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Cognitive Load Reduces the Effects of Optic Flow on Gait and 2 Electrocortical Dynamics During Treadmill Walking 3

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1 **Cognitive load reduces the effects of optic flow on gait and**
2 **electrocortical dynamics during treadmill walking**

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47 **ABSTRACT**

48 While navigating complex environments the brain must continuously adapt to both
49 external demands such as fluctuating sensory inputs, as well as internal demands, such as
50 engagement in a cognitively demanding task. Previous studies have demonstrated changes in
51 behavior and gait with increased sensory and cognitive load, but the underlying cortical
52 mechanisms remain unknown. Here, in a Mobile Brain/Body Imaging (MoBI) approach sixteen
53 young adults walked on a treadmill with high density EEG and 3D motion capture tracked
54 kinematics of the head and feet. Visual load was manipulated with the presentation of optic flow
55 with and without mediolateral perturbations, and the effects of cognitive load were assessed by
56 the performance of a Go/No-Go task on half of the blocks. During increased sensory load,
57 participants walked with shorter and wider strides, which may indicate a more cautious pattern
58 of gait. Interestingly, cognitive task engagement attenuated these effects of sensory load on
59 gait. Using an Independent Component Analysis (ICA) and dipole-fitting approach, neuro-
60 oscillatory activity was evaluated from source-localized cortical clusters. Significant modulations
61 in spectral power in the theta (3-7Hz), alpha/mu (8-12Hz), beta (13-30Hz), and gamma (31-
62 45Hz) frequency bands were observed over occipital, parietal and frontal source clusters, as a
63 function of optic flow and task load. These findings provide insight into the neural correlates of
64 gait adaptation, and may be particularly relevant to older adults who are less able to adjust to
65 ongoing cognitive and sensory demands while walking.

66

67

68 **Keywords:** EEG, Mobile Brain/Body Imaging (MoBI), dual-task design, Independent
69 Component Analysis (ICA), power spectral density

70

71 INTRODUCTION

72 Although we typically take walking for granted, the neural systems that regulate it
73 perform many complex functions (Rossignol, Dubuc, & Gossard, 2006). Human locomotion
74 involves the combination of descending pathways from the brainstem to the spinal cord
75 (Duysens & Van de Crommert, 1998), the involvement of the cerebellum and basal ganglia to
76 modulate posture and balance (Grillner, Wallen, Saitoh, Kozlov, & Robertson, 2008) and more
77 recent evidence points to the direct involvement of cortical motor areas in the control of leg
78 muscle activity (Artoni et al., 2017; T. H. Petersen, Willerslev-Olsen, Conway, & Nielsen, 2012).
79 Moreover, when navigating new or unpredictable environments, cortical resources must be
80 recruited to integrate numerous sources of sensory information including visual, vestibular,
81 somatosensory and proprioceptive inputs (Jacobs & Horak, 2007; Varraine, Bonnard, &
82 Pailhous, 2002), or to attend to cognitively demanding secondary tasks (De Sanctis, Butler,
83 Malcolm, & Foxe, 2014). Behavioral studies have previously explored adaptations in gait in
84 response to various manipulations in sensory and cognitive load, however much remains
85 unknown about the cortical underpinnings of sensorimotor mechanisms during locomotion.
86 Here, the aim was to assess the effects of both increased environmental (deployed with optic
87 flow stimuli and visual perturbations) and cognitive load (either engaging in performing a
88 cognitive task or not) on gait and neuro-oscillatory activity.

89 Vision provides a fundamental source of information for the control of goal-directed
90 movements (Lappe, Bremmer, & van den Berg, 1999; W. H. Warren, Jr., Kay, Zosh, Duchon, &
91 Sahuc, 2001). Optic flow, the visual motion we experience as a result of traveling through our
92 environment (Lappe & Grigo, 1999) is a powerful signal that can be used to control the
93 parameters of our movements. Prior studies have introduced perturbations and experimentally
94 manipulated visual inputs to better understand how changes in the visual environment
95 contribute to locomotion. For example, changing the speed of optic flow causes people to
96 modulate their walking speed accordingly (Prokop, Schubert, & Berger, 1997) and the direction

97 of optic flow influences heading direction (Butler, Smith, Campos, & Bulthoff, 2010; Lappe et al.,
98 1999; W. H. Warren & Hannon, 1988). Other studies have employed sinusoidally oscillating
99 visual scenes and observed anisotropic changes in gait parameters, i.e. measures such as step
100 width variability increased in magnitude in accordance with the degree and direction of visual
101 perturbations (O'Connor & Kuo, 2009). Walking in a virtual reality (VR) environment matched to
102 the speed of the treadmill, Hollman et al. (2006) found that young adults took shorter strides and
103 wider steps, with increased variability in stride velocity and step width, compared to walking in a
104 non-VR (visually static) environment, modulations they characterize as reflecting gait instability
105 (Hollman, Brey, Robb, Bang, & Kaufman, 2006). Furthermore, likely due to the fact that humans
106 are more unstable in the ML direction during locomotion (Bauby & Kuo 2000, Donelan 2004,
107 Dean 2007, O'Connor & Kuo 2009) previous studies have noted an increased sensitivity to
108 environmental perturbations in the mediolateral (ML) direction, opposed to those in the anterior-
109 posterior (AP) direction. For example, ML oscillations introduced into a VR environment in the
110 form of continuous but unpredictable (pseudo-random) perturbations resulted in participants
111 taking shorter and wider steps with increased gait variability compared to the no-perturbation
112 condition, but this effect was not observed with perturbations in the AP direction (McAndrew,
113 Dingwell, & Wilken, 2010; McAndrew, Wilken, & Dingwell, 2011). These gait adaptations were
114 interpreted as assuming a more cautious pattern of walking.

115 When navigating complex real-world environments, another challenge to walking
116 behavior is the accommodation of an attentionally demanding secondary task, such as following
117 directions or responding to a text message. Dual-task walking situations have been widely
118 studied in various laboratory settings and with different populations, for reviews see (Al-Yahya
119 et al., 2011; Woollacott & Shumway-Cook, 2002). Depending on the type and complexity of the
120 secondary cognitive task, young adults have typically shown none or minimal costs in the
121 maintenance of postural control (Woollacott & Shumway-Cook, 2002) and gait stability.
122 However some reported changes in young adults' walking as a result of dual-tasking have

123 included reduced gait velocity (Hollman, Kovash, Kubik, & Linbo, 2007; Springer et al., 2006),
124 reduced cadence and stride length, increased stride time and stride time variability (Al-Yahya et
125 al., 2011) and decreased step width variability (Grabiner & Troy, 2005). These modulations
126 have been interpreted as an adoption of a more conservative gait pattern, and may be
127 explained in terms of a capacity sharing model of attentional trade-off (Pashler, 1994; Tombu &
128 Jolicoeur, 2003) in which cortical resources recruited to maintain steady gait become taxed
129 under conditions of increased cognitive load (O'Shea, Morris, & Iansek, 2002). Additional
130 support for this model comes from findings in older adults who often exhibit greater costs in the
131 form of gait instability, when engaged in a cognitive task (Yogev-Seligmann, Hausdorff, & Giladi,
132 2008). Finally, in nondemented older adults, declines in executive function were correlated with
133 decreased walking speed (Ble et al., 2005) and increased gait variability (Springer et al., 2006).
134 These studies provide indirect evidence of the presence of shared cortical resources for
135 cognitively demanding tasks and the maintenance of gait stability.

136 Research using functional magnetic resonance imaging (fMRI) and motor imagery
137 (Bakker et al., 2008), as well as functional near-infrared spectroscopy (fNIRS) (Harada, Miyai,
138 Suzuki, & Kubota, 2009; Miyai et al., 2001) and transcranial magnetic stimulation (TMS) (N. T.
139 Petersen et al., 2001; N. T. Petersen, Pyndt, & Nielsen, 2003), have provided additional
140 evidence of cortical involvement in walking behavior. But due to the slow temporal resolution of
141 hemodynamic measures, EEG, with its portability, relative low cost and excellent temporal
142 resolution, has emerged as the method of choice to assess electrocortical functioning during
143 active movements. When utilized in concert with body motion tracking this approach is referred
144 to as Mobile Brain/Body Imaging (MoBI) (Gramann et al., 2011; Gramann, Jung, Ferris, Lin, &
145 Makeig, 2014; Makeig, Gramann, Jung, Sejnowski, & Poizner, 2009). Recent MoBI studies from
146 our group and others have established the viability (De Sanctis, Butler, Green, Snyder, & Foxe,
147 2012; Gramann, Gwin, Bigdely-Shamlo, Ferris, & Makeig, 2010) and long-term reliability
148 (Malcolm et al., 2017) of recording event-related potentials (ERPs) reflecting cognitive

149 processes during treadmill and outdoor (De Vos, Gandras, & Debener, 2014) walking.
150 Additional studies have employed the MoBI approach to measure differences in electrocortical
151 activity and gait at varying levels of walking speed (De Sanctis et al., 2014; Kline, Poggensee, &
152 Ferris, 2014), and have assessed age-related differences in behavior, gait and ERPs during
153 dual-task walking (Malcolm, Foxe, Butler, & De Sanctis, 2015).

154 Other MoBI studies have provided evidence of suppression of oscillatory activity over
155 motor cortex during walking (Cevallos et al., 2015; Presacco, Goodman, Forrester, & Contreras-
156 Vidal, 2011) compared to standing, signifying increased activations during motion compared to
157 at rest (Wagner et al., 2012), in agreement with prior research showing that efficient motor
158 actions are associated with alpha (8-12Hz) and beta band (13-35Hz) rhythms, such as local
159 field desynchronizations over sensorimotor and parietal cortex (Pfurtscheller, 2000;
160 Pfurtscheller, Gramann, Huggins, Levine, & Schuh, 2003). Recording EEG activity during
161 locomotion, researchers have also begun to investigate the cortical dynamics associated with
162 different phases of the gait cycle during steady-state walking (Gwin, Gramann, Makeig, & Ferris,
163 2011; T. H. Petersen et al., 2012; Severens, Nienhuis, Desain, & Duysens, 2012), walking in
164 synchrony with pacing tones (Wagner, Makeig, Gola, Neuper, & Muller-Putz, 2016) and when
165 experiencing a loss of balance (Sipp, Gwin, Makeig, & Ferris, 2013). In an effort to explore the
166 neurophysiological correlates of active vs. passive locomotion, Wagner et al. (2012) recorded
167 EEG while participants walked in a Lokomat machine for robotic-assisted walking rehabilitation
168 (Wagner et al., 2012). Spectral patterns over sensorimotor cortical areas revealed significant
169 modulations in mu (8-12Hz), beta (18-21Hz) and gamma (25-40Hz) band frequencies, as well
170 as activity that may represent the transition from stance to swing phase of the gait cycle.

