# **Motion P3 Demonstrates Neural Nature of Motion ERPs**

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*Abstract*—The technical challenges of recording electroencephalographic (EEG) data during motion are considerable, but would enable the possibility of investigating neural function associated with balance, motor function and motion perception. The challenges include finding a reliable method of motion stimulus reproduction, removing artifacts, and ensuring that the recordings retain sufficient EEG signal for proper interpretation. This study details the use of the P3 waveform to validate the concept of motion-based EEG data, and discusses some potential future uses in experimental and clinical settings.

#### I. INTRODUCTION

The proper perception of self-motion is important in everyday function - walking, standing and sitting rely on knowing whether and how much the body is in motion. The vestibular system is very important in the perception of selfmotion, as it detects the body's acceleration. Heading, or direction, is a fundamental feature of motion and recent studies have shown that the vestibular system plays a pivotal role in the perception of self-motion heading in humans (e.g. [1, 2]). There have been a number of advances in the understanding of the neuronal processing of visual and vestibular signals for self-motion perception in non-human primates [3-6]. However to date, there has only been very few studies investigating the neural correlates of self motion perception in humans. Electroencephalography (EEG) is the most suitable candidate for these purposes as invasive recordings are often impossible, and other non-invasive methods such as function magnetic resonance imaging (fMRI) or positron emission tomography (PET) require

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Heinrich H. Bülthoff is with Department Bülthoff, Max Planck Institute for Biological Cybernetics, Tübingen, Germany, and the Department of Brain and Cognitive Engineering, Korea University, Seoul, South Korea (email: heinrich.buelthoff@tuebingen.mpg.de). bulky equipment which does not lend itself to motion studies.

These studies have primarily investigated rotational selfmotion [7]-[10]. These have been seen to have some potential for clinical research [11] but problems such as electrooculogram (EOG) and electromyogram (EMG) artifacts from rotational stimuli are frequently reported [8, 9, 11]. Advances in signal processing methodologies such as independent component analysis (ICA) now allow the attenuation of such artifacts. Previously we have shown that it is feasible to acquire EEG responses to auditory stimuli during continuous linear self-motion on a Stewart platform without noticeable electromagnetic or EMG interference [12]. Building upon these results, here we investigate the possibility of recording EEG responses to a linear motion stimulus.

We have chosen heading as the motion feature to manipulate, and the P3 event-related potential (ERP) component as the EEG feature to evoke. To do this we have adapted the classical two-stimulus oddball paradigm [13], in which participants are presented a stream of frequent standard stimuli (80%) and infrequent target stimuli (20%) and are instructed to respond to the target stimuli. The target stimulus elicits the P3 component, which is a response to a new stimulus. The P3 component is a well-studied component evoked by observing change in the experimental environment; it has been observed in multiple stimulus modalities: visual, auditory, somatosensory [14] and olfactory [15].

In this study, participants were seated upon a Stewart motion platform while EEG was acquired using a 128-channel system. The standard and target heading stimuli were forward translations at a 45° angle to the left or right of straight ahead. The goal of our study was to investigate the neural correlates of self-motion perception using high density EEG to characterize vestibular processing of heading change.

#### II. METHODS

#### A. Subjects

Sixteen subjects with normal or corrected-to-normal vision participated in the experiment. The age range was 22 - 35 (mean 28.1 ± standard deviation 3.9). Subjects gave their informed consent before taking part in the experiment, which was performed in accordance with the ethical standards laid down in the 1964 Declaration of Helsinki. One subject's data

was particularly noise-contaminated and was removed from the study.

## B. Apparatus

A Maxcue 600 platform manufactured by Motion-Base PLC [16] was employed. This is a 6-legged Stewart platform with 6 degrees of freedom, which are rotation and translation about 3 axes. A fixation cross was displayed on a projection screen, with a field of view of  $86^{\circ} \times 65^{\circ}$ , a resolution of  $1400 \times 1050$  and a refresh rate of 60 frames per second. Subjects wore Sennheiser HD600 headphones with one-way communication capability while white noise was played to mask the sound of the platform. EEG data were recorded at 512Hz using a Biosemi ActiveTwo<sup>TM</sup> 128 channel EEG system with 7 supplementary electrodes recording EOG and reference channels. Data were referenced to the average reference.

