Effects of sensitive electrical stimulation based cueing in Parkinson’s disease: a preliminary study
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Effects of sensitive electrical stimulation based cueing in Parkinson's disease: a preliminary study

Benoît Sijobert (1), Christine Azevedo-Coste (1), David Andreu (1), Claudia Verna (2), Christian Geny (2)

(1) INRIA - LIRMM Université de Montpellier, Montpellier, France; (2) CHU Montpellier, Montpellier, France

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Abstract

This study aims to investigate the effect of a sensitive cueing on Freezing of Gait (FOG) and gait disorders in subjects suffering from Parkinson’s disease (PD). 13 participants with Parkinson’s disease were equipped with an electrical stimulator and a foot mounted inertial measurement unit (IMU). An IMU based algorithm triggered in real time an electrical stimulus applied on the arch of foot at heel off detection. Starting from standing, subjects were asked to walk at their preferred speed on a path comprising 5m straight, u-turn and walk around tasks. Cueing globally decreased the time to achieve the different tasks in all the subjects. In “freezer” subjects, the time to complete the entire path was reduced by 19%. FOG events occurrence was lowered by 12% compared to baseline before and after cueing. This preliminary work showed a positive global effect of an electrical stimulation based cueing on gait and FOG in PD.

Key Words: Freezing of gait, Parkinson’s disease, electrical stimulation, cueing, gait analysis, inertial sensors
where they otherwise had problems with FOG. From the previous statements, we decided to design a protocol based on electrical stimuli cueing.

In both on and off conditions (under medication or not), turning phase has been demonstrated as the most frequent trigger of freezing of gait in Parkinson’s disease. This is also what we observed in our experiments. Plotnik et al. tried to explain this occurrence by the asymmetric nature of the task which would increase interlimb synchronization difficulties. Crenna et al. showed it could possibly be related to head rotation. Patients in the early stage of the disease starting their head rotation later than controls while turning. In Nieuwboer et al. work, authors chose to focus their work on different cueing modalities on turn speed only. In addition, as shown in, evaluating freezing of gait poses difficulties as its likelihood to happen highly relies on environmental triggers, cognitive input and medication. In order to increase FOG occurrence during experimentations, we designed an experimental path including a maximum of turning phases. Numerous studies confirmed Parkinson’s disease motor deficits are associated with proprioceptive impairment. In Vaugoyeau et al. work, the authors subjected standing subjects to small angular sinusoidal perturbations applied to a supporting platform and asked them to maintain verticality. In the absence of visual cues, the PD subjects were clearly unable to use proprioceptive information as feedback to control their body verticality and stabilize the body segments, resulting in blocking head and shoulders. The same strategies have been observed during their gait. The authors concluded sensorimotor integration deficits partly account for the postural and locomotion impairments observed in PD. Using muscle vibration on the trajectories of voluntary dorsiflexion movements of the ankle joint, Khudados et al. showed that proprioceptive regulation of voluntary movement is disturbed in PD. El-Tamawy et al. used augmented proprioceptive cues during gait on thirty levodopa-dependent PD subjects. They applied vibratory stimuli to the feet plantar surfaces (below the heel and forefoot) through miniature hidden vibrating devices that sent rhythmic vibrations to the skin synchronized with the step in the push off-phase of the gait. Results demonstrated a significant improvement in gait kinematics and angular excursion of lower limb joints. Similarly, Kleiner et al. applied mechanical stimulation (AMPS: Automated Mechanical Peripheral Stimulation Treatment) on four specific target areas in patient’s feet while the subject was laid down and reported a 15% improvement in gait velocity after treatment.

**Materials and Methods**

**Stimulation**

Inspired by Spaich work on hemiparetic gait, we stimulated the arch of the foot as shown in Fig. 1. The stimulation pattern consists in five 1 ms-wide biphasic pulses delivered at 200 Hz, repeated 4 times at 15 Hz. Current amplitude was adjusted in order for the subject to feel the stimulation without any discomfort. Patients with advanced Parkinson’s disease are usually subject to altered gait patterns which makes difficult to segment and to reliably detect gait events or compute

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**Fig 1.** Electrodes location used for electrical stimulation. To deliver stimulation, a self-adhesive electrode (2.6 cm²) is placed as cathode on arch of the foot and a large common anode on dorsum of the foot.

