Initial circulatory response to active standing in Parkinson’s disease without typical orthostatic hypotension

Resposta circulatória inicial após ortostatismo ativo na doença de Parkinson sem hipotensão ortostática típica

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ABSTRACT
While the circulatory response to orthostatic stress has been already evaluated in Parkinson's disease patients without typical orthostatic hypotension (PD-TOH), there is an initial response to the upright position which is uniquely associated with active standing (AS). We sought to assess this response and to compare it to that seen in young healthy controls (YHC). Method: In 10 PD-TOH patients (8 males, 60±7 years, Hoehn and Yahr ≤3) the changes in systolic blood pressure (SBP) and heart rate that occur in the first 30 seconds (sec) of standing were examined. Both parameters were non-invasively and continuously monitored using the volume-clamp method by Peñáz and the Physiocal criteria by Wesseling. The choice of sample points was prompted by the results of previous studies. These sample points were compared to those of 10 YHC (8 males, 32±8 years). Results: The main finding of the present investigation was an increased time between the AS onset and SBP overshoot in PD-TOH group (24±4 vs. 19±3 sec; p<0.05). Conclusion: This delay might reflect a prolonged latency in the baroreflex-mediated vascular resistance response, but more studies are needed to confirm this preliminary hypothesis.

Keywords: Parkinson's disease, autonomic nervous system, orthostatic hypotension, hemodynamics, cardiovascular physiological Phenomena.

RESUMO
Apesar da resposta circulatória ao estresse ortostático já foi estudada em pacientes com doença de Parkinson sem hipotensão ortostática típica (PD-TOH), não há uma resposta inicial que é exclusivamente associada com o ortostase ativa (AS). Portanto, buscou-se avaliar esta resposta e compará-la à observada em jovens saudáveis (YHC). M étodo: Em 10 PD-TOH pacientes (8 homens, 60±7 anos, Hoehn e Yahr ≤3) as mudanças na pressão arterial sistólica (PAS) e da frequência cardiaca que ocorrem nos primeiros 30 segundos (seg) de pé foram examinados. Ambos parâmetros foram monitorizados continuamente através do método Peñáz e os critérios de Wesseling. Os pontos de amostragem foram escolhidos com base em estudos anteriores. Estes pontos foram comparados com os de 10 YHC (32±8 anos). Resultados: O principal achado deste estudo foi o aumento do tempo entre o início de AS e rebote sistólica no grupo PD-TOH (24±4 vs 19±3 seg, p<0.05). Conclusão: Este atraso pode refletir uma latência prolongada na resposta da resistência vascular mediado pelo barorreflexo, mas outros estudos são necessários para confirmar esta hipótese preliminar.

Palavras-chave: doença de Parkinson, sistema nervoso autônomo, hipotensão ortostática, hemodinâmica, fenômenos fisiológicos cardiovasculares.

There is an initial circulatory or hemodynamic response (first 30 seconds) to the upright position which is solely associated with active standing (AS)1. This response has been described in both young and elderly healthy subjects1-4. When orthostatic hypotension occurs within 15 seconds (sec) of standing, it is termed initial orthostatic hypotension.
(IOH) and is attributed to a mismatch between cardiac output and peripheral vascular resistance (PVR). The aforesaid mismatch is due to a reduction in PVR and moreover, this decrease is clearly associated with the depth of the blood pressure (BP) drop. Symptoms of cerebral hypoperfusion are likewise present during this drop.

Orthostatic symptoms (OS) occur when cerebral perfusion is sufficiently impaired and, in turn, hypoperfusion develops when cerebrovascular autoregulation (CVA) fails to cope with the BP reduction. This is why, in typical orthostatic hypotension (TOH), these symptoms take place more readily in patients with severe cerebrovascular autoregulatory failure. Nevertheless, autonomic dysfunction in Parkinson’s disease (PD) does not seem to impair the CVA nor its response to orthostatic stress. These CVA differences could account for the reduced reporting of OS in PD.

Autonomic dysfunction in PD is associated with a generalized sympathetic denervation. Recently, it has been reported that PD patients with TOH (PD+TOH) have a lower basal leg vascular resistance, which might suggest that sympathetic denervation is more pronounced in PD+TOH than in PD patients without TOH (PD-TOH). Yet, Oka et al. documented that latent cardiac and vasomotor sympathetic dysfunction occurs in PD-TOH patients.

