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# Suppressing movements with phantom limbs and existing limbs evokes comparable electrophysiological inhibitory responses

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# Abstract

Phantom limb is a common sensation in amputees, who often report vivid experiences of voluntarily moving their phantom. Previous studies showed that phantom movement can be functionally disentangled from imagined movement comparable to the actual movement of an intact limb. How and to what extent phantom movement and real movement share similar physiological mechanisms? Here, we focused on a specific aspect of motor control, the motor inhibition, and we asked whether inhibitory physiological responses are implemented when a phantom movement has to be suppressed. Sixteen twohanded controls and two left upper-limb amputees (with and without phantom movement) underwent a Go/Nogo paradigm, while event-related potentials (ERPs) were recorded. The task was performed with both the right (intact) and the left (phantom) hand, either in real or imagery conditions. Opposite results between the moving-phantom case and the static-phantom case were found. In the real condition, moving-phantom case showed the classical motor-inhibition related ERP pattern, with large P300 inhibitory wave when the movements of both (right) intact and (left) phantom limbs have to be suppressed. This inhibitory response was not different from that found in controls (who performed the task with an existing hand; real condition), but, crucially, it was significantly different from the imagery condition of controls. Contrariwise, in the static-phantom case, the ERP responses to Nogo trials during real condition were different from the real condition in controls but were not different from their imagery conditions. Importantly, in real condition, Nogo-ERP responses were significantly different between the two phantom cases. Taken together, these findings provide compelling evidence that phantom movements share the same neurophysiological correlates of real movements, not only when an action has to be executed, but also when it should be inhibited.

# Keywords: phantom movement; motor inhibition; ERPs; phantom limb; motor imagery

# **1. Introduction**

People with a body amputation commonly continue to perceive the presence of their missing limb, as if 1 it was still present (Herta Flor, Nikolajsen, & Staehelin Jensen, 2006). Amputees report that their 2 phantom limb has certain sensory properties like touch (Melzack, 2006) and up to 80% of these phantom 3 sensations are experienced as painful (Kikkert et al., 2017; Weeks, Anderson-Barnes, & Tsao, 2010). 4 5 Interestingly, some patients with phantom limbs claim they can voluntary move their phantom 6 (Garbarini, Bisio, Biggio, Pia, & Bove, 2018; Raffin, Giraux, & Reilly, 2012; Raffin, Mattout, Reilly, & 7 Giraux, 2012; Ramachandran & Hirstein, 1998). They experience sensations of reaching out to grab an 8 object, making a fist, or moving their fingers individually (Ramachandran & Hirstein, 1998). It has been described that phantom movement involves some aspects of "real" motor execution. For example, 9 voluntary movements of the phantom hand trigger characteristic patterns of distal movements in the 10 residual limb muscle activity, which differed from the activity recorded in the same muscle groups during 11 12 proximal movements, suggesting that the brain's motor areas can be differentially activated according to the movement the patients intend to perform with their phantom (Reilly, Mercier, Schieber, & Sirigu, 13 2006). Furthermore, an ischemic nerve block applied to the residual limb abolished not only 14 electromyographic (EMG) signal of the residual limb, but also the ability to voluntarily move the 15 phantom (Reilly et al., 2006). 16

Many neuroimaging studies, examining the cerebral activity associated with phantom limb movements (Hugdahl et al., 2001; Lotze, Flor, Grodd, Larbig, & Birbaumer, 2001; Roux et al., 2003), showed a maintained representation of the missing hand in the primary sensorimotor missing hand cortex (Kikkert, Johansen-Berg, Tracey, & Makin, 2018; Makin et al., 2013) and an increased activity in motor areas, such as the contralateral primary motor cortex (M1) and the supplementary motor area (SMA), in a similar way as real/actual movements do (Raffin, Giraux, et al., 2012; Reilly et al., 2006). Even using a different neuroimaging technique, such as the electroencephalography (EEG) time-frequency analysis, it has been shown that, in a case with congenital absence of arm, similar sensorimotor rhythms were present
in both real movements and phantom movements (Walsh, Long, & Haggard, 2015). These neuroimaging
findings suggest that, when amputees "perform" voluntary movements with their phantom limb, they
recruit a similar neurophysiological brain activity as that employed during the actual movements of an
existing limb.

Importantly, amputees, when asked to perform real or imagined movements with their phantom, report 29 30 different perceptions and sensations during each task. Some studies tried to disentangle motor imagery from motor execution both with behavioral (Garbarini et al., 2018; Raffin, Giraux, et al., 2012) and 31 neuroimaging measures (Hugdahl et al., 2001; Lotze et al., 2001; Raffin, Mattout, et al., 2012; Roux et 32 33 al., 2003). From a behavioral point of view, it has been observed that motor execution with the phantom is slower than with the intact limb, whereas the time to imagine a movement is similar for both the 34 phantom and intact limbs, suggesting that amputees limb loss differentially affects these two tasks 35 (Raffin, Giraux, et al., 2012). In a recent study (Garbarini et al., 2018), the intermanual transfer of 36 sequence learning, occurring when one hand motor skills improve after training with the other hand, was 37 used to dissociate motor execution from motor imagery with a phantom limb. It has been reported that 38 only after an active training with the phantom limb, and not with an imagery training, the moving-39 phantom case showed a faster performance of the intact hand, as in two-handed controls actually 40 41 performing the training with an existing hand (Garbarini et al., 2018). Neuroimaging studies on real and imagery phantom movements showed that two distinct brain networks are activated during motor 42 execution (i.e. hand area of the contralateral M1, hand area of the contralateral primary somatosensory 43 44 cortex – S1, contralateral dorsal premotor areas, bilateral medial premotor areas, ipsilateral anteromedial part of the cerebellum) and motor imagery (i.e. contralateral dorsal premotor areas, bilateral medial 45 premotor areas, bilateral inferior areas) (Raffin, Mattout, et al., 2012). Interestingly, these two networks 46 partially overlap (i.e. parts of SMA, the dorsal premotor cortex, the posterior parietal cortex and the 47