**Fig 2.** Example of stimulation triggering based on Equ 2 from an experimental record. In green and red are respectively the start and stop actual stimulation events triggered by the non-stationary period detection algorithm. Heel off (HO) and swing (SW) phases were manually added to the figure from literature for reference.
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Gait parameters compared to healthy subjects. Different methods have been proposed based on wearable and non-wearable systems using multiple sensors, but our aim has been to conceive a ready-to-use patient-oriented solution, using a minimum amount of sensors and not requiring individual calibration or threshold parameterization for correctly triggering stimulation. In Moore et al. work, the investigators monitored during 75 min subjects suffering from Parkinson. For detecting locomotor activity, they defined periods where the RMS vertical acceleration was greater than 0.4 m/s² above baseline. Hundza et al. proposed a method for accurately and reliably detect gait cycle in PD. They used gyroscope angular rate reversal to identify the start of each gait cycle during walking. By interpolating zero-crossing of angular rate from a foot mounted IMU, they detect the termination of forward swing (TOFS) and consider it as physically close to heel strike. They define it as the start and end point of the stride time for each gait cycle. In our case, the strategy was to determine the feasibility of using an inertial sensor as a heel switch alternative, in order to trigger and adapt the stimulation. We aimed to detect stationary periods from the foot mounted IMU combining accelerometer and gyrometer measurements. For defining lowest sensibility thresholds, we had to firstly filter raw inertial data. As the latency was a crucial parameter, we chose to use an Exponential Moving Average (EMA, low pass, Infinite Impulse Response - IIR) filter. At any time, output of the filter is a weighted sum of the new sensor value and the old filter output. Filter coefficient controls the filtering effects:

\[ gX_{\text{filt}} = (1 - \alpha) \cdot gX_{\text{filt-old}} + \alpha \cdot gX_{\text{raw}} \]  

(1)

with \( \alpha \in [0,1] \)

Our data being processed at 100Hz, we designed a low pass filter of order 1 type butterworth with an attenuation of 3dB at a cutoff frequency of 5Hz using \( \alpha = 0.1367 \) with only one sample late.

In motionless situation, the acceleration norm is supposed to be around \( \text{acc}_{\text{th}} = 9.81 \text{ m.s}^{-2} \) (depending on sensor quality and calibration). From gyrometer angular profile, we determined a magnitude threshold, which is the limit between foot flat phase and heel off phase (\( \varpi_{\text{th}} \approx 30 \text{ deg/s} \)). Based on Equ. 2, stimulation was triggered when a non-stationary period was detected.

\[
\begin{align*}
\text{if } & \left\{ \begin{array}{l}
\text{norm(accel)} \leq \text{acc}_{\text{th}} \\
\text{and} \\
\text{norm(gyro)} \leq \varpi_{\text{th}}
\end{array} \right. \\
\Rightarrow & \text{stationary state}
\end{align*}
\]  

(2)

As shown in Fig 2, post-processing of angular speed combined with stimulation start-stop events clearly demonstrates that stimulus was sent during heel-off phase, which was the initial goal.

**Fig 3.** The subject is equipped with an inertial measurement unit (a) and a wirelessly programmable stimulator (b) connected through a PC.

**Fig 4.** Five meters experimentation path. The subject starts in the middle of the walkway, walks 2.5m, performs a u-turn at the line, goes back 5m, walks around a cone, walks back 2.5m then stops where he started.
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Experimental Protocol

13 subjects with Parkinson’s disease (10 male, 3 female; Age range: 60 to 82 years) participated to the study. The protocol has been approved by the local ethical committee (international identification number NCT02317289). Participants were recruited at the Neurology (Chauliac Hospital) and Gerontology (Balmes Center) departments of Montpellier hospital (CHU Montpellier). All subjects gave their informed written consent. Subjects started from standing in the middle of a gait carpet, equipped with one HikoB© Fox (HikoB© Villeurbanne, France) inertial measurement unit strapped to the foot and a wirelessly programmable electro-stimulator (Phenix© Neo Usb, Montpellier, France) strapped around the shank. Two electrodes were set up on the foot as shown in Fig. 1. After a short familiarization to walk under stimulation, subjects were instructed to walk towards a line draw on the ground, then do a U-turn, walk 5 meters, walk around a cone and keep walking to the start-stop line in the middle of the carpet. The test was repeated five times under the following conditions: no cueing (C0), stimulation cueing (C1), no cueing (C0bis). C0 is considered as the baseline. For eliminating learning bias, we asked the subject to perform a 10-min ecological path (random walk in the hospital) between C1 and C0bis conditions.

Results

In each condition, we analyzed the last three repetitions. Each FOG event, u-turn execution times, 5-meters execution times and times to walk around the cone were quantified from video recordings. In a previous study, we observed that subjects scored with a high occurrence of FOG in daily life were not necessarily those who were the most subject to FOG during clinical experimental protocol. In table 1, we can see subject 2 and 4 were reported as frequent freezers in their daily life meanwhile they did not freeze in C0. Thus we chose to classify the results between subjects who did experienced FOG in C0 and those who did not “freeze”.

