impact of the different iTBS protocols on trained and untrained hand 457 458 excitability, we compared the normalized excitability (relative to pre-test) of cM1, iM1 and 459 sham groups before and after the application of iTBS. As shown in figure 4, iTBS reduced normalized excitability of circuits projecting to the trained hand from 1.50 (± 0.26) at pre-460 iTBS to 1.23 (± 0.24) at post-iTBS when delivered to the motor cortex contralateral to the 461 trained hand (cM1 group), but had little effect when delivered to the ipsilateral motor cortex 462 463 (iM1 group; pre-iTBS: 1.51 ± 0.30 ; post-iTBS: 1.41 ± 0.21) or the vertex (sham group; preiTBS: 1.40 ± 0.30 ; post-iTBS: 1.58 ± 0.43). In contrast, iTBS *increased* normalized excitability 464 465 of the untrained hand from 0.98 (± 0.19) at pre-iTBS to 1.38 (± 0.39) at post-iTBS in the iM1

466	group, whereas normalized excitability of the untrained hand was less affected by iTBS
467	delivered to the cM1 (pre-iTBS: 1.04 \pm 0.24; post-iTBS: 1.14 \pm 0.28) or to the vertex (sham
468	group; pre-iTBS: 1.03 \pm 0.18; post-iTBS: 1.13 \pm 0.14). ANOVA conducted on nMEP values
469	revealed a significant time x hand x group interaction, $F(2,33) = 4.31$, $p = .02$, $\eta_p^2 = .21$,
470	indicating that the hand- and group-specific effects described above were statistically
471	significant. Posthoc pairwise comparisons revealed the <i>decrease</i> in excitability in the cM1
472	groups' trained hand ($p = .04$, $d = .57$) and the <i>increase</i> in excitability in the iM1 groups'
473	untrained hand ($p < .001$, $d = .69$) following iTBS to be significant, while no other pairwise
474	comparisons reached significance (all $p > .17$, $d < .26$).
475	Correlation analyses revealed that trained and untrained hand excitability gains were
476	not significantly related to each other following the first training block at pre-iTBS (r = .12, p
477	= .50) or following iTBS at post-iTBS (r = .29, p = .09). For the iM1 group, there was a
478	marginal positive correlation between the extent of iTBS-induced change in performance in
479	the untrained hand, and the extent of iTBS-induced change in excitability (i.e., at post-iTBS)
480	in the untrained motor cortex ($r = .52$, $p = .08$), but not for the trained motor cortex and
481	hand ($r =29$, $p = .36$). Also there were no such associations between performance and
482	excitability changes following iTBS for the cM1 group's trained ($r = .09$, $p = .78$) or untrained
483	hand ($r = .33$, $p = .29$) at post-iTBS.
484	

485

"Figure 4 about here"

486

487 iTBS-induced effects on subsequent motor training

The effect of the second training block was to increase trained hand excitability
(relative to values observed at post-iTBS) in all groups. Specifically, normalized excitability of

490	the trained hand increased to 1.51 (\pm 0.48) in the cM1 group, to 1.10 (\pm 0.19) in the iM1
491	group and to 1.17 (\pm 0.22) in the sham group. Normalized excitability of the untrained hand
492	(relative to post-iTBS) at post-test was 0.97 (\pm 0.18) in the cM1 group, 1.01 (\pm 0.17) in the
493	iM1 group and 1.23 (\pm 0.28) in the sham group. Despite the apparent differences between
494	groups and hands described qualitatively above, ANOVA conducted to assess the effect of
495	the second training bout (i.e., post-test excitability normalized to post-iTBS excitability)
496	revealed no significant differences between groups (p = .50). The main effect of hand,
497	$F(1,33) = 3.45$, $p = .07$, $\eta_p^2 = .10$, and the interaction of hand and group were marginal,
498	$F(2,33) = 3.05$, $p = .06$, $\eta_p^2 = .16$. Post-hoc tests showed that the marginal interaction was
499	driven by the greater excitability gain in cM1 group's trained hand at post-test as compared
500	to their untrained hand ($p = .005$).
501	Correlation analyses revealed a significant relationship between the extent of
502	excitability increases in cM1 group's trained hand induced as a result of the second training

period and the extent of the previous iTBS-induced change (reduction) in excitability (r = -

504 0.74, p = .003). That is, the greater the reduction in excitability induced by iTBS, the greater 505 the subsequent increase in excitability as a result of motor learning.

