

1 **The role of extra-striate areas in conscious motor behavior: a**  
2 **registered report with Fast-Optical Imaging**

3 Elisabetta Colombari<sup>a</sup>, Giorgia Parisi<sup>a</sup>, Sonia Mele<sup>a</sup>, Chiara Mazzi<sup>a</sup> and Silvia  
4 Savazzi<sup>a</sup>

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6 <sup>a</sup> Perception and Awareness (PandA) Laboratory, Department of Neuroscience, Biomedicine and  
7 Movement Sciences, University of Verona, Strada le Grazie 8, Verona, Italy

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## 10 **Abstract**

11 Disclosing the brain areas responsible for the emergence of visual awareness and their timing of  
12 activation represents one of the major challenges in consciousness research. In particular, isolating  
13 the neural processes strictly related to consciousness from concurrent neural dynamics either related  
14 to prerequisites or post-perceptual processing has long engaged consciousness research. In this  
15 framework, the present study aims at unravelling the spatio-temporal dynamics underlying conscious  
16 vision by adopting a distinctive experimental design in which both awareness and motor response are  
17 manipulated, allowing the segregation of neural activity strictly related to awareness from response-  
18 related mechanisms. To this aim, we will employ a GO/NOGO detection task, in which participants  
19 will respond or withhold responding according to the experimental condition. Critically, during the  
20 performance of the task, participants' brain activity will be recorded by means of Event-Related  
21 Optical Signal (EROS) technique, which provides accurate information about brain functions both  
22 from the temporal and spatial point of view, simultaneously. The combination of this experimental  
23 design with EROS recording will enable us to pinpoint the neural correlates underlying conscious  
24 vision and to disentangle them from processes related to the response. In addition, by coupling  
25 conventional EROS analysis with Granger Causality analysis, we will be able to clarify the potential  
26 interplay between consciousness-related extra-striate areas and response-related motor areas.

## 27 **1. Introduction**

28 Consciousness, namely the set of subjective experiences we have when we are awake, is one of the  
29 most intriguing topics debated in neuroscience research. In particular, the search for its neural  
30 correlates (NCC) has permeated the literature in recent decades. In broad strokes, one of the most  
31 widely used approaches to assess such NCCs involves contrasting brain activity occurring when a  
32 visual stimulus enters consciousness with brain activity occurring when the same stimulus does not  
33 reach awareness. This renowned paradigm is known as *contrastive analysis* (Aru *et al.*, 2012) and  
34 has been frequently combined with electrophysiological recording or functional neuroimaging,  
35 leading to numerous and dissimilar results (Förster *et al.*, 2020). Indeed, the interpretations of spatio-  
36 temporal dynamics underlying conscious vision are among the most disparate. ERP studies propose  
37 two possible electrophysiological markers as correlates of visual awareness: an earlier occipito-  
38 temporal negative deflection (i.e., Visual Awareness Negativity – VAN) detectable 200 ms after the  
39 presentation of the stimulus, and a later positivity (i.e., Late Positivity – LP) widespread over centro-  
40 parietal regions, peaking 300-500 ms after the stimulus onset (Koivisto & Revonsuo, 2010). However,  
41 the electrophysiological signature/s characterizing conscious vision has still to be elucidated. This  
42 may be attributed to one of the main limitations of the contrastive analysis, which is represented by  
43 its ineffectiveness in dissociating the true NCC (i.e., the set of neural correlates necessary and  
44 sufficient to enable consciousness) from concurrent neural dynamics either related to prerequisites or  
45 post-perceptual processing (Aru *et al.*, 2012). In most prior studies aiming at identifying such NCCs,  
46 participants were asked to make judgments about their experience. However, such an operation could  
47 lead to confounding neural processes related to the task, not strictly to awareness per se.

48 For this reason, in an effort to disentangle the proper correlates of consciousness from neural activity  
49 related to the response, no-report paradigms have been employed. In this framework, no-report  
50 paradigms, where participants are not requested to perform any tasks or to provide any judgments  
51 about their perceptual experience, represent an advantageous tool to dissociate the neural processes  
52 strictly related to consciousness from subsequent processes related to the required response (Tsuchiya  
53 *et al.*, 2015; Hatamimajoumerd *et al.*, 2022). Studies employing this kind of paradigm with different  
54 techniques such as EEG and fMRI concluded that LP is highly modulated by several different  
55 cognitive processes occurring at later stages of processing (Mazzi *et al.*, 2020; Schlossmacher *et al.*,  
56 2020; Dembski *et al.*, 2021; Kronemer *et al.*, 2022), as well as by the task relevance of the stimulus  
57 (Makeig & Jung, 2000; Pitts *et al.*, 2014; Shafto & Pitts, 2015; Schelonka *et al.*, 2017; Dellert *et al.*,  
58 2021; Hense *et al.*, 2024). By contrast, the role of response requirements, as well as that of attention,  
59 on the VAN are still debated as different studies have reported both positive (e.g., Bola &  
60 Doradzińska, 2021; Dellert *et al.*, 2021; Doradzińska & Bola, 2024) and negative (e.g., Koivisto *et*

61 *al.*, 2006; Cohen *et al.*, 2020; Dellert *et al.*, 2022; Ciupińska *et al.*, 2024) results. Interestingly, in a  
62 study published in 2016 by Koivisto and colleagues (Koivisto *et al.*, 2016), authors successfully  
63 dissociated ERP correlates of visual awareness from those related to post-perceptual mechanisms,  
64 disclosing that VAN was not modulated by response requirements. The authors adopted a particular  
65 partial-report paradigm in which participants were sometimes asked to provide a report by pressing a  
66 response button when they were aware of the stimulus and sometimes to withhold responding in case  
67 of awareness. They found that, while the amplitude of LP was modulated by the response (i.e., it was  
68 greater in trials where participants were asked to respond in case of awareness, compared to the Aware  
69 condition where they were asked to withhold responding), VAN did not change depending on task  
70 requirements. This allowed Koivisto and colleagues to advocate for an early onset of visual  
71 awareness: the phenomenal content of a visual experience, indeed, takes place before LP, more  
72 specifically in the temporal window of VAN.

73 Several pieces of evidence are consistent in considering VAN as the electrophysiological signature  
74 of phenomenal consciousness (Koivisto *et al.*, 2008; Railo *et al.*, 2015), while the localization of its  
75 neural generator still remains open. In this regard, previous MEG source localization studies (Vanni  
76 *et al.*, 1996; Liu *et al.*, 2012) identified the Lateral Occipital Complex (LOC), an extra-striate visual  
77 areas traditionally associated with objects recognition, as the generator of VAN. **Moreover, previous**  
78 **no-report studies using both EEG and fMRI (Dellert *et al.*, 2021; Kronemer *et al.*, 2022) have also**  
79 **found awareness effects in LOC and linked it to VAN.** The same result was achieved in a recent work  
80 aimed at unravelling the spatio-temporal dynamics underlying conscious vision (Colombari *et al.*,  
81 2024). In such study, participants were asked to perform a discrimination task on the orientation of a  
82 tilted Gabor patch while their brain activity was recorded first with EEG and then with Fast Optical  
83 Imaging. This allowed authors to identify the exact temporal window of VAN and LP and then, by  
84 taking advantage of the peculiarity of Fast Optical Imaging of achieving both temporal and spatial  
85 accurate information (Gratton & Corballis, 1995; Gratton & Fabiani, 2010; Baniqued *et al.*, 2013), to  
86 investigate the spatio-temporal unfolding of brain activity occurring in these predetermined time  
87 windows. Authors contrasted activity of Aware trials (i.e., trials in which participants reported to  
88 perceive the orientation of the stimulus) with activity of Unaware ones and observed a sustained  
89 activation of LOC in the VAN temporal window, consistently with the above-mentioned MEG  
90 studies. More interestingly, they observed that, only when the stimulus crossed the threshold of  
91 consciousness, activity in extra-striate visual areas triggered subsequent activation of motor areas,  
92 although motor response was required in both Aware and Unaware conditions. Authors tried to  
93 interpret this unexpected finding by ascribing it to the selection of the correct response, that could be  
94 provided in the Aware trials only where participants consciously perceived the stimulus. Indeed, in