171 A major issue in obtaining informative event-related cortical activity during walking is the
172 presence of movement-related artifacts. While gait-specific activity has been identified in low
173 EEG frequencies <10Hz (Castermans, Duvinage, Cheron, & Dutoit, 2014; Gwin, Gramann,
174 Makeig, & Ferris, 2010; Kline, Huang, Snyder, & Ferris, 2015; Presacco, Forrester, & Contreras-

175 Vidal, 2012), several different groups have investigated different approaches that may be
176 employed to successfully isolate and remove head movement and gait artifacts from cortical
177 signals (Gwin et al., 2010; Kline et al., 2015; Nathan & Contreras-Vidal, 2015; Severens et al.,
178 2012; K. L. Snyder, Kline, Huang, & Ferris, 2015). One approach involves using independent
179 components analysis (ICA), already shown to be effective at separating eye and muscle-related
180 noise from EEG signals obtained during seated tasks (Delorme, Sejnowski, & Makeig, 2007;
181 Jung et al., 2000), combined with dipole fitting procedures that model independent components
182 (ICs) as equivalent current dipoles (Oostenveld & Oostendorp, 2002), to accurately localize the
183 resulting neural sources. Snyder et al. (2015) recently tested this tactic by recording EEG over a
184 silicone swim cap, thus blocking all real electrophysiological activity, and demonstrated that ICA
185 and dipole fitting procedures accurately isolated 99% of non-neural sources by location (outside
186 of the brain) or by a lack of dipolar characteristics (K. L. Snyder et al., 2015).

187 Here, we employed a MoBI approach with young adult participants to evaluate the
188 effects of environmental load, in the form of three different visual conditions (consistent optic
189 flow, optic flow with visual perturbations and static) as well as cognitive load, on gait and
190 electrocortical dynamics. Spatiotemporal measures of gait and variability in head movement
191 were captured with kinematics recordings. Spectral power was obtained from high-density EEG
192 using an ICA and dipole fitting procedure. Independent Component cluster models were then
193 used to identify modulations in average spectral activity across participants as a result of optic
194 flow and cognitive task load. We hypothesized that increased load during walking would lead to
195 a more conservative and more variable pattern of gait. Furthermore, previous literature has
196 reported increased cortical excitability exhibited in the form of power reductions, or
197 desynchronization, in the alpha frequency band over occipital regions during visual processing
198 (Pfurtscheller & Lopes da Silva, 1999), as well as in the mu and beta bands before and during
199 movements (Pfurtscheller & Klimesch, 1991; Wagner et al., 2012). In line with these findings,
200 we predicted that increased sensory load (optic flow vs. static) and cognitive load (processing

201 letters vs. not processing letters) would lead to a decrease in alpha power over occipital regions,
202 and decreased mu and beta power over sensorimotor cortex. Finally, based on literature linking
203 increased alpha power over parietal regions to attentional mechanisms used to suppress task-
204 irrelevant information (Foxye & Snyder, 2011), we predicted that sensory load, particularly
205 unreliable visual scene motion in the form of mediolateral perturbations, would result in
206 increased alpha power over parietal cortex.

207

208

209 **METHODS**

210 *Participants*

211 Eighteen healthy young adults participated in the experiment. Data from two participants
212 were excluded due to technical issues; therefore results reported here were derived from
213 sixteen individuals (five females) with a mean age of 25.6 years (SD = 4.5 years). All individuals
214 reported normal or corrected-to-normal vision and were free from any neurological, psychiatric
215 or locomotor disorders, by self-report. Participants were recruited from the lab's existing subject
216 pool and from flyers posted at the Albert Einstein College of Medicine. The Institutional Review
217 Board of the Albert Einstein College of Medicine approved the experimental procedures and all
218 participants provided their written informed consent. All procedures were compliant with the
219 principles laid out in the Declaration of Helsinki for the responsible conduct of research.

220

221 *Stimuli and procedure*

222 While walking on the treadmill, participants were presented with a full-field visual display
223 consisting of a star field (200 randomly placed white dots projected onto a black background). In
224 the two dynamic optic flow conditions, the stars emanated outward from a central focus of
225 expansion point, either moving steadily with no visual perturbations (NOP) or oscillating with
226 continuous perturbations in the mediolateral direction (MLP). Optic flow was programmed from:

227

228

$$D(t) = A \times \sin(0.4 \times 2\pi t)$$

229

230 Where $D(t)$ was the translation distance (m), A was the amplitude of displacement and t was
231 time (sec). Sinusoidal perturbations in the ML direction were applied at amplitudes of 0 (NOP
232 condition) or 0.12 m (MLP condition). The frequency selected (0.4 Hz) was within the range of
233 those used in previous studies of human gait (McAndrew et al., 2010; O'Connor & Kuo, 2009).
234 The star field moved in this manner throughout the duration of a three-minute walking block. A
235 static condition was also employed in which the same number of stars were randomly presented
236 across the visual field projection but did not move, i.e., no optic flow. Participants were
237 instructed to keep their eyes fixed on a central fixation cross.

238 In addition to these three visual conditions, participants were presented with a Go/No-Go
239 response inhibition task. Stimuli consisting of letters were shown in the center of the visual field,
240 not interfering with the optic flow. During 'Task' blocks, participants were instructed to engage in
241 the cognitive task by responding quickly and accurately to the frequently-occurring Go trials by
242 clicking a wireless mouse button following the presentation of the letter 'O', while withholding
243 responses during infrequent No-Go trials, designated by the presentation of the letter 'X.' The
244 probability of Go and No-Go trials was 0.80 and 0.20, respectively. The duration of each letter
245 was 400ms with a random stimulus-onset-asynchrony (SOA) ranging from 600-800ms.
246 Response inhibition performance was assessed by a participant's percentage of Correct
247 Rejection (CR) trials, defined as when a response was correctly withheld following a No-Go
248 stimulus. In order to evaluate the effect of cognitive task load on gait and EEG spectral power,
249 half of the experiment was performed as 'No-Task' blocks, in which the Go/No-Go stimuli were
250 shown but participants were instructed not to respond to the task, or to cognitively engage in the
251 task. Images were projected centrally (InFocus XS1 DLP, 1024 x 768 pixel) onto a black wall

252 approximately 1.5m in front of the participant. The stimulus display was programmed with
253 Presentation software version 18.1 (Neurobehavioral Systems, Berkeley, CA).

254 With the two factors of visual condition (static, optic flow with no perturbation, optic flow
255 with mediolateral perturbation) and cognitive task (task performance or no-task performance),
256 this design resulted in a total of six different experimental conditions. Each participant performed
257 three blocks of each condition, resulting in a total of 18 blocks, each lasting three minutes. All
258 conditions were conducted in a pseudo-random order, counterbalanced across participants, and
259 a practice block was performed before undertaking the main experiment. Several rest breaks
260 were provided in between blocks. Participants self-selected their walking speed at the beginning
261 of the experiment and maintained that speed throughout. Average walking speed was 3.9 km/hr
262 (range: 3.2–4.5 km/hr). All subjects walked while wearing comfortable shoes and a safety
263 harness. See Figure 1 for a representation of the recording set-up. No specific task prioritization
264 instructions (i.e., walking versus cognitive task) were given, other than for participants to direct
265 their gaze towards the central fixation cross (and presentation of task-relevant letters) during no-
266 task as well as task blocks.

267
268 -----
269 Insert Figure 1 Here
270 -----

271
272 *Kinematics recording*

273 Three-dimensional kinematic data were collected at 100Hz using a 9-camera Optitrack
274 infrared motion capture system and Arena v.1.5 acquisition software (Natural Point). Each
275 participant wore 10 reflective markers: four were placed on the head (attached to the EEG cap,
276 right and left sides, front and back), and three markers were placed on each foot. These were
277 placed over the participants' shoes, on the calcanei, the second and the fifth distal metatarsals.

278

279 *Electrophysiological recording*

280 Continuous EEG was recorded with a 72-channel BioSemi ActiveTwo system (digitized
281 at 512Hz; 0.05 to 100 Hz pass-band, 24 dB/octave). Stimuli from Presentation software were
282 transmitted to BioSemi ActiView via a parallel cable. Time-synchronized acquisition of stimulus
283 triggers, behavioral responses, EEG and rigid body motion tracking was conducted with Lab
284 Streaming Layer software (Swartz Center for Computational Neuroscience, UCSD, available at:
285 <https://github.com/sccn/labstreaminglayer>).

286

287 **Data Analysis**

288 All EEG and kinematic data analyses were performed using custom MATLAB scripts
289 (MathWorks, Natick, MA) and EEGLAB (Delorme & Makeig, 2004).

290

291 *Kinematics*

292 Heel strikes were computed from the heel marker trajectory, using an automated peak-
293 picking function (MATLAB custom scripts) and confirmed by manual inspection, to identify the
294 point where the heel marker was at the most anterior point in the anterior-posterior direction
295 (Dingwell, John, & Cusumano, 2010; Zeni, Richards, & Higginson, 2008). Individual strides were
296 defined as consecutive heel strikes of the same foot. Responses to visual optic flow and
297 cognitive task load on the gait cycle were assessed by three dependent measures. Stride time
298 (ST) was defined as the time between consecutive heel strikes of the same foot, while Stride
299 length (SL) was calculated as the sum of each pair of consecutive step lengths that made up
300 each stride (Alton, Baldey, Caplan, & Morrissey, 1998; Dingwell & Cusumano, 2015). Step width
301 (SW) was computed as the lateral distance between the two heel markers at the time of right
302 heel strike (Kang & Dingwell, 2008; Kline et al., 2014; Owings & Grabiner, 2004). The means
303 and standard deviations of each of these measures were calculated for each block of each
304 condition, for each participant. Finally, the mean SD of the head markers in the mediolateral and

305 anterior-posterior directions was used as a measure of postural stability and overall variability in
306 movement position on the treadmill. The SD was calculated for each block separately, and then
307 averaged over conditions, then subjects.

