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² Motor Preparation of Step Initiation: Error-related Cortical Oscillations

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Abstract—Gait initiation can vary as a function of the available and engaged attentional resources. Conflict resolution can disrupt movement preparation and lead to "errors" in motor programming. These "errors" are physiologically useful by enabling us to adapt our motor behavior to situations with conflicting information. The objective of the present study was to analyze the patterns of cortical activation associated with motor programming errors and the corresponding error corrections. Incongruent flankers around a target arrow were used to trigger errors in anticipatory postural adjustments (APAs) prior to gait initiation; i.e. perturbed motor programming but normal execution. Thirty healthy adults performed a gait initiation task. The event-related potentials (ERPs) and event-related desynchronization (ERD) after target presentation were analyzed according to the presence or absence of an APA error. The ERP was similar in both conditions, except that the Ne and P300 peak latencies were longer for APA errors. Motor programming errors during gait initiation were characterized by longer, less intense low-beta-band ERD over the sensorimotor cortex and alpha ERS followed by stronger alpha ERD during errors. APA errors were associated with a specific alpha/beta oscillation profile over the sensorimotor cortex; these beta oscillations might be sensitive markers of non-conscious motor error and correction monitoring.

Key words: cortical activation, gait initiation, posture, inhibition, attention.

INTRODUCTION

Gait initiation is a motor program characterized by the 11 transition from a static stable stance to a continuously 12 unstable posture during locomotion. The characteristics 13 of gait initiation can vary as a function of the available 14 attentional resources. Indeed, gait initiation can be 15 modulated when the subject is obliged to deal with 16 conflicting information (Uemura et al., 2012). Gait is initi-17 ated in two phases: a motor preparation phase (corre-18 sponding to anticipatory postural adjustments, APAs) 19 and then an execution phase (corresponding to the time 20 interval between "toe-off" and "heel strike" for the swing 21 leg). During standard gait initiation (i.e. in the absence 22 of external or internal stimuli requiring modulation of the 23 motor program), healthy subjects display a stereotypical 24 APA pattern. Foot-off of the swing leg is preceded by a 25 shift in body weight that displaces the center of pressure 26

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Abbreviations: APAs, anticipatory postural adjustments; CoP, center of pressure; CRN, Correct-Related Negativity; EEG, electroencephalogram; ERD, event-related desynchronization; ERD/S, event-related (de)synchronization; ERN, Error-Related Negativity; ERPs, event-related potentials; Pe, Positive error-related.

(CoP) backward and toward the swing leg. Next, the CoP 27 is displaced forward and toward the stance leg. Hence, 28 APAs create the conditions required for progression 29 (Brenière and Do, 1991). Furthermore, APAs along the 30 mediolateral axis are predictive of postural stability 31 (McIlroy and Maki, 1999). However, it is known that self-32 triggered gait initiation is not always preceded by an 33 APA (Delval et al., 2014; Lu et al., 2017). The lack of a lat-34 eral or posterior APA was nevertheless infrequent (in 2% 35 of the trials) in healthy elderly controls during externally 36 triggered rapid stepping (Delval et al., 2014). Conversely, 37 the absence of APAs can be frequently observed in 38 patients with freezing of gait and an increased risk of falls 39 (Delval et al., 2014). The occurrence of APA errors can 40 also perturb the gait initiation program by delaying the 41 onset of movement execution (Cohen et al., 2011). This 42 corresponds to the correction of an APA when the initial 43 direction of postural adjustment is not appropriate (for 44 example, when the CoP moves inappropriately toward 45 the stance leg and is then appropriately moved first 46 toward the swing leg and only then toward the stance 47 leg). This APA error corresponds to a motor program 48 error, which is efficiently corrected and prevents incorrect 49 step initiation from taking place. It is known that APA 50 errors are more frequent in conditions modulated by 51 attention (especially in the presence of incongruent stimuli 52

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(Uemura et al., 2013) or with invalid cues (Tard et al., 2013)) than in conditions with congruent step initiation stimuli. However, it is not known if and how these APA 55 errors are modulated in the cortex.

Cortical areas involved in gait initiation include the 57 sensorimotor cortex, premotor cortex, basal ganglia and 58 brainstem structures. It was initially suggested that the 59 60 motor programs underlying the elicitation of gait initiation were stored in subcortical structures, and could be 61 elicited by a startling stimulus or a decision for action 62 (Takakusaki, 2008; Queralt et al., 2010; Delval et al., 63 2012; Watanabe et al., 2016a). However, studies in 64 patients with focal lesions of the supplementary motor 65 66 area and studies in patients with Parkinson's disease (Viallet et al., 1992; Gantchev et al., 1996) have shown 67 68 that APAs can be modulated at the supraspinal level. since the supplementary motor area, the basal ganglia 69 and the pontomedullary reticular formation are linked by 70 neural networks. Moreover, inhibitory repetitive transcra-71 nial magnetic stimulation over the supplementary motor 72 area shortens the APA duration for a brief period, i.e. 73 for the first stepping trial after stimulation (Jacobs et al., 74 2009). The output of this pathway is located in the mid-75 76 brain locomotor region (which may correspond in part to 77 the cuneiform nucleus and the dorsal part of the peduncu-78 lopontine nucleus), which is connected to limbic struc-79 tures and the basal ganglia (Pahapill and Lozano, 2000).