95 Aware trials participants had to press a specific button on the response box (to provide the correct  
96 answer about the orientation of the Gabor patch), while when the stimulus was unseen (i.e., Unaware  
97 trials) they had to respond randomly, by pressing indifferently one of the two response buttons.  
98 However, the employed experimental paradigm did not allow the authors to thoroughly investigate  
99 this issue. Thus, in order to clarify the interplay between extra-striate areas and motor regions in  
100 awareness, in the present study we will adopt a go/no-go detection task (similar to that adopted by  
101 Koivisto *et al.*, 2016), while recording participants' brain activity by means of Fast Optical Imaging.  
102 Specifically, Event-Related Optical Signal (EROS) technique will be employed. This technique, by  
103 shedding near-infrared light through the brain tissues, is able to detect changes in the light scattering  
104 properties that are known to be directly related to neural activity, thus providing accurate information  
105 about brain functions both from the temporal and spatial point of view, simultaneously (Gratton *et*  
106 *al.*, 1997; Gratton & Fabiani, 1998, 2001). Critically, the study will adopt a peculiar **distinctive**  
107 paradigm manipulating both awareness and response. The latter, indeed, will be provided sometimes  
108 in the Aware condition (condition Aware-GO/Unaware-NOGO) and sometimes in the Unaware one  
109 (condition Aware-NOGO/Unaware-GO). This double manipulation will enable us to unravel the  
110 spatio-temporal unfolding of awareness-related activity, by disentangling neural activity related to  
111 awareness from response-related mechanisms. Indeed, in the present study, we can investigate the  
112 NCCs both when the motor response is required and when no task is performed, thus allowing to  
113 isolate consciousness effects from the effects related to the task. Importantly, the experimental  
114 paradigm adopted will enable us to elucidate the interplay between extra-striate visual areas and  
115 motor areas. Indeed, in addition to conventional EROS analyses, we will perform Granger Causality  
116 analysis, in order to disclose the relationship existing among the investigated areas. In broad strokes,  
117 Granger analysis allows to move beyond the classical identification of cortical activation provided by  
118 EROS analysis by disclosing functional circuits underpinning the investigated brain function (Seth *et*  
119 *al.*, 2015). When coupled with EROS, Granger Causality analysis represents a powerful tool to  
120 highlight predictive relationship between activations in the investigated regions of interest (ROI) at  
121 different time-points (Parisi *et al.*, 2020).

122 Based on previous literature suggesting that VAN is independent from subjective report (Koivisto *et*  
123 *al.*, 2016; Ye *et al.*, 2024) and LOC represents the cortical generator or VAN (Liu *et al.*, 2012;  
124 Colombari *et al.*, 2024), we expect Aware trials to elicit early greater activation of LOC,  
125 independently of the response requirement. Moreover, by combining EROS conventional analysis  
126 with Granger Causality analysis, and manipulating both awareness and motor response, we aim to  
127 highlight potential interplay between consciousness-related extra-striate areas and response-related  
128 motor areas both when the motor response is required and when it has to be inhibited.

## 129 **2. Methods**

### 130 **2.1 Ethics Information**

131 The study is approved by the local Ethics Committee (Prog.171CESC) and it will be conducted in  
132 accordance with the principles laid down in the 2013 Declaration of Helsinki ~~and~~. Participants will  
133 be recruited from the University of Verona community, by means of printed flyers displayed on notice  
134 boards at different University of Verona sites and through advertisements on social media. Each  
135 participant will be fully informed about the modalities of the study before taking part in the  
136 experiment and written informed consent will be signed. In addition, participants will receive  
137 compensation for their participation and will be debriefed after the conclusion of the experiment.

### 138 **2.2 Participants**

139 We will recruit healthy adults, right-handed (as assessed by means of the standard handedness  
140 inventory *Edinburgh Handedness Questionnaire*; Oldfield, 1971) and aged between 18 and 50 years  
141 old. All of them will have to report normal or corrected-to-normal vision, no history of neurological  
142 or psychiatric disorders and no contraindications to MRI. The study will be conducted at the Panda  
143 lab of the University of Verona (Italy).

#### 144 *2.2.1 Sample size estimation*

145 The estimate of the sample size for the current study is based on our previous EROS study (Colombari  
146 et al., 2024), in which a similar paradigm was adopted, and similar analyses on similar ROIs were  
147 performed. Specifically, EROS data from the ROI of LOC were extracted, and significant time-points  
148 were averaged within participants so to have one value for each of them. Then, a one-sample t-test  
149 was performed ( $t(23) = 2.99$ ,  $p = .006$ , Cohen's  $d = .611$ ), and the resulting Cohen's  $d$  was employed  
150 to compute the sample size estimation for the current study. Specifically, the estimated sample size  
151 for research questions Q1 (i.e., "Can we replicate Colombari et al., 2024 findings showing that LOC  
152 is an NCC?") and Q2 (i.e., "Is the activity in LOC independent from the response?") was calculated  
153 with G-Power software (v. 3.1.9.7), with a power of 90% and a level of significance of 2%. The  
154 estimated sample size resulted in 32 participants (critical  $t = 2.143$ ; actual power = 0.900). Considering  
155 that the estimated sample size for this study ( $n = 32$ ) is more than double the typical sample size of  
156 EROS studies present in literature, the same estimated sample size seems to be also adequate to  
157 answer research questions Q3 (i.e., "Does consciousness modulate the activation of motor areas in a  
158 detection task?") and Q4 (i.e., "Does consciousness modulate the activation of motor areas in  
159 ABSENCE of motor response?"). For a review of the existing EROS literature, see Supplementary  
160 Table 1 at

161 [https://osf.io/ebfu3/?view\\_only=9ec2e6bf32ba4a8bb8b858639ec40a59](https://osf.io/ebfu3/?view_only=9ec2e6bf32ba4a8bb8b858639ec40a59)) from which emerges that,  
162 on average, EROS studies employ experimental samples composed of about 13 participants (mean  
163 12.944; SD 7.008).

164 ~~Since in EROS literature no previous studies report the effect size because of technical constraints of~~  
165 ~~the employed dedicated software, an a priori statistical sample size estimation for the present study~~  
166 ~~is not achievable based on EROS data. For this reason, we first based our sample size estimation on~~  
167 ~~a review of the existing EROS literature (see Supplementary Table 1 at~~  
168 ~~[https://osf.io/ebfu3/?view\\_only=9ec2e6bf32ba4a8bb8b858639ec40a59](https://osf.io/ebfu3/?view_only=9ec2e6bf32ba4a8bb8b858639ec40a59)) (GRATTON *et al.*, 1995;~~  
169 ~~Gratton *et al.*, 1997, 2000, 2001; Gratton & Fabiani, 2003; Wolf *et al.*, 2003; Low *et al.*, 2006; Tse~~  
170 ~~& Penney, 2007; Medvedev *et al.*, 2008, 2010; Proulx *et al.*, 2018; Toscano *et al.*, 2018; Parisi *et al.*,~~  
171 ~~2020; Tse *et al.*, 2021; Knight *et al.*, 2024), from which emerges that, on average, EROS studies~~  
172 ~~employ experimental samples composed of 13 participants (mean 12.944; SD 7.008). Moreover, we~~  
173 ~~decided to estimate the sample size for the present study basing on a previous EEG study employing~~  
174 ~~the same experimental design adopted in the present study (Koivisto *et al.*, 2016). The estimated~~  
175 ~~sample size for research questions Q1 (i.e., “Can we replicate Colombari *et al.*, 2024 findings showing~~  
176 ~~that LOC is an NCC?”) and Q2 (i.e., “Is the activity in LOC independent from the response?”) was~~  
177 ~~calculated with G Power software (v. 3.1.9.7), with a power of 90% and a level of significance of~~  
178 ~~2%. To estimate the power needed to detect the effect of awareness (aware vs. unaware trials), we~~  
179 ~~considered the significant main effect of awareness of a within subjects repeated measures ANOVA~~  
180 ~~( $F(1,14)=17.06$ ,  $P=0.001$ ,  $\eta_p^2=0.55$ ) carried out in Koivisto *et al.*(2016). The estimated sample~~  
181 ~~size resulted in 15 participants (critical  $F=6.887$ ; actual power=0.918). Since EROS signal to noise~~  
182 ~~ratio is lower than that of EEG, we will increase our final sample to 26 participants. Considering that~~  
183 ~~the estimated sample size for this study ( $n=26$ ) is the double of the typical sample size of EROS~~  
184 ~~studies present in literature, the same estimated sample size seems to be also adequate to answer~~  
185 ~~research questions Q3 (i.e., “Does consciousness modulate activation of motor areas in a detection~~  
186 ~~task?”) and Q4 (i.e., “Does consciousness modulate activation of motor areas in ABSENCE of motor~~  
187 ~~response?”).~~

### 188 2.2.2 Exclusion Criteria

189 As better specified in section 2.3, before getting involved in the study, participants will undergo a  
190 perceptual threshold assessment, in order to identify the proper stimulus to be employed in the main  
191 experiment. To be enrolled in the study, participants will have to successfully complete this session.  
192 The criterion used is that one of the stimuli presented during the threshold assessment will have to  
193 be acknowledged as perceived a minimum of 25%, a maximum of 75%, or closest to the 50% of the

194 times (i.e., at perceptual threshold level). If no stimulus results at the threshold level, the participant  
195 should not be enrolled in the study.