While the hemodynamic response to orthostatic stress has been already evaluated in PD-TOH, IOH can be documented only by continuous beat-to-beat BP monitoring during AS. Consequently, in this study, we assessed beat-to-beat the initial hemodynamic response to AS in PD-TOH and compared this response to that seen in young healthy controls (YHC).

METHOD

Study population

We enrolled 20 non-institutionalized patients with mild to moderate PD (16 males, mean age 60±7 years, mean duration of disease 3.5±2 years, Hoehn and Yahr stage 2[1-3])10. All of them fulfilled the UK Brain Bank Clinical Criteria for PD. PD+TOH patients were excluded from the present study; the foregoing condition was defined as a fall in BP of at least 20mmHg systolic (SBP) and 10mmHg diastolic (DBP) within 3 minutes (min) in the upright position. BP measurement was performed using standard sphygmomanometry. These patients were all investigated after 12 hours off medication. We also included ten YHC (8 males, mean age 32±8 years) as control group, since the transient hypotension seen during AS does not increase with age. Their health status was determined by a thoughtful medical history and a directed physical examination. This study was approved by the Ethical Committee of the National Institute of Medical Sciences and Nutrition and an informed written consent was obtained from all participants.

Protocol

All evaluations were performed in the morning. Participants in both groups were instructed to avoid alcohol, caffeinated beverages and over-the-counter medications after 22:00 on the night before the evaluation. Finger arterial pressure (FAP) was non-invasively and continuously (beat-to-beat) monitored using the volume-clamp method by Penõáz and the Physiocal (physiological calibration) criteria by Wesseling. SBP, DBP and mean BP (MBP) were then obtained using a slightly modified version of the method already employed by Imholz et al.15 Statistical analysis was performed with STATISTICA for Windows (v.5.1). Prior to group analyses, individual data were tested for normality (Shapiro-Wilk test). Comparisons between groups were performed by either using Student’s t-test or Mann-Whitney U test. A p-value below 0.05 (p<0.05) was considered significant. All results are expressed as mean ±SD or median (interquartile range).
RESULTS

All data were visually inspected prior to analysis. Five patients were excluded either because of tremor-induced artifact (n=2) or asymptomatic TOH (n=3). Besides, the characteristic hemodynamic pattern seen during AS was almost completely lost in five patients, thus hampering attempts to locate the sample points and further comparison between groups. Data from these patients were hence eliminated from study analysis (n=5).

Initial hemodynamic response (Tables 1 and 2). In the ten remaining patients (8 males, mean age 60±7 years), basal SBP and HR were, respectively, 99±8 millimetres of mercury (mmHg) and 64±8 beats per min (bpm). On standing SBP increased suddenly from 107±10 (t0) to 133±18 mmHg (SBPa) and then decreased from that value to 75±20 mmHg (SBPb). An overshoot was evident in seven patients during the recovery phase of SBP (SBPos, 127±18 mmHg). These changes reached its maximum (SBPa) after 3±1 sec (tSBPa) and its minimum (SBPb) after 11±2 sec (tSBPb). For its part, in the seven aforementioned patients, SBPos was reached after 24±4 sec (tSBPos). Thereafter, at the end of the initial hemodynamic response (t30), SBP was 113±23 mmHg (Figure 1).

Standing was accompanied by an absence of a true HRa in all patients and, as a consequence, also by an absence of HRb. The awaited transition from the HR surge to an increscent further rise was likewise not precisely definable. In this manner, HR increased gradually from 66±7 (t0) to 89±8 bpm (HRc) upon standing and subsequently dropped from that value to 70±6 (HRd). These changes reached its maximum (HRc) after 13±4 sec (tHRc) and its minimum (HRd) after 23±5 sec (tHRd). HR was lastly 72±8 bpm at t30 (Figure 1).

Patients were slower to stand up than YHC (11.7 [8.8-14] vs. 4.9 [4.2-5] sec; p<0.001). SBPmax was evident in nine YHC during the recovery phase of SBP (115±10 mmHg). While its value in mmHg was not significantly different between groups, its timing (tSBPos) was delayed in the PD-TOH group (24±4 vs. 19±3 sec; p<0.05). The remaining SBP parameters did not significantly differ from those found in YHC (Figures 1 and 2). HRa (or its corresponding aforedescribed transition) was present in nine YHC (86±5 bpm) at 2±1 sec (tHRa); HRb was afterwards identified in only seven YHC (80±9 bpm) at 4±1 sec (tHRb). HR was smaller in the PD-TOH group than in YHC (89±8 vs. 103±11 bpm; p<0.05), howbeit its timing (tHRb) did not significantly differ between groups. No significant differences were found between groups for the rest of the HR parameters (Figures 1 and 2).