80 Attentional control can modulate gait initiation - either directly by involving brainstem structures (for example. 81 the alert process induced by a loud stimulus can 82 product a start-react effect) or indirectly via a cortical 83 loop that includes more complex attentional networks 84 (Delval et al., 2012; Tard et al., 2013). Cortical movement 85 preparation can be measured through electroencephalo-86 gram (EEG) features like event-related potentials (ERPs) 87 and event-related (de)synchronization (ERD/S). In the 88 frequency domain, ERD (mainly in the alpha- and beta-89 90 bands) is the cortical marker of movement intention. It has been demonstrated that gait initiation is associated 91 92 with desynchronization of sensorimotor rhythms related to sensorimotor cortex activation (Pfurtscheller and 93 Andrew, 1999). If the EEG is response-locked (i.e. locked 94 to the motor response), a movement-related cortical 95 potential (MRCP) is present before gait initiation or when 96 97 imagining gait initiation (Vidailhet et al., 1993, 1995). For 98 the gait initiation task, if the EEG is target-locked, the 99 early ERP components are probably influenced by the different physical characteristics of the stimuli (Rektor et al., 100 2006) and a posterior P300 can be found, higher in case 101 of stimulus driven attention for example (Tard et al., 102 103 2013), whereas late components reflect motor preparation (Hamano et al., 1997). More recently, combined 104 ERP and ERD/S recordings via an EEG brain-computer 105 106 interface were used to detect gait initiation (Jiang et al., 2015; Sburlea et al., 2015). 107

ERPs are also used to monitor cognitive control of 108 action. During error recognition, a negativity (named the 109 "Error-Related Negativity" (ERN or Ne)) and then a 110 "Positive error-related wave" (Pe) can be observed (for 111 a review, see Wessel and Aron (2017)). The functional 112 significance of ERN was associated with error detection 113

(Falkenstein et al., 1991). Alternatively, the ERN was pro-114 posed to reflect conflict resolution due to a finding of the 115 "Correct-Related-Negativity" (CRN) (Vidal et al., 2000; 116 Meckler et al., 2011). However, errors during gait initiation 117 are mostly non-conscious and the presence of an ERN or 118 Pe during an APA error in healthy subjects remains uncer-119 tain. For example, Watanabe found similar frontal ERN 120 and CRN in trials with or without APA errors during gait 121 initiation (Watanabe et al., 2016b). The significance of 122 these potentials remains discussed. Indeed, it has been 123 demonstrated that the ERN occurs also after "partial 124 errors", i.e., incorrect activities that are not sufficient to 125 produce overt errors (Carbonnell and Falkenstein, 126 2006), which is observed during spontaneous correction 127 of APA errors. To date, the cortical areas involved in gait 128 initiation errors have not been extensively studied and the 129 focus was only on Fz, FCz, and Cz (Watanabe et al., 130 2016b). Indeed, the human sensorimotor system needs 131 to be able to rapidly correct for errors in an ongoing motor 132 command brought about by sudden, unexpected changes 133 in the movement environment (such as conflicting infor-134 mation, for example) (Krigolson et al., 2008). The present 135 study was designed to evaluate the cortical changes 136 induced by these adaptive reactions called APA errors. 137 The study's primary objective was to use a combined 138 ERP and time-frequency analysis to evaluate cortical 139 activation during correct gait initiation (i.e. with no APA 140 errors) and during disturbed step initiation (i.e. with APA 141 errors). Our starting hypothesis was that APA errors 142 would be associated with ERP modulations featuring 143 error-related potentials (for example, error-related nega-144 tivity/positivity (Ne/Pe) (Falkenstein et al., 2000)) and/or 145 changes in beta-band ERS, for example increased beta 146 ERS, as observed in stop-signal paradigms for move-147 ments requiring motor inhibition (Duque et al., 2017). 148 Modulations in lower bands (delta-theta) have also been 149 attributed to error monitoring in children, young and 150 elderly adults (Kolev et al., 2001, 2005; Albrecht et al., 151 2009). 152

EXPERIMENTAL PROCEDURES

Participants

Thirty healthy adult volunteers (16 females, 14 males; 29 right-handed; mean \pm standard deviation (SD) age: 39.4 \pm 14.2 years) participated in the study after providing written, informed consent. None of the participants had history of medication use (neuroleptics, а 159 benzodiazepines, etc.) or disease (neurological, orthopedic or psychiatric) that could have interfered with gait. The mean ± SD Montreal Cognitive Assessment score (Nasreddine et al., 2005) was 28.5 ± 2. Participants with a score of less than 26 out of 30 were excluded. The study was approved by the local independent ethics committee (CPP Nord-Ouest, Lille, France; reference: 2015-A00013-46).