196 In addition, participants who will not complete all the experimental sessions, as well as participants  
197 reporting a level of Awareness superior to 75% or inferior to 25% at the end of the experiment will  
198 be excluded from analyses. This is to maintain comparable the number of trials in the two  
199 experimental conditions (i.e., Aware and Unaware) and to ensure a reliable EROS activity (because  
200 of its relatively low signal-to-noise ratio, EROS needs a high number of trials per condition, in  
201 order to compute statistics). Moreover, participants whose behavioral performance will be affected  
202 by biases related to the behavioral response (as assessed by catch trial analysis, explained more in  
203 detail below) will be excluded from the analyses (see below –*Section 2.8.1 Behavioral data* for  
204 more detailed information on the analysis of catch trials). Finally, participants whose EROS signal  
205 could not be detected properly during the experiment (for example because of too dark hair or  
206 technical issues) will not ~~also~~ be included in the analyses **as well**. In particular, the opacity value  
207 (i.e., the product of the scattering and absorption coefficients) will be estimated for each participant.  
208 Based on this value, it is possible to judge the quality of the signal for each participant,  
209 independently from the experimental condition. Opacity values of all participants will be averaged  
210 together providing the absorption coefficient to be used when running statistical analysis.  
211 Participants whose opacity value is equal to 0 or exceeds three standard deviations of the mean will  
212 be excluded from statistical analyses.

213 Importantly, each participant who will be excluded due to the previously mentioned exclusion criteria,  
214 will be replaced with the recruitment of another participant. Thus, the number of participants to be  
215 recruited will be increased to reach a total of **32** ~~26~~ analyzed subjects, as specified in section 2.2.1.

## 216 **2.3 Stimuli**

217 Stimuli will be created by means of a custom-made Matlab script (version R2022b; the MathWorks,  
218 Inc., Natick, MA) and resized by means of Photoshop (Adobe Photoshop CC, v2014.0.0). As shown  
219 in Figure 1, they will be gray circles (.85 .85 .85 RGB), presented on a black background, with 8 radii  
220 equally distanced one from another. One radius (the first one, clockwise) can be slightly thicker than  
221 the others (critical trials) or not (catch trials). The thickness of the radius for critical stimuli will be  
222 individually assessed for each participant on the basis of a subjective perceptual threshold assessment  
223 that will be held before the main experiment.

224 Both in the perceptual threshold assessment and in the main experiment, the stimulus will be  
225 presented in the lower right quadrant of the screen, specifically at an eccentricity of  $3.5^\circ$  from the  
226 fixation cross along the vertical meridian and of  $2^\circ$  along the horizontal one. This is to allow a left-



227 lateralized EROS montage, as a full-head montage is not achievable in our lab due to technical  
228 constraints (i.e., insufficient probes). Moreover, since EROS technique is sensitive to depth, a right-  
229 lateralized stimulus ensures that it elicits activity in the left portion of the primary visual cortex, which  
230 is known to be anatomically closer to the skull compared to the right one (Zhao *et al.*, 2022), thus  
231 ensuring a better penetration of near-infrared light through brain tissues.

## 232 **2.4 Perceptual Threshold Assessment**

233 Before starting the experiment, participants will undergo a perceptual threshold assessment, with the  
234 aim of identifying, for each participant, the level of thickness of the critical radius so that it results to  
235 be perceived as thicker 50% of the times. To this aim, stimuli with different levels of radius thickness  
236 will be randomly presented and the subjective perceptual threshold will be measured using the method  
237 of constant stimuli. Specifically, 9 levels of radius thickness will be presented. The range of stimuli  
238 to be used in the perceptual threshold assessment will be selected based on the results of a pilot  
239 experiment in which participants were asked to perform the same task employed in the perceptual  
240 threshold assessment while presented with a wider range of radius thickness. This will allow us to  
241 identify a smaller range of optimal stimuli to be presented thus excluding a range of stimuli whose  
242 thickness was almost never or always reported by participants. Each level of radius thickness will be  
243 presented 5 times per block, for a total of 8 blocks. Thus, all the stimuli, as well as the catch stimulus,  
244 will be presented 40 times each. Participants will be asked to press the spacebar as soon as they detect  
245 the stimulus with a thicker radius. The stimulus identified as perceived a minimum of 25%, a  
246 maximum of 75%, and closest to 50% of the times at the end of the subjective perceptual threshold  
247 assessment will be used in the experimental task, together with the catch. The perceptual threshold  
248 assessment, as well as the main experiment, will be conducted in a dimly illuminated room and  
249 participants will be sitting in front of a 17 in. LCD monitor (resolution 1920x1080, refresh rate of  
250 144 Hz) placed at a viewing distance of 57 cm. Their head will be held in place by means of an  
251 adaptable chin rest so that eyes are aligned with the center of the screen. Both the perceptual threshold  
252 assessment and the main experiment will be programmed and administered using E-Prime 3.0  
253 software (E-Prime Psychology Software Tools Inc., Pittsburgh, PA, USA). Before starting the  
254 perceptual threshold assessment, participants will undergo a fixation training (Leung *et al.*, 2009), in  
255 order to ensure they will maintain their gaze on the central fixation cross correctly.

## 256 **2.5 Experimental Procedure**

257 The experiment will be composed of two identical sessions lasting approximately 3 hours each  
258 performed on different days. The first session will be preceded by the assessment of the subjective

259 perceptual threshold, which, in turn, will last around 20 minutes. The two experimental sessions will  
 260 be identical except for the EROS montages, specifically devised to obtain better coverage of the brain  
 261 areas of interest. The order of the montages will be counterbalanced across participants, as well as  
 262 the order of conditions (see below for more detailed information).

263 The task will be a two-conditions go/no-go detection task, similar to that adopted by Koivisto *et al.*,  
 264 2016, in which participants have to respond in different ways according to the experimental condition  
 265 (Table 1). In condition “Aware-GO”, they will be asked to press the spacebar on the keyboard as soon  
 266 as they perceive the thicker radius, and withhold responding when they do not perceive any difference  
 267 among radii. On the contrary, in condition “Aware-NOGO”, participants will be asked to withhold  
 268 responding when they perceive a thicker radius, and press the response button when they do not  
 269 perceive any difference. Each trial will begin with the presentation of a central fixation cross,  
 270 followed 500 ms later by a sound (1000Hz) presented for 100 ms, notifying participants of the  
 271 subsequent onset of the stimulus. After a random interval ranging from 500 to 600 ms, the stimulus  
 272 will be presented for 100 ms in the lower right quadrant of the screen. After that, participants will be  
 273 asked to respond according to the experimental condition. Each experimental session will be  
 274 composed of 24 blocks: 12 blocks for condition Aware-GO/Unaware-NOGO and 12 blocks for  
 275 condition Aware-NOGO/Unaware-GO, counterbalanced across participants according to the order  
 276 depicted in Table 1. Each block will consist of 50 critical trials and 15 catch trials. The whole  
 277 experiment will be composed of 48 blocks per participant, for a total of 2400 critical trials and 720  
 278 catch trials per participant.