In “freezers” group (n=9), we observed that cueing globally decreased of 12% FOG occurrence compared to baseline without cueing (Fig. 5). Table 2 shows cueing’s effects in relation to baselines for the total group and in each subgroup during the different experimental path phases. In every tasks on both “freezers” and “non-freezers” subjects, Table 2 shows that cueing improved gait performances. Considering

![Fig 5. Number of Freezing of Gait events compared between baseline (C0), stimulation (C1) and control baseline (C0bis) on all subjects.](image)

**Table 1 Clinical profiles of subjects who participated in the study.**

<table>
<thead>
<tr>
<th>ID</th>
<th>AGE</th>
<th>DISEASE DURATION</th>
<th>STAGE (H&amp;Y)</th>
<th>AGE OF ONSET</th>
<th>*UPDRS 3.11: FREEZING / 3.10: WALK / GLOBAL PART III</th>
<th>Freezing (Occasional / Frequent)</th>
<th>Falls (Y/N)</th>
<th>*MOCA</th>
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</thead>
<tbody>
<tr>
<td>1</td>
<td>71</td>
<td>5</td>
<td>2</td>
<td>57</td>
<td>1/1/28</td>
<td>O</td>
<td>N</td>
<td>26</td>
</tr>
<tr>
<td>2</td>
<td>63</td>
<td>7</td>
<td>3</td>
<td>53</td>
<td>1/1/28</td>
<td>F</td>
<td>Y</td>
<td>30</td>
</tr>
<tr>
<td>3</td>
<td>71</td>
<td>18</td>
<td>3</td>
<td>53</td>
<td>2/2/40</td>
<td>na</td>
<td>na</td>
<td>na</td>
</tr>
<tr>
<td>4</td>
<td>74</td>
<td>22</td>
<td>3</td>
<td>52</td>
<td>1/2/23</td>
<td>F</td>
<td>Y</td>
<td>25</td>
</tr>
<tr>
<td>5</td>
<td>72</td>
<td>7</td>
<td>3</td>
<td>65</td>
<td>2/2/na</td>
<td>F</td>
<td>Y</td>
<td>27</td>
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<tr>
<td>6</td>
<td>74</td>
<td>8</td>
<td>3</td>
<td>48</td>
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<td>F</td>
<td>Y</td>
<td>12</td>
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<td>7</td>
<td>60</td>
<td>13</td>
<td>3</td>
<td>47</td>
<td>3/na/na</td>
<td>F</td>
<td>Y</td>
<td>25</td>
</tr>
<tr>
<td>8</td>
<td>66</td>
<td>3</td>
<td>4</td>
<td>63</td>
<td>na</td>
<td>F</td>
<td>Y</td>
<td>23</td>
</tr>
<tr>
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<td>76</td>
<td>7</td>
<td>3</td>
<td>69</td>
<td>na</td>
<td>F</td>
<td>N</td>
<td>23</td>
</tr>
<tr>
<td>10</td>
<td>74</td>
<td>10</td>
<td>3</td>
<td>64</td>
<td>2/3/35</td>
<td>F</td>
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</tr>
<tr>
<td>11</td>
<td>66</td>
<td>14</td>
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<td>52</td>
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<td>na</td>
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<td>3</td>
<td>61</td>
<td>2/2/41</td>
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<tr>
<td>13</td>
<td>82</td>
<td>15</td>
<td>3</td>
<td>47</td>
<td>1/3/47</td>
<td>F</td>
<td>Y</td>
<td>26</td>
</tr>
</tbody>
</table>

*UPDRS: Unified Parkinson’s Disease Rating Scale, from 0(normal) to 4(inability)

*MOCA: Montreal Cognitive Assessment, the total possible score is 30 points; a score of 26 or above is considered normal.
all participants, we observed a reduction of 15% in turning time, 14% in 5-m covering duration and 19% in time needed to walk-around the cone. In “freezers” subgroup, turning time is improved by 21%, time to walk-around the cone is reduced of 25% and the duration needed to cover the 5-m walk decreased of 18%. The entire path is completed 19% shorter than baseline.