The experimental setting

The participant was told to stand in a stable, comfortable, 169 natural posture on a force platform, with his/her feet 170

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171 parallel and with a gap of a few centimeters between the feet. A computer display screen was placed at head 172 height 1 meter in front of the participant. The attentional 173 task was an adaptation of the attentional network test 174 (Fan et al., 2002) (Fig. 1). The participant was instructed 175 to initiate a forward step after presentation of the visual 176 target (an arrow pointing to the right or to the left, which 177 was visible for 1500 ms). If the arrow pointed to the left. 178 the participant had to initiate gait with the left foot; con-179 versely, if the arrow pointed to the right, the participant 180 had to initiate gait with the right foot. The balance weight 181 shift between the 2 feet was controlled online (position of 182 the CoP between the 2 feet visualized by the Nexus soft-183 184 ware). The present study only assessed the condition with incongruent flankers (i.e. flankers pointing in the opposite 185 direction to the target arrow), in which the frequency of 186 APA errors is reportedly higher (Uemura et al., 2013). 187 Indeed, differences in ERP amplitudes have been 188 reported between congruent and incongruent conditions 189 in a similar study design in seated condition (Neuhaus 190 et al., 2010). A total of 144 incongruent trials (out of a total 191 of 300) were available for each participant. 192

193 Motion analysis

Data were collected with a three-dimensional motion 194 analysis system (VICON 370®, Oxford Biometrics, 195 Oxford, UK), using eight infrared cameras and a 196 sampling frequency of 100 Hz. The CoP was measured 197 with two force platforms (the ORG-5 model from 198 199 AMTI®, Watertown, MA, USA) at a sampling frequency 200 of 1000 Hz. Reflective markers were placed on precise, reproducible, anatomic landmarks on each foot; the toe 201



Fig. 1. Attentional Network test. A schematic diagram of the attentional task. S1: cue; S2: target. The target appeared for 1500 ms and was surrounded by flankers (four arrows: two on each side) that were either congruent or incongruent. The targets could be preceded by valid or invalid spatial cues, i.e. asterisks indicating (or not) the direction of the arrow. Four blocks of 75 trials were administered. The blocks were separated by short breaks of variable duration. The cue and target conditions were presented in a pseudo-random order: 156 congruent trials, 144 incongruent trials, 49 trials with no cue, 61 trials with a central cue, and 190 trials with a spatial cue (142 valid and 48 invalid cues). Only incongruent trials was 16.3%, the proportion of neutral cue trials was 20.3%, the proportion of valid cue trials was 47.3%, and the proportion of invalid cue trials was 16%.

(the head of the second metatarsal), the lateral malleolus, and the heel. The data were then computed by the same operator using an in-house MATLAB® routine (The MathWorks, Natick, MA, USA).

The direction of the APA was considered to be normal 206 if the CoP moved backward and sideways toward the 207 swing foot. Conversely, the direction of the APA was 208 considered to be abnormal (i.e. an APA error) if the CoP 209 moved first toward the stance foot and only then toward 210 the swing foot (see Fig. 2). The reaction time (RT) was 211 defined as the time interval between the appearance of 212 the target (S2) and the beginning of the APA or T_0 An 213 RT < 100 ms was classified as a false start and was 214 excluded from further analyses. Incorrect starts (i.e. 215 starts with the wrong foot) were also excluded. An in-216 house MATLAB® routine detected changes in CoP 217 velocity > mean + 3 SD of the baseline period (-1500)218 to 1000 ms before target stimulus), the experimenter 219 then chose the start of the APAs according to the 220 curves in X and Y axis. Toe-off was detected visually 221



Fig. 2. Normal Anticipatory Postural Adjustments (APA) and APA error. Step initiation with the right foot by a study participant. Top panel: a normal APA, where the center of pressure (CoP) shifts to the swing leg (right) and then to the stance leg (left) (lateral CoP: red line). Bottom panel: an APA error: the CoP shifts toward the stance leg (left) but the trajectory is corrected. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

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(from the toe marker trajectory in the sagittal plane) and then reported on the CoP curve. It corresponded to the time the CoP shifted forward. The APA duration was assessed by subtracting T_0 from the toe-off time. For APA errors, the correction time was defined as the time interval between T_0 and the sideways corrective shift (i.e. the beginning of the APA in the correct direction).

229 Acquisition of electroencephalographic data

The electroencephalogram (EEG) was recorded with an 230 Ag/AgCl 128-scalp-electrode cap (Waveguard®, ANT 231 The Netherlands), Neuro. Enschede, positioned 232 according to the 10/05 international system (Oostenveld 233 and Praamstra, 2001). Data were acquired with ASA™ 234 software (ANT Neuro), using a 0.01- to 100-Hz band-235 236 pass filter, a sampling frequency of 512 Hz, and impedances below 20 kΩ. The data were pre-processed with 237 ASA[™] software in order to reject ocular artifacts and apply 238 a 50-Hz notch filter to the recordings. Next, interpolation 239 was performed for artifact-affected electrodes, with a 240 241 maximum interpolation rate of 10% (n = 13) for the whole set of 128 scalp electrodes. 242

The EEG data were then analyzed with Brain Vision 243 Analyzer 2.0 software (BrainProducts). Muscle artifacts 244 were manually removed from the EEG layout; 245 thereafter, we segmented the EEG data into 2500-ms 246 epochs that were time-locked with respect to target 247 onset (1500 ms before and 1000 ms afterward). The 248 median (min-max) number of epochs selected (after 249 artifact rejection) per participant was 49 (20-98) for APA 250 errors and 69 (38-107) for normal APAs. 251

ERP analyses. ERP were analyzed with the EEGLAB 252 toolbox (Delorme and Makeig, 2004), using a baseline 253 254 from 1500 ms before the target onset to 1000 ms for 255 target-locked ERP and from 1500 ms before the APA onset to 650 ms for response-locked ERP. The time win-256 dow analysis was from S2 to 1000 ms after target onset. 257 We first analyzed ERP scalp distribution maps. Next, 258 the ERP wave's characteristics were assessed and col-259 260 lected by grand-averaging over the main central elec-261 trodes (Fz, Cz, Pz).

