Effect of clinical autonomic dysfunction on cognitive functions in Parkinson’s disease

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Received: 27.10.2016; Revised: 16.06.2017; Accepted: 27.07.2017

Abstract
Objective: Parkinson’s disease (PD) is a chronic progressive neurodegenerative disorder characterized by tremor, rigidity, bradykinesia, and postural instability. PD also involves nonmotor manifestations such as autonomic failure, cognitive disorders, and sleep disorders. These clinical characteristics are not identical in severity, frequency, and onset time in all PD patients. We assessed whether there is a negative effect on cognition of clinical autonomic dysfunction in PD patients.

Methods: This prospective study includes 37 PD patients with autonomic failure. From each patient, a questionnaire (SCOPA-AUT) including symptoms associated with clinical autonomic dysfunction such as constipation, urinary incontinence, orthostatic hypotension, and hyperhidrosis was obtained and the patient’s clinical condition was rated on the Hoehn and Yahr (H-Y) scale in the ON-medication state. The patients’ cognitive function was assessed by the Mini-Mental State Examination (MMSE), Blessed score, Frontal Assessment Battery (FAB), and Digid Span Test (DST) (forward, reverse). Cognitive test scores were compared with SCOPA-OUT scores.

Results: Mean age was 66.5±11.2 years. There was no correlation between cognitive test scores and SCOPA-OUT scores (p>0.05). However, H-Y scores were negatively correlated with the DSTf, DSTr, FAB, and MMSE scores (p50.005, r=-0.451; p50.025, r=-0.367; p50.040, r=-0.340; p50.044, r50.333, respectively).

Conclusions: According to our results, clinical autonomic dysfunction did not seem to have an effect on cognition. In addition, severity of cognitive dysfunction showed a strong negative correlation with the stage of disease.

Keywords: Blessed score, Clinical autonomic dysfunction, Cognitive test, Parkinson’s disease, SCOPA-AUT

DOI: 10.5798/dicletip.338976
Parkinson hastalarındaki klinik otonomik disfonksiyonun kognitif fonksiyonlar üzerine etkisi

Özet

Yöntemler: Bu prospektif çalışmaya otonomik yetmezliği olan 37 PH dahil edildi. Her bir hasta konstipasyon, uriner inkontans, ortostatik hipotansiyon ve hiperhidrozis gibi klinik otonomik disfonksiyon ile ilişkili semptomları içeren SCOPA-AUT anketi ve hastaların on durumundaki klinik durumun Hoehn Yahr (H&Y) skalası ile elde edildi. Hastaların kognitif fonksiyonu mini mental durum muayenesi (MMSE), blessed skoru, frontal değerlendirme ölçeği (FAB), ileri (DSTf) ve geri (DSTr) sayı testi ile değerlendirildi. Kognitif test skorları SCOPA-AUT skoru ile karşılaştırıldı.

Bulgular: Yaş ortalaması 66,5±11.2 yıldı. Kognitif test skoru ve SCOPA-AUT skoru arasında istatistiksel bir ilişki yoktu (p>0.05). Fakat H-Y skoru ile DSTf, DSTr, FAB ve MMSE skoru arasında negatif bir korelasyon vardı (p=0.005, r:-0.451; p=0.025, r:-0.367; p=0.044, r:-0.333, sırasıyla).

Sonuçlar: Bizim sonuçlarımızına göre, klini otonomik disfonksiyonun kognisyon üzerine bir etkisi olmadığıdır. İlave olarak kognitif fonksiyonun ciddiyeti hastalığın evresi ile güçlü negatif bir korelasyon gösterdi.

Anahtar kelimeler: Blessed skoru, klinik otonomik disfonksiyon, kognitif test, Parkinson hastalığı, SCOPA-AUT

