Overnight Monitoring of Turnover Movements in Parkinson’s Disease Using A Wearable Three-Axis Accelerometer

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Rec date: Mar 15, 2016; Acc date: May 30, 2016; Pub date: June 2, 2016

Abstract

Background: In patients with Parkinson’s disease (PD), the impairment of voluntary and involuntary movement during sleep might affect their natural sleep and quality of life; however, there is no reliable method to evaluate turnover movements during sleep. We aimed to clarify whether overnight monitoring of turnover movements in bed using a wearable three-axis accelerometer is a feasible and reliable tool for evaluating the impact of motor complications during sleep in PD.

Methods: The number of turnover movements in bed was counted based on the graphic pattern in the X, Y, and Z axis using threshold values in each axis to discriminate turnover movements from other movements mainly associated with respiration or cough. These threshold values were defined by the recordings of various turnover movements in normal volunteers. Overnight monitoring of turnover movements in bed from 9:00 pm to 7:00 am was performed in 7 normal volunteers and 5 patients (mean age, 76.4 ± 4.6 yrs, Hoehn-Yahr stage, 3.6 ± 0.5; duration of disease, 8.8 ± 5.6 yrs). In patients with PD, monitoring was performed before and after adjustment of anti-Parkinson medications.

Results: The number of turnover movements was significantly more restricted in PD before drug control than in control subjects (p=0.005). The median number of overnight turnover movements increased from 0 to 5 times after drug control in patients with PD. The number of overnight turnover movements increased significantly in all 5 patients after adjusting their anti-Parkinson medications (p=0.041).

Conclusion: Overnight monitoring of turnover movements in bed using a wearable three-axis accelerometer is feasible. Further studies are warranted to evaluate the impact of movement disorders during sleep on patients with PD.

Keywords: Parkinson’s disease; Turnover movement; Wearable three-axis accelerometer

Introduction

Turnover movements in bed generally involve elevation of the upper limbs and their movement towards the shoulder, movements of the head and trunk, and raising of one leg [1]. In patients with Parkinson’s disease, movement disorders during sleep might seriously affect their quality of life.

Disability in bed or adjusting bed clothes are evaluated in the Unified Parkinson’s Disease Rating Scale (UPDRS) Part II [2], but these evaluations are based solely on patients’ subjective answers. There is no reliable method to evaluate turnover movements during sleep, and no information is available about the impact of overnight movement disorders in handicapped patients, including those with Parkinson’s disease.

Recently, a wearable three-axis accelerometer has been developed, with which acceleration or angular velocity can be continuously recorded for analysis of human motion, such as three-axis gait analysis in Parkinson’s disease [3,4].

Our goal was to demonstrate the feasibility of evaluating overnight movement disorders using a wearable motion recorder equipped with three-axis acceleration sensors in patients with Parkinson’s disease.

Methods

Measurement of turnover movements in bed

A wearable motion recorder (75 mm × 50 mm × 20 mm, 120 g) equipped with three-axis acceleration sensors (Mimamori-Gait system, MG-M1110-HW, LSI Medience Corporation, Tokyo, Japan) was used to record the motion from the three-axis acceleration data. The motion recorder was fixed on the centre of the abdomen at the navel using a special belt.

Three-axis acceleration data as shown in Figure 1 were collected 100 times a second and serially stored in the micro SD in the recorder through A/D conversion. These data were analysed offline using special commercially available software (MG-M1100-PC, LSI Medience Corporation).
Typical turnover movements in bed are shown in Figure 2. Because a turnover movement in bed is a movement around the Y axis, acceleration in the Y axis direction remains essentially zero. In the 90° turnover movements, i.e., from supine to recumbent position, simultaneous changes in the X and Z axes are observed.

The graphic patterns can be useful to determine the direction of the turnover movements (Figures 2A and 2B). The Y axis value maintained 1.0 gravity (G) during sitting, standing, or walking.

Once the turnover movement occurs, body position remains stable for a variable duration from a few minutes to hours which was defined as turnover interval. Therefore, turnover movements in bed can be counted based on the typical turnover movements followed by a stable period (Figure 3).

However, there is a great variation in the angle of each turnover movement, and slight drift may be observed because of the movement of the abdomen usually associated with respiration or cough. In this study, three-dimensional acceleration data were obtained in various turnover angles at 30°, 60°, and 90°, in 7 normal volunteers (mean age, 28.6 ± 2.4 years old) to define cut-off values to discriminate turnover movements from noise.

Overnight monitoring

Overnight monitoring of turnover movements in bed from 9:00 pm to 7:00 am was performed in the 7 normal volunteers (mean age, 62.1 ± 21.7 years old) and 5 patients with Parkinson’s disease (mean age, 77.0 ± 3.5 years old; Hoehn-Yahr stage, 3.8 ± 0.4, duration of disease, 10.6 ± 5.2 yrs). Then, the number of turnover movements in each recording was calculated, and total bed resting time (minutes) was also calculated by defining the Y axis value as <0.324. Parkinson’s disease was diagnosed according to the United Kingdom Parkinson’s disease Society Brain Bank clinical diagnostic criteria [5].