		Awareness	
		yes	no
Response	no	Aware-NOGO	Unaware-NOGO
	yes	Aware-GO	Unaware-GO

279 **Table 1. Experimental conditions.** Both Awareness and Response are manipulated: Awareness is  
 280 experimentally manipulated by employing a threshold stimulus, so that sometimes it is consciously perceived  
 281 (Aware) and sometimes not (Unaware). Response is manipulated by the task: in condition GO participants are  
 282 asked to respond by pressing a key, while in condition NOGO they are asked to withhold responding. The  
 283 combination of these two manipulations gives rise to the 4 experimental conditions depicted in the table.

284

Participants	Day 1		Day 2	
	EROS montage 1	Task	EROS montage 2	Task

1	A	GNGG	B	NGGN
2	B	GNGG	A	NGGN
3	A	NGGN	B	GNGG
4	B	NGGN	A	GNGG

285 **Table 2. Counterbalancing of montages and task conditions across participants.** Both EROS montages  
286 and task conditions (G = Aware-GO/Unaware-NOGO; N = Aware-NOGO/Unaware-GO) will be  
287 counterbalanced across participants. In the column “Task”, each letter represents 6 blocks of task. Thus, each  
288 day, participants will perform 12 blocks per condition, for a total of 24 blocks of task per day.

## 289 2.6 Optical Recording

290 Three synchronized Imagent frequency domain systems (ISS, Inc., Champaign, IL) will be used to  
291 record continuous fast optical data throughout experimental sessions. Each system is equipped with  
292 4 photo-multiplier tubes detectors, for a total of 12 detectors. Near-infrared light (830 nm) will be  
293 delivered from 48 laser diodes on participants’ scalp and it will be modulated at 110 MHz. Each of  
294 12 detectors will receive light from sets of 16 light emitters, multiplexed every 25.6 ms, resulting in  
295 a sampling rate of 39.0625 Hz.

296 To avoid cross-talk between channels, the array of source-detector pairs (i.e., the montage) will be  
297 created by means of a specific program (NOMAD, Near-Infrared Optode Montage Automated  
298 Design) implemented in Matlab, useful to place sources and detectors at optimal distances. In this  
299 experiment, we will set the minimal distance to 17.5 mm and the maximum distance to 50 mm, in  
300 order to ensure an extensive coverage of the brain regions of interest both from the spatial and the  
301 depth point of view. The distance between the source and the detector of a channel, in fact, determines  
302 the depth of the light pathway (Gratton *et al.*, 2000), thus corresponding to the depth of the  
303 investigation: namely, longer channels can investigate deeper layers and shorter channels can  
304 examine shallower regions.

305 Both light emitters and detectors will be placed on participants head using a custom-built helmet. To  
306 minimize interferences, before placing the optical fibers on the head, the hair will be carefully moved  
307 with cotton buds, so that the fibers can reach the scalp directly. In order to better adhere to the head  
308 of the participant, we will employ two helmets of different sizes: one 55-56 cm large, and one 57-58  
309 cm large. For each helmet, we will develop two different montages, so that to provide a dense  
310 coverage of the regions of interest (i.e., the left occipital, temporal and parietal cortices, see Figure  
311 2). Each montage will consist of the combination of 12 detectors and 48 light emitters, resulting in a  
312 total of 192 channels per montage. As mentioned before, each montage will be recorded in a separate  
313 session, and the order will be counterbalanced across participants.

314 At the end of each EROS session, the scalp location of each source and detector will be digitized in  
315 relation to four fiducial points (i.e., nasion, inion and pre-auricular points) with a neuro-navigation

316 software (SofTactic, E.M.S., Bologna, Italy) combined with a 3D optical digitizer (Polaris Vicra, NDI,  
317 Waterloo, Canada). Afterwards, the digitized scalp locations will be co-registered with each  
318 participant's individual MRI, using a dedicated software package (OCP, Optimized Co-registration  
319 Package, Matlab code developed by Chiarelli and colleagues (Chiarelli *et al.*, 2015).

320 For this reason, participants will undergo a structural MRI at the Azienda Ospedaliera Universitaria  
321 Integrata of Verona (AOUI).

## 322 **2.7 MRI Acquisition**

323 Participants' individual structural MRI will be acquired by means of a 3 Tesla Philips Ingenia scanner  
324 with a 32-channel head RF receive coils. A whole brain high-resolution 3D T1-weighted image (T1w)  
325 Turbo-field echo image (1mm-isotropic TE/TR=3.8/8.4 ms, TI=1050 ms) will be acquired.

326 The T1w field of view (240 x 240 x 180 mm) will be large enough to allow for the ears and the entire  
327 scalp to be fully included in the image to facilitate later and accurate co-registration with functional  
328 data.

## 329 **2.8 Data Analysis**

### 330 *2.8.1 Behavioral data*

331 Raw data will be processed by means of custom scripts created on Matlab (the MathWorks, Inc.,  
332 Natick, MA). Data will be divided into the 4 experimental conditions (i.e., Aware-GO, Unaware-  
333 NOGO, Aware-NOGO, Unaware-GO). For each participant, trials with reaction times lower than 150  
334 ms and higher than 3 standard deviations from the mean will be excluded from the analysis. Data will  
335 be successively analyzed using Jamovi (version 2.3.28): first, the percentage of Aware and Unaware  
336 trials will be calculated, in order to assess that a sufficient amount of trials is present for each  
337 condition. Participants presenting more than 75% or less than 25% of Awareness will be discarded  
338 from the sample. This is because, in that case, the number of Unaware (or Aware) trials would be  
339 insufficient for statistical EROS analysis. EROS technique, indeed, although having a high  
340 localization power from both the spatial and temporal point of view, has a relatively low signal-to-  
341 noise ratio. For this reason, a high number of trials is required for statistical analysis. Subsequently,  
342 reaction times (RTs) will be analyzed for the "GO" conditions, thus paired sample t-tests (two-tailed)  
343 will be applied to compare the mean RTs between Aware-GO and Unaware-GO conditions. Finally,  
344 to verify that participants are performing the task accurately and that there are no biases related to the  
345 response, catch trials will be analyzed. As mentioned above, catch trials are those trials in which all  
346 the radii of the stimulus are equally thick, thus no differences in the stimulus are present. In case of  
347 catch trials, the participants' task will be different according to the condition: in the Aware-GO  
348 condition, they are expected to withhold responding, while in the Aware-NOGO condition, they are

349 expected to respond. Thus, catch trials will be analyzed separately for the two conditions (GO and  
350 NOGO) by means of a paired sample t-test (two-tailed), in order to ensure that the behavioral  
351 performance follows the above-mentioned trend. Paired sample t-tests (two-tailed) will indeed be  
352 performed to test whether catch trials performance is significantly different from critical trials.