Discussion

Through this study, we investigated the feasibility of using electrical stimulation as a cueing method in Parkinson’s Disease. The aim was to investigate the capability of this cueing modality to prevent or at least reduce FOG events and to improve gait performances. As partly related to environmental triggers, freezing of gait is difficult to evaluate during a clinical protocol. Based on previous observations and literature, we designed a protocol including turning phases in order to increase FOG events occurrence. This hypothesis has been validated as FOG repartition on all trials was four times more frequent during turning phases than when walking in a straight line. Results show a global positive effect on gait performances, as the time needed to achieve the protocol was considerably shorter with stimulation cueing. “Freezer” subjects tend to be more responsive to cueing, with a turning time improved by 21%. We also observe a 12% decrease in FOG occurrence compared to baseline. However, our subject population was too small for showing a statistically significant effect. Subjects reported no discomforts with electrical stimulation sensation. Some of them expressed an interest in such a possibility to be helped while walking in their daily-life and seemed to accept the additional technological equipment coming with it. We noticed an important range between subjects regarding minimum electrical intensity needed for feeling the stimulus. Such as auditory and visual stimuli in other studies, the electrical cueing responsivity seemed also to be clearly disparate depending on the subjects. Among our 13 participants, stimulation had a strong significant effect on two (respectively 70% of FOG events reduction and a 5-m path 45% shorter compared to baseline), while it did not affect at all some others. In the last case, stimulation cueing never worsen performances or FOG occurrence. In this protocol, the use of an inertial sensor based trigger did not offer much more functionality than a basic heel switch. However, having access to gait kinematics data and to path information from only one sensor could be useful to real-time adapt cueing, when for example a turning phase or a FOG event is detected by the sensor. We could also modulate stimulation or dynamically change the trigger timing. Many other triggering strategies could be investigated and some technical aspects need to be improved for getting rid of some latency problems we experienced during the trials.

In conclusion, this study suggests a new sensitive cueing modality based on electrical stimulation for Parkinson Disease population. Experimental results brought to light a favorable effect in both gait performances and FOG occurrence. Improvements of our cueing strategy have still to be investigated and discussed, but this preliminary work demonstrated encouraging results. A larger study is also required to statistically support these findings and to compare them to other cueing methods usually applied in PD.

Contributions

BS is the principle author and responsible for the algorithm and software design and data analysis. CA and BS carried out the experiments. DA provided his electrical stimulation devices technical expertise for implementing the experimental protocol. CG was the medical doctor. Helped by CV, they were in charge of designing the protocol accepted by the ethical committee, subject evaluation and inclusion. All the

Table 2. Durations (standard deviation) of U-turn, 5-meters and Walk-around phases compared between baseline 1 (C0), stimulation (C1) and baseline 2 (C0bis) for the total group (n=13) and subgroups (freezers and non-freezers in C0).

<table>
<thead>
<tr>
<th></th>
<th>All (N=13)</th>
<th>Non Freezers (n=4)</th>
<th>Freezers (n=9)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>U-Turn Time (s)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline 1 (C0)</td>
<td>3.0 (1.6)</td>
<td>1.9 (0.6)</td>
<td>4.1 (3.0)</td>
</tr>
<tr>
<td>Stimulation (C1)</td>
<td>2.6 (1.1)</td>
<td>1.8 (0.6)</td>
<td>3.4 (2.7)</td>
</tr>
<tr>
<td>Baseline 2 (C0bis)</td>
<td>3.2 (1.3)</td>
<td>2.2 (0.9)</td>
<td>4.1 (3.0)</td>
</tr>
<tr>
<td></td>
<td>Walk Around Time (s)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline 1 (C0)</td>
<td>4.7 (3.5)</td>
<td>2.2 (0.3)</td>
<td>7.2 (3.5)</td>
</tr>
<tr>
<td>Stimulation (C1)</td>
<td>3.8 (2.3)</td>
<td>2.1 (0.4)</td>
<td>5.4 (4.1)</td>
</tr>
<tr>
<td>Baseline 2 (C0bis)</td>
<td>4.7 (3.5)</td>
<td>2.2 (0.3)</td>
<td>7.2 (6.5)</td>
</tr>
<tr>
<td></td>
<td>5m Time (s)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline 1 (C0)</td>
<td>6.7 (1.8)</td>
<td>5.4 (1.3)</td>
<td>7.9 (4.2)</td>
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<tr>
<td>Stimulation (C1)</td>
<td>5.8 (1.3)</td>
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<td>6.7 (2.1)</td>
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<tr>
<td>Baseline 2 (C0bis)</td>
<td>6.6 (1.6)</td>
<td>5.4 (0.8)</td>
<td>7.7 (3.3)</td>
</tr>
</tbody>
</table>
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authors read and approved the final version of the manuscript.

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Conflict of Interest
The authors declare no potential conflict of interests.

Corresponding Author
Benoît Sijobert, INRIA - LIRMM Université de Montpellier, Montpellier, France.
E-mail: benoit.sijobert@inria.fr
E-mails of coAuthors
Christine Azevedo: christine.azevedo@inria.fr.
David Andreu: david.andreu@inria.fr
Claudia Verna: c-verna@chu-montpellier.fr
Christian Geny: c-geny@chu-montpellier.fr

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