INTRODUCTION
Parkinson’s disease (PD) is a chronic progressive neurodegenerative disease which includes motor signs and symptoms such as rigidity, bradykinesia, rest tremor and balance disorder1. Parkinson’s disease is a neurodegenerative disease which courses especially with the loss of dopaminergic cells in substantia nigra (SN). A great number of mechanisms have been put forward in its pathogenesis1,2. Non-motor symptoms of this disease are not well defined as motor symptoms. Non-motor symptoms can be seen in the early stages of the disease or years before motor symptoms start1. Non-motor features include neuropsychiatric disorder, sleep disorder, autonomic dysfunction, pain and sensory impairment2. As the disease progresses, the variety of these symptoms increase3. As in motor symptoms, non-motor symptoms significantly affect life quality and physical activity4. While motor symptoms respond well to dopaminergic treatment, non-motor symptoms do not show an obvious improvement. Although there are studies in literature which show a directly proportional association between motor response and PD dementia and autonomic dysfunction and advanced PD, there are also studies which support the opposite5,6. There are limited numbers of studies in literature which show the effect of autonomic deficiency on dementia. To the best of our knowledge, there are no studies directly assessing the association between autonomic dysfunction, cognitive function and dementia clinically. In this sense, we believe that our study will contribute to literature. In our study, we evaluated the effects of autonomic dysfunction clinically found in Parkinson’s disease patients on cognitive functions.

METHODS
The hospital research ethics boards approved this study, and informed written consent was obtained from all participants.
60 Parkinson’s patients were assessed for the study. 23 were excluded and 37 patients were included in the study. The diagnosis of Parkinson’s disease was made by a neurologist who is an expert in the field and the related tests were made by another neurologist who is also experienced in the field.

The study included patients who were registered in ***, Neurology Parkinson’s and Motion Sicknesses Polyclinic with a diagnosis of Parkinson’s disease and who were found to have clinical autonomic dysfunction with the Scales for Outcomes in Parkinson’s disease - Autonomic (SCOPA-AUT) questionnaire. Cognitive tests and SCOPA-AUT questionnaire were administered in the pre-stage (within the six hours following dopaminergic drug intake). Clinical presence and severity of autonomic involvement was determined with SCOPA-AUT questionnaire. Hoehn and Yahr (H&Y) scale was administered to assess the disease severity. The patients who had moderate and severe depression according to Geriatric Depression Scale, those who had visual hallucinations, patients who were taking medication that can affect autonomic function and those who had 4 and 5 Hoehn-Yahr staging were excluded from the study.

Mini Mental State Examination test: 30 points in total (30 is the best), the test consists of items that measure time and space orientation (10 points), memory (3 points for recording and 3 points for remembering, 6 points in total), attention (5 points), language (8 points) and visual-spatial functions (1 point). Blessed test: In order to reach a total of 28 points (28 is the worst), error numbers in each item are multiplied with pre-determined weighted coefficients. Following the time orientation items (year and month), an imaginary name-surname and address of 5 elements are given to be memorized as memory item. After a time orientation (hour of the day) and two attention items (counting down from 20 one by one and counting the months from backwards) are placed within the delay time, the address is scored with free recall.

Frontal Evaluation Test: It is a test which can be administered in a short time and it is used in the assessment of frontal lobe functions. It consists of 6 sub-divisions that give a total possible score between 0 and 18. Sub-divisions are similarities (conceptualization), word fluency (mental flexibility), motor series (programming), contradicting instructions (sensitivity to interference), do-don’t do (inhibitory control) and catch behavior (environmental autonomy). High scores taken from the test mean better performance.

Geriatric Depression Scale: A score of over five can correspond to depression; it should be evaluated with the patient’s clinic.

Number sequence test: It is the most commonly used attention/short-term memory test. It consists of two parts as number range, forward and backward counting range. In both, the subject is read random numbers with breaks of one second in-between, the numbers increase at each try and the subject is asked to repeat with the same order. For both parts, the number of the first series out of the two consequent series the subject fails forms the number range. For normal individuals, the lower limit is generally accepted as 6 forward and 4 backward.

RESULTS

Of the 37 patients included in the study, 18 were male and 19 were female and their averages of age were 66.53±11.2 (46-79). Average age for disease onset was 60.61±6.14 and the average duration of disease was
4.87±1,91 years. Mean MMSE score and Blessed score were 23.6 (15–30) and 15.8 (3-26), respectively. H&Y clinical stage averages of the cases was 1.37±0.49. Mean total SCOPA-AUT score was 12.8 and H&Y stages did not correlate with SCOPA-AUT (p < 0.05) (Table 1). There was no correlation between cognitive test scores (MMSE, FAB, Blessed score, DSTf, DSTr) and SCOPA-OUT scores (p>0.05) (Table 2). However, H-Y scores negatively correlated with the DSTf, DSTr, FAB, and MMSE scores (p=0.005, r:-0.451; p=0.025, r:-0.367; p=0.040, r:-0.340; p=0.044, r=-0.333, respectively) (Table 3). In addition, there was a positive correlation between the Blessed score with age (r = 0.41, p < 0.01), disease duration (r = 0.45, P < 0.05) and H&Y score (r = 0.37, p < 0.05). No significant association was found between the side on which the disease started and SCOPA-AUT and cognitive tests. Autonomic dysfunction may be present from disease onset, whereas the rate of cognitive decline increases with disease progression.