cerebellum as well as M1), and they are similar to those activated during imagination and execution with 48 the intact limb (Hugdahl et al., 2001; Lotze et al., 2001; Raffin, Mattout, et al., 2012; Roux et al., 2003). 49 However, when considering effective connectivity between brain areas, it has been shown that, as in 50 healthy controls (Bruno, Fossataro, & Garbarini, 2018; Kasess et al., 2008; Solodkin, Hlustik, Chen, & 51 Small, 2004; Tak, Kempny, Friston, Leff, & Penny, 2015), real and imagery tasks with the phantom limb 52 had an opposite effect on the SMA-M1 network: while during motor execution, SMA exerts an 53 54 excitatory effect on M1, since the movement has to be performed; during motor imagery, SMA exerts an inhibitory effect on M1, since the movement has to be suppressed (Raffin, Giraux, et al., 2012). 55

Taken together, these previous findings suggest that, when a movement has to be executed, real and 56 57 phantom limbs share common behavioral and physiological mechanisms. But, what happens when a phantom movement has to be suppressed? Are inhibitory physiological responses fully implemented, as 58 during the suppression of a real movement? The ability to suppress, withhold, delay or interrupt ongoing 59 or planned actions is a fundamental aspect of motor control in everyday life. Thus, if controlling phantom 60 and real movements rely on common mechanisms, we should expect that suppressing movements with 61 existing and phantom limbs evokes comparable inhibitory responses. The most common task employed 62 in the study of motor inhibition is the Go/Nogo task. During this task, participants are required to respond 63 to frequent imperative stimuli and they must withhold the response to other infrequent alternatives 64 (Donders, 1969). Several electrophysiological studies on the Go/Nogo task described an enhanced 65 frontocentral negativity occurring around 140-300 ms, as well as an enhanced central positivity 66 occurring around 300-600 ms, following the presentation of a Nogo stimulus (Falkenstein, Hoormann, 67 68 Christ, & Hohnsbein, 2000; Falkenstein, Koshlykova, Kiroj, Hoormann, & Hohnsbein, 1995; Pfefferbaum, Ford, Weller, & Kopell, 1985; Veen & Carter, 2002). These peaks are referred to as N200 69 and P300 respectively, and they have been interpreted as indexes of response inhibitory process in the 70

frontal lobe (Bokura, Yamaguchi, & Kobayashi, 2001; Cojan, Archimi, Cheseaux, Waber, &
Vuilleumier, 2013; Kok, 1986; Smith, Johnstone, & Barry, 2008).

The purpose of the present study was to investigate whether and to what extent phantom limbs share 73 similar physiological mechanisms with actual limbs, not only when a movement has to be performed, 74 but also when it has to be inhibited. To this aim, we recorded event-related potentials (ERPs) during a 75 76 Go/Nogo task in two upper-limb amputees who have already participated in a previous study (Garbarini 77 et al., 2018). Both the amputees reported vivid sensations on their phantom, but in different ways: one reported the ability to move her phantom (moving-phantom case), the other reported vivid phantom limb 78 79 sensation without phantom movement, which, in turn, he perceived as paralyzed (static-phantom case). 80 See details in section 2.1. The ERPs collected in amputees were compared to those collected in twohanded controls. All participants underwent two different conditions of the task: the real condition, 81 82 during which a real key press in response to Go trials was requested (either with the intact or with the 83 phantom hand) and an imagery condition, during which a mental simulation of the key press in response to Go trials was requested (either with the intact or with the phantom hand). If a/the? phantom movement 84 has similar physiological properties as the real movement, we should expect that, during the Go/Nogo 85 task, the ERP pattern of the moving-phantom case will be comparable to that found in two-handed 86 controls, with the typical inhibitory ERP response associated to Nogo trials. In two-handed controls, the 87 88 inhibitory ERP response was expected to be greater in real condition than in imagery condition and, coherently, a similar pattern was expected in the moving-phantom case. On the contrary, the static-89 phantom case, who did not experience kinematical sensations with his phantom, was expected to use 90 91 motor imagery to perform the real Go/Nogo task with his phantom limb, thus showing a similar ERP pattern as controls during the imagery task. 92

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# 94 2. Materials and methods

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#### 95 2.1 Participants

Two left upper-limb amputees took part in this study. Both of them reported phantom limb sensations, 96 but in different ways. The first phantom case was a 31 years old female, with 18 years of education. Her 97 left upper limb was amputated under the shoulder as a consequence of arm paralysis due to brachial 98 99 plexus injury after a car accident 11 years ago. Her phantom limb sensation started immediately after the accident and continues to persist, although it may vary daily depending on various factors (e.g., stress). 100 101 At the time of testing, on a visual analog scale (VAS) aiming at assessing pain intensity on her phantom limb ranging from 0 (absolutely not painful) to 10 (absolutely painful), she reported a score of 0, 102 103 indicating that she did not perceive painful sensation on her phantom (even if she reported phantom pain 104 in the past, for about two years after the accident). She reported to be able to move her phantom at will, and her perceived movement control is limited to relatively simple movements (e.g., grasping, pressing, 105 pointing). We named her "moving-phantom case". The second phantom case was a 53 years old male, 106 107 with 13 years of education. He got the amputation of the left upper limb after an accident at work when he was 35. The amputation was made above the left elbow and he was not using any prosthesis. At the 108 109 time of testing, he did not report painful sensations on his phantom (even if he reported phantom pain in the past), but sometimes he felt it as ticklish. He reported a score of 0 on the VAS about pain intensity 110 on his phantom limb. He did not report the ability to move his phantom, which, in turn, described as rigid 111 112 and contracted. We named him "static-phantom case". Sixteen two-handed volunteers (6 men; mean age  $\pm$  sd: 23.9  $\pm$  1.3 years; mean of educational years  $\pm$  sd: 17.6  $\pm$  0.8) were recruited as controls. All 113 participants (i.e. the two phantom cases and controls) were right-handed, according to the Standard 114 115 Handedness Inventory (Oldfield, 1971). Neither amputees nor controls had suffered a brain lesion or had a history of neurological or psychiatric illness. All participants were naïve to the purpose of the study 116 and gave written informed consent according to the declaration of Helsinki. The Ethical Committee of 117 the University of Turin gave approval to the project (prot. n. 125055, 12/07/16). 118