For target-locked ERP, we focused on P300 over Pz. 262 For response-locked ERP, we analyzed ERN/CRN over 263 Fz and P300/Pe over Pz. The amplitude of potentials 264 was measured as the difference between the maximum 265 peak of the ERP waveform and the mean baseline 266 voltage (which occurs prior to the stimulus). Latency 267 268 was defined as the interval between target presentation and the point of highest positive amplitude in the time 269 270 window of the potential. For target-locked ERP, time 271 window of P300 ranged from 250 to 500 ms after target presentation. For response-locked ERP, the time 272 273 window of the ERN/CRN ranged from -50 to 200 ms 274 after APA start and from 0 to 400 ms for the posterior component. 275

ERD/ERS analyses. Time-frequency analysis
requires computing the power spectrum over a sliding
latency window. ERD data were analyzed using
EEGLAB software (Delorme and Makeig, 2004) with a

500-ms baseline (between 1500 and 1000 ms before 280 the target's appearance, target-locked and response-281 locked). To characterize event-related EEG oscillations 282 like ERD and ERS, we applied a time-frequency analysis 283 based on a continuous wavelet transform. We used a ver-284 sion of sinusoidal wavelets in which the number of cycles 285 increases slowly with frequency (e.g. 1.5 cycles at 4 Hz, 286 and 5.6 cycles at 30 Hz) with a window width of 213 sam-287 ples (416 ms). This procedure has been described in 288 (Delorme and Makeig, 2004), and similar approaches for 289 time windows of around 2500 ms have been described 290 in (Fan et al., 2007). Time-frequency analyses were per-291 formed between 4-7 Hz (the theta-band), 8-12 Hz (the 292 alpha-band), and 13-30 Hz (the beta-band, divided in a 293 low-beta-band (13-20 Hz) and a high-beta-band (20-294 30 Hz)). 295

Cortical sources. A realistic head model was built by 296 segmenting a template MRI data with Freesurfer 297 software (Dale et al., 1999). The lead field matrix was 298 then computed for a cortical mesh with 15.000 vertices. 299 using Brainstorm software (Tadel et al., 2011) and Open-300 MEEG software (Gramfort et al., 2010). The weighted 301 minimum-norm estimate was then used to reconstruct 302 the cortical sources (using Brainstorm toolbox (Tadel 303 et al., 2011)) in the time window corresponding to motor 304 preparation. 305

Statistical analyses

Characteristics of APAs were compared using a one-way 307 ANOVA after checking normality of the distributions. To 308 evaluate differences in cortical activation (ERPs, ERD/S 309 and source localization) in trials with an APA error vs. 310 trials with a normal APA, we used a non-parametric 311 permutation (randomization) test to obtain the p-value 312 for each electrode and each time point (for ERP and 313 ERD/S) and for each source reconstruction. The false 314 discovery rate (FDR) method was used to correct for 315 multiple comparisons (Genovese et al., 2002), and 316 enabled us to determine which electrodes differed 317 between the two conditions at the different time points 318 (scalp maps) and to compare ERD/ERS maps between 319 both conditions. These analyses were performed with 320 the EEGLAB toolbox (Delorme and Makeig, 2004), which 321 includes MATLAB statistical routines at this purpose. For 322 comparisons between source localizations, we used the 323 scripts included in the Brainstorm toolbox (Tadel et al., 324 2011)). Peak amplitudes and latencies (for P300 on Pz, 325 for example) were compared in a t-test (in SPSS 17 for 326 Windows) after checking the normality of distribution in 327 a Kolmogorov-Smirnov test. The threshold for statistical 328 significance was set to p < 0.05 for all analyses. 329

RESULTS

Behavioral data

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In trials with incongruent flankers, the APA error rate was 332 41.0%. 333

The false start rate. The false start rate (i.e. 334 RTs < 100 ms) was 11.9%. These trials were excluded 335

from further analysis, since they did not correspond to 336 APA errors. 337

The error step rate. A start with the wrong foot was 338 rare, since it occurred in only 0.56% of the trials. These 339 trials were excluded from the analysis because they did 340 not correspond to correct error monitoring. The low 341 number of these events prevented us from analyzing 342 them separately. 343

The mean \pm SD RT was 0.27 \pm 0.08 s for normal 344 APAs and 0.23 ± 0.06 s for APA errors (p < 0.001). 345 The mean ± SD APA duration was longer for APA 346 errors $(0.64 \pm 0.13 \text{ s})$ than for normal APAs (0.47)347 \pm 0.10 s) (p < 0.001). The mean \pm SD correction time 348 349 for APA errors was 0.20 ± 0.07 s (i.e. 0.43 ± 0.07 s after target presentation, on average). 350

Erp 351

352 Target-locked ERP: as shown in Fig. 3, the ERP scalp 353 distribution maps revealed an early anterior component 354 (N2, see discussion), a late central negative component (corresponding to preparation of movement) and a 355 posterior positive component (at the same scalp sites as 356 the P300). There were no differences in the ERP maps 357 between the "APA error" and "normal APA" conditions 358 except for P300 component (see Figs. 3 and 4). 359

The mean ± SD P300 peak latency (Pz electrode) 360 was longer for the APA error condition than the normal 361 APA condition $(0.50 \pm 0.08 \text{ s} \text{ vs. } 0.47 \pm 0.08 \text{ s},$ 362 respectively; p < 0.01). No differences in P300 peak 363 amplitude were observed. 364