506

507 Intracortical inhibition

Average SICI ratios at pre-test were 0.56 ± 0.13 , 0.71 ± 0.18 and 0.66 ± 0.17 for the right FDI and were 0.64 ± 0.19 , 0.72 ± 0.17 and 0.72 ± 0.25 for the left FDI for cM1, iM1 and sham group participants respectively. There were no differences in SICI at pre-test between the groups (p > .45). ANOVA conducted on non-normalized SICI ratios of trained and untrained hands (before and after the first training block) revealed a marginal main effect of time, F(1,33) = 3.17, p = .08, $\eta_p^2 = .09$, indicating a small (and non-significant) increase in the

level of inhibition (averaged over all groups and both hands) as a result of the first training block (pre-test: 0.67 ± 0.11; pre-iTBS: 0.61 ± 0.09). All other main effects and interactions were not statistically significant (all p > .17).

517

518 Multiple Regression analysis to elucidate predictors of cross-limb transfer

519 Averaged across the three groups (n=36), performance of the untrained, left hand 520 increased by 56.6% (± 13.3%) as a result of unilateral, right hand motor training from pretest to pre-iTBS; this is equivalent to $61.2 \pm 28.6\%$ of the gains observed in the *trained* hand 521 522 (i.e., untrained hand normalized performance gains relative to trained hand gains following 523 the first block of motor training). To identify the main predictors of (normalized) untrained 524 hand performance gains at pre-iTBS and to assess their relative predictive strength, we employed a multiple regression analysis. Initially, we entered predictor variables that were 525 526 directly related to the excitability change and dynamics of the muscle bursts in *untrained* 527 hand (i.e., nMEP_{untrained}, nEMG_{untrained} and EMG_{mirror}, respectively). A second model also 528 included predictor variables that were related to the *trained* hand performance, muscle 529 activity and excitability changes (i.e., nACC_{training}, nMEP_{trained}, nEMG_{trained},) to additionally 530 account for the impact of adaptations in the trained hand on adaptations in the untrained limb. 531

532 Untrained hand performance gains at pre-iTBS (i.e., cross-limb transfer) were 533 significantly predicted by model 2 ($\Delta R^2 = 0.49$, $\Delta F(3,29) = 10.06$, $\Delta p < 0.001$), but not by 534 model 1 (adjusted $R^2 = -0.06$, F(3,32) = .38, p = 0.77). Normalized performance gains of the 535 trained hand during training (nACC_{training}, $\beta = 0.66$, t(35) = 4.71, p < .001) and training-536 induced excitability changes of the trained hand (nMEP_{trained}, $\beta = 0.37$, t(35) = 2.86, p = .008) 537 explained 43.6 % and 13.7 % of the variance in untrained hand performance gains following

538	unilateral practice, respectively, when controlled for the other variables in the equation. The
539	analysis also revealed a marginal (unique) contribution of the training-induced excitability
540	changes in the untrained hand (nMEP _{untrained} , β = 0.25, $t(35)$ = 1.89, p = .07) explaining at
541	least 6.4 % of the variance in normalized untrained hand performance following motor
542	practice at pre-iTBS. Partial regression plots for the variables that have been shown to
543	explain significant (marginal) portions of variance in untrained hand performance gains at
544	pre-iTBS are displayed in Figure 5.

- 545
- 546

"Figure 5 about here"

547

548 Control Experiment

The results of the main experiment showed that both performance and MEPs 549 550 increased in the untrained hand after iTBS to the untrained M1, but it is unclear whether this 551 performance gain reflected a general improvement in motor function due to enhanced 552 excitability, or an interaction with the recently transferred motor skill. We therefore 553 analyzed the iTBS-induced change in excitability and motor performance (normalized values 554 relative to pre-test) at post-iTBS (i.e. following iTBS over right M1) in a control group that performed no prior motor training. Normalized excitability of circuits projecting to the right 555 and left hands at post-iTBS was 0.95 (\pm 0.12) and 1.13 (\pm 0.18) respectively (see Figure 6A). 556 557 Normalized motor performance following iTBS was $0.93 (\pm 0.11)$ for the right hand and was 558 $0.86 (\pm 0.07)$ for the left hand (see Figure 6B). One-sample t-Tests (against pre-test level, i.e. 1) revealed the decrease in left hand performance to be significant, t(11) = -3.71, p = .003, 559 but not the increase in left hand (right M1) excitability, t(11) = 1.407, p = .18. Thus, the 560 561 expected LTP-like effect of iTBS was not statistically significant for the entire group due to