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# 120 2.2 Experimental design

Each participant underwent an EEG recording session while she/he performed a Go/Nogo task (see details in the next paragraph). During the same session, participants were asked to perform the task in both *real* and *imagery* condition. The only difference between the two conditions was that during the *imagery* condition, the subjects had to imagine the hands' movements (i.e. key press, see details in the next paragraph) with a kinesthetic motor imagery, (Bisio et al., 2017; Bruno et al., 2018; Jeannerod, 1995; Piedimonte, Garbarini, Rabuffetti, Pia, & Berti, 2014) instead of moving the hand as in *real* conditions.

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# 129 2.3 Go/Nogo task and procedure

A similar version of the Go/Nogo task used by Cojan and colleagues, aiming at comparing motor 130 131 inhibition mechanisms responsible for paralysis during hypnosis and those recruited by voluntary inhibition (Cojan et al., 2013) was used. Participants were seated on a chair in front of a 21-inches Sony 132 CRT screen placed at a distance of 55 cm, in a dimly illuminated room. Visual stimuli were presented 133 on the computer screen. Each stimulus represented the dorsal view of a hand (left or right) and it could 134 be colored in grey, green, or red. Each trial started with a fixation cross (jittered interval 6000-8000 ms), 135 136 followed by a preparation cue (i.e. Preparation) which represented a grey hand, either left or right (jittered interval 1000-1200 ms). The grey hand instructed the participant to prepare to press a key on the keyboard 137 with the corresponding hand. Then, the grey hand could turn either green (i.e. Go stimulus) or red (i.e. 138 139 Nogo stimulus) (fixed duration of 750 ms). During the real condition, participants had to press the key as quickly as possible when the hand turned green (75%), and to withhold the prepared response if the 140 hand turned red (25%). During the imagery condition, participants had to imagine to press the key when 141 the hand turned green and to withhold the prepared (imagined) response if the hand turned red. After 142

each imperative stimulus (Go or Nogo), the fixation cross re-appeared (Figure 1A). It is important to note 143 that the task, both in real and imagery condition, required that both phantom-limb cases performed it 144 with their left/phantom and their right/intact hand. While the moving-phantom case was able to perform 145 the task with her phantom, the static-phantom case was asked to "try to perform" the task with his 146 147 phantom. The presentation of stimuli was pseudorandomized, in a way that more than two sequential Nogo stimuli never appeared. Stimuli display and reaction times (RTs) collection were controlled by E-148 prime v.2 (Psychology Software Tools, http://www.pstnet.com). Before starting the experiment, all the 149 participants (i.e. the two phantom-cases and controls) performed a short 10 trials practice to familiarize 150 with the task. Six blocks of 40 trials (half right hand, half left hand) were performed per condition (i.e. 151 152 real; imagery), resulting in a total of 480 stimuli: 90 real Go left, 90 real Go right, 30 real Nogo left, 30 real Nogo right, 90 imagery Go left, 90 imagery Go right, 30 imagery Nogo left, 30 imagery Nogo right. 153 All the participants (i.e. the two cases and the two-handed controls) performed alternately a block of real 154 155 and a block of imagery condition. Half of the controls started with a real block, the other half with an imagery one. The two phantom-cases started both with a real block (Figure 1B). Each block lasted about 156 6 minutes. A 2 minutes break was performed between each block. 157

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- 159

--- Figure 1 about here ---

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# 161 *2.4 Electroencephalogram recording and processing*

Participants were seated in a comfortable chair in a silent, temperature-controlled room. They were asked to focus on the task, keep their eyes open and try to avoid blinking when stimuli appeared. Continuous EEG activity was acquired from 32 channels (HandyEEG, SystemPlus Evolution, Micromed, Treviso, Italy) by using tin electrodes mounted in an elastic cap according to the International 10-20 system and referenced to the nose. Eye movements (electrooculogram, EOG) were recorded from two surface 167 electrodes, one placed over the right lower eyelid and the other placed lateral to the outer canthus of the 168 right eye. Electrode impedances were kept below 5 k $\Omega$ . Signal was digitized at a sampling rate of 1,024 169 Hz (SD32; Micromed, Treviso, Italy). Data were continuously streamed to a laptop connected to a second 170 computer generating the stimuli. These two computers interfaced via a serial port for precise 171 synchronization.