308

309 *EEG and power spectral density*

310 EEG data were first high-pass filtered at 1Hz using a zero phase FIR filter (order 5632)
311 (Winkler, Debener, Muller, & Tangermann, 2015). Then all blocks for each subject were
312 concatenated into one dataset. Noisy channels were identified and removed by visual inspection
313 and by automatic detection of channels with signals more than five times the standard deviation
314 of the mean across all channels. The remaining channels were re-referenced to a common
315 average reference. Continuous data were then subjected to a manual visual inspection resulting
316 in the rejection of any sequences that contained large or non-stereotypical artifacts. An
317 extended Independent Components Analysis (ICA) decomposition was performed on the
318 remaining data using default training mode parameters (Makeig, Bell, Jung, & Sejnowski, 1996).
319 ICA separates various generators of task-evoked cortical activity to help distinguish and
320 separate from artifactual sources such as electrical noise, eye blinks, neck muscles and
321 walking-related artifacts such as cable sway (Jung et al., 2000).

322 The resulting Independent Components (ICs) were then coregistered with a standard
323 MNI (Montreal Neurological Institute) boundary element head model and fit with single
324 equivalent current dipole models using the DIPFIT toolbox in EEGLAB (Delorme, Palmer,
325 Onton, Oostenveld, & Makeig, 2012; Oostenveld & Oostendorp, 2002). Only ICs for which the
326 estimated dipole model was located within the brain and explained > 85% of the variance of the
327 IC scalp map were retained (Gwin et al., 2011). These were then examined and any that were
328 clearly artifactual were rejected; these could have included activity originating from eye blinks,
329 bad electrodes and muscle noise. Rejection criteria were based on topography, spectra,
330 component activation time course, and dipole location (Jung et al., 2000). Following this

331 procedure there were an average of 10 brain related ICs per participant (ranging from 6 to 16
332 ICs) for use in further analyses. Presumably, these ICs reflect activity generated in cortical
333 sources close to the location of their equivalent dipole model (Akalin Acar & Makeig, 2013).
334 Remaining ICs were then clustered across participants with EEGLAB clustering routines using
335 the parameters of 3-D dipole location, scalp topography and power spectra (3-45Hz) (Onton &
336 Makeig, 2006). Using principal components analysis, these feature vectors were reduced to 10
337 principal components and clustered using a k-means algorithm implemented in EEGLAB. *K*-
338 means is a well-known clustering algorithm that requires no prior information about the
339 associations of data points with clusters. ICs that were further than three standard deviations
340 from any of the resulting cluster centers were identified as outliers. Finally, only clusters that
341 included ICs from at least half of the participants were retained, resulting in the eight clusters
342 reported below.

343 For the spectral analysis, we chose to look at the neural oscillatory pattern resulting from
344 component activations, in comparison to the data from separate channels, since independent
345 components may help to explain the activity underlying a specific cognitive function. whereas
346 channel activations are the result of summed potentials volume-conducted from different parts
347 of the brain (Onton, Westerfield, Townsend, & Makeig, 2006). Even though EEG does not have
348 the spatial resolution of fMRI, this technique has been shown to provide a spatial resolution of
349 around a few centimeters (Mullen, Acar, Worrell, & Makeig, 2011). Power spectral density
350 (PSD) was computed using Welch's method, separately for each IC and for each of the six
351 experimental conditions. Periodograms were obtained in windows of 512 samples (1 sec), an fft
352 length of 1024, with 50% overlap, and windowed with a Hamming window of the same length as
353 the segment. Similar parameters were used to calculate spectra for ICs in a previous MoBI
354 study (K. L. Snyder et al., 2015). The resulting periodograms were averaged over the ICs in
355 each cluster to produce an estimation of the absolute PSD for four frequency bands of interest:
356 theta (3-7Hz), alpha (8-12Hz), beta (13-30Hz) and gamma (31-45Hz).

357

358 *Statistical analyses*

359 Cognitive task performance was analyzed with a one-way repeated-measure ANOVA,
360 with the factor of visual load (static, no perturbation optic flow and ML optic flow). Gait and
361 posture data were analyzed with 2 (Task Load) x 3 (Visual Load) repeated measures ANOVAs.
362 Because walking speed has a direct relationship with stride length and stride time (Dingwell et
363 al., 2010; Kang & Dingwell, 2008) walking speed was included as a covariate in the analysis of
364 these gait parameters. The covariate was mean-centered, i.e., deviations from the mean speed
365 were used instead of the raw values, to avoid interfering with the test of the main effects
366 (Delaney & Maxwell, 1981). For the analysis of power spectral density (PSD), separate two-
367 factor (task load, visual condition) repeated-measures ANOVAs were performed for each IC
368 cluster and frequency band of interest. Greenhouse-Geisser corrections were applied when
369 appropriate, but original degrees of freedom have been reported. All statistical analyses were
370 performed using IBM SPSS (v. 24).

371

372

373 **RESULTS**

374 *Cognitive Task Performance*

375 Figure 2 shows the percentage of Correct Rejections (CRs) for each visual condition
376 (static, no perturbation optic flow and optic flow with ML perturbations). No differences were
377 found for response inhibition performance as a function of the visual condition employed, $F_{2, 30} =$
378 0.27 , $p = .76$, indicating that participants were able to perform the Go/No-Go task equally well
379 regardless of the dynamic state of the star field.

380

381

382

Insert Figure 2 Here

383

384

385 *Gait and Posture*

386 Average and mean SD of stride time, stride length and step width for all six conditions
387 are presented in Figure 3.

388 **Stride Time:** For the parameter of average stride time, there was a main effect of task load, $F_{1, 14} = 8.51$, $p = .01$, and an interaction between task load and visual condition, $F_{2, 28} = 3.99$, $p =$
389 $.03$. Follow-up paired comparisons showed that, averaged over all visual conditions, participants
390 took significantly faster strides when engaged in the task (Mean = 1179ms, SD = 75) compared
391 to the no-task blocks (Mean = 1189ms, SD = 78), $t_{15} = 3.02$, $p = .009$. Furthermore, for the no-
392 task conditions, participants exhibited increasingly faster strides with increasing levels of visual
393 load. The slowest strides were observed for the static no-task condition, closely followed by the
394 no perturbation optic flow condition. On the other hand, during the task blocks, there was
395 minimal difference in average stride time according to visual stimuli. Averaging across task
396 conditions revealed significantly longer strides for the no-perturbation optic flow condition in
397 comparison to the presentation of ML perturbations ($p = .02$). For the measure of average stride
398 time variability, no significant effects were found.

400 **Stride Length:** There was a main effect of visual condition on average stride length, $F_{2, 28} =$
401 3.59 , $p = .04$, as well as a significant effect of task load, $F_{1, 14} = 11.85$, $p = .004$, and an
402 interaction was observed between these two factors, $F_{2, 28} = 5.43$, $p = .01$. In line with the
403 findings outlined above for Stride Time, participants took significantly shorter strides when
404 engaged in the task (Mean = 1423mm, SD = 115) compared to not performing the task (Mean =
405 1438mm, SD = 122), $t_{15} = 3.50$, $p = .003$. The effect of visual condition on average stride length
406 exhibited the greatest difference between the static star field condition in which participants took
407 overall longer strides, compared to the ML visual perturbations ($p = .04$). This effect also
408 appeared to be most prominent for the no-task blocks, as stride length progressively decreased

409 with the dynamic optic flow and even more so as perturbations were applied to the star field. For
410 stride length variability no effects reached the level of significance, though interestingly, strides
411 tended to be more variable when participants observed the dynamic optic flow and were not
412 engaged in the cognitive task.

413 **Step Width:** There was a significant effect of the visual condition on average step width, $F_{2, 28} =$
414 7.14, $p = .003$, reflecting the fact that compared to the static visual condition, participants
415 walked with wider steps during the no perturbation optic flow blocks ($p = .002$) as well as with
416 ML perturbations ($p = .02$), regardless of whether they performed the cognitive task. Average
417 step width variability exhibited a robust effect of task load, $F_{1, 14} = 11.77$, $p = .004$, with more
418 variable step widths across all visual conditions when not performing the cognitive task (Mean =
419 16.2mm, SD = 4.8), compared to during task blocks (Mean = 14.5mm, SD = 3.7).

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421 -----
422 Insert Figure 3 Here
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424
425 **Mean SD of head position:** Figure 4 shows the mean SD of head position in the mediolateral
426 (ML) direction (left) and anterior-posterior (AP) direction (right). For head position variability in
427 the ML direction, there was a main effect of task load, $F_{1, 15} = 8.56$, $p = .01$, indicating decreased
428 variability in head position in the lateral direction when performing the cognitive task (Mean =
429 26.7mm, SD = 9.2) in contrast to walking without engaging in the task (Mean = 29.6mm, SD =
430 10.8).

431 For the average variability in head position in the AP direction, there was also a main
432 effect of cognitive task load, $F_{1, 15} = 10.12$, $p = .006$, as well as an interaction between cognitive
433 load and visual condition, $F_{2, 30} = 7.33$, $p = .003$. This effect was indicative of increased
434 variability on the no-task blocks (Mean = 44.8mm, SD = 19.2) compared to performing the task

435 (Mean = 33.5mm, SD = 18.0), and while the different task blocks were shown to maintain a
436 similar level of variability, the no-task conditions showed a decrease in variability from the static
437 visual condition, to the no perturbation optic flow, and then even more so with the introduction of
438 ML perturbations.

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Insert Figure 4 Here

443

444 *Power spectral density*

445 Table 1 lists the specifics (number of ICs and subjects included in each cluster and the
446 approximate anatomical location (Brodmann area and Talairach coordinates) of cluster
447 centroids) of the eight clusters that were localized to cortical areas and composed of ICs from at
448 least half of the participants. Figure 5 shows the clusters of electrocortical sources localized to
449 occipital, parietal and frontal cortical areas.