562	inter-subject variability (as has been reported previously, Hamada et al., 2013; Hinder et al.,
563	2014). We therefore looked at the subset of six control participants who showed the largest
564	MEP changes for the left hand, to be sure that an iTBS-induced increase in MEP amplitude
565	does not change motor performance. Average normalized excitability of the sub-sample was
566	0.98 (\pm 0.13) and 1.37 (\pm 0.23) for circuits projecting to right and left hands respectively (see
567	Figure 6C). Average normalized performance following iTBS for the subset of best
568	'responders' was 0.98 (\pm 0.16) for the right hand and was 0.88 (\pm 0.08) for the left hand (see
569	Figure 6D). One-sample t-Tests revealed both the increase in left hand (right M1) excitability,
570	t(5) = 2.84, $p = .04$ and the decrease in left hand performance to be significant, $t(5) = -2.85$,
571	p = .04. Moreover, changes in MEP amplitude and motor performance following iTBS were
572	not associated, neither across the entire group of 12 subjects (left hand: r = .05, p = .89; right
573	hand: r =38, p = .22), nor in the subset of participants that exhibited the largest MEP
574	changes in the left hand following iTBS (left hand: r = .03, p = .96; right hand: r =11, p =
575	.84). Taken together, the data imply that there was no tendency towards increased motor
576	performance simply as a result of increased excitability produced by iTBS in the absence of
577	training.
578	

579

Discussion

The present study used non-invasive brain stimulation to probe the neural mechanisms underpinning motor learning and cross-limb transfer. The major novel finding was that when applied following an initial period of motor learning, brain stimulation that induces LTP-like plasticity in the resting-state motor cortex (iTBS) had unilateral effects on motor performance and corticospinal excitability, the nature of which differed depending on which cortex was stimulated. Specifically, iTBS applied to the trained cortex (cM1 group)

586	resulted in statistically significant <i>reductions</i> of both prior training-induced performance
587	gains (Figure 3A) and corticospinal excitability increases (Figure 4A) in the trained hand and
588	motor cortex, without affecting performance in, or corticospinal projections to, the
589	untrained hand. The reversal of the typical facilitatory effect of iTBS on corticospinal
590	excitability (Huang et al. 2005) is consistent with homeostatic plasticity (see Müller-Dahlhaus
591	and Ziemann 2015 for a review), whereas the reduction in training-induced performance
592	gains suggests that non-invasive brain stimulation interfered with circuits involved in storage
593	or retrieval of the new motor memory (Muellbacher et al. 2002). In contrast, iTBS applied to
594	the untrained hemisphere (iM1 group), resulted in <i>improved</i> motor performance (Figure 3B)
595	and increased corticospinal excitability (Figure 4B) in the untrained hand and motor cortex
596	without affecting the performance or projections to the trained hand (see Figure 1B for that
597	prediction). Moreover, these changes in performance and excitability seem functionally
598	related; the extent of performance transfer to the untrained hand predicted the magnitude
599	of excitability increases. The <i>distinct</i> effects of iTBS on performance in the trained
600	(performance decrements) and untrained (performance gains) cortices is highly suggestive
601	that different mechanisms mediate motor learning and cross-limb transfer. Importantly, the
602	observed differences in the manner in which iTBS affected performance in the trained and
603	untrained hands appeared despite the fact that both hands had exhibited increases in
604	performance following the initial unilateral motor learning.
605	

606 Homeostatic versus non-homeostatic processes in the trained and untrained M1s

The interaction between the mechanisms underpinning motor learning in the trained
hand and iTBS is consistent with the notion of homeostatic metaplasticity. In this instance,
rather than LTP-like plasticity from motor learning and iTBS accumulating, the prior motor