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173 **3. Data analysis** 

#### 174 Electrophysiological data

ERPs were pre-processed and analyzed offline using Letswave v.6 (https://www.letswave.org/, Mouraux 175 176 & Iannetti, 2008). The analyses were performed on both Go and Nogo trials (stimulus-locked). Epochs were selected from 500 ms prior to onset to 1000 ms after the onset of the imperative stimulus (total 177 epoch duration: 1500 ms). All the epochs were band-pass filtered (1-30 Hz) using a fast Fourier 178 transform filter. Each epoch was baseline corrected using the 500 ms pre-stimulus recording period as 179 reference. Artifacts due to eye blinks or eye movements were subtracted using a validated method based 180 on an Independent Component Analysis (Jung et al., 2000). Only correct responses to Go and Nogo 181 imperative stimuli were analyzed. Blinks were found to be the most frequent cause of rejection. Epochs 182 belonging to the same experimental condition were averaged time-locked to the onset of the stimulus. 183 184 Thus, separately for each hand (left/phantom hand, right/intact hand), for each condition (Go, Nogo) and for each task (real, imagery), 8 average waveforms (Go real left/phantom, Go real left/phantom, Go 185 imagery left/phantom, Go imagery left/phantom, Nogo real left/phantom, Nogo real left/phantom, Nogo 186 187 imagery left/phantom, Nogo imagery left/phantom) were obtained for each subject.

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189 *3.1. Group analysis* 

190 We firstly performed a group analysis on the two-handed controls to investigate: i) if our task was able to reproduce the classical electrophysiological pattern associated with motor inhibition (Bokura et al., 191 2001); ii) if there were any differences in ERPs between real and imagery conditions. Mean ERP 192 responses to Go and Nogo stimuli were analyzed by a point-by-point repeated-measure ANOVA with 193 194 three within-subject factors: Condition (two levels: Go; Nogo), Side (two levels: left; right) and Task (two levels: real; imagery). Cluster-based permutation testing approach (1000 random permutations 195 196 testing across all 32 channels) was employed to correct for multiple comparisons across different time points (cluster threshold was set at the 95th percentile of the cluster magnitude distribution; i.e. p<0.05, 197 Maris & Oostenveld, 2007). This yielded seven distinct waveforms, representing the significance of the 198 main effect of each of the three factors across time, as well as their first order and second-order 199 interactions. To further explore the interactions, we then performed for each contrast of interest two-200 tailed paired T-tests, by means of cluster-based permutation testing approach (1000 random permutations 201 202 testing across all 32 channels) to correct for multiple comparisons across different time points (cluster threshold was set at the 95th percentile of the cluster magnitude distribution; i.e. p<0.05, Maris and 203 Oostenveld 2007). 204

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#### 206 *3.2 Single-subject analyses*

Based on the results of the group-analysis in two-handed controls, showing a crucial difference between real condition and imagery condition in the amplitude of the Nogo-P300 wave, we focused our singlesubject analyses of the phantom limb cases on this component. In each participant (i.e. including amputees and two-handed controls), the mean amplitude of Nogo-P300 peak at Fz was measured for each Nogo stimulus of the left/phantom and right/intact hand, during both real and imagery condition. P300 wave was defined as the most positive deflection following the onset of Nogo stimulus. In particular, the amplitude of P300 was computed for each individual subject in a 100 ms time window centered on her/his average peak in Nogo trials (P300 range: 280–380 ms). Then, Nogo-P300 peaks were
normalized in z-scores, calculated within-subject across all conditions (i.e. real, imagery, left, right). Zscores peaks were used as dependent variables.

of SingleBayes ES.EXE 217 By means program 218 (http://homepages.abdn.ac.uk/j.crawford/pages/dept/Single Case Effect Sizes.htm) we compared, in 219 separated analyses, P300 amplitudes of both experimental cases (moving-phantom and static-phantom) to the P300 amplitudes recorded in the control group. The program is specifically devised to test whether 220 an individual's score is significantly different from a control or normative sample (Crawford, Garthwaite, 221 and Porter 2010). It provides a point estimate of the effect size for the difference between the case and 222 223 controls (Z-CC) with an accompanying 95% credible interval and it provides a point and interval estimate of the abnormality of the case's score (i.e. it estimates the percentage of the population that would obtain 224 a lower score, together with a 95% credible interval on this percentage) (Crawford, Garthwaite, and 225 226 Porter 2010). This test was used to perform four comparisons, in which the amplitude of the left Nogo-P300 of each phantom limb case, during the real task, was compared with that of controls, either during 227 the real task or the imagery task. The  $\alpha$  value was set at 5%, and Bonferroni correction for multiple 228 229 comparisons was applied (p-value: 0.05/4=0.0125).

The above described Crawford's test was used also to investigate the presence of both a side effect and 230 231 a task effect in the two cases. With respect to the side effect, to analyze potential differences between left (phantom) and right (intact) hand during Nogo trials of the real task, we computed for all the participants 232 a delta (right minus left) where positive values indicate a greater P300 amplitude in response to right 233 234 than left stimuli. The resulting delta of each phantom-case was compared with the same delta of controls. The  $\alpha$  value was set at 5%, and Bonferroni correction for multiple comparisons was applied (p value: 235 0.05/2=0.025). With respect to the task effect, to analyze potential differences between real and imagery 236 conditions in the left/phantom side, we computed for all the participants a delta (real minus imagery) 237

where positive values indicate greater P300 amplitude in real condition than in imagery condition. The resulting delta of each case was compared with the same delta of two-handed controls. The  $\alpha$  value was set at 5%, and Bonferroni correction for multiple comparisons was applied (p-value: 0.05/2=0.025).