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Insert Table 1 Here

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Insert Figure 5 Here

458

459 Three clusters were located over occipital cortex. Scalp topography, dipole location (blue
460 dots indicate the location of each IC, red dots represent the cluster centroid) and average power
461 spectral density (PSD) for these clusters are presented in Figure 6. For the cluster located over
462 medial occipital cortex, no significant modulations were found in the PSD of any frequency

463 bands according to task load or visual condition. In contrast, for the right occipital cluster, robust
464 differences in spectral power attributable to the presentation of the three different visual
465 conditions were found in theta, $F_{2, 22} = 8.94$, $p = .008$ and alpha, $F_{2, 22} = 21.50$, $p < .001$,
466 frequencies, with a smaller effect observed in the beta range, $F_{2, 22} = 4.18$, $p = .05$. The same
467 pattern emerged for the lower frequencies (theta and alpha) in that there was on average higher
468 spectral power for the static conditions compared to both dynamic optic flow conditions (p 's <
469 .05), but no difference between the two optic flow conditions (p 's > .50). For the beta range,
470 increased power was observed during the static conditions compared to the no perturbation
471 optic flow ($p = .006$), but no significant differences were found between the static and ML optic
472 flow or between the two dynamic conditions (p 's > .10). Furthermore, there was a strong effect
473 of task load on alpha spectral power, $F_{1, 11} = 14.15$, $p = .003$, with higher power over all three
474 no-task conditions compared to when participants performed the cognitive task. Finally, a
475 significant interaction between task load and visual condition was found for gamma band power,
476 $F_{2, 22} = 4.96$, $p = .02$, indicating that while spectral power remained relatively consistent across
477 the visual conditions when participants engaged in the cognitive task, when they did not perform
478 the task gamma power remained high for the static visual condition but decreased greatly during
479 the no perturbation optic flow blocks, and decreased to a lesser extent with ML perturbations in
480 optic flow.

481 For the IC cluster located in left occipital cortex, modulations in spectral power were only
482 observed in the alpha frequency range. There was a robust effect of visual condition, $F_{2, 26} =$
483 10.76 , $p = .004$, with higher alpha power observed for the static visual condition compared to
484 both dynamic conditions (p 's = .005) but no difference apparent between the two dynamic
485 conditions ($p = .88$). There was also a trend towards higher alpha power on no-task blocks, $F_{1, 13}$
486 $= 4.52$, $p = .053$, compared to blocks when participants engaged in the cognitive task.

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489 Insert Figure 6 Here

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491

492 Figure 7 shows scalp topography, dipole location and power spectra for one cluster over
493 left temporal cortex and two clusters located over right parietal cortex. For the cluster localized
494 to the left superior temporal gyrus, the different visual conditions had a significant effect on both
495 theta band ($F_{2, 22} = 7.00, p = .02$) and alpha band ($F_{2, 22} = 9.65, p = .001$) spectral power. Both
496 frequency bands showed significantly greater power during the static visual blocks compared to
497 both dynamic optic flow conditions (p 's < .05), with no differences between the two dynamic
498 conditions (p 's > .05). Additionally, no differences were apparent at higher frequencies. For the
499 cluster localized to the right inferior parietal lobule, the only significant difference in spectral
500 power was observed for the factor of visual condition in the alpha frequency range, $F_{2, 18} = 5.94,$
501 $p = .01$. Again, regardless of task load, there was higher alpha power during the static star field
502 blocks in comparison to both of the optic flow conditions (p 's < .02), but there was no difference
503 in power between the two dynamic star field displays ($p = .66$).

504 Significant modulations in spectral power as a result of visual condition were observed in
505 the cluster of ICs located over medial parietal cortex, localized to the precuneus. This effect
506 occurred across all frequency bands of interest: theta ($F_{2, 28} = 20.09, p < .001$), alpha ($F_{2, 28} =$
507 $15.63, p = .001$), beta ($F_{2, 28} = 10.33, p < .001$), and gamma ($F_{2, 28} = 3.66, p = .04$). For
508 frequencies in the theta, alpha and beta bands, significantly greater power was observed for the
509 static condition compared to both the no perturbation optic flow (p 's < .01), and the ML
510 perturbation condition (p 's < .01), but there was no difference between the two optic flow
511 conditions (p 's > .05). In the gamma range only a significant difference between static and ML
512 perturbations was observed ($p = .05$) but there was no difference between static and no
513 perturbation ($p = .22$) or between the two optic flow conditions ($p = .24$). For the alpha and beta
514 frequency bands, differences were also observed in spectral power linked to cognitive task

515 engagement: alpha ($F_{1, 14} = 21.07, p < .001$), beta ($F_{1, 14} = 13.16, p = .003$). For both, overall
516 higher power was found for the no-task blocks compared to when participants performed the
517 task. Finally, for frequencies in the alpha range there was an interaction between task load and
518 visual condition, $F_{2, 28} = 5.94, p = .007$, indicating that while either performing the cognitive task
519 or not, there was a desynchronization in alpha power between the static visual condition to the
520 no-perturbation optic flow condition, whereas a different result was observed with the
521 introduction of ML perturbations. When performing the task, average power continued to
522 decrease when perturbations were introduced into the optic flow, but when not engaged in the
523 task, alpha power actually increased with the ML perturbations.

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526 Insert Figure 7 Here
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528
529 The final two clusters located over frontal cortical areas including the supplementary
530 motor area and the anterior cingulate are depicted in Figure 8. For the cluster located over
531 supplementary motor area, significant changes in spectral power were found in the theta range
532 linked to task performance, $F_{1, 22} = 9.41, p = .006$, representing the effect that average spectral
533 power was higher when subjects performed the cognitive task compared to when they did not.
534 Significant effects of the visual condition on spectral power were observed in the theta ($F_{2, 44} =$
535 $5.01, p = .02$), alpha ($F_{2, 44} = 29.52, p < .001$) and beta ($F_{2, 44} = 17.20, p < .001$) bands. For alpha
536 and beta this was reflected in the fact that higher spectral power was observed for the static
537 condition compared to both no perturbations (p 's $< .001$) and ML perturbations (p 's $< .001$),
538 whereas no difference occurred between the two dynamic conditions (p 's $> .05$). For the theta
539 range, spectral power during the ML perturbation conditions were significantly lower compared
540 to presentation of the static star field ($p = .006$), as well as the no perturbation optic flow ($p =$

541 .04), while there was no difference between the static and no perturbation conditions ($p = .19$).
542 Additionally, for frequencies in the alpha range, there was a significant interaction between task
543 load and visual condition, $F_{2, 44} = 5.77$, $p = .01$, indicating that the average spectral power
544 remained approximately the same between task conditions for both the static and no
545 perturbation star field displays, however with ML visual perturbations spectral power increased
546 during no-task blocks but decreased with task engagement.

547 Lastly, for the IC cluster located to anterior cingulate cortex, significant changes in
548 spectral power were observed only in lower frequencies. The visual conditions significantly
549 affected spectral power in both the theta ($F_{2, 28} = 10.16$, $p < .001$) and alpha ($F_{2, 28} = 10.13$, $p <$
550 $.001$) frequency ranges. This effect was indicative of greater power for the static visual condition
551 compared to both the no perturbation optic flow (p 's $< .005$) and the ML perturbation optic flow
552 (p 's $< .005$), but no difference was apparent between the two dynamic conditions (p 's $> .10$).
553 Additionally, in the theta range, spectral power significantly increased during performance of the
554 cognitive task in comparison to no-task blocks, across all three visual conditions, $F_{1, 14} = 11.61$,
555 $p = .004$.

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561

562 **DISCUSSION**

563 *Effects of optic flow and cognitive load on gait*

564 The objective of the current experiment was to examine changes in gait and cortical
565 network activity in response to the presence of optic flow stimuli, as well as the engagement or
566 lack of engagement in a cognitive task. During two dynamic visual conditions a pattern of optic

567 flow created by the movement of a star field radiating outwards generated a sense of forward
568 movement. The optic flow either moved steadily (no perturbations) or oscillated with continuous
569 mediolateral (ML) perturbations. Sensory load, presented here in the form of optic flow, did not
570 result in decrements in task performance, i.e., there were no costs in the behavioral domain as
571 a result of the different visual conditions. Conversely, both sensory and cognitive load had
572 significant effects in the motor domain. Participants took shorter strides as cognitive and
573 sensory load increased. Average step width also increased with visual load, with wider steps
574 during both optic flow conditions compared to the static star field. However, the interactions
575 observed between cognitive and sensory load for the measures of stride time/length and head
576 position in the AP direction reveal that optic flow modulates gait more so when participants are
577 disengaged from the cognitive task. For example, participants made faster and shorter strides
578 during both optic flow conditions compared to the static condition, when they were not engaged
579 in the task. Also, during no-task blocks, average head position variability in the anterior-posterior
580 (AP) direction decreased as the amount of visual load increased, i.e., in the presence of optic
581 flow and even more so with the introduction of ML perturbations. These findings likely indicate
582 the engagement of a more conservative pattern of gait with increased load: shorter and wider
583 steps, and the maintenance of a more consistent position along the length of the treadmill. This
584 may be indicative of increased allocation of sensorimotor resources in order to accommodate
585 potentially destabilizing sensory load.

586 The primary effect of increased cognitive load on gait appeared to be a reduction in
587 walking variability. When performing the inhibitory control task, participants exhibited decreased
588 variability in step width and head position in both directions. These findings suggest that
589 cognitive task engagement actually led to a more consistent pattern of motor behavior.
590 Participants adopted a more stereotyped manner of walking, with less stride-to-stride
591 fluctuations when attention was bound to the Go/No-Go task. Prior dual-task walking (DTW)
592 findings seem to go along with these results, as Grabiner & Troy (2005) also observed

593 decreased step width variability and more conservative gait under cognitive load (Grabiner &
594 Troy, 2005). Additionally, Lovden et al. (2008) observed that when young adults performed a
595 moderately difficult cognitive task, gait variability decreased (Lovden, Schaefer, Pohlmeier, &
596 Lindenberger, 2008). They contend that an external focus of attention is beneficial to motor
597 performance, and that there will be no cross-domain competition, resulting in costs, as long as
598 cognitive load remains moderate (Lovden et al., 2008).

599 Interestingly, when the ML perturbations were applied to the optic flow, participants often
600 did not exhibit increased movement in that direction, in relation to the no-perturbation optic flow.
601 This was somewhat surprising considering that other studies have observed effects such as
602 changes in posture and increased variability in gait and dynamic stability (McAndrew et al.,
603 2011) in response to ML visual perturbations. One possible reason that this may have occurred
604 is that participants in this study walked in a safety harness and wore an EEG cap, with
605 electrodes tethered to an overhead platform. Another explanation for the relative lack of
606 modulation in body position in the ML direction is that participants may have become
607 accustomed to the perturbations over time and were able to 'entrain' their walking behavior to
608 accommodate them. Because the ML visual oscillations were constant for the duration of each
609 three-minute block, in contrast to some studies that have employed pseudo-random
610 perturbations (McAndrew et al., 2010; McAndrew et al., 2011), participants here may have
611 unconsciously come to predict the effect that oscillations may have had on body position and
612 adjusted their gait accordingly. For example, Brady et al (2009) applied continuous ML
613 perturbations to the treadmill surface and observed that within five minutes people showed
614 adaptation in the form of entrainment and began to time their steps to occur in line with the
615 phase of oscillation applied (Brady, Peters, & Bloomberg, 2009). Also, in a very recent study,
616 young adults quickly adapted to continuous mediolateral optic flow perturbations by taking
617 shorter, wider and more variable steps, until after approximately three minutes step length and
618 width returned to normal (unperturbed) levels, while variability did not (Thompson & Franz,

619 2017). The authors attribute these results to visuomotor adaptation processes – the return of
620 step length and width to normal levels as visual perturbations continued likely reflects a
621 deprioritization of visual inputs while presumably other inputs, such as vestibular and
622 proprioceptive modalities were up-regulated. On the other hand, they point to the sustained
623 increase in variability as indicative of a necessary, reactive step-to-step balance control
624 strategy.