### 353 2.8.2 EROS data

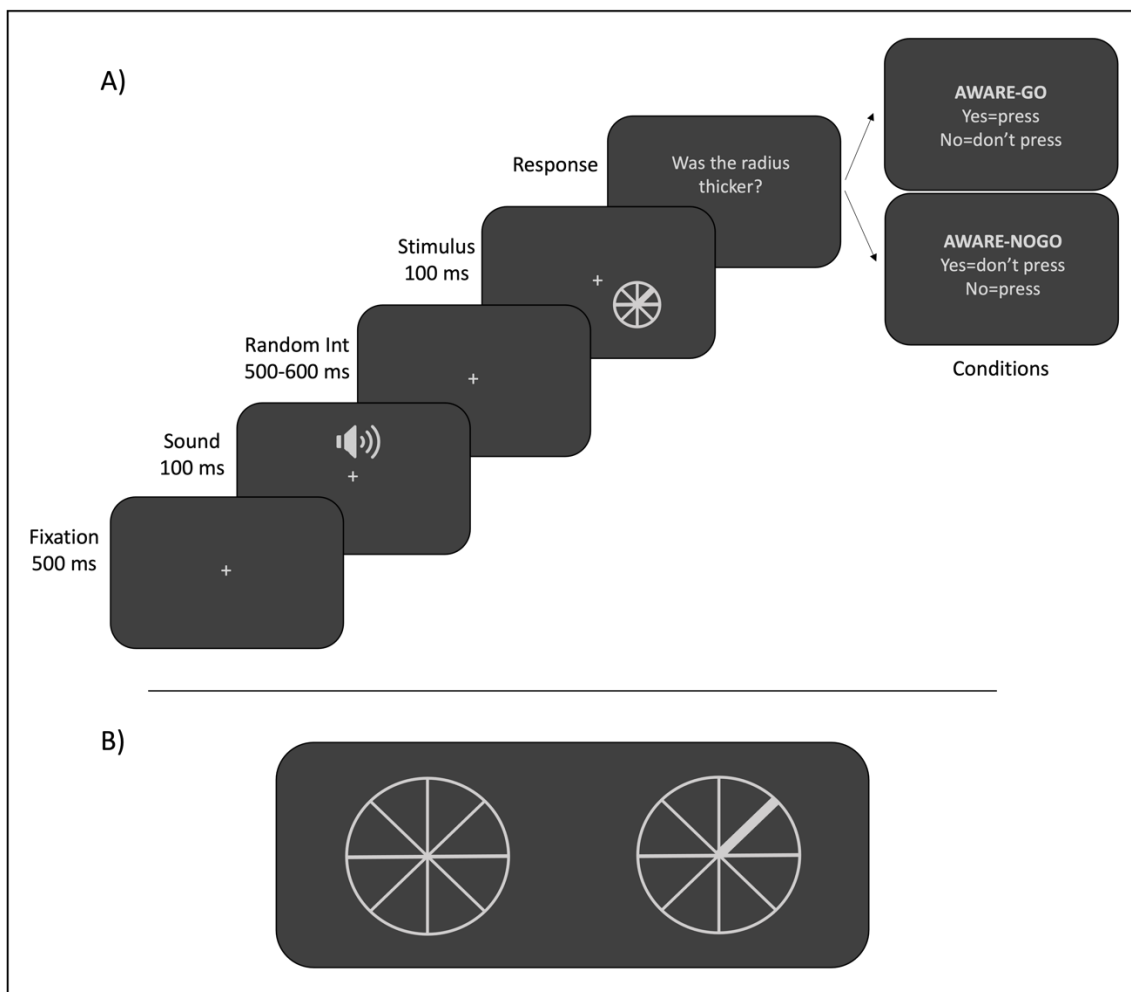
354 Pre-processing of continuous phase delay (i.e., time-of-flight) data will be computed by means of a  
355 dedicated in-house software, P-POD (Pre-Processing of Optical Data, run in Matlab, version  
356 R2013b). Thus, raw data will be normalized (i.e., corrected for phase wrapping and de-trended to  
357 remove low-frequency drifts), ~~baseline-corrected~~ **demeaned** and filtered by means of a 6<sup>th</sup> order  
358 Butterworth band-pass filter which allows frequencies between 0.5 Hz and 15 Hz. Pulse artifact will  
359 be removed by using a regression algorithm (GRATTON *et al.*, 1995). After that, data will be  
360 averaged separately for each subject, condition, and channel and segmented into epochs time-locked  
361 to the onset of the stimulus. Each epoch will comprise a period from 486 ms before the stimulus onset  
362 to 998 ms following the stimulus onset, resulting in an epoch lasting 1484 ms. Subsequently,  
363 statistical analyses will be computed with an in-house software package (Opt-3d; (Gratton, 2000)),  
364 which provides statistical spatial maps of fast optical data.

365 To perform statistics, data from channels whose diffusion paths intersect a given voxel will be  
366 combined (Wolf *et al.*, 2014). Phase delay data will be spatially filtered with an 8-mm Gaussian  
367 kernel **and baseline corrected using a 204 ms time-window preceding the stimulus onset**. Within each  
368 ROI, t-Statistics will be calculated at group level, converted into Z-scores and corrected for multiple  
369 comparisons using random field theory (Worsley *et al.*, 1995; Kiebel *et al.*, 1999). Then, Z-scores  
370 will be weighted and orthogonally projected onto the surface of an MNI template brain, according to  
371 the physical homogenous model (Arridge & Schweiger, 1995; Gratton, 2000).

372 In order to investigate the neural dynamics related to conscious vision and to disentangle the role of  
373 the motor areas, the following contrasts between conditions will be computed: 1) Aware-GO versus  
374 Unaware-GO and 2) Aware-NOGO versus Unaware-NOGO. These contrasts allow to investigate the  
375 research questions the proposed study aims at answering (see Section 3 for a detailed description of  
376 the planned analysis). **Importantly, both frequentist and Bayesian statistics (with default priors) will  
377 be computed, to test both positive and negative effects.**

378 Moreover, Granger Causality analysis will be computed. Granger Causality analysis allows to explore  
379 the predictive interactions between different brain areas at different time-points. Specifically, this  
380 approach requires a region of interest (ROI) to be used as a “seed” and investigating whether the

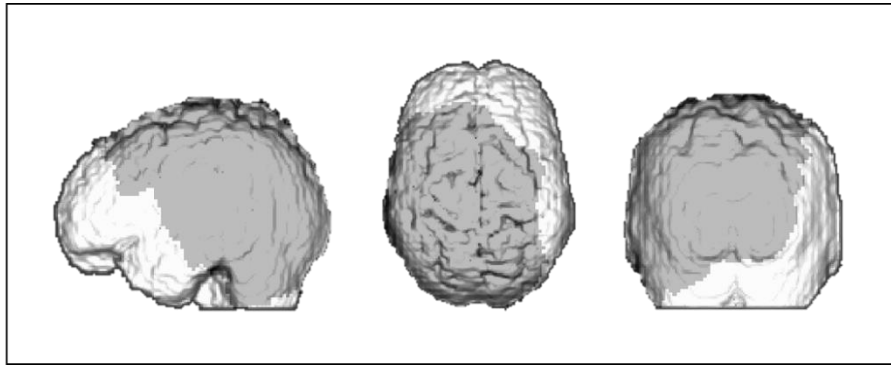
381 activity of this seed predicts activity in the other ROIs at a later time-lag, by deriving statistical maps  
 382 from t-statistics computation (then transformed into z scores) for each lag.  
 383 Statistical functional analysis will be computed within specific predetermined regions of interest  
 384 (ROIs) and time intervals. ROIs will be defined by a 2-dimensional box-shaped structure, covering  
 385 an area of 20x20 millimeters. Critical ROIs will be selected on the basis of the results obtained in the  
 386 above-mentioned experiment (Colombari *et al.*, 2024) and ~~by visual inspection of functional data.~~  
 387 ~~In particular,~~ they will be located in the occipital and in the left parietal and temporal lobes, specifically  
 388 over the primary visual cortex (V1, Brodmann Area 17), the left lateral occipital cortex (LOC,  
 389 Brodmann Area 19), the left supplementary motor area (SMA, Brodmann Area 6), the left premotor  
 390 area (PM, Brodmann Area 6) and the left primary motor cortex (M1, Brodmann Area 4). Statistical  
 391 analysis will be computed within specific temporal windows of interest selected on the basis of the  
 392 results obtained by Colombari *et al.*, 2024. This is to reduce the risk of false positives, as Opt3d does  
 393 not offer the possibility to correct data for multiple comparisons in the temporal domain. The specific  
 394 time windows tested for each hypothesis are listed in Table 3.



395  
 396 **Figure 1. Trial procedure and stimuli:** A) Experimental procedure: the trial begins with a fixation cross  
 397 persisting at the center of the screen for 500 ms. After that, an acoustic tone lasting 100 ms will be presented,

398 followed by a random interval ranging from 500 to 600 ms. Then, the stimulus will be presented for 100 ms  
399 and participants will be asked to respond according to the experimental condition (i.e., Aware-GO or Aware-  
400 NOGO). **B)** Example of stimuli: on the left is shown the catch stimulus, with all the radii equally thick; on the right  
401 is depicted the critical stimulus, with the first radius, clockwise, thicker than the others.

402



403

404 **Figure 2. Covered area.** The gray area represents the area covered by the EROS montages (combined  
405 together) from the sagittal, axial and coronal point of view.