Response-locked: in both APA conditions, early 365 negativity (in the time window of ERN or CRN) 366 occurred, later in case of APA error. Late positivity 367 (P300 or Pe, see discussion) also occurred in both 368 conditions, later in case of APA error on posterior 369 regions. The mean ± SD ERN/CRN peak latency (Fz 370 electrode) was longer for the APA error condition than 371 the normal APA condition $(0.12 \pm 0.05 \text{ s vs.} 0.08)$ 372 \pm 0.05 s, respectively; p < 0.001). No differences in 373 peak amplitude were observed. The mean ± SD Pe/ 374 P300 peak latency (Pz electrode) was longer for the 375 APA error condition than the normal APA condition 376 $0.19 \pm 0.06 \, \mathrm{s},$ respectively; $(0.26 \pm 0.07 \, s)$ 377 VS. p < 0.001). No differences in peak amplitude were 378 observed. 379

ERD data 380

As shown in Figs. 5 and 6, we observed similar theta-381 band ERS (between 200 and 600 ms, target-locked; 382 383 starting at T0, response-locked) in both conditions.

384 Alpha ERS was significantly more pronounced in trials with an APA error (starting around 300 ms after S2, 385 target-locked; during APA, response-locked) and was 386 followed by a stronger alpha ERD (response-locked). 387

Central beta ERD was observed, starting 200 ms after 388 S2 (target-locked); or just before T0 (response-locked). 389 This feature lasted significantly longer over Cz in trials 390 with an APA error (Fig. 6). Moreover, beta ERD over 391 the sensorimotor cortex was more attenuated in the low-392 beta-band (i.e. 13-20 Hz) than in the high-beta-band 393

(20-30 Hz) in trials with an APA error (relative to trials 394 with a normal APA) in target-locked analysis (Figs. 5 395 and 6). We can observe that this beta ERD was present 396 during both normal APAs and APA errors but was more 397 prolonged (response-locked and target-locked) in case 398 of error. 399

Cortical sources of changes in the EEG signal during motor programming

Cortical sources in the 0- to 600-ms time interval (target-402 locked) are shown in Fig. 7. Occipital and temporoparietal 403 regions were activated at 200 ms, and then the 404 sensorimotor cortex and the frontal dorsolateral cortex 405 were activated during both normal APAs and APA errors. 406

DISCUSSION

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Our present results showed that an error in motor 408 programming during gait initiation in healthy subjects 409 was not associated with obvious differences in ERPs. 410 We only observed a longer P300 peak latency in trials 411 with an APA error. However, we observed extended 412 beta ERD over the sensorimotor cortex, and more 413 pronounced alpha ERS followed by an ERD in trials with 414 an APA error. 415

Are APA errors low-level errors?

It is known that the motor program can be adjusted during 417 APAs. This process might involve rapid, direct 418 sensorimotor loops via visual afferences for stimulus 419 detection and proprioceptive afferences for the ongoing 420 APA. Hence, healthy subjects are able to adjust the 421 motor program after it has started by delaying foot lift 422 until the correct motor program has been selected. 423 Response inhibition - the ability to rapidly cancel an 424 action - is a critical component of executive function. In 425 gait initiation, response inhibition guickly corrected APAs 426 initiated in the wrong direction (around 200 ms after the 427 start of the APA error (Tard et al., 2015)). This means that 428 subjects can react to the perception of conflicting informa-429 tion and quickly reorient ongoing actions. Many research-430 ers have investigated the neural substrates of behavioral 431 inhibition by applying laboratory tasks based on the stop-432 signal paradigm and that require a planned action to be 433 stopped (Duque et al., 2017); however, these tasks 434 require complete inhibition of the motor program, rather 435 than just correction (as in APA errors). 436

Moreover, the different types of errors described in the 437 literature appear to have different neural bases: Hill and 438 Raab (Hill and Raab, 2005) first distinguished the correc-439 tion of errors induced externally and internal errors gener-440 ated by the subject itself. Another distinction has been 441 made between low-level errors (i.e. non-conscious, 442 quickly corrected errors) involving posterior regions of 443 the brain, and high-level errors (i.e. conscious errors that 444 are not always corrected) involving the medial frontal lobe 445 (Krigolson and Holroyd, 2007). The errors in our study 446 would be classified as internal, low-level, since the partic-447 ipants were not aware of them; although some partici-448



500-600 ms

Fig. 3. Event-related potentials in normal Anticipatory Postural Adjustments (APA) and APA error. Top view of topographic voltage maps for each condition (i.e. a normal APA-N- or an APA error-E-). 0 ms corresponds to either the target presentation (S2), target-locked (T), or to the start of the APAs (response-locked:R). Cold and hot colors correspond to negative and positive ERPs, respectively. Red dots correspond to electrodes with differences between conditions (whatever the direction), as indicated by permutation tests with FDR correction. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

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pants described "start hesitation" in a few trials, most 449 were unaware of the quickly corrected motor program. 450

Cortical activations during unexpected events have 451 recently been reviewed (Wessel and Aron, 2017). Most 452 of the relevant studies were based on the use of stop-453 signal paradigms and fMRI to identify the underlying 454 motor inhibition system. The right inferior frontal cortex, 455 pre-supplementary motor area and subthalamic nucleus 456 (STN) of the basal ganglia are all involved, with down-457 stream effects on the pallidum, thalamus, and primary 458 459 motor cortex. Indeed, the STN's role has been emphaand then appropriate movement execution. With regard to response checking, the participant had to recognize the engaged motor program (the left or right foot) usually a non-conscious process - and determine whether or not it corresponded to the appropriate response. This process is much the same in APA errors and normal APAs.