625 There were a few gait parameters in which a significant difference was found between
626 the two optic flow conditions (e.g., shorter strides and less variability in AP head position with
627 added ML perturbations). These findings are consistent with other studies that have employed
628 constant, sinusoidal oscillations and have still observed changes in measures such as step
629 width (O'Connor & Kuo, 2009) and stride length variability. It is possible that when people come
630 to predict the environmental perturbations, they will exhibit consistent changes in gait to better
631 accommodate them. For example, that participants walked with faster and shorter strides and
632 increased step width could be interpreted as a more cautious gait approach in response to
633 sensory load. Furthermore, walking on a treadmill requires the strict regulation of both walking
634 speed and position, but Dingwell (2015) showed that young adults regulated stride-to-stride
635 fluctuations in walking by prioritizing speed maintenance, not their position in the anterior-
636 posterior direction, therefore letting themselves drift to the front and back of the treadmill before
637 correcting (Dingwell & Cusumano, 2015). This notion may explain the current finding of
638 increased influence of visual flow on head position variability only in the AP direction, not the ML
639 direction. When walking without additional cognitive load, participants may have allowed
640 themselves to drift forwards and backwards on the treadmill, especially during the static no-task
641 condition (perhaps the least attentionally demanding).

642

643 *Modulations in power spectral density*

644 Following an Independent Components Analysis (ICA) and dipole-fitting procedure,
645 neuro-oscillatory activity was evaluated from eight source-localized clusters of Independent
646 Components (ICs). To identify modulation in the power content across different frequencies as a
647 function of visual load and cognitive task engagement, power spectral density (PSD) was
648 calculated for each condition in each IC, and then averaged over all ICs in each cluster. These
649 results provide new information about the frequency-related effects of optic flow stimulation and
650 task load on brain activity during locomotion. The locations of the IC clusters reported here were
651 similar to locations cited in other mobile EEG studies (Gwin et al., 2011; Kline, Huang, Snyder,
652 & Ferris, 2016; Sipp et al., 2013; Wagner et al., 2012): three clusters were located over occipital
653 cortical areas, one cluster localized to the left superior temporal gyrus, right inferior parietal
654 lobule, the precuneus in the parietal lobe, and two frontal clusters over supplementary motor
655 area and anterior cingulate cortex. Results indicate a widely distributed cortical network
656 exhibiting task-specific fluctuations in spectral power.

657 Occipital Region

658 Other than the IC cluster over medial occipital cortex where no significant modulations
659 were observed, all other clusters exhibited significant changes in the spectral power of lower
660 frequencies (theta and alpha) linked to visual presentation and optic flow. For the right occipital
661 cluster, average spectral power was reduced in the theta, alpha and beta ranges, with increased
662 sensory load. In the left occipital cluster, this effect was also seen, but only in the alpha range.
663 Additionally, decreased alpha power was also observed in the right occipital cluster as
664 participants processed the Go/No-Go task letters; with a trend towards this effect in the left
665 occipital cluster as well. Alpha band oscillations have long been shown to play an important role
666 in directing attention, for a review see (Foxe & Snyder, 2011). Desynchronization in the alpha
667 band over occipital regions is assumed to reflect cortical excitation related to various stages of
668 stimulus processing (Pfurtscheller, Stancak, & Neuper, 1996), thus the reduction in alpha power

669 when presented with increased visual demands in the form of optic flow. Furthermore, EEG
670 studies of parieto-occipital alpha band activity have revealed a more sophisticated role as a
671 mechanism involved in selectively attending to relevant information in the environment (Fuxe,
672 Simpson, & Ahlfors, 1998; Fuxe & Snyder, 2011; Kelly, Lalor, Reilly, & Fuxe, 2006; A. C. Snyder
673 & Fuxe, 2010; Worden, Fuxe, Wang, & Simpson, 2000). Presumably this paradigm required
674 participants to selectively disengage from processing the distracting optic flow information when
675 they were performing the task, reflected as synchronization in alpha power over cortical regions
676 dedicated to optic flow processing. Subsequently, during the blocks in which they saw the
677 Go/No-Go letters but were instructed to not engage cognitively, an effortful, top-down
678 recruitment strategy would likely be employed in order to ignore the letters, i.e., synchronization
679 in alpha to inhibit processing, but at the same time suppress any potentially destabilizing
680 information from the dynamic star field. Gait results indicate the adoption of a progressively
681 more conservative manner of walking with increased visual input but no task engagement.
682 Consequently, it seems that even though the optic flow lent no meaningful information to
683 walking behavior, on some level participants did pay attention to and process this information.
684 Thus, these findings may indicate a flexible deployment of enhanced alpha band activity to
685 selectively suppress to-be-ignored aspects of this complex environment (Dockree, Kelly, Fuxe,
686 Reilly, & Robertson, 2007; Fuxe & Snyder, 2011; Worden et al., 2000). This pattern of results is
687 consistent with alpha desynchronization not simply due to visual stimulation but being
688 specifically task driven (Kelly et al., 2006; Klimesch, 2012), a theory that goes along with the
689 current finding of a greater desynchronization in alpha power when individuals also engaged in
690 the task.

691 There was also an interaction between cognitive and sensory load in the gamma range
692 (31-45Hz) in the right occipital cluster. Here, gamma power increased while participants were
693 engaged in the Go/No-Go task, then decreased as they disengaged during no-task conditions,
694 though only while exposed to optical flow. Sustained attention requires ongoing activation of

695 task-relevant regions and evidence links gamma in sensory cortices as a mechanism to
696 enhance processing of task-relevant sensory inputs (Clayton, Yeung, & Cohen Kadosh, 2015).
697 Previous studies have also reported enhancement of gamma band activity during visuospatial
698 attention tasks (Siegel, Donner, Oostenveld, Fries, & Engel, 2007) and gamma power has been
699 associated with task complexity (Fitzgibbon, Pope, Mackenzie, Clark, & Willoughby, 2004).
700 However, if sustained gamma power in this region is indeed related to sustained task
701 engagement, it is an open question as to why the static no-task condition maintained a higher
702 average spectral power in relation to the other no-task conditions.

703

704 Left Superior Temporal Gyrus

705 A main effect of visual condition was observed for theta and alpha activity in this region,
706 with reductions in spectral power associated with increased optical flow input. Animal studies
707 have shown that this area is involved in processing optic flow and visual motion information
708 generated from environmental stimuli (Duffy & Wurtz, 1991). Therefore the current findings may
709 indicate increased activation in this region when presented with more computationally
710 demanding visual environments.

711

712 Parietal Region

713 The IC cluster localized to the right inferior parietal lobule showed a significant decrease
714 in alpha spectral power for both dynamic flow conditions compared to the static visual condition.
715 The precuneus cluster exhibited a similar pattern with decreased spectral power in theta, alpha
716 and beta for optic flow relative to static, while gamma power was higher for the static condition
717 relative only to visual ML perturbations. Thus, if we are to assume that participants invest more
718 resources to counteract unreliable proprioceptive information (generated by ML visual
719 perturbations), evidenced by their engagement in a more conservative pattern of gait, higher

720 gamma power during the static condition may be acting to increase reliance on proprioceptive
721 information via enhanced sensory processing (Clayton et al., 2015).

722 The precuneus also showed modulations in spectral power as a result of cognitive task
723 load, with activity in both alpha and beta bands ramping up during no-task blocks, possibly as a
724 mechanism to inhibit and down-regulate visual load (Banerjee, Snyder, Molholm, & Foxe, 2011;
725 Foxe et al., 1998; Foxe & Snyder, 2011). Furthermore, an interaction between visual and
726 cognitive load within the alpha-band reveals that power decreases with visual load as
727 participants are engaged in the cognitive task, but increases as participants disengage from the
728 task, particularly while exposed to mediolateral perturbations. Considering precuneus
729 connections with sensorimotor regions (Cavanna & Trimble, 2006), the latter finding might
730 indicate an alpha-band mediated gating/suppression mechanism of unreliable information to
731 sensorimotor regions. Interestingly, the IC clustering approach produced two parietal clusters,
732 sensitive to attentional demands resulting from cognitive task engagement and the radiating star
733 field, that were both localized to the right hemisphere. This finding fits nicely with several reports
734 in the literature. The precuneus has been linked to the processing of scenes, with previous
735 imaging studies reporting middle parietal cortex to be involved in visuospatial processing (Harris
736 et al., 2000), and specifically the right hemisphere to be more spatially oriented to the
737 surrounding environment (Joseph, 1988). Topographic mapping of high-density EEG recorded
738 in a line-bisection task revealed a right hemisphere dominant network with activation spreading
739 from right parieto-occipital scalp, to regions over right superior cortices (Foxe, McCourt, & Javitt,
740 2003). The right hemisphere may also control shifts in attention when viewing a scene - fMRI
741 studies have reported right-lateralized fronto-parietal activity during shifts in visual attention
742 (Corbetta, Kincade, Ollinger, McAvoy, & Shulman, 2000). Furthermore, a recent EEG study
743 found increased processing of optic flow speed over right parietal recording sites (Vilhelmsen,
744 van der Weel, & van der Meer, 2015). And in an older study that used positron emission
745 tomography (PET), the right precuneus was cited as one of three areas that showed increased

746 cerebral blood flow specifically in response to optic flow stimulation (de Jong, Shipp, Skidmore,
747 Frackowiak, & Zeki, 1994). These authors claim that both dorsal and ventral pathways are
748 involved in the processing of optic flow stimuli, based on their finding of occipito-parietal as well
749 as occipito-temporal activation patterns (de Jong et al., 1994). This claim is in line with a recent
750 proposal that the inferior parietal lobe does not fit into the traditional dorsal-ventral visual
751 processing stream dichotomy, and that specifically the right inferior parietal lobe plays an
752 important role in maintaining attention while also responding to salient new information (Singh-
753 Curry & Husain, 2009).