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610	learning reversed, or occluded, the 'expected' effects of a subsequent LTP-inducing protocol
611	(in this case, iTBS; see Di Lazzarro et al. 2008; Huang et al. 2005) applied to the trained
612	hemisphere (see Müller-Dahlhaus and Ziemann 2015 for a review; Rosenkranz et al. 2007;
613	Stefan et al. 2006; Stöckel et al. 2015; Ziemann et al. 2004).
614	In contrast, the increases in corticospinal excitability observed in the untrained
615	hemisphere following iTBS applied to the untrained M1 (iM1 group), reflect an apparent LTP-
616	like effect. This is consistent with iTBS effects observed when applied in isolation (Huang et
617	al. 2005). Conceivably, because the behavioral gains in the untrained hand (following motor
618	training) were not accompanied by increases in excitability of the untrained hemisphere
619	(Figure 4), the iTBS protocol was still able to act in the 'expected' direction and induce
620	facilitation of MEPs.
621	
622	Effects of iTBS on subsequent performance and learning
623	Because NBS is a potential candidate to augment neuro-rehabilitation (Müller-
624	Dahlhaus and Ziemann 2015; Ridding and Rothwell 2007), it is important to consider its
625	effects on subsequent motor performance and learning. Previous work shows that learning
626	can be enhanced in the trained limb when iTBS is applied to the contralateral M1 (cM1;
627	trained hemisphere) prior to practice (Teo et al. 2010; c.f. Agostino et al. 2008 for a
628	contradictory report, note that their experiments involved either a short training protocol
629	that caused limited learning, or a small sample of n = 5). In the current study, performance

was reduced when assessed without feedback immediately after iTBS to the contralateral

M1, but rapidly increased during the second learning bout such that final performance was

influence of iTBS on subsequent learning was complicated in the current study by the fact

no different from a group that received sham stimulation. However, disentangling the

634	that motor practice was also performed prior to iTBS delivery. The rapid recovery of
635	performance during the first few trials of T2 could be viewed as an increase in learning rate
636	following iTBS (as per Teo et al. 2010), or the dissipation of a homeostatic interaction
637	between iTBS and prior training (Stöckel et al. 2015).
638	More importantly, we were interested in the influence of iTBS to the ipsilateral M1
639	(iM1; untrained hemisphere) on subsequent performance and <i>transfer</i> from the trained limb,
640	as to our knowledge, this effect has not been previously investigated. The increase in
641	ipsilateral excitability that we observed after iTBS to the ipsilateral M1 appeared to drive
642	further performance gains in the untrained hand. Because neither excitability nor
643	performance were significantly increased following iTBS in the non-training control group,
644	we propose that the effects of prior training with the opposite limb interact with iTBS
645	delivered to the untrained M1. In particular, it appears in this case that the NBS-induced
646	facilitatory effect summated with the transfer-induced performance gains. Similar to the
647	results for the trained hand, however, final performance measured in the untrained hand
648	after T2 was not different between groups. This indicates that the immediate performance
649	benefit conferred by iTBS to the ipsilateral M1 failed to improve subsequent performance
650	gains due to transfer from the opposite limb.
651	

652 What type of ipsilateral adaptations mediate untrained hand performance?

Unlike previous research demonstrating bilateral increases in corticospinal
excitability following unilateral, ballistic motor practice (Carroll et al. 2008; Lee et al. 2010;
Hinder et al. 2011), substantial performance improvements in the untrained hand were not
accompanied by increased excitability of corticospinal projections to the untrained hand in
the current study. Transfer of performance without changes in excitability of the untrained

658	cortex is consistent with evidence from sequencing tasks (Camus et al. 2009; Pascual-Leone
659	et al. 1995; Perez et al. 2007). Moreover, previous work has also shown that transfer of
660	ballistic motor skill can even be accompanied by decreases in excitability of the untrained
661	hemisphere (Duque et al. 2008). A likely mechanism that contributes to enhanced
662	performance in the untrained limb is reduced inter-hemispheric inhibition, which is reduced
663	from the trained to the untrained M1 after various types of sequence learning (Camus et al.
664	2009; Perez et al. 2007), and after strength training (Hortobagyi et al. 2011; see Ruddy and
665	Carson 2013 for a review). Thus, while ballistic motor training reliably potentiates
666	corticospinal excitability in the trained M1 (cf. Liepert et al. 1998; Muellbacher et al. 2001),
667	untrained, left hand performance gains following unilateral practice (i.e., as a result of cross-
668	limb transfer) are not necessarily accompanied by overt changes in excitability in the
669	untrained, right M1. However, the fact that iTBS applied to the untrained hemisphere
670	amplified untrained hand performance gains in the current study suggests that some form of
671	adaptation occurred within the untrained M1 which mediated performance improvements
672	in the untrained hand. In support of this view there is evidence from neuroimaging data on
673	the encoding of (sequential) single finger movements (Diedrichsen et al. 2013; Wiestler et al.
674	2014) demonstrating similar (mirrored) representation patterns in both motor cortices (and
675	sensory motor cortices) that include the same fine-grained details of the movement, but
676	with suppressed BOLD signals (relative to resting baseline) in the motor cortex ipsilateral to
677	the active hand. This raises the possibility that multiple processes may influence ipsilateral
678	cortical function, including generalised suppression of activation (which might underlie the
679	lack of corticospinal excitabitly we observed) and patterned activation specifically associated
680	with task performance (which might underlie transfer of performance to the untrained limb).