Finally, the performance of the two cases was directly compared by means of the C CTC.exe program

242 (http://homepages.abdn.ac.uk/j.crawford/pages/dept/Compare\_Two\_Cases.htm). This program uses Bayesian statistical methods to draw inferences concerning the difference between the test scores of two 243 single cases by referring their scores to a control sample. The p-values for this test are used to determine 244 whether we can reject the null hypothesis that the difference between the two cases is an observation 245 from the distribution of differences between pairs of controls. Point and interval estimates of the effect 246 247 size for the difference (Z-PCC) between the two cases are reported (Crawford, Garthwaite, & Wood, 2010). This test was used to perform four comparisons, in which the two cases was compared on the 248 following variables: left Nogo-P300 real, left Nogo-P300 imagery, Nogo-P300 delta right-left, Nogo-249 P300 delta real-imagery. It is to note that these programs are all implemented Bayesian statistics, which 250 is more adequately suited for supporting null results. See Supplementary materials for other analyses 251 including both Go and Nogo trials and both N200 and P300 waves. 252

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#### 254 **Behavioral data**

The statistical analyses on behavioral data were performed on mean RTs in response to Go stimuli. The analyses were performed on all the RTs collected in response to right/intact hand stimuli during the real task, because i) no behavioral data were collected for the two phantom-cases with respect to left/phantom hand stimuli during the real task; ii) no behavioral data were collected during the imagery task. Analyses were performed only on correct responses (i.e. incorrect responses corresponded to a wrong key press or a delayed response). Trials with RTs faster than 150 ms or deviating more than 2 standard deviations from individual mean RT were also excluded. As for electrophysiological data, Crawford's tests were
used to compare each phantom-case with controls and the two phantom-cases between each other.

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264 **4. Results** 

# 265 Electrophysiological data

266 *4.1 Group analysis* 

Grand average waveforms are depicted in Figure 2 and 3. To note, ERPs evoked by Nogo stimuli in real 267 condition presented the typical neurophysiological markers of motor inhibition responses, with the 268 emerge of N200/P300 complex, replicating previous studies using similar paradigms (Bokura et al., 269 2001). The point-by-point ANOVA revealed different significant effects. As expected, a significant main 270 effect of Condition was found. At Fz, this factor was a significant source of variance within three different 271 time intervals: 165-256 ms [coinciding with the latency of N200 ( $F_{1,15}=49.8$ , p<0.0000038)]; 294-415 272 273 ms [coinciding with the latency of P300 ( $F_{1,15}=31.9$ , p<0.0005)]; and 487-650 ms [coinciding with the latency of the negative shoulder following P300 wave ( $F_{1,15}=28.3$ , p<0.0001)], with greater responses 274 after Nogo than in Go stimuli (Figure 2A). Importantly, although a similar pattern of activity was present 275 during imagery condition, the point-by-point ANOVA showed a significant main effect of Task at Fz 276 within these intervals: 258-345 ms [coinciding with the latency of P300 ( $F_{1,15}$ =39.3 p=0.000028)], 443-277 523 ms [coinciding with the latency of the negative shoulder of P300 (F1,15=11.4, p=0.004)], with 278 significant lower amplitudes during imagery with respect to real task (Figure 2B). Furthermore, the 279 ANOVA found a significant Condition\*Task interaction. At Fz, this factor was a significant source of 280 281 variance within two different time intervals: 126-215 ms [coinciding with the latency of N200 wave  $(F_{1,15}=34.2, p<0.00032)$ ]; and 254-359 ms [coinciding with the latency of P300 wave  $(F_{1,15}=40.1, p<0.00032)$ ]; 282 p=0.000013)]. Two-tailed paired T-tests revealed that larger Nogo-N200 - and Nogo-P300 were elicited 283 in real with respect to the imagery task (p always <0.001) (Figure 3). 284

285	Figure 2 about here
286	Figure 3 about here

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# 288 *4.2 Single-subject analyses*

Our main goal was to examine electrophysiological activity associated with the inhibition of motor action in phantom limb syndrome with and without phantom movement. Voluntary motor inhibition was probed by correct responses to Nogo stimuli in the normal condition, which required an interruption of the prepared action. The waveforms in response to right/intact and left/phantom Nogo stimuli of the movingphantom case, controls, and static-phantom case are depicted in Figure 4A.

294 The moving-phantom case showed the typical neurophysiological pattern of motor inhibition (Bokura et al., 2001), with the emerge of the classical ERP pattern not only in response to right/intact stimuli but, 295 crucially, also in response to left/phantom Nogo stimuli, in a way similar to two-handed controls. In 296 297 particular, when the amplitude of the left Nogo-P300 of the moving-phantom case, during the real task, was compared with the left Nogo-P300 amplitude of the two-handed controls performing the real task, 298 Crawford's test did not find any significant differences (p=0.3) (Figure 4B). However, when it was 299 300 compared with the left Nogo-P300 amplitude of the two-handed controls performing the imagery task, a significant difference was found (Z-CC=2.880; p=0.006); i.e. the Nogo-P300 amplitude was significantly 301 greater when the moving-phantom case performed the real task than when two-handed controls 302 performed the imagery task (see Figure 4B). 303

On the contrary, the static-phantom case showed the classical ERP pattern in response to Nogo stimuli only when performing the task with the right/intact hand. In particular, when the amplitude of the left Nogo-P300 of the static-phantom case, during the real task, was compared with the left Nogo-P300 amplitude of the controls performing the real task, Crawford's test found a significant difference (Z-CC=-2.654; p=0.010), with significant greater Nogo-P300 amplitudes in two-handed controls than in the static-phantom case (Figure 4B). However, when it was compared with the left Nogo-P300 amplitude of
the two-handed controls performing the imagery task, no significant difference was found (see Figure
4B), suggesting a similar performance between the imagery task of the two-handed controls and the real
task of the static-phantom case (see Figure 4B). Coherently, when directly comparing P300 amplitude in
response to left/phantom Nogo stimuli during the real task of the two cases, a significant difference was
found (Z-PCC=2.488; p=0.0125), corresponding to a greater P300 in moving- than in static-phantom
case (Figure 4B).