754

755 Supplementary motor area (SMA)

756 The SMA has been implicated in an enormous variety of motor functions including
757 planning and gait initiation (Mihara, Miyai, Hatakenaka, Kubota, & Sakoda, 2007) and
758 coordinating more demanding walking tasks (Kurz, Wilson, & Arpin, 2012), as well as cognitive
759 control functions (Nachev, Kennard, & Husain, 2008). In a recent MoBI study employing
760 connectivity analysis based on fluctuations in spectral power between cortical IC clusters, the
761 authors proposed a cortical network underlying both active and viewed limb movements driven
762 by the right premotor cortex and SMA, but also including cingulate and parietal areas (Kline et
763 al., 2016). In another MoBI paradigm, brain-to-muscle connectivity was assessed by measuring
764 heel-strike related spectral perturbations and electromyographic recordings (Artoni et al., 2017).
765 They found evidence of unidirectional drive from contralateral motor cortex to leg muscles in the
766 swing phase, with stronger modulations in mu, beta and gamma bands for clusters over motor
767 areas compared to non-motor areas. And motor regions, including the cingulate motor cortex,
768 supplementary motor area, and primary foot motor cortex were among the cortical areas with
769 maximal influence on lower limb muscles during stereotyped walking (Artoni et al., 2017).

770 Therefore it does not come as a surprise that we also observed modulatory activity
771 resulting from both sensory and cognitive processing in this cluster. A desynchronization was

772 observed in both alpha and beta bands associated with increased visual input (optic flow),
773 compared to the static condition. Additionally, theta power was significantly reduced with MLP,
774 in comparison to the other visual conditions. Furthermore, an interaction was observed for alpha
775 frequencies, as the presentation of ML perturbations resulted in a different pattern of spectral
776 modulation depending on whether one was engaged in the task or not. Finally, in relation to
777 cognitive load, theta power was higher when participants performed the task; in line with
778 findings showing theta power is sensitive to the recruitment of executive control in interference
779 situations (Nigbur, Ivanova, & Sturmer, 2011).

780

781 Anterior cingulate cortex (ACC)

782 According to fMRI studies, the ACC is thought to monitor ongoing mental processes and
783 signal the need for increased attentional focus (Fassbender et al., 2009; O'Connell et al., 2007;
784 Simoes-Franklin, Hester, Shpaner, Foxe, & Garavan, 2010). We observed significant increases
785 in theta and alpha spectral power in this cluster, observed across approximately 3-12Hz
786 frequencies, for the static visual condition compared to both dynamic optic flow conditions.
787 Additionally, theta power showed a significant increase for task performance, in line with the
788 results observed in the SMA cluster above, and points to the role of theta oscillations in
789 executive control processes during increased task load (Clayton et al., 2015). Cognitive-task
790 related modulations in this cluster likely reflect processing demands dedicated to the Go/No-Go
791 task, as the ACC has frequently been cited for recruitment in processing error detection and
792 correction (O'Connell et al., 2007; Walton, Croxson, Behrens, Kennerley, & Rushworth, 2007)
793 as well as evidence from a Go/No-Go ERP study implicating this area in conflict monitoring and
794 attentional allocation (Dias, Foxe, & Javitt, 2003; Fallgatter, Bartsch, & Herrmann, 2002).

795

796 In conclusion, by utilizing an ICA and clustering approach to isolate cortical sources
797 supporting dual-task walking activity, we have demonstrated that the MoBI technique is capable

798 of distinguishing subtle modulations in gait and spectral power attributed to sensory and
799 cognitive load. Future investigations will examine event-related spectral perturbations (ERSPs)
800 to determine if the timing of spectral power fluctuations is associated with specific phases of the
801 gait cycle. This will add to the literature as cortical involvement in gait is already being explored
802 in the context of steady-state and robotic-assisted treadmill walking (Gwin et al., 2011; Presacco
803 et al., 2012; Seeber, Scherer, Wagner, Solis-Escalante, & Muller-Putz, 2014; Wagner et al.,
804 2016; Wagner et al., 2012). In future MoBI protocols, the utilization of spatially-filtered EEG
805 signals during active movements may provide insight into the neural dynamics underlying gait
806 adaptation. This area of research is especially relevant for applications such as
807 neurorehabilitation, for example to decode user intentions from EEG in brain-computer
808 interfaces (Kilicarslan, Prasad, Grossman, & Contreras-Vidal, 2013; Wagner et al., 2012).
809 Additionally, valuable information may be gained in relation to monitoring the neural correlates
810 underlying disease progression and rehabilitation in diseases such as Multiple Sclerosis and
811 Parkinson's (Boyd, Vidoni, & Daly, 2007). Finally, older adults often have difficulty adapting to
812 increased cognitive load during locomotion and show evidence of declines in proprioceptive,
813 vestibular and somatosensory processing (Goble, Coxon, Wenderoth, Van Impe, & Swinnen,
814 2009; Hay, Bard, Fleury, & Teasdale, 1996), factors that may increase fall risk (Ayers, Tow,
815 Holtzer, & Verghese, 2014; Setti, Burke, Kenny, & Newell, 2011). MoBI approaches in virtual
816 reality environments (e.g., visual perturbations) could be employed in combination with gait
817 training strategies to successfully challenge people's walking ability, with the aim of reducing fall
818 risk in vulnerable populations.

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850 **Figure 1:** Representation of recording apparatus: a participant walking on the treadmill wearing
851 an EEG cap and motion capture markers, facing the optic flow display.

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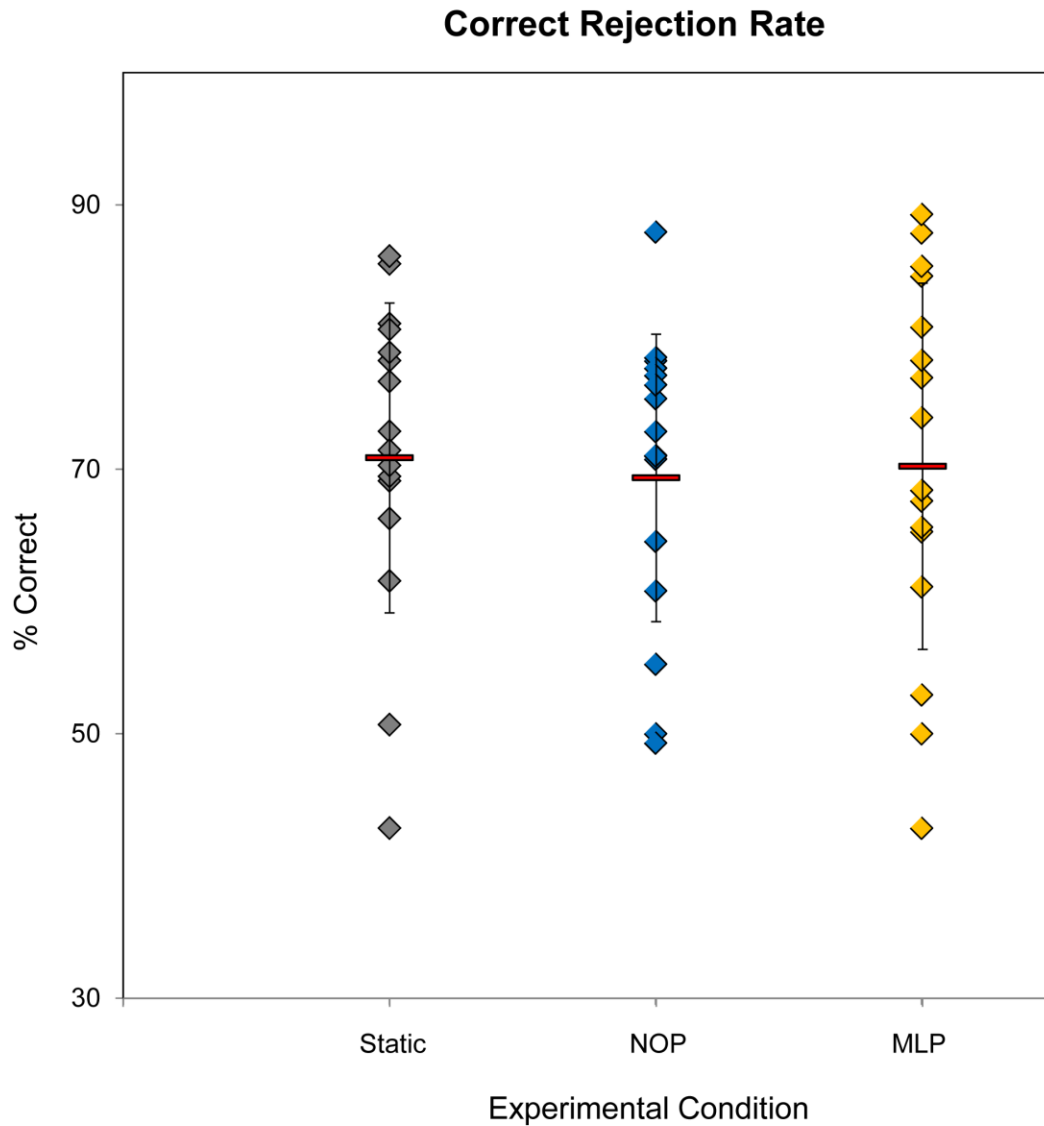
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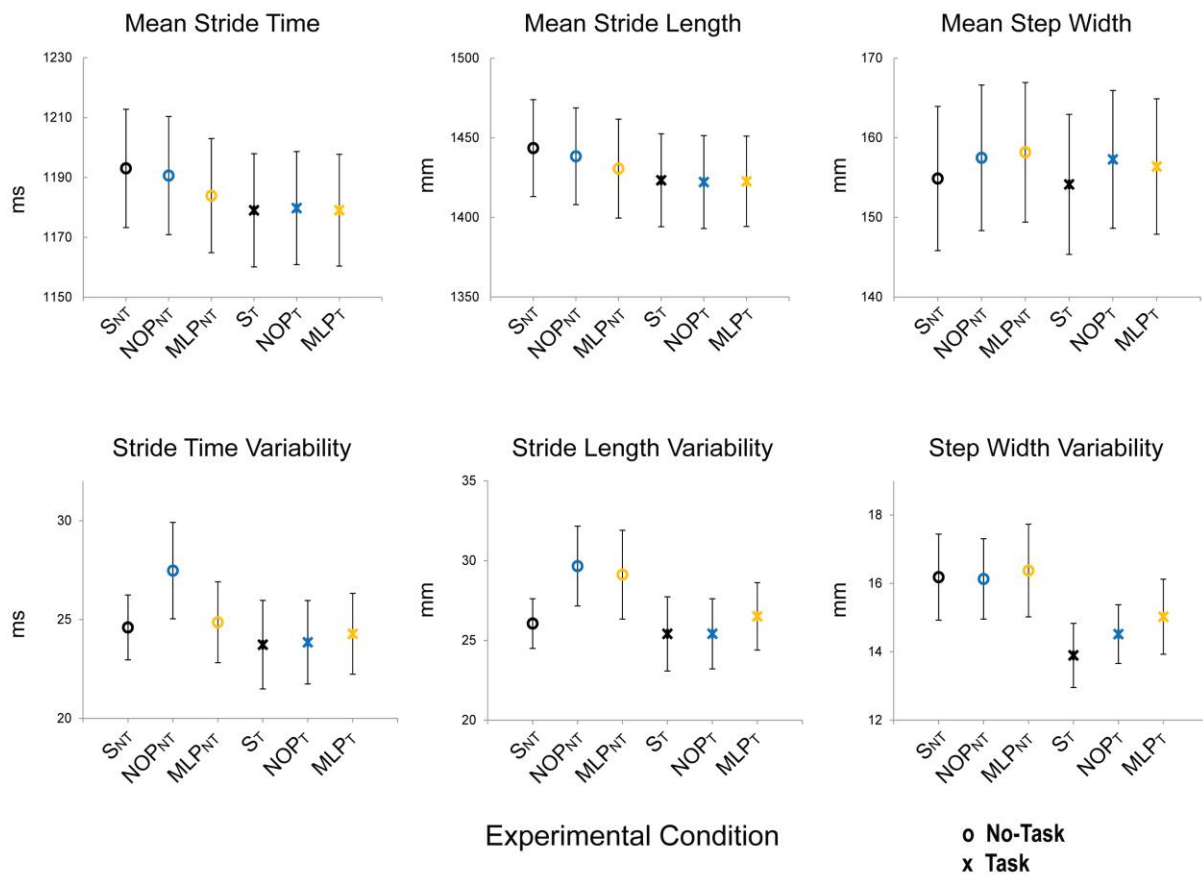
865 **Figure 2:** Response inhibition performance on the Go/No-Go task. From left to right: percentage
866 of correct rejections (CRs) for all 16 participants during static visual field, dynamic optic flow with
867 no perturbation (NOP) and optic flow with ML perturbation (MLP). Red markers indicate the
868 means for each condition, with vertical bars representing standard deviations.
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874 **Figure 3:** Average and mean variability for stride time, stride length and step width. Top row
 875 represents means, bottom row is mean SD, for stride time (left column), stride length (center)
 876 and step width (right column) for all six experimental conditions. Open circles represent the No-
 877 Task conditions, while crosses represent Task blocks. S_{NT} = Static No Task, NOP_{NT} = No
 878 perturbation No Task, MLP_{NT} = Mediolateral perturbation No Task, S_T = Static Task, NOP_T = No
 879 perturbation Task, MLP_T = Mediolateral perturbation Task.