All patients responded well to dopamine replacement therapy. All monitoring was performed during hospital admission. After completion of overnight monitoring, additional anti-Parkinson medication was started for two to four weeks to control their movement disorders. Then, the overnight monitoring was repeated to assess the effects of treatment on turnover movements in bed. Additional doses of anti-Parkinson medication were expressed as levodopa equivalent doses [6].

All study protocols were approved by the Bioethics Committee of St. Marianna University School of Medicine. Informed consent to participate in the investigation was obtained from all participants after explaining the purposes of the study.

Statistical Analyses

The characteristics of the subjects are given as means and standard deviation (SD), unless otherwise indicated. The Wilcoxon signed-rank test and the Mann-Whitney test were used for data that were not normally distributed. Values of $p < 0.05$ were considered significant.
statistical analyses were performed using SPSS version 21 (IBM SPSS Statistics for Windows; IBM Corp, Armonk, NY).

Results

Turnover movements in normal volunteers

Recording of turnover movements in bed at 30°, 60°, and 90° was performed using a wearable three-axis accelerometer in 7 normal volunteers. Each session consisted of 5 recordings. The mean values and 95% range of acceleration values for the 3 axes are shown in Table 1. Based on the highest value of the range in the Y axis, the threshold value of <0.324 G was adopted as the requirement for resting in bed.

While the Y axis value remained below 0.324 G, turnover movement was defined as the presence of simultaneous changes in both X and Z axes of at least 0.580 G and 0.200 G, respectively, for further studies of overnight monitoring.

<table>
<thead>
<tr>
<th>Turnover Angle</th>
<th>Acceleration Change (G, Absolute Value)</th>
<th>Mean (95% Range)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>X-axis</td>
<td>Y-axis</td>
</tr>
<tr>
<td>30°</td>
<td>0.74</td>
<td>0.178</td>
</tr>
<tr>
<td></td>
<td>(0.580-0.900)</td>
<td>(0.089-0.267)</td>
</tr>
<tr>
<td>60°</td>
<td>0.932</td>
<td>0.189</td>
</tr>
<tr>
<td></td>
<td>(0.828-1.036)</td>
<td>(0.086-0.292)</td>
</tr>
<tr>
<td>90°</td>
<td>1.113</td>
<td>0.245</td>
</tr>
<tr>
<td></td>
<td>(1.014-1.212)</td>
<td>(0.166-0.324)</td>
</tr>
</tbody>
</table>

Table 1: Acceleration changes in turnover movements of 7 control subjects.

Overnight monitoring

The numbers of turnover movements and turnover intervals in 7 normal volunteers during a 10-hour monitoring period from 9:00 pm to 7:00 am are shown in Table 2 and a typical overnight recording in a normal volunteer is shown in Figure 4. Total bed resting time varied from 364 minutes to 596 minutes. Total number of turnover movement during the bed resting time also varied from 4 times to 49 times.

<table>
<thead>
<tr>
<th>Case</th>
<th>Age</th>
<th>Sex</th>
<th>Total bed resting time (minute)</th>
<th>Total number of turnover movement</th>
<th>Median of turnover interval (second)*</th>
<th>Total time of standing or sitting position (minute)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>29</td>
<td>M</td>
<td>364</td>
<td>9</td>
<td>1013 (9-2794)</td>
<td>236</td>
</tr>
<tr>
<td>2</td>
<td>28</td>
<td>M</td>
<td>546</td>
<td>49</td>
<td>232 (5-3128)</td>
<td>54</td>
</tr>
<tr>
<td>3</td>
<td>79</td>
<td>M</td>
<td>530</td>
<td>19</td>
<td>771 (225-3153)</td>
<td>70</td>
</tr>
<tr>
<td>4</td>
<td>73</td>
<td>F</td>
<td>579</td>
<td>27</td>
<td>714 (180-2908)</td>
<td>21</td>
</tr>
</tbody>
</table>

*Median values (minimum-maximum).

Table 2: Parameters of turnover movements in control subjects.

Figure 4: Overnight recording of three-axis acceleration in a normal volunteer.

A 73-year-old woman, a total of 27 turnover movements are observed during the 10-hour recording period.

Figure 5: A 77-year-old patient with Parkinson's disease (Case #4 – showing the three-axis locus chart before and after drug control). Overnight monitoring of a Parkinson's disease patient with a levodopa equivalent dose of 450 mg/day, Hoehn & Yahr Stage 4, duration of disease: 3 years, 77-year-old woman (Table 3, Case #4). Figure 5A shows the three-axis locus chart before drug control, and Figure 5B shows the locus chart after drug control. An increase in the number of turnover movements from before drug control (3 times, LVED 450 mg) to after drug control (5 times, LVED 600 mg) is seen.