With respect to the side effect in the real task, the delta right-left did not differ between the movingphantom case and controls (p=0.3), suggesting that similar P300 amplitudes were evoked irrespective of the side (Figure 4C). Conversely, the delta right-left of the static-phantom case was significantly different with respect to both controls (Z-CC=2.889; p=0.006) and moving-phantom case (Z-PCC=-2.553; p=0.011), with significantly higher delta values for the static-phantom case, suggesting a lower P300 amplitude in response to left than right Nogo stimuli (Figure 4C).

With respect to the task effect in the left/phantom side, we found that the delta real-imagery did not differ 322 between the moving-phantom case and controls (p=0.3), indicating that they showed similarly different 323 ERP patterns between the real than in the imagery tasks (Figure 4D). Contrariwise, the delta real-imagery 324 of the static-phantom case was significantly different with respect to both controls (Z-CC=-2-211; 325 p=0.024) and moving-phantom case (Z-PCC=2.084; p=0.027), with significant lower values for the 326 static-phantom case, indicating a similar P300 amplitude evoked in the real and in the imagery task 327 (Figure 4D) (P300 mean amplitude z-scores; *moving-phantom case*: real left/phantom= 0.43; delta real 328 329 right/intact-left= -0.08; delta left/phantom real-imagery= 0.86; controls: real left/phantom=  $0.29\pm0.16$ ; imagery left/phantom=  $-0.29\pm0.25$ ; delta real right/intact-left=  $0.05\pm0.18$ ; delta left/phantom real-330 imagery= 0.58±0.38; static-phantom case: real left/phantom=-0.14; delta real right/intact-left= 0.57; 331 delta left/phantom real-imagery= -0.26). 332

333 See Supplementary materials for the results on the differential response between Nogo and Go stimuli334 with respect to both N200 and P300.

- 335
- --- Figure 4 about here ---
- 337

# 338 Behavioral data

The analysis on RTs relative to the right/intact hand in response to Go trials showed no difference (p always > 0.4) between each of the two cases with respect to controls (RTs; mean±sd: *moving-phantom case*: 361.57; *controls*: 317.32±46.17), and no difference between the two cases (p=0.93), suggesting that potential differences (e.g. in age or skills) between the two cases and controls did not affect the behavioral performance during the Go/Nogo task. See Supplementary Figure 4 in Supplementary materials.

344

# 345 **5. Discussion**

In the present study, the phantom limb syndrome was investigated from an electrophysiological 346 perspective, aiming at describing a neural counterpart of the vivid experience, sometimes reported by 347 amputees, of being able to perform voluntary movements with their phantom. In particular, we focused 348 on a specific aspect of the motor control, the movement inhibition (i.e. the ability to suppress, withhold, 349 350 delay or interrupt ongoing or planned actions) and we asked what happens when a phantom movement has to be suppressed. To this aim, we recorded ERPs during a Go/Nogo tasks, where, according to a 351 consolidated literature in healthy subjects (e.g. Kok 1986; Bokura et al. 2001; Smith et al. 2008), larger 352 353 inhibition-related ERP components were expected in EEG responses to Nogo stimuli, as compared to EEG responses to Go stimuli. In our experimental design, the effects of "actual" (real key press during 354 Go trials) and mental (imagery key press) Go/Nogo task were investigated in two phantom cases (one 355 with and one without phantom movement) and in 16 healthy controls. First of all, we found that, in 356

healthy controls, suppressing real and imagined movements produces a discriminative EEG pattern, with
significantly greater inhibitory ERP responses in the former task relative to the latter. More crucial for
the present study, in the moving-phantom case, suppressing "real" movements with the existing limb and
the phantom limb evoked comparable inhibitory ERP responses to Nogo stimuli, which were clearly
distinguishable from those evoked by imagined movements. On the contrary, in the static-phantom case,
who did not experience kinematical sensations with his phantom, inhibitory ERP responses to Nogo
stimuli during real and imagery tasks were indistinguishable.

364

365 Disentangling inhibitory responses during the suppression of real and imagined movements in two-366 handed controls

As expected, a main effect of Condition showed the classical EEG pattern for a Go/Nogo task, with greater amplitude of N200 and P300 components in Nogo trials than in Go trials, irrespective of Side (left/right) and Task (real-imagery). See Figure 2A. Furthermore, a Task effect suggested that, irrespective of Side and Condition (Go/Nogo), the amplitude of P300 component was greater in real than in imagery task. See Figure 2B. More interestingly, a Condition by Task interaction suggested that, irrespective of Side, the Nogo-Go differential response in N200 and P300 components was significantly greater in real than imagery task. See Figure 3.

With respect to N200 and P300, several studies on Go/Nogo task showed that these two ERP components are typically associated with motor inhibition and that they are generally elicited by Nogo stimuli (Bokura et al., 2001; Cojan et al., 2013; Falkenstein et al., 1995). Some authors suggested that N200 Nogo effect may reflect a non-motoric stage of inhibition, or recognition of the need for inhibition, while the Nogo P300 may overlap with a positive movement-related potential occurring specifically on trials where overt motor responses must be inhibited (Smith et al. 2008). More specifically, when comparing Go and Nogo stimuli, it has been reported that P300 component in response to Nogo stimuli has a larger amplitude (Bruin et al. 2001; Schmajuk et al. 2006; Nakata et al. 2010; Cojan et al. 2013) and a more anterior
distribution (Smith et al. 2008; Barry et al. 2010) than that elicited by Go stimuli. This anterior
distribution has been linked to the inhibition of a motor response when a Nogo stimulus is presented, a
sort of index of an active inhibitory process (e.g. Woodward et al. 1991; Rockstroh et al. 1992; Smith et
al. 2008; Randall and Smith 2011).