#### C. Motion Paradigm

The classical oddball paradigm, in which an infrequent target is identified in a sequence of frequent standard stimuli [17] was adapted for a bi-directional motion stimulus.

Subjects were presented with sequential angled forwards motions; the angles were either 45 degrees left or 45 degrees right. Left and right were presented in an oddball manner, with random presentation probabilities of 0.8 for standard and 0.2 for target. The use of left and right for the standard and target was balanced across runs. The return to center after each stimulus was of sub-threshold amplitude. There was a random interval of 1.5 - 2.5 s after the return. There were a total of 8 blocks, each consisting of 50 trials – 40 standards and 10 targets – yielding a total of 320 standard stimuli and 80 target stimuli per subject. Each block lasted approximately 5 minutes. All subjects were given two breaks lasting 5-10 minutes.

The ideal motion displacement profile was:

$$s(t) = 0.49 \frac{(2\pi t - \sin(2\pi t))}{4\pi^2}$$
  $0 < t \le 1s$ 

where t is time. It had a maximum displacement, velocity and acceleration of 0.078 m,  $0.156 \text{ ms}^{-1}$  and  $0.49 \text{ ms}^{-2}$  respectively, lasting 1s, which are above the detection thresholds reported in [18].

# D. EEG Processing

All data were high-pass filtered at 1 Hz and low-pass filtered at 95 Hz for processing, with a bandstop filter at 47 - 53 Hz. The FASTER processing method [19] was used to preprocess data and remove artifacts. Epochs of 1000ms with 500ms pre-stimulus baseline were extracted from the continuous data; there was a mechanical delay of approximately 200ms from the trigger onset to the onset of the motion.

#### E. Analysis

The goal of this study was to determine whether motionbased EEG responses were valid. The P3 is known to be centered on the midline between electrode sites Cz and Pz, depending on the modality. It is known to occur



Figure 1. (Left) The outside of the Stewart platform. (Right) A subject with EEG electrodes in place, ready to perform the task. During this study, a cloth sheet covered the platform to mask visual motion cues, and a Biosemi ActiveTwo EEG system was installed to record EEG data.

approximately 300 - 400 ms after stimulus onset in healthy adults. Thus the spatial regions of interest were taken as electrode sites Pz, Cz and Fz along the midline. Temporal regions of interest were pre-identified as 300ms – 800ms after stimulus onset, to allow for the non-discrete nature of the motion stimulus. Due to the wide peak observed from the ERPs, the target ERP was also epoched relative to the button press to remove some of the inter-subject temporal variability [20].

#### III. RESULTS

# A. Behavioral Results

Reaction times to the target stimulus were measured, along with hit rates.

Measure	Value (mean ±	Intra-participant
	SD)	SD
Reaction time, overall (ms)	876 ± 214	57 ± 24
Reaction time, right (ms)	872 ± 223	67 ± 34
Reaction time, left (ms)	886 ± 221	61 ± 51
Hit rate, right (%)	$95.14\pm5.60$	
Hit rate, left (%)	$93.91\pm9.36$	

Table 1. The group average and standard deviation of the reaction times and hit rate for the target. Hit rates were calculated over all runs rather than per-run, and so there is no intra-participant SD.

A paired t-test of left vs. right values showed no significant difference between left (mean=93.91%, SD=9.36%) or right (95.14%  $\pm$  5.60%) headings for true positives (p=0.38), false positives (p=0.76), or left (886  $\pm$  221) and right (872  $\pm$  223) reaction times (p=0.14).

### B. EEG Results



Figure 2. (Left) Group average standard (gray) and target (black) ERPs from (A) Fz, (B) Cz and (C) Pz after epoching relative to motion onset. (Right) Group average target ERPs from (A) Fz, (B) Cz and (C) Pz after epoching relative to the button response.