681	The finding that untrained hand performance gains following motor training are only
682	affected (i.e. up-regulated) by iTBS applied to the untrained, but not the trained, M1 is
683	strongly suggestive of a contribution of the ipsilateral M1 to cross-limb transfer. In line with
684	the cross-activation hypothesis (Cramer et al. 1999; Dettmers et al. 1995; Parlow and
685	Kinsbourne 1989) and previous experimental work using the same motor learning task (Lee
686	et al. 2010), our results suggest that task-related information stored in the untrained
687	hemisphere during motor learning is subsequently retrieved to drive cross-limb adaptations
688	when the task is undertaken with the untrained limb. The fact that training related
689	improvements in the untrained hand are positively correlated with subsequent iTBS-induced
690	improvements in this same hand is further evidence to suggest that cross-limb
691	improvements (if governed by LTP-like processes) and iTBS interact in a non-homeostatic
692	manner. Alternatively, because the untrained limb improvements were not associated with
693	excitability increases, it may be that transfer itself is not driven by LTP-like effects, and hence
694	the subsequent iTBS LTP-inducing protocol was able to act without being affected by a prior
695	history of LTP. Although our evidence strongly suggests that the ipsilateral, untrained M1 is
696	involved in transfer, the lack of change in MEP, compared with MEP increases in the
697	contralateral, trained M1, implies either that 1) the majority of neurons that contribute to
698	the peripheral responses to TMS (i.e. MEPs) are not involved in transfer, or 2) other non-
699	primary areas also contribute substantially to transfer. Finally, it should be noted that the
700	current cohort consisted of right handed adults who trained with their dominant hand. Thus,
701	although we feel it unlikely, the possibility exists, that influences of hand dominance and
702	hemisphere-specific effects of iTBS in the right and left M1 (irrespective of which
703	hand/hemisphere had trained) may have had a small effect on the results.

704	With respect to potential non-primary contributions to transfer, it has indeed been
705	demonstrated that a broad neural network is involved in cross-limb transfer (Gerloff and
706	Andres 2002; Rizzolatti et al. 1998; Ruddy and Carson 2013), with evidence for bilateral
707	changes in different secondary motor areas (Hardwick et al. 2013; Wiestler and Diedrichsen
708	2013). As such, neural adaptations in other brain regions beyond M1 ipsilateral to the
709	trained limb could account for cross-limb transfer effects observed in the current study. In
710	this case, the stimulation protocol used in the present study would only have affected a part
711	of the acquired skill representation for the untrained limb, and thereby limiting the
712	conclusions drawn here to features of the task represented within M1. For example, Romei
713	and colleagues (Romei et al. 2009) provided evidence that M1 contributes to intrinsic (i.e.,
714	knowledge represented in body-centred coordinates; muscle and joint based) but not
715	extrinsic components (i.e., world-centred coordinates; movement features in external space)
716	of motor skill learning. Therefore, future studies should examine the relative contribution of
717	a more extensive brain network in the untrained cortex to cross-limb adaptations following
718	unilateral practice of different motor tasks (e.g., ballistic, sequential, or reaction time tasks).
719	

720 Conclusions

In sum, the present study suggests that, while occurring simultaneously, motor learning and cross-limb transfer represent distinct neural adaptation processes which interact differently with iTBS. The typical effect of an LTP-like inducing brain stimulation protocol were reversed in the hemisphere projecting to the trained hand, consistent with the suggestion that LTP contributes to ballistic motor learning. That is, motor learning resulted in subsequent iTBS having a LTD-like effect in the trained M1. In contrast, LTP-like effects following iTBS were observed in the hemisphere projecting to the transfer hand, suggesting

- either that LTP within the untrained M1 does not underlie cross limb transfer, or that the
- 729 majority of neurons that contribute to the peripheral responses to TMS (applied to the
- vultrained M1) are not involved in transfer. Importantly, iTBS had a unilateral effect on both
- the training and transfer process, offering further support that transfer is governed by the
- ross-activation hypothesis (Cramer et al. 1999; Dettmers et al. 1995; Lee et al. 2010).