386 When ERP responses to Nogo stimuli were considered individually (as in the post hoc comparisons of 387 the Task by Condition interaction), the amplitude of both N200 and P300 components was significantly larger in real than in imagery task (Figure 4, lower part). However, in the Task effect, the overall 388 difference between real and imagery tasks was found only in P300 component (Figure 2B), suggesting 389 390 that this component is more relevant in disentangling when a real or imagined movement has to be suppressed. In particular, as mentioned above spazio Go-P300 has been described as a motoric stage of 391 392 inhibition, while Nogo-N200 as a non-motoric motoric stage that is more related to a general need for 393 inhibition (e.g. Smith et al. 2008). Thus, the stronger result on P300 component suggest that real, more than imagined movements, reflect the motoric stage of inhibition; the weaker results on N200 component 394 suggest that the more cognitive stage of inhibition can be less relevant to discriminate between real and 395 imagined movements. 396

Our Task effect is in agreement with previous data on motor imagery (Burle, Vidal, & Bonnet, 2004; 397 398 Galdo-Alvarez, Bonilla, González-Villar, & Carrillo-de-la-Peña, 2016; González-Villar, Bonilla, & Carrillo-de-la-Peña, 2016), showing that the overt and covert execution of the task seems to be sufficient 399 to produce a discriminative pattern in the EEG activity. Although these previous studies reported that the 400 401 imagined performance of a stop-signal task produces a pattern of brain electrical activity with diminished amplitude with respect to that associated with real performance of the task, they focused more on the 402 similarities between real and imagery EEG pattern rather than on their differences, supporting the view 403 404 of a functional equivalence of imagined and real performances (Jeannerod, 2001). On the one hand, our results confirm this functional equivalence, showing a similar effectiveness of real and imagery task in
inducing an ERP modulation during Go/Nogo paradigm; on the other hand, they clearly show that the
EEG discriminative patter is greater during actual than covered execution of the task, with respect to
P300 component.

409

Suppressing phantom movements induces inhibition-related ERP responses comparable to those induced 410 by suppressing real movements and different from those induced by suppressing imagined movements 411 Crucially for the purpose of the present study, we compared inhibitory motor responses in the two 412 phantom limb cases, with and without phantom movements. Based on the results we found in two-handed 413 414 controls, we focused on the Nogo-P300 component, which is crucial in the distinction between real vs. imagined movement suppression. We found that, when the moving-phantom case has to "actually" 415 perform the task, she showed the classical ERP pattern associated with motor inhibition, comparable to 416 417 that found in healthy controls, who performed the task with two existing hands. On the contrary, the static-phantom case, who did not experience kinematical sensations with his phantom, showed this ERP 418 response only for the (right) intact hand, while, for the (left) phantom hand, it was not present. 419 420 Furthermore, in our results, phantom movement can be functionally disentangled from motor imagery, so that, as in healthy controls, ERP responses were dramatically different when the moving-phantom 421 case was asked to "actually" perform the Go/Nogo task (i.e. to "press" the key with her phantom) or to 422 imagine doing it. 423

Interestingly, single-subject analyses revealed a strong modulation on the P300 component, in which opposite results were found in the two cases. In moving-phantom case, the P300 modulation during real condition was comparable to that found in controls during the same condition, but, crucially, it was significantly different from that found during their imagery condition (see Figure 4B). Contrariwise, in the static-phantom case, the amplitude of the P300 component during real condition was significantly

different from real condition of controls and it was comparable to their imagery conditions (see Figure 429 430 4B). Importantly, a significant difference was found when the P300 was compared between the two phantom cases during the real task (see Figure 4B). In addition, when comparing the differential ERP 431 response to Nogo stimuli with the right/intact hand and the left/phantom hand during the real task, no 432 433 difference emerged between the moving-phantom case and controls, suggesting that the moving-phantom case showed a comparable response when the movement has to be suppressed with her intact (right) hand 434 435 and with her phantom (left) hand, similarly as two-handed controls (Figure 4C). On the contrary, the static-phantom case showed a significant difference with respect to both controls and moving-phantom 436 case, with greater differential P300 between the right/intact hand and the left/phantom hand during the 437 438 real task, suggesting the presence of genuinely motoric inhibition only when the movement has to be suppressed with his intact (right) hand and not when it has to be suppressed with his phantom (left) hand 439 (Figure 4C). Interestingly, single-subject analyses on the delta real-imagery response show that, while 440 441 the moving-phantom case shares with controls similar ERP pattern, the static-phantom case presents significant differences with respect to both controls and moving-phantom case (see Figure 4D). When 442 the movement has to be suppressed with his phantom, the static-phantom case showed no difference 443 during the real and the imagery task contrary to what showed by controls and moving-phantom case, 444 which presented a clearly distinct ERP pattern between real execution and motor imagery with their 445 446 left/phantom hand. We can speculate that, since the static-phantom case cannot "move" his phantom 447 limb, then he could only imagine to move it, resulting therefore in a similar ERP pattern during the real and the imagery task. 448

According to previous behavioral (Garbarini et al., 2018; Raffin, Giraux, et al., 2012) and fMRI (Raffin, Mattout, et al., 2012) studies, these results strongly support the view that phantom movements and imagery movement are functionally disentangled, also extending this concept to the context of motor inhibition. Indeed, the present results suggest that a similar amount of inhibitory cerebral activity is