### 3. Study design

Question	Hypothesis	Sampling Plan	Analysis Plan	Rationale for deciding the sensitivity of the test for confirming or disconfirming the hp	Interpretation given different outcomes
<p><b>Q1:</b> Can we replicate Colombari et al., 2024 findings showing that LOC is an NCC?</p>	<p><b>H1:</b> We hypothesize to replicate Colombari et al., 2024 results: greater activity in LOC in an early temporal window (i.e., 150-350 ms post stimulus onset) is observed when contrasting Aware and Unaware trials in the condition in which the response is required (i.e., GO condition)</p> <p><b>Expected outcome:</b> LOC aware-GO&gt;LOC unaware-GO, as measured by EROS activity</p>	<p><del>Because of technical constraints of the dedicated EROS software, the effect size for EROS data is not computable and thus sample size cannot be determined basing on previous EROS findings. For this reason, the sample size estimation for the present question is basically determined according to two strategies:</del></p> <p>1) a systematic review of existing EROS literature revealing that the typical sample size used is 13 participants (see Supplementary Table 1)</p> <p>2) sample size estimation based on a previous EEG study of Koivisto et al., 2016, in which authors employed the same experimental paradigm adopted in the present study and aware trials were compared to unaware trials. Sample size calculation was thus performed with G-Power software (v. 3.1.9.7);</p>	<p><b>A1:</b> The goal is to replicate the results of Colombari et al., 2024, in which the manual response was required for both Aware and Unaware conditions. Here, in order to perform the same analysis, early LOC activity in Aware-GO and Unaware-GO trials will be compared by using a paired-sample one-tailed t-test, computed with the EROS dedicated analysis software “Opt3d”.</p> <p><b>Contrast to be computed:</b> AWARE GO VS UNAWARE GO</p> <p><b>ROI to be tested:</b> LOC</p> <p><b>Time interval of interest:</b> 150-350ms after stimulus onset</p>	<p>Effect size for EROS data is not computable. This is because the existing software dedicated to statistical EROS analysis (i.e., Opt-3d) does not allow to calculate this measure. However, we estimated our sample size basing of the effect size of a previous EEG study (Koivisto et al., 2016) employing the same experimental design and based on the sample used in EROS literature.</p>	<p><b>O1.1:</b> A significant t-test within the interval of interest will be interpreted as a successful replication of previous findings, supporting the involvement of LOC in NCC.</p> <p><b>O1.2:</b> The absence of this effect will not confirm the hypothesis, suggesting that LOC is not involved in the conscious detection of a stimulus property.</p>



		<p>with a power of 90% and a level of significance of 2%, resulting in 15 participants. However, since EROS signal to noise ratio is lower than that of EEG, we will increase our final sample to 26 participants, which is almost the double of the estimated sample size.</p> <p>Since the present research question aims at replicating the results of Colombari et al., 2024, sample size estimation is based on those EROS data. Sample size calculation was thus performed with G-Power software (v. 3.1.9.7), with a power of 90% and a significance level of 2%, resulting in 32 participants.</p>			
<p><b>Q2:</b> Is the activity in LOC independent from the response?</p>	<p><b>H2:</b> We hypothesize that LOC activity is independent from response requirement: when contrasting activity elicited by Aware-NOGO trials with activity elicited by Unaware-NOGO trials, we expect to find the same activation of LOC found in the Aware-GO vs Unaware-GO contrast.</p>	<p>Since research question Q2 involves the same analyses of research question Q1 (but for the NOGO condition), the sampling plan for Q2 is the same as for Q1.</p>	<p><b>A2.1:</b> A paired-sample one-tailed t-test will be computed in order to compare early activity in LOC in the NOGO condition. Thus, activity in Aware-NOGO and Unaware-NOGO trials will be contrasted.</p> <p>Both frequentist and Bayesian statistics (with default prior) will be computed.</p>	<p>As above</p>	<p><b>O2.1.1:</b> A significant t-test in the time window of interest will suggest that LOC activity is independent from response, since its activity is observed even when no response is required (NOGO conditions).</p> <p><b>O2.1.2:</b> If greater activity in LOC in the time window of interest is not observed, then it means that LOC activity is somehow related to the motor response.</p>

	<p><b>Expected outcome:</b></p> <p>LOC aware-NOGO&gt;LOC unaware-NOGO, as measured by EROS activity</p> <p>(LOC aware-GO&gt;LOC unaware-GO)=(LOC aware-NOGO&gt;LOC unaware-NOGO)</p>		<p><b>Contrast to be computed:</b></p> <p>AWARE NOGO VS UNAWARE NOGO</p> <p><b>ROI to be tested:</b></p> <p>LOC</p> <p><b>Time interval of interest:</b></p> <p>150-350ms after stimulus onset</p>		
			<p><b>A2.2:</b> The interaction effect between awareness and motor response will be tested by means of a paired-sample one-tailed t-test computed between contrast Aware-GO VS Unaware-GO and contrast Aware-NOGO VS Unaware-NOGO</p> <p><b>Contrast to be computed:</b></p> <p>(AWARE GO VS UNAWARE GO) - (AWARE NOGO VS UNAWARE NOGO)</p> <p><b>ROI to be tested:</b></p> <p>LOC</p> <p><b>Time interval of interest:</b></p> <p>150-350ms after stimulus onset</p>		<p><b>O2.2.1:</b> Significant interaction effect will suggest that activity in LOC depends from response requirement</p> <p><b>O2.2.2:</b> The absence of a difference between the two effects will suggest that motor response does not affect awareness-related activity in LOC</p>

<p><b>Q3:</b> Does consciousness modulate activation of motor areas in a detection task?</p>	<p><b>H3:</b> When a motor response is required, consciousness modulates activation of motor areas (MA), as activity in motor areas is triggered by LOC (Colombari et al., 2024)</p> <p><b>Expected outcome:</b></p> <p>MA aware-GO &gt; MA unaware-GO, as measured by EROS activity</p> <p>LOC activity predicts MA activity (investigated by means of Granger Causality Analysis)</p>	<p>Considering that the estimated sample size for this study (n=26 32) is <b>more than twice the double</b> of the typical sample size of EROS studies present in literature (see <b>Supplementary Table 1, where a systematic review of existing EROS literature revealing that the typical sample size used is 13 participants is depicted</b>), the same estimated sample size <b>of Q1 and Q2</b> seems to be also adequate to answer research questions Q3 and Q4.</p>	<p><b>A3.1</b> A paired-sample one-tailed t-test will be computed in order to compare early activity in Motor Areas in the GO condition. Thus, activity in Aware-GO and Unaware-GO trials will be contrasted.</p> <p><b>Contrast to be computed:</b></p> <p>AWARE GO VS UNAWARE GO</p> <p><b>ROI to be tested:</b></p> <p>Motor areas</p> <p><b>Time interval of interest:</b></p> <p>Based on mean RTs, with a time window of <math>\pm 1.5</math> sd around the mean</p> <p><b>A3.2:</b> In order to further investigate the flow of activity occurring in the investigated brain areas,</p>	<p>As above</p>	<p><b>O3.1.1:</b> A statistically significant difference between the two conditions will suggest that, even in a detection task, response related motor activity is stronger in the Aware condition compared to the Unaware one.</p> <p>In Colombari et al., 2024 this difference was observed. Importantly, in this previous study a <i>discrimination</i> task was employed and participants were asked to provide two different responses in case of Awareness (intentional) or Unawareness (random). Instead, in this study participants are asked to perform a <i>detection</i> task, in which the motor behavior made to provide the response, when required, is the same for both Aware and Unaware condition and thus no response selection is required.</p> <p><b>O3.1.2:</b> If no difference between the tested conditions is observed, it will suggest that in a detection task there is no difference in the motor activity related to the response.</p> <p><b>O3.2.1:</b> Significant predictive interactions between LOC and motor areas will suggest that, when the stimulus enters consciousness,</p>

			<p>Granger Causality Analysis will be performed. In the present study, we will perform Granger analysis on the “Aware-GO VS Unaware-GO” contrast, since we are interested in investigating whether activity in motor areas is predicted by previous activity in LOC, when a motor response is required (i.e., in the GO condition). Thus, LOC will be used as seed ROI and later activity in motor areas will be investigated.</p> <p><b>Contrast to be computed:</b></p> <p>AWARE GO VS UNAWARE GO</p> <p><b>ROI to be tested:</b></p> <p>LOC (as seed ROI) Motor areas as predicted areas</p> <p><b>Time interval of interest:</b></p> <p>LOC: 150-350 ms after the stimulus onset</p> <p>MA: based on mean RTs</p>		<p>awareness-related activity in LOC predicts subsequent activity in motor areas. This (expected) outcome will suggest that consciousness modulates subsequent response-related motor activity, by directly triggering activation of motor areas, as observed in Colombari et al., 2024</p> <p><b>O3.2.2:</b> If no significant interactions between LOC and MA will be highlighted, then it would mean that activity in motor areas is not predicted by LOC. Specifically, it could be surmised that in a <i>detection</i> task, consciousness does not modulate activation of motor areas, as observed in Colombari et al., 2024, where a <i>discrimination</i> task was employed. The difference in the two tasks, indeed, consists in the type of motor response required: in the case of the discrimination task, the participant is asked to press one button or another according to the response. Conversely, in a detection task, the participant has to press a key when the target stimulus is detected. Thus, no selection of the response is needed. This difference could play a role in the relationship between consciousness and motor areas.</p>
<b>Q4:</b> Does consciousness modulate	<b>H4:</b> Consciousness modulates activation of motor areas, even if the	As Q3	<b>A4.1:</b> A paired-sample one-tailed t-test will be computed in order to	As above	<b>O4.1.1:</b> A statistically significant t-test will suggest that, when a motor response is not provided, the inhibition