Baroreflex and vasoconstriction (Table 3). BRI and VCI were determined in seven patients and in almost all YHC (n=9). BRI was lower in PD-TOH than in YHC. VCI was also decreased, although not significantly (p=0.09), in the former group (Figure 3). BRS was measured in all the participants and did not significantly differ between groups.

BP abnormalities. Three patients and six YHC met the criteria for asymptomatic IOH. In one patient, SBP recovery phase was noticeably prolonged (28 sec) but nevertheless it does not fulfilled the TOH criteria, since BP after standing for 3 min was not altered. This matter will be addressed in detail in a subsequent communication.

Table 1. Initial HR response to active standing.

<table>
<thead>
<tr>
<th></th>
<th>PD-TOH (n=10)</th>
<th>YHC (n=10)</th>
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</thead>
<tbody>
<tr>
<td>MHHR0 (bpm)</td>
<td>64±8</td>
<td>63±7</td>
</tr>
<tr>
<td>HRt0</td>
<td>64±8</td>
<td>61±5</td>
</tr>
<tr>
<td>HRt5</td>
<td>64±9</td>
<td>63±6</td>
</tr>
<tr>
<td>HRt10</td>
<td>66±7</td>
<td>65±6</td>
</tr>
<tr>
<td>HRt9</td>
<td>86±5</td>
<td>80±9‡</td>
</tr>
<tr>
<td>HRt8</td>
<td>89±8*</td>
<td>103±11</td>
</tr>
<tr>
<td>HRt7</td>
<td>70±6</td>
<td>69±7</td>
</tr>
<tr>
<td>HRt6</td>
<td>72±8</td>
<td>73±9</td>
</tr>
<tr>
<td>tHRa (sec)</td>
<td>2±1‡†</td>
<td></td>
</tr>
<tr>
<td>tHRb</td>
<td>4±1†‡</td>
<td></td>
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<tr>
<td>tHRc</td>
<td>13±4</td>
<td>12±3</td>
</tr>
<tr>
<td>tHRd</td>
<td>23±5</td>
<td>20±2</td>
</tr>
</tbody>
</table>

All results are expressed as mean±standard deviation. PD-TOH: Parkinson’s disease without typical orthostatic hypotension; YHC: young healthy controls; MHHR0: mean heart rate (HR) during a 5-min supine rest; bpm: Beats per minute; HRt0: HR at 10 seconds (sec) before active standing (AS); HRt5: HR at 5 sec before AS; HRt10: HR at the start of AS; HRt9: HR at first peak; HRt8: HR at first valley; HRt7: HR at second peak or maximum; HRt6: HR at second valley; HRt5: HR at 30 sec after AS; tHRa: Time at HRt0; tHRb: Time at HRt5; tHRc: Time at HRt9; tHRd: Time at HRt7; tHRmax: Time at HRt5; tHRmin: Time at HRt9.

*p<0.05 PD-TOH vs. YHC. †n=9, ‡n=7.

Table 2. Initial SBP response to active standing.