734 Acknowledgements

- 735 The research was supported by a fellowship within the Postdoc-Program of the German
- 736 Academic Exchange Service (DAAD) to TS and an Australian Research Council DECRA
- 737 fellowship awarded to MRH. We also would like to acknowledge NHMRC funding
- 738 (APP1050261) awarded to JJS and MRH. We thank Paola Reissig for assisting in data
- 739 collection.
- 740 The authors declare no competing financial interests.

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866	Figure captions
867	
868	Figure 1. Predictions of the study. Panel A) predicts that, if learning and transfer are both
869	mediated by LTP-like processes, then the effect of a subsequent intervention that induces
870	LTP-like effects in the absence of training (here iTBS) should be reduced or reversed due to
871	homeostatic plasticity in both the trained or untrained hemispheres. Panel B) predicts that, if
872	learning relies on LTP-like processes, then it should cause a reduction or reversal in the
873	expected LTP-like effects induced by iTBS to the contralateral M1 (cM1; trained hemisphere).
874	However, if the processes that mediate transfer are not LTP-like, then the typical LTP-like
875	effect of iTBS to the ipsilateral M1 (iM1; untrained hemisphere) should be observed. (\downarrow
876	indicates reduction of iTBS-induced LTP-like effect due to homeostatic interaction of
877	learning/transfer with iTBS; \leftrightarrow indicates maintenance of the expected iTBS-induced LTP-like
878	effects).
879	
880	Figure 2. Experimental procedure. Transcranial magnetic stimulation (TMS) and motor
881	performance (MP) measures were assessed before a first training block, before and after
882	applying intermittent theta-burst stimulation (iTBS) and after a second training block for right
883	and left hands and motor cortices. Depending on group affiliation, iTBS was induced over the
884	trained or untrained motor cortex or over the vertex.
885	
886	Figure 3. Normalized performance relative to pre-test. Following a first training block of
887	ballistic right hand practice (150 trials), iTBS was applied to the contralateral, trained (cM1)
888	[panel A], the ipsilateral, untrained motor cortex (iM1) [panel B], or over the vertex (sham
889	group) [panel C]. Performance was tested in participants trained (right) hand (circles) and
890	untrained (left) hand (triangles) before the first training block, before and after iTBS and after

- a second training block. Performance values were normalized to the corresponding pre-test
 performance for that hand (as indicated by the X-axis). Error bars indicate 95% Cl.
 Figure 4. Average normalized MEPs relative to pre-test for cM1, iM1 and sham group
- participants. MEP amplitudes were tested on the trained (right) hand (circles) and on the
 untrained (left) hand (triangles) before a first training block (T1), before and after iTBS and
 after a second training block (T2). MEP amplitudes were normalized to pre-test values. Error
 bars indicate 95% CI and the X-axis represents pre-test values.
- 899

900 Figure 5. Partial regressions. Multiple regression analysis was performed to identify main 901 predictors of cross-limb transfer (i.e., normalized untrained hand performance) following 902 unilateral practice (i.e., at pre-iTBS) and to study their relative strength. Displayed are partial 903 regressions of the (A) normalized performance of the trained hand during the first training 904 block, (B) normalized MEPs of the trained hand following the first training block, and (C) 905 normalized MEPs of the untrained hand at pre-iTBS; all of which contributed unique variance 906 to the regression model, i.e. were identified as potential predictors of cross-limb transfer. 907 Relations between the measures are displayed by linear trend lines. X- and Y-axes represent 908 respective pre-test values.

909

Figure 6. Control experiment data summary. (A) MEP and (B) performance changes in right
and left hands following right M1 iTBS averaged across all participants of the no-training
control group (n = 12). (C) MEP and (D) performance changes in the right and left hands
averaged across the 6 best responders to iTBS (out of the 12 no-training control group
participants). MEP amplitudes and performance values were normalized to pre-test values
(before iTBS). Error bars indicate 95% CI.



B) Learning and Transfer represent distinct neural

A) Learning and Transfer represent similar neural mechanisms