Group average ERPs computed from the standard (grey) and target (black) conditions with respect to motion onset and button press are shown in Figure 2. As participants did not respond during the standard condition, there is no standard condition for the button press ERPs. Inspection of the group average ERP-waveforms, in Figure 3, showed a clear positive response at approximately 600ms at the parietal midline site (Pz).

Statistical analysis of the amplitudes at electrode sites Fz, Cz and Pz, identified from the P3 literature, showed that the target peaks at ~600ms differ significantly from standard peaks (t(14) = 3.89, t(14) = 3.22, t(14) = 4.61, respectively, p < 0.01 for all).

Figure 3 shows butterfly plots of the group average standard (A), target (B) and button press (C) waveforms. These figures display each channel's ERP overlaid, and scalp topographies at three time periods, including the largest positive peak. These topographies represent two time periods of similarity between standard and target waveforms at 300 ms and 350 ms, and the P3 peak at 600ms which is seen to differ between conditions.

The results indicate that the elicited response contains the P3 waveform.



Figure 3. (Left) Butterfly plots repicting (A) Standard, (B) Target and (C) Target after epoching relative to button response. (Right) Target ERPs from (A) Fz, (B) Cz and (C) Pz after epoching relative to the button response.

#### IV. DISCUSSION

In this study we investigated the feasibility of recording motion-based EEG responses using a basic oddball paradigm. The results display a typical P3 topography (see [21]) with significant peak amplitude increase at 600ms. This provides strong evidence that we have successfully recorded a P3 response to a motion stimulus without significant EMG or EOG interference.

While this result may initially seem straight forward, it should be emphasized that there were a number of potential factors which may have interfered with proper EEG recording. The foremost of these is EMG interference – the possibility that the ERPs recorded were in fact capturing muscle movement of the neck to stabilize head in response to

the full-body motion via the vestibulo-collic reflex. However, the P3 component occurs only in the target stimuli which are directionally different but of the same amplitude, and so the increase in amplitude cannot be considered to be related to neck muscle movement. Furthermore, frequency analysis showed no increase in high-frequency activity which would be characteristic of EMG responses. Topographically, such stabilizing EMG responses would be likely to occur temporally, while here the P3 is seen to be active in the parieto-central regions. For these reasons, we conclude that the ERPs are not driven by EMG activity.

Due to the vestibular-ocular reflexes, the possibility of contamination of the EEG by EOG signals was also a possibility. Such signals are characterized by high-amplitude frontal topographies, which are not seen the ERP responses recorded here. Furthermore, it has been seen that FASTER, the artifact reduction method employed in this study, is highly effective in removing ocular contamination [19].

A further result of the study arises when considering the differences between the target ERP after epoching with respect to motion onset and after epoching with respect to the button-press. Due to the sinusoidal nature of the stimulus and the inherent ramp-up of intensity that comes with it, detection of motion onset is not instant. The wide peak seen in the P3 in the motion-onset epoched ERP is considerably sharper in the button-press epoched ERP. This implies that the inter-subject variability of motion detection threshold is high, as the sharper P3 peak from each subject is spread across time when averaged, resulting in the wide peak. This concept is supported by the behavioral data where the inter-subject variability in response time is 214ms, while the intra-subject variability is only 57ms. This observation should be taken into account in further motion-based EEG studies.

The consequences of these results are far-ranging. As the vestibular system contributes strongly to motion perception [1], the possibility of testing vestibular function experimentally, to enhance understanding of the neural correlates of vestibular function, and clinically, for aiding diagnosis in vestibular dysfunctions. Furthermore, the investigation of the multisensory contribution of visual, somatosensory and auditory senses to motion perception – which is currently poorly understood – is also a possibility. This may be particularly important in research in aging – falls are the most common injury for elderly people [22], and as multisensory integration has been seen to become inefficient with age and may be implicated in falls [23], investigating the multisensory elements of motion perception may lead to better prevention of falls.

In conclusion, this study has demonstrated a P3 component elicited using a motion-based stimulus, without interference from reflexive EMG or EOG. This provides a strong grounding for further EEG-based studies into the processing of full-body motion.

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