The total bed rest times were similar among the control subjects (median; 546 minutes, min.-max.; 364-596 minutes) and patients with Parkinson's disease before (median; 568 minutes, min.-max.; 427-598 minutes) and after the drug control (median; 556 minutes, min.-max.;
480-557 minutes). Numbers of turnover movements in patients with Parkinson’s disease before and after the drug control were significantly fewer than that in the normal volunteers, p=0.005 and p=0.018 respectively (Mann-Whitney test Figure 6). The number of overnight turnover movements in patients with Parkinson’s disease increased significantly after drug control (p=0.041, Wilcoxon signed-rank test, Figure 6).

<table>
<thead>
<tr>
<th>Case</th>
<th>Age</th>
<th>Sex</th>
<th>Disease</th>
<th>Hoehn-Yahr</th>
<th>Levodopa equivalent dose (mg)</th>
<th>Number of movements</th>
<th>Bed resting time (minutes)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Before</td>
<td>After</td>
<td>Before</td>
</tr>
<tr>
<td>1</td>
<td>80</td>
<td>M</td>
<td>17</td>
<td>4</td>
<td>0</td>
<td>116</td>
<td>0</td>
</tr>
<tr>
<td>2</td>
<td>68</td>
<td>M</td>
<td>4</td>
<td>3</td>
<td>660</td>
<td>820</td>
<td>0</td>
</tr>
<tr>
<td>3</td>
<td>76</td>
<td>M</td>
<td>6</td>
<td>3</td>
<td>0</td>
<td>420</td>
<td>8</td>
</tr>
<tr>
<td>4</td>
<td>77</td>
<td>F</td>
<td>3</td>
<td>4</td>
<td>450</td>
<td>600</td>
<td>3</td>
</tr>
<tr>
<td>5</td>
<td>81</td>
<td>F</td>
<td>14</td>
<td>4</td>
<td>1400</td>
<td>1535</td>
<td>0</td>
</tr>
</tbody>
</table>

Table 3: Change in number of turnover movements in patients with Parkinson’s disease before and after drug control.

Discussion

Our study demonstrated that the overnight monitoring of turnover movements using a wearable three-axis accelerometer is feasible. Turnover movements are complex, and although not all turnover movements with various rotational angles are captured, with the present method, at least the turnover movements determined by a change in the accelerometer that records a turnover movement of ≥30° are captured. Numbers of turnover movements in normal volunteers varied from 4 to 49, implying that cut-off value for the diagnosis of abnormally restricted turnover movements could not be determined. However, the patients with Parkinson’s disease (Hoehn and Yahr stages III to IV) in the present study made significantly fewer turnover movements at night than healthy individuals. It is of great importance that no turnover movement was observed in three patients with advanced Parkinson’s disease (Hoehn-Yahr stage IV) and turnover movements were recorded after the adjustment of anti-Parkinson medications. Our methods could be utilized to improve motor complications during sleep in patients with advanced Parkinson’s disease.

Previous research on turnover movements reported that rotational movement in the Y-axis was poor in tetraplegic patients, and turnover movements were difficult in patients with bradykinesia caused by extrapyramidal damage [7]. Moreover, muscle spindles are abundantly distributed in the sub-occipital muscle groups [8], and we expected that, in Parkinson’s disease patients, there would be not only impaired bending of the neck due to rigidity of the sub-occipital muscle groups, but also greater tension in the back muscles, which would hinder flexible movement of the spinal cord and make turnover movements difficult. The presence of glenohumeral rhythm across the entire movable region is also essential for movement of the shoulder joint complex [9]. Thus, we predicted that establishing the rhythm for the turnover movements would also be impaired in Parkinson’s disease patients. Therefore, capturing turnover movements as representing the status of overnight turnover movements can become a new method for evaluating movement disorders in Parkinson’s disease patients.

The present study had several limitations. First, research has shown that there is considerable variation in turnover movements [1], and that there is a bimodal component of either ≤10 seconds or ≥100 seconds in the turnover movements of healthy individuals, but these components disappear in patients with neurodegenerative conditions [10]. Further studies are needed to investigate whether our method of capturing and measuring turnover movements as simple body-axis rotational movements can accurately quantify and serve as a clinically significant evaluation of night-time movement disorders. Second, this study had a small sample size, potentially resulting in insufficient power to detect significant differences.
In conclusion, the present study demonstrated that overnight monitoring of turnover movements in bed by a wearable three-axis accelerometer is feasible. This method could be used to evaluate the impact of movement disorders during sleep on patients with Parkinson's disease.

References