453 needed to suppress real and phantom movements, while the amount of inhibition necessary to block the 454 imagined movement is lower and, consequently, the inhibitory-related ERP components appear reduced. Thus, only in moving-phantom case, who experienced the ability to voluntarily move her phantom, 455 inhibitory responses are fully implemented during the real task, when frontal areas exert, via premotor 456 457 cortex and/or SMA, an inhibitory control over the primary motor cortex. These results can contribute to 458 the debate about cortical reorganization (H. Flor et al., 1998; Herta Flor et al., 2006; Lotze et al., 2001; Ramachandran, Rogers-Ramachandran, & Stewart, 1992) and maintained cortical representation 459 (Kikkert et al., 2016, 2018; Makin et al., 2013) in phantom limb syndrome. The cortical reorganization 460 model (H. Flor et al., 1998; Herta Flor et al., 2006; Lotze et al., 2001; Ramachandran et al., 1992) suggests 461 462 that the deprivation of sensory inputs in the S1 missing hand cortex leads to cortical reorganization, where the deprived cortex becomes responsive to inputs from its cortical neighbors. Specifically, it has 463 been showed that referred sensations in the phantom (that is, painful and non-painful phantom sensations 464 465 that can be elicited by stimulating body areas adjacent to but also far from the amputated limb) are a perceptual correlate of reorganizational processes in the S1 cortex (Ramachandran et al., 1992). 466 However, other studies showed instead that multiple factors interact to maintain local structural and 467 functional representations (Makin et al., 2013), with a maintained representation of the phantom hand in 468 the S1 cortex following amputation (Bogdanov, Smith, & Frey, 2012; Raffin, Mattout, et al., 2012) and 469 470 comparable patterns of movement-related activity when amputees moved their phantom hand compared 471 with two-handed controls moving their non-dominant hand (Makin et al., 2013). Our results seem to support the maintained cortical organization hypothesis, at least in the moving-phantom case, who 472 473 showed an ERP pattern of motor inhibition with her phantom limb comparable to that found in twohanded controls. In particular, in agreement with previous studies, our results suggest that the 474 475 representation of the amputated limb is functionally active, just like a real hand.

476	Although this study was performed only on two phantom limb cases, these findings provide <b>a</b> compelling
477	evidence that phantom movements share the same neurophysiological correlates of real movements, not
478	only when an action has to be executed, but also when it should be inhibited.
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623 Captions to figures:

Figure 1. A. Experimental task. Stimuli were visual, and they represented the dorsal view of a right or
a left hand. Gray hands: preparation cue; green hands: Go signal; red hands: Nogo signal. B.
Experimental procedure. Each participant (i.e. controls; moving-phantom case and static-phantom
case) underwent an EEG session during which she/he performed the Go/Nogo task. Each EEG session
comprised both blocks of real and imagery task.

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Figure 2. Group analysis results: main effects of Condition and Task. A) Main effect of Condition. 630 Mean ERPs in response to Go and Nogo signals, irrespective of the side (left, right) and of the task (real; 631 632 imagery). Data are displayed in microvolts as a function of time post-cue onset, for Fz electrode (referenced to the nose). The green waveform represents Go stimuli and the red waveform represents 633 Nogo stimuli. Point-by-point F value is represented below. Time intervals where the difference between 634 635 conditions was significant are highlighted in gray and the corresponding scalp topographies are shown. B) Main effect of Task. Mean ERPs in response to real and imagery task, irrespective of the side (left, 636 right) and the condition (Go; Nogo). Data are displayed in microvolts as a function of time post-cue 637 onset, for Fz electrode (referenced to the nose). The solid gray waveform represents ERPs in response to 638 the real task and the dashed gray waveform represents ERPs in response to the imagery task. Point-by-639 640 point F value is represented below. Time intervals where the difference between the tasks was significant are highlighted in gray and the corresponding scalp topographies are shown. 641

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Figure 3. Group analysis results: Condition\*Task interaction. Mean ERPs in response to Go real, Go imagery, Nogo real and Nogo imagery, irrespective of the side (left, right). Data are displayed in microvolts as a function of time post-cue onset, for Fz electrode (referenced to the nose). The solid green waveform represents Go stimuli during the real task, the dashed green waveform represents Go stimuli

during the imagery task, the solid red waveform represents Nogo stimuli during the real task, the dashed 647 red waveform represents Nogo stimuli during the imagery task. Point-by-point F value is represented 648 below. Time intervals where the difference between conditions was significant are highlighted in gray 649 and the corresponding scalp topographies are shown. Two-tailed paired T tests results of the contrast of 650 651 interest (Nogo real vs Go real; Nogo imagery vs Nogo imagery; Nogo real vs Nogo imagery; Go real vs Go imagery) are displayed below the F value. Time intervals during which the contrasts of interest were 652 significantly different are colored in blue and pink, such as the topographical distribution of T values 653 reflecting the statistical comparison. 654

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656 Figure 4. A. ERP waveforms of the moving-phantom case, controls and the static-phantom case. Mean ERPs of the moving-phantom case (left), controls (center) and static-phantom case (right) Nogo 657 stimuli, both with the left/phantom (upper part) and with the right/intact (lower part) limb, during the real 658 task (solid lines) and during imagery task (dashed lines). Data are displayed in microvolts as a function 659 of time post-cue onset, for Fz electrode (referenced to the nose). B. Single-subject analyses results. 660 Mean z-scores of the P300 amplitude in response to Nogo stimuli of the moving-phantom case (left), 661 controls (center) and the static-phantom case (right). C. Side effect: Mean z-scores of the P300 amplitude 662 of the delta right/intact-left/phantom of the moving-phantom case (left), controls (center) and the static-663 phantom case (right) performing the real task. D. Task effect: Mean z-scores of the P300 amplitude of 664 the delta real-imagery of the moving-phantom case (left), controls (center) and the static-phantom case 665 (right) in response to Nogo stimuli requiring the suppression of left/phantom movements. -  $*p \le 0.05$ , 666 667 \*\*p<0.01, \*\*\*p<0.001. Error bars represent standard error of the mean of two-handed controls.