<p>activation of motor areas in ABSENCE of motor response?</p>	<p>motor response is not required</p> <p><b>Expected outcome:</b></p> <p>MA aware-NOGO&gt;MA unaware-NOGO</p> <p>LOC predicts MA (investigated by means of Granger Causality Analysis)</p>		<p>compare activity in Motor Areas in the NOGO condition. Thus, activity in Aware-NOGO and Unaware-NOGO trials will be contrasted.</p> <p><b>Contrast to be computed:</b></p> <p>AWARE NOGO VS UNAWARE NOGO</p> <p><b>ROI to be tested:</b></p> <p>Motor areas</p> <p><b>Time interval of interest:</b></p> <p>Based on mean RTs, , with a time window of <math>\pm 1.5</math> sd around the mean</p>		<p>required to withhold responding is stronger when the visual characteristic of the stimulus is consciously perceived, compared to when no difference is perceived.</p> <p><b>O4.1.2:</b> If no difference between the tested conditions is observed, this will suggest that i) no inhibition is required to withhold responding, both in the Aware and Unaware condition, or ii) the inhibition is equally strong for the two conditions.</p>
			<p><b>A4.2:</b> With the aim of investigating the flow of activity occurring in the investigated brain areas also in the condition where no response is required, Granger Analysis will be performed on the “Aware-NOGO VS Unaware-NOGO” contrast. This will allow to investigate whether activity in motor areas is triggered by previous activity in LOC, even when a motor response is not required.</p>		<p><b>O4.2.1:</b> If significant predictive interactions between LOC and motor areas will be observed, then consciousness modulates subsequent activity in motor areas also in absence of a motor response. This could be due to inhibition of the response processes.</p> <p><b>O4.2.2:</b> If no significant interactions between LOC and MA will be highlighted, then LOC does not predict activity in motor areas in absence of motor response.</p>

			<p>Thus, LOC will be used as seed ROI and later activity in motor areas will be investigated.</p> <p><b>Contrast to be computed:</b></p> <p>AWARE NOGO VS UNAWARE NOGO</p> <p><b>ROI to be tested:</b></p> <p>LOC (as seed ROI)</p> <p>Motor areas as predicted areas</p> <p><b>Time interval of interest:</b></p> <p>LOC: 150-350 ms after the stimulus onset</p> <p>MA: Based on mean RTs</p>		
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## 408 **4. Pilot study**

409 In order to test the experimental paradigm, we pilot-tested the task.

410 A total of 10 right-handed participants (5 females and 5 males; mean age  $\pm$  standard deviation: 21  
411 years  $\pm$  1.0) took part in the pilot study. They all reported normal or corrected-to-normal vision and  
412 no history of neurological or psychiatric disorders. All of them provided written informed consent  
413 before starting the experiment.

414 After the first session, two participants dropped out the experiment, hence data from 8 participants  
415 were included in the statistical analyses. Moreover, in order to maintain an equal number of trials in  
416 both the conditions (i.e., Aware and Unaware), the percentage of Aware and Unaware trials was  
417 calculated and data from participants reporting a proportion of awareness equal or superior to 80%  
418 (i.e., 3 participants) were discarded from subsequent analysis. For this pilot study, we decided to raise  
419 the awareness threshold of acceptance to 80% (instead of 75%, that will be used in the experiment)  
420 in order to be more inclusive, given the low number of participants.

421 Thus, in total, data from 5 participants were included in the behavioral and functional analyses.

### 422 **4.1 Preliminary Results**

#### 423 *4.1.1 Behavioral results*

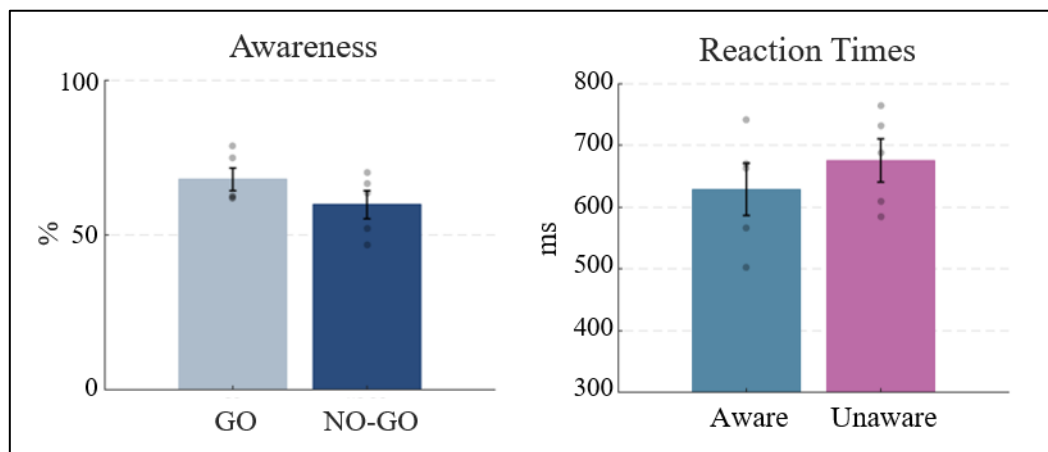
424 Raw data were processed by means of scripts created on Matlab (version R2017b; the MathWorks,  
425 Inc., Natick, MA). According to the participants' responses, trials were sorted into the four  
426 experimental conditions (i.e., Aware-GO, Unaware-NOGO, Aware-NOGO and Unaware-GO).  
427 Aware trials were those trials in which the participant reported to perceive the thicker radius, while  
428 Unaware trials were those trials in which participants could not perceive that the radius was thicker.  
429 As specified in Section 2.8, trials with RTs lower than 150 ms or higher than 3SD from the mean  
430 were removed. After removal, we had on average 830.6 trials for the Aware-GO condition, 389.2 for  
431 the Unaware-NOGO condition, 738.8 trials for the Aware-NOGO condition and 491.4 for the  
432 Unaware-GO.

433 Subsequently, once assessed the normality of RTs and Awareness distributions (Shapiro-Wilk test.  
434 RTs distribution:  $W=0.824$ ,  $p=0.125$ ; Awareness distribution:  $W=0.817$ ,  $p=0.112$ ), the percentage of  
435 Awareness for the two conditions was calculated: in the GO condition, Aware trials represented on  
436 average 68.02% of the trials, while in the NOGO condition, Aware trials constituted the 59.82% of  
437 the trials. Paired sample (two-tailed) t-test performed with Jamovi (version 2.3.28) highlighted that  
438 there was no significant difference between the two conditions ( $t_{(4)} = 1.88$ ,  $p = .134$ , Cohen's  $d =$   
439 ~~.839~~), suggesting that they are comparable. Similarly, mean RTs for Aware and Unaware trials in the  
440 GO condition were contrasted and the statistical analysis (Paired sample two-tailed t-test) revealed

441 that mean RTs for the Aware condition (628.530 ms) and the Unaware condition (675.317 ms) were  
442 not statistically different ( $t(4) = -1.77$ ,  $p = .152$ , Cohen's  $d = -.791$ ). This indicated that there was no  
443 difference in the responsiveness between the two conditions. The behavioral results are depicted in  
444 Figure 3.