We did not observe any differences in the ERN/CRN amplitudes. The amplitudes of CRN and ERN were also similar for the stepping task in (Watanabe et al., 2016b) that used a Simon task to elicit APA errors. They pro-

sized by several fMRI studies 460 (Aron and Poldrack, 2006; Li 461 et al., 2008); activity in the STN is 462 greater during both stop successes 463 and stop errors than in "go" trials, 464 and greater for stop errors than 465 for stop successes (Li et al., 466 2008). These findings suggest that 467 the STN has a role in suppressing 468 thalamocortical output, which 469 thereby blocks motor response 470 execution via a hyperdirect path-471 way (Aron and Poldrack, 2006). 472 The mechanisms in our paradigm 473 were less clear, since error correc-474 tion re-oriented a movement rather 475 than stopping it completely as in 476 stop-signal paradigms. Subjects 477 were able to shift their weight 478 toward the swing leg to correct 479 the error. This could correspond 480 to "partial errors", i.e., incorrect 481 activities which are not sufficient 482 produce overt errors to 483 (Carbonnell and Falkenstein, 484 2006). These latter could produce 485 both ERN and CRN. However, we 486 would have expected more ample 487 negativity over frontal regions in 488 case of APA errors. 489

Cortical markers of error monitoring

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Evaluating the precise timing of 492 cortical activations requires 493 electrophysiological recordings 494 based on local field potentials or 495 EEG. It is generally thought that 496 an anterior component (ERN/Ne) 497 reflects error inhibition (Kopp 498 et al., 1996), conflict detection 499 (Carter et al., 1998) or the 500 comparison (response checking) 501 of the neural representation of the 502 actual (erroneous) response and 503 the representation of the required 504 (i.e. correct) response. In our 505 paradigm, however, successful 506 error inhibition was followed by a 507 motor programming correction 508



Fig. 4. Event-related potentials in normal Anticipatory Postural Adjustments (APA) and APA error in midline derivations. ERP for Fz, Cz and Pz sites in two conditions: blue: grand average of normal APA vs. green, grand average of APA error. 0 ms corresponds to either the target presentation (S2), target-locked, or to the start of the APAs (response-locked). Target-locked: P300 occurred later in case of APA error. Response-locked: negative components (?Ne and CRN) were observed, later in case of APA error. P300 or Pe occurred later in case of APA error. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

520 posed different interpretations: the more convincing one in our opinion is that APA errors are brief, covert, and 521 are likely corrected by initiating a step with the appropriate 522 leg. As a consequence, ERN amplitude may have 523 become smaller because APA errors were not recognized 524 as definite errors. Once again, this error is not conscious. 525 The late, posterior, positive ERP component observed 526 after presentation of the target is more difficult to interpret. 527 Considering target-locked responses, the ERP peaked in 528 the parietal cortex at about 400 ms (target-locked) and 529 100-200 ms (response-locked). It might therefore be a 530 531 P300 component. It is very similar to the responselocked posterior component. The latency (but not the 532 533 amplitude) differed according to the presence or not of an APA error. The absence of a difference in amplitude 534 was not very surprising because P300's amplitude is pri-535 marily modulated by the stimulus's rarity (as in the oddball 536 paradigm). Here, only incongruent targets were consid-537 ered. They elicited a large P300, regardless of the forth-538 coming motor preparation (Neuhaus et al., 2010; Deiber 539 et al., 2013). Alternatively, the ERP component might cor-540 respond to Pe. The latter is thought to reflect (i) error cor-541 rection, (ii) a delayed parietal P300 (since it is present in 542 correct trials) or (iii) additional error processing or post-543 error processing (for a review, see Falkenstein, 2010). 544 Here, the distribution is posterior and not anterior. More-545 546 over, in our paradigm, there were no amplitude differ-547 ences between trials with and without APA errors. It 548 must be borne in mind that the variability in Pe depends on error detectability: the larger the difference between 549 the representations (i.e. the easier the error is to detect), 550 the larger and/or earlier the Pe. There are several possi-551

ble explanations for the lack of difference in the amplitude of Pe. The participant was not given any information about APA error monitoring. Indeed, both Ne and Pe are closely related to conscious perception of the error (Nieuwenhuis et al., 2001; Charles et al., 2013). In fact, in the work by Charles et al., the ERN was absent only when subjects reported that they did not see the target. That was not the case in our study since subjects well identified the target (no error of step side).