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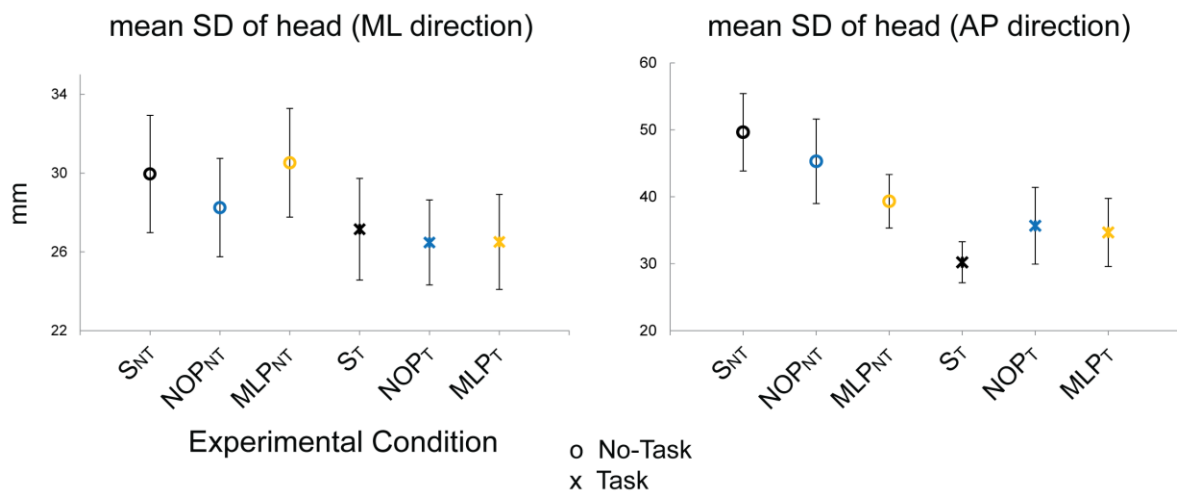
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887 **Figure 4:** Average variability in head position, in ML (left) and AP (right) directions for all six
888 experimental conditions. Open circles represent the No-Task conditions, while crosses
889 represent Task blocks. S_{NT} = Static No Task, NOP_{NT} = No perturbation No Task, MLP_{NT} =
890 Mediolateral perturbation No Task, S_T = Static Task, NOP_T = No perturbation Task, MLP_T =
891 Mediolateral perturbation Task.

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Table 1: Clusters of Independent electrocortical sources (ICs). Description and approximate location (Brodmann area and Talairach coordinates) of cluster centroids for all clusters located in the cortex and containing ICs from more than half of the participants.

Functional Area	Brodmann Area	Talairach coordinates (x,y,z)	No. of subjects (S) and ICs
Medial occipital lobe, lingual gyrus	BA17	11, -94, -10	12 S, 14 ICs
Right occipital	BA19	47, -73, -1	11 S, 12 ICs
Left occipital	BA19	-43, -71, 14	12 S, 14 ICs
Left superior temporal gyrus	BA22	-47, -17, -6	10 S, 12 ICs
Right inferior parietal lobule	BA40	43, -34, 36	8 S, 10 ICs
Parietal lobe, precuneus	BA7	12, -62, 34	11 S, 15 ICs
Supplementary motor area	BA6	-6, -16, 45	14 S, 23 ICs
Limbic lobe, anterior cingulate	BA24	1, 25, 22	14 S, 15 ICs

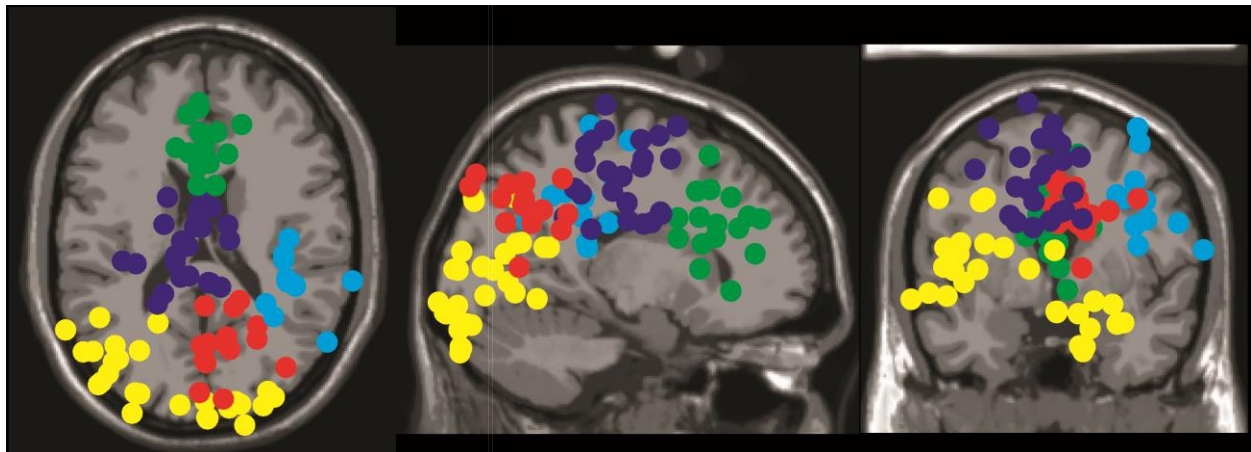
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917 **Figure 5:** Clusters of electrocortical sources localized to occipital cortex (yellow), parietal cortex
918 (inferior parietal lobule: cyan, precuneus: red) and frontal cortex (Supplementary Motor Area:
919 purple, Anterior Cingulate Cortex: green).