445 Moreover, in order to verify that the employed paradigm works as planned and that participants  
446 performed the task accurately, analysis on catch trials was performed as described in Section 2.8.1  
447 *Behavioral data*. As specified above, catch trials were those trials in which all the radii of the stimulus  
448 are equally thick. Hence, in those cases, participants should report not to see the thicker radius. As  
449 expected, they correctly reported not seeing the thicker radius on average the 96.47% of times  
450 ( $sd=2.49$ ) in the Aware GO condition and the 98.36% ( $sd=1.89$ ) in the Aware NOGO condition.  
451 Paired sample (two-tailed) t-test revealed no significant difference between the two conditions.

452



453

454 **Figure 3. Behavioral results.** The percentage of Awareness was calculated for both “GO” and “NOGO”  
455 conditions (on the left). Mean reaction times were calculated for Aware and Unaware trials only for the “GO”  
456 condition (on the right). No significant differences were observed. Error bars represent SEM and gray dots  
457 represent individual data points showing the data distribution.

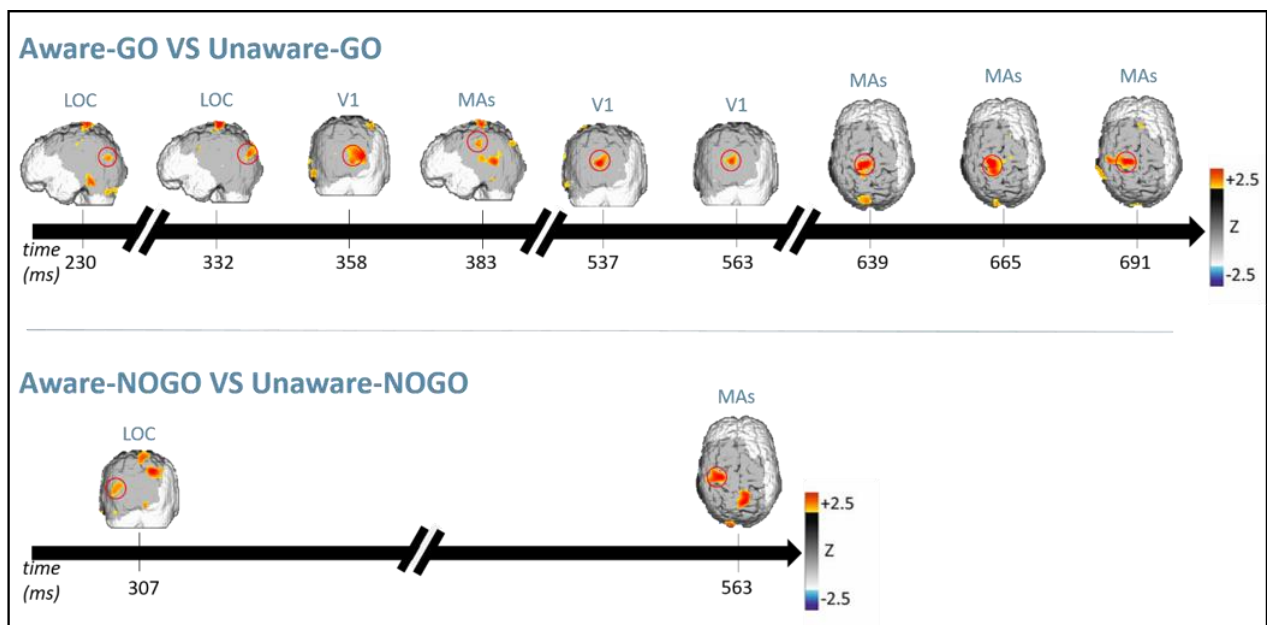
#### 458 4.1.2 EROS results

459 EROS data were pre-processed with a dedicated in-house software, P-POD (Pre-Processing of Optical  
460 Data, run in Matlab, version R2013b), as described in Section 2.8. Subsequently, we computed  
461 statistical analyses on pre-processed data by means of the dedicated in-house software package Opt-  
462 3d.

463 For this pilot study, participants' individual structural MR images could not be acquired, so an  
464 estimated MR-based head model was individually created using the Softaxic Optic system (Softaxic,  
465 E.M.S., Bologna, Italy) combined with a 3D optical digitizer (Polaris Vicra, NDI, Waterloo, Canada).  
466 EROS data were thus co-registered with the estimated MRI using a specific procedure performed in  
467 OCP software package (as specified above). Finally, co-registered data were transformed into MNI  
468 space for subsequent analyses.



469 For both GO and NOGO conditions, Aware and Unaware trials were contrasted. As shown in Figure  
 470 4, the Aware-GO vs Unaware-GO contrast replicated the results obtained by Colombari et al., 2024  
 471 ~~under review~~. In this contrast, indeed, we compared conditions in which the motor response was  
 472 required, thus replicating the task carried out in the previously mentioned experiment. Also in this  
 473 case, Aware trials elicited a sustained activation of LOC (230 and 332 ms after the stimulus onset),  
 474 followed by the recurrent activation of the primary visual cortex (V1) and the motor areas (MA) at  
 475 later stages of stimulus processing.  
 476 Similarly, contrasting Aware and Unaware trials in the condition where the motor response was not  
 477 required (i.e., the NOGO condition), greater activation of LOC was elicited in a timing comparable  
 478 to that of the contrast just mentioned above (i.e., 307 ms after the stimulus presentation). Interestingly,  
 479 also in this case awareness-related processing elicited activity in the motor areas, 563 ms after the  
 480 stimulus onset, despite in this condition no response was required, possibly suggesting an inhibition  
 481 to respond for the NOGO trials.



482  
 483 **Figure 4. EROS results.** Statistical parametric maps of the z-score difference computed contrasting Aware  
 484 and Unaware trials in the GO (upper panel) and NOGO condition (lower panel). Each map represents a 25.6  
 485 ms interval.

#### 486 4.2 Preliminary Discussion

487 The aim of the present pilot study was to assess whether the task and the experimental procedure were  
 488 suitable to investigate the study's research questions.

489 As described in Section 4.1, the pilot study successfully replicated the trend of activations observed  
 490 by Colombari et al., 2024, suggesting that the proposed study proves to be feasible in terms of  
 491 methodology. For the sake of clarity, it is important to point out that the preliminary results reported  
 492 here do not reach the statistical level of significance. This outcome was expected as data from only 5

493 participants were included in the analysis. For the same reason, we decided not to perform Granger  
494 Causality analysis as for this kind of analysis results from 5 participants would have been  
495 uninformative. Nevertheless, it was possible to observe that the proposed task could elicit a pattern  
496 of activation similar to that observed by Colombari et al., 2024, suggesting that the experimental  
497 paradigm proposed to investigate the research questions is suitable.  
498

## 499 **Data availability**

500 Upon acceptance of the Stage 2 registered report, we will share all raw and processed anonymized  
501 data as well as study materials publicly available as open data. Pilot raw and processed data can be  
502 found on this link: [https://osf.io/ebfu3/?view\\_only=9ec2e6bf32ba4a8bb8b858639ec40a59](https://osf.io/ebfu3/?view_only=9ec2e6bf32ba4a8bb8b858639ec40a59)

## 503 **Code availability**

504 All analysis codes will be made publicly available upon acceptance of the Stage 2 registered report.

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511 Resilience Plan (NRRP), project MNESYS (PE0000006)—"A Multiscale integrated approach to the  
512 study of the nervous system in health and disease" (DN. 1553 11.10.2022), CM is supported by MIUR  
513 D.M. 737/2021—"Neural correlates of perceptual awareness: from neural architecture to the  
514 preservation of conscious vision in brain tumor patients".

## 515 **Author contributions**

516 **EC** Conceptualization, Methodology, Software, Validation, Formal analysis, Investigation, Data  
517 Curation, Writing - Original Draft, Visualization, Funding Acquisition; **GP** Methodology, Formal  
518 Analysis, Investigation, Data Curation, Writing - Review & Editing; **SM** Methodology, Investigation,  
519 Writing - Review & Editing **CM** Methodology, Software, Data Curation, Writing - Review & Editing,  
520 Supervision; **SS** Conceptualization, Methodology, Resources, Writing - Review & Editing,  
521 Supervision, Project administration, Funding acquisition.

522 **Competing interests**

523 The authors declare no competing interests.

524 **References**

- 525 Arridge, S.R. & Schweiger, M. (1995) &lt;title&gt;Sensitivity to prior knowledge in optical  
526 tomographic reconstruction&lt;/title&gt; In Chance, B. & Alfano, R.R. (eds), *Optical*  
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