Instructing the participant to pay attention (or not) to the error stimulus (Ramautar et al., 2006) can also amplify the ERP. In Ramautar et al.'s study, Pe was much more pronounced for perceived errors than for unperceived errors. We suggest that these ERPs reflect cognitive processing of the stimulus (i.e. incongruent flankers surrounding the arrow) more than perception of the APA error. These scalp ERPs did not seem to be relevant for studying the non-conscious monitoring of

an ongoing action, when the error was corrected online before the possible erroneous outcome (i.e. initiation with the wrong foot). According to Krigolson and Holroyd, the P300 component has a role in the online control process for low-level errors (Krigolson and Holroyd, 2007). In a corrective limb adjustment task using a joystick (in which the target's location changed unexpectedly following movement onset, in order to elicit errors), the researchers concluded that if P300 arises after behavioral changes associated with the online control of movement, then it cannot be involved in the evaluation of target errors (Krigolson et al., 2008). Indeed, the P300 started after the participants had begun to adjust their motor output to accommodate the target perturbation. Moreover, Krigolson and Holroyd did not observe a difference in amplitude according to the presence or absence of correction. As suggested by Krigolson et al. (Krigolson et al., 2008), we hypothesize that P300 reflects the updating of an internal model of the movement environment processing of flankers, for example (Donchin and Coles, 1988).

Time–frequency analysis and motor programming

Non-phase-locked (induced) changes can be studied in a 604 time-frequency analysis, which highlights the cortical 605 oscillations related to an external or internal event 606 (Rektor et al., 2006). Indeed, motor-related cortical oscil-607 lations are generally assessed by guantifying increases or 608 suppressions in spectral power. For example, increases 609 in amplitude of the cortical oscillations in the delta-band 610 (2-4 Hz) and the gamma (bands 60-200 Hz) are 611

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Fig. 5. Event-related desynchronization/synchronization (ERD/ERS) in normal Anticipatory Postural Adjustments (APA) and APA error. Top view of spectral maps (in dB) in the 4–7 Hz, 8–12 Hz, 13–20 Hz and 20–30 Hz bands for normal APAs (N) and APA errors (E). Target-locked (T) and response-locked (R) are shown. 0 ms correspond either to the target presentation or to T0 (start of the APAs). The color at each image pixel indicates the power (in dB) of a given frequency band. Hot colors correspond to an increase in power relative to the baseline, and cold colors correspond to a decrease in power. Theta, alpha ERS were observed in both conditions after target presentation. Beta ERD was observed in both conditions but was more prolonged for APA errors. The low-beta ERD was less intense for APA errors (target-locked only). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)



Fig. 6. Event-related desynchronization/synchronization (ERD/ERS) in normal Anticipatory Postural Adjustments (APA N) and APA error (E) over Cz. ERD/ERS in the different frequency bands over Cz in trials with a normal APA (N top row) or an APA error (E second row). The line indicates the target presentation time (target-locked: T) or T0 (start of the APAs, response-locked: R). ERD is shown in blue, and ERS is shown in red in dB. We observed alpha ERS during APAs more intense for APA errors, beta ERD during the normal APA and during the APA error. This ERD was longer (targetlocked, response-locked) and less intense (target-locked) in the low-beta-band for APA errors. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

observed during both the planning and execution of 612 613 movement (Combrisson et al., 2017). The initiation of voluntary movements has also been linked to desynchro-614 nization of cortical activity in the alpha-band (8-12 Hz) 615 and the beta-band (13-30 Hz) in electrocorticography 616 and scalp EEG recordings (and then decrease of ampli-617 tude of oscillations in the corresponding frequency band) 618 over the motor and premotor cortex (Pfurtscheller, 1981; 619 Neuper et al., 2006). 620

Firstly, we observed theta synchronization and a more 621 622 pronounced alpha synchronization in case of error. Secondly, beta ERD was observed over sensorimotor 623 cortex. 624

Theta-band ERS is linked to an alert effect (Luu et al., 625 2004) and to stimuli monitoring and discrimination (Wang 626 et al., 2005) engaged in attentional processes (Luu et al., 627 2004; Fan et al., 2007; Song et al., 2014). Alpha ERS 628 (coupled with theta ERS) at the start of APAs was fol-629 630 lowed by stronger posterior alpha ERD in case of APA error. This particular pattern has been previously 631 described during different variants of the Simon task dur-632 ing errors in an upper limb task (van Driel et al., 2012) and 633 634 seems consistent. These patterns according to the vari-635 ants of the attentional task (in terms of amplitude, loca-636 tion, coupling between different cortical regions) were influenced by performance monitoring (theta ERS), error speed processing (posterior alpha ERD) (van Driel et al., 2012). Increasing alpha ERS is supposed to facilitate the goal-directed behavior (Dockree et al., 2007) by reflecting active mechanisms of sensorv suppression for irrelevant-task stimulus (Foxe and Snyder, 2011) which could be particularly useful for proper motor execution in case of APA errors triggered by incongruent stimuli. Then, alpha ERD (response-locked) is strongly associated to visual perception and involved in task anticipation to modulate the excitability in human parieto-occipital cortex (Capotosto et al., 2017). Alpha ERD is therefore stronger in trials with APA errors in order to facilitate the modulation of the sensorimotor cortex with stimulus' perception. It signs the further orientation and maintenance of visual attention (posterior predominant. engagement of parieto-occipital areas (Foxe et al., 1998)).