<table>
<thead>
<tr>
<th></th>
<th>PD-TOH (n=10)</th>
<th>YHC (n=10)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MSBP0 (mmHg)</td>
<td>99±8</td>
<td>100±9</td>
</tr>
<tr>
<td>SBP-10</td>
<td>108±7</td>
<td>102±9</td>
</tr>
<tr>
<td>SBP-5</td>
<td>107±8</td>
<td>102±9</td>
</tr>
<tr>
<td>SBP+5</td>
<td>107±10</td>
<td>102±7</td>
</tr>
<tr>
<td>SBP+0</td>
<td>133±18</td>
<td>123±17</td>
</tr>
<tr>
<td>SBP+1</td>
<td>75±20</td>
<td>70±16</td>
</tr>
<tr>
<td>SBP+2</td>
<td>127±18*</td>
<td>115±10§</td>
</tr>
<tr>
<td>SBP+3</td>
<td>113±23</td>
<td>108±15</td>
</tr>
<tr>
<td>ΔSBP0–SBP+3 (mmHg)</td>
<td>31±17</td>
<td>31±13</td>
</tr>
<tr>
<td>tSBP0 (sec)</td>
<td>3±1</td>
<td>2±0.8</td>
</tr>
<tr>
<td>tSBP+1</td>
<td>11±2</td>
<td>10±1</td>
</tr>
<tr>
<td>tSBP+3</td>
<td>24±4*</td>
<td>19±3†</td>
</tr>
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All results are expressed as mean±standard deviation. PD-TOH: Parkinson’s disease without typical orthostatic hypotension; YHC: young healthy controls; MSBP0: Mean systolic blood pressure (SBP) during a 5-min supine rest; mmHg: Millimetres of mercury; SBP+1: SBP at 10 seconds (sec) before active standing (AS); SBP+0: SBP at 5 sec before AS; SBP+2: SBP at the start of AS; SBP+3: SBP at first peak; SBP+1: SBP at valley; SBP0–SBP+3: SBP at overshoot; HRt30: SBP at 30 sec after AS; ΔSBP0–SBP+3: Absolute change from SBP0 to SBP+3; tSBP0: Time at SBP0; tSBP+1: Time at SBP+1; tSBP+3: Time at SBP+3.

*p<0.05 PD-TOH vs. YHC. †n=7, ‡n=1.
DISCUSSION

The main finding of the present investigation was an increased time (tSBPos) between the start of AS (t0) and the overshoot (SBPos) in PD-TOH group. When compared between groups, SBP (in mmHg) was not significantly different at tSBPos; notwithstanding it could not be determined in three patients. Peripheral sympathetic vasoconstriction, in part baroreflex-mediated, is responsible for the rebound of BP from SBPb to SBPos following AS1,18. Local reflexes are involved as well in this PVR increment8,19,20. Except for the venoarteriolar response (VAR) in those aged over 75 years, both central and local responses are independent of age in healthy subjects1,19,21. Thus, we suggest that an increased tSBPos may be caused either by an impaired VAR or by an abnormal baroreflex response. Indeed, these mechanisms are not mutually exclusive22.

With respect to the first possibility, Andersen et al. found that, in PD patients without symptoms or signs of autonomic dysfunction, local reflexes did not differ from those in YHC.
in age-matched healthy controls\(^1\). This finding was corroborated by Fusina et al.\(^2\) and more recently by Groothuis et al.\(^3\). Regarding the second possibility, it is noteworthy that baroreflex function assessment has concentrated on baroreceptor-HR reflex, even though PVR control is of greater importance during orthostatic stress. In the same manner, this attention has focused on the sensitivity of the baroreflex response and not in its latency. Gulli et al. reported, in connection with the foregoing, a prolonged delay in the baroreflex-mediated PVR response, after baroreceptor unloading (simulation of BP drop), in patients with poor orthostatic tolerance\(^4\). Additionally, it must be noted that the temporal response to baroreceptor unloading is similar in young and older healthy subjects\(^5\). We therefore deemed the latter possibility as more likely in the present case.

VCI was proposed as an indicator of PVR function\(^6\) and in our study, it was found decreased in PD-TOH compared to YHC (although not significantly). The lack of significant difference may be due to the way in which it is calculated (i.e., from the ratio between MBP recovery and its corresponding time)\(^7\), inasmuch as this calculus takes into account both the magnitude (mmHg) and the duration (sec) of the recovery and, as in the case of SBP, the former did not significantly differ between groups (data not shown). Even PD-TOH patients have baroreflex abnormalities, albeit more subtle than those experienced by PD +TOH patients\(^8\). In our study, baroreceptor-HR reflex function was assessed using two different procedures. While it is true that BRS can be determined more easily than BRI in PD-TOH, its results were inconsistent with those obtained by BRI, since it did not detect blunted baroreceptor-HR reflex function.

Getting out of the bed is often difficult for PD patients\(^9\). This situation was patent when our patients assumed the upright posture from the supine and were comparatively slower than YHC. We also found that the bimodal pattern of the initial HR response to AS was entirely lost in all patients, this absence could be attributed to the aforesaid motor slowness\(^1\). For its part, HR\(_c\) attenuation in PD-TOH group might be explained by the age effect\(^1\).

In conclusion, the present study demonstrates that, in PD-TOH patients, there is a delay between the AS onset and SBP overshoot. This delay possibly reflects a prolonged latency in the baroreflex-mediated PVR response, but more studies are needed to confirm this preliminary hypothesis.

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