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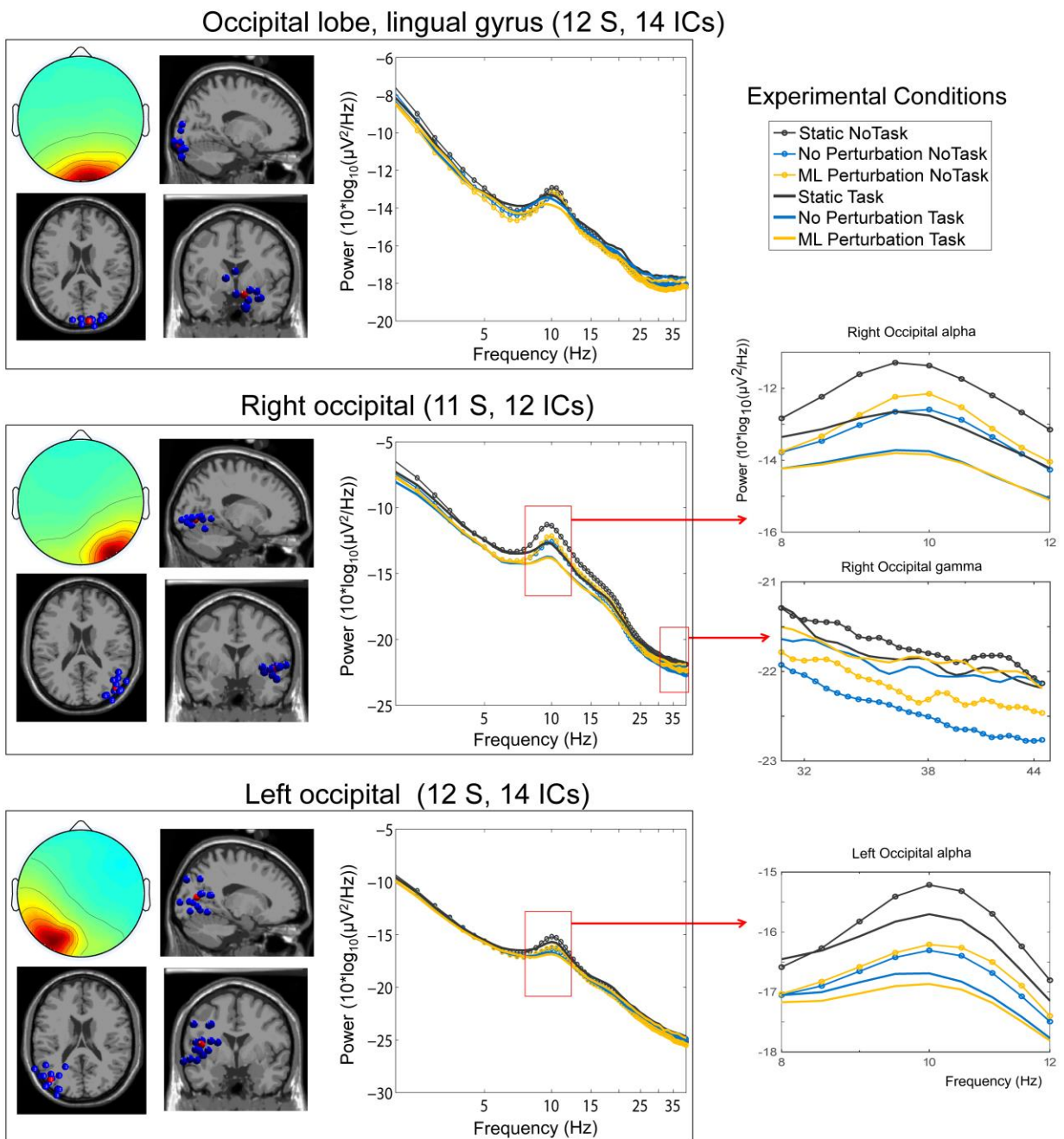
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936 **Figure 6:** Occipital cortex clusters. Scalp topography, dipole location (blue dots indicate the
937 location of each IC, red dots represent the cluster centroid) and average power spectral density.

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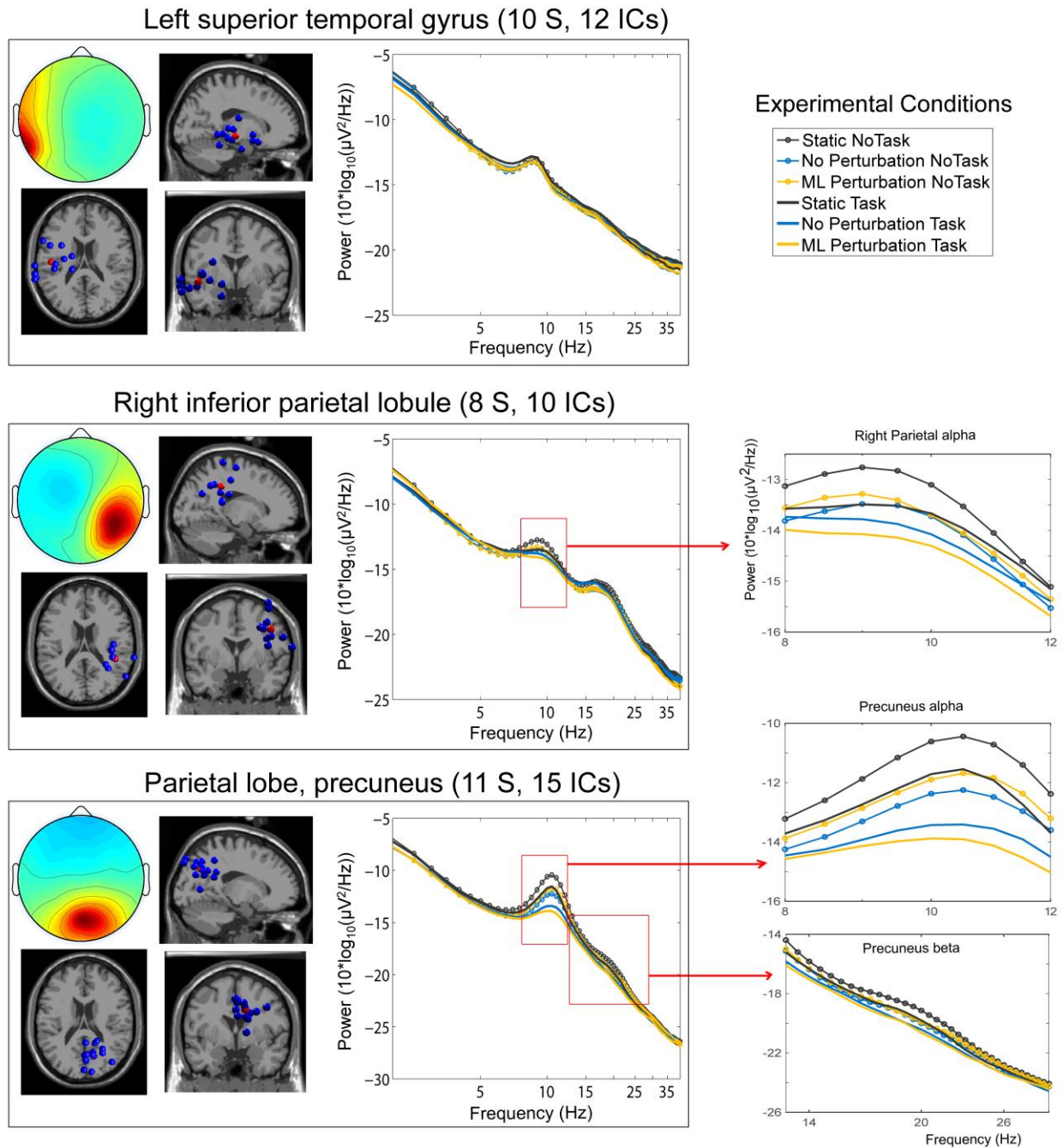
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942 **Figure 7:** Clusters located over temporal and parietal cortex. Scalp topography, dipole location
943 (blue dots indicate the location of each IC, red dots represent the cluster centroid) and average
944 power spectral density.

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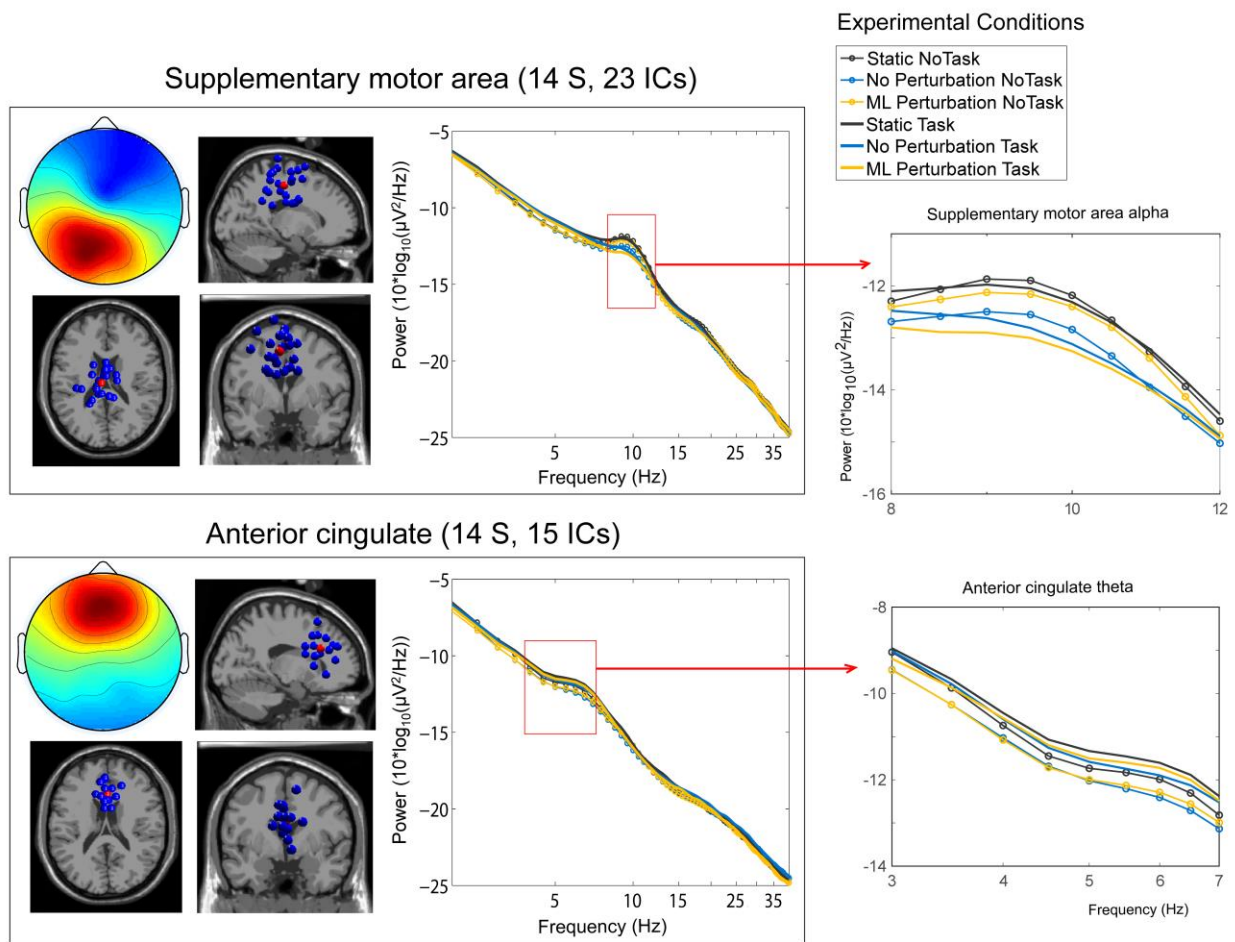
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948 **Figure 8:** Clusters located over frontal cortical areas. Scalp topography, dipole location (blue
949 dots indicate the location of each IC, red dots represent the cluster centroid) and average power
950 spectral density for the cluster localized to the supplementary motor area (top panel) and the
951 anterior cingulate (bottom panel).

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