Beta ERD is thought to reflect the activation of regions engaged in visuospatial attention or motor execution (Pfurtscheller and Lopes da Silva, 1999). During tasks requiring attention, significant increases in the delta-, thetaand gamma-bands have been

reported during the planning phase and especially during execution. In contrast, alpha, beta and low-gamma power falls after an execution cue (Combrisson et al., 2017). In our paradigm, the occurrence of beta ERD following the appearance of the target was consistent with this pattern. Oscillations in the beta- and gamma-bands during motor preparation have been studied in paradigms that compare successful stops with unsuccessful stops (Swann et al., 2009, 2012). Overall, brief beta ERS is followed by longer beta ERD. The latter is more pronounced in successful stop trials. These findings provide insight into our results - even though our study did not feature successful vs. unsuccessful stops. In trials with APA errors, we observed prolonged beta ERD over the sensorimotor cortex; this probably reflected the fact that movement preparation was longer when an APA error occurred (Cohen et al., 2011). We also observed less intense low-beta ERD during APA errors. In a study evaluating a shifting cognitive task during gait (Wagner et al., 2016), two different beta oscillations were noted: beta ERD (expressing motor execution and motor readiness related to gait movements) and a frontal beta ERS (related to cognitive top-down control on gait). The less intense ERD in the APA error condition might be due to the summation of concomitant ERD

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Fig. 7. Source localization of cortical activity in the 0- to 600-ms time window N: normal APAs; E: APA errors. The source localizations were the same in both conditions.

(for movement preparation) and ERS (for the cognitive 698 load, with a complicated process for conflict resolution 699 and correction of the engaged motor program). Further-700 more, beta frequency oscillations are associated with 701 702 the maintenance of the "status quo", such as holding a fixed position. In the event of an APA error, the status 703 quo is disrupted. For example, stabilized gait on treadmill 704 is characterized by beta-band desynchronization that 705 lasts for at least 15 steps (Wagner et al., 2016). Here, 706 the presence of differences in low-beta ERD during an 707 APA error suggested that the status quo had been dis-708 rupted (Engel and Fries, 2010). However, low-beta mod-709 ulations have been studied more frequently in cognitive 710 tasks (including memory tasks) than in motor tasks. Other 711 spectral properties (such as phase and phase-amplitude 712 coupling) have also been shown to carry information with 713 714 regard to the oscillatory dynamics underlying motor pro-715 cesses, and an analysis of these variables might be of 716 value in studying network dynamics during low-level errors (Combrisson et al., 2017). There was a contradic-717 tion between the findings of ERN and theta/alpha ERD/ 718 S. Contrary to ERP, alpha synchronization appeared in 719 this study to be more sensitive, probably because of its 720

time resolution and the relative 721 specificity of the different fre-722 quency bands to monitor brief cov-723 ert errors such as APA errors. 724 Indeed, previous studies (Kolev 725 et al., 2005, 2005; Albrecht et al., 726 2009) found modulations in low-fre-727 quency band (delta, theta) and 728 ERN in different paradigms (choice 729 reaction task, flankers...) provok-730 ing overt errors in different popula-731 tions but once again, APA errors 732 are of different nature. 733

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Limitations

There is a strong relationship between the ERP and ERD/ERS. However, ERD and ERS are not systematically time-locked to the target ERP. Theta- and deltaband ERSs correspond most closelv to N200 P300. and respectively (Huster et al., 2013). In fact, an increase in the amplitude of P300 is invariably associated with an increase in power in the low-frequency bands - as observed in the present study. Removing or not the ERP signal of the time-frequency analysis is still subject to debate. In a nonpresented analysis, we removed the ERP signal from the EEG before performing event-related spectral analysis in the beta-band; this enabled us to study the induced response alone and not the evoked response: no clear dif-

ferences (topographical or statistical maps) were observed after the removal.

The attention network task enabled the study of 760 different components of attention (Fan et al., 2002). How-761 ever, trials with cues might influence the RTs or the pro-762 portion of pre-APAs (i.e. APAs occurring between the 763 cue and the target presentation but not followed by a step, 764 with a return to the baseline posture at the moment of tar-765 get presentation (Tard et al., 2015)); however, the propor-766 tions of valid, invalid cued and uncued trials did not differ 767 after removal of trials with artifacts. The proportion of no 768 cue trials was 17.8% (normal APA) vs. 16.7% (APA error) 769 vs 16.3% (before removal of trials with artifacts), the pro-770 portion of neutral cue trials was 19.3% (normal APA) vs. 771 18.2% (APA error) vs. 20.3% (before removal of trials with 772 artifacts), the proportion of valid cue trials was 47.5% 773 (normal APA) vs. 47.4% (APA error) vs. 47.3% (before 774 removal of trials with artifacts), the proportion of invalid 775 cue trials was 15.4% (normal APA) vs 17.6% (APA error) 776 vs. 16% (before removal of trials with artifacts). Further-777 more, we excluded false start trials (i.e. those with an 778 RT < 100 ms). The median number of trials by subject / 779 condition after removal of artifacts was 49 (20-98) for 780

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APA errors and 69 (38–107) for normal APAs. One subject has a very low number of trials taken into account but the ERP was clearly identifiable in this particular case.
The subject was thus not excluded.

Our present results highlighted a cortical marker of 785 gait initiation APA errors in healthy subjects. Differences 786 in sensorimotor activation (reflected by differing alpha/ 787 788 beta-band ERS/ERD patterns) were observed during APA errors. It remains to be seen how these cortical 789 oscillations are influenced by cortical-subcortical loops. 790 Future research should consider the role of the basal 791 ganglia (and specifically the STN) in movement 792 inhibition. In contrast to ERP analysis, time-frequency 793 794 methods are useful for monitoring non-conscious errors. These methods could also be used to monitor motor 795 programming errors (in patients with dysexecutive 796 syndrome, for example), and could be implemented in 797 brain-computer interface algorithms. 798

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800 Capotosto et al. (1991).

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