<table>
<thead>
<tr>
<th>Table 1: Clinical characteristics of the patients, mean ± SD</th>
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<tbody>
<tr>
<td>Number of patient (M,F)</td>
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<tr>
<td>Mean Age</td>
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<tr>
<td>Mean disease duration</td>
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<td>Mean disease onset age</td>
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<td>Hoehn-Yahr stage</td>
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<td>SCOPA-AUT test</td>
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<td>MMSE test</td>
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<td>Blessed score</td>
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<td>Digid span forward</td>
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<td>Digid span reverse</td>
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<td>Frontal assesment battery</td>
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M:Male; F:Female; SCOPA-AUT: the Scales for Outcomes in Parkinson’s disease - Autonomic

**DISCUSSION**

Although the mechanisms in the pathogenesis of Parkinson’s disease are not fully understood, it is thought to occur as a result of chain of events that emerge with disorders in the protein processing triggered by genetic factors and environmental factors, oxidative stress, mitochondrial disorder, excitotoxicity, inflammation and changes in the immune system. Initially, genetic factors were not thought to be effective in PD, it was thought to be a sporadic disease. With the findings in studies that in 20% to 25% of sporadic PD cases at least one of the first degree relatives had PD, PD risk is increased in individuals whose first degree relatives had PD, genetic factors in twins were important for early onset PD and the discovery of genes related with familial PD, genetic factors gained importance.

Intracytoplasmic Lewy body accumulation and neuron dysfunction starts from olfactory bulb and dorsal vagal nucleus and then involves the cortex. Clinical findings are known to start after the disappearance of a specific rate of dopaminergic cells which include melanin in SN. Cognitive dysfunction and dementia develop with the accumulation of Lewy bodies in limbic cortex and frontal neocortical regions and the decrease in frontal dopaminergic cortical projection. Executive functions are influenced based on basal ganglia being affected and the decrease in the dopaminergic input to prefrontal cortex. Autonomic dysfunction develops based on the sympathetic ganglion and peripheral autonomous nerve system as well as synuclein pathology in dorsal vagal nucleus, hypothalamus and neurons in the intermediolateral nucleus of the spinal cord. In addition, cardiac sympathetic fiber exposure can cause orthostatic hypotension and postural hypotension while parasympathetic fiber exposure can cause changes in heart rate. One study showed...
degeneration in vagal nerve, spinal cord and celiac ganglion. This explained the clinical variety in motor and non-motor findings\(^5\). Another study showed that non-motor symptoms were seen frequently and especially in the advanced periods of the disease and they had serious effects on the quality of life. The same study reported that 21% of the patients were admitted with non-motor symptoms\(^5\). Non-motor features include neuropsychiatric disorder (depression, apathy/anhedonia, anxiety, frontal executive malfunction, dementia, psychosis), sleep disorder (insomnia, RUDB, daytime sleep), autonomic dysfunction (orthostatic hypotension, constipation, urogenital dysfunction, extreme sweating), pain and sensory impairment\(^2\).

Typical clinical feature of cognitive dysfunction is an executive dysfunction in the foreground (problem solving, attention, planning) and disorders in verbal fluency and visual-spatial skills since early periods. Memory impairment in early periods is not limbic\(^12\).

Different studies have reported the reasons which increase the development of dementia in PD cases as mild cognitive dysfunction, postural instability, visual hallucinations, orthostatic hypotension, REM sleep behavioral disorder, speech and swallowing difficulty, symmetrical parkinsonism findings, gastrointestinal or urological dysfunction, male gender, depression, akinetic-rigid Parkinson and advanced age\(^12,14\).

With neuropsychological tests in PD, cognitive exposure was reported in rates between 20-83\(^%\)\(^15\). In one study, cognitive dysfunction was found in 84% of Parkinson's patients and as a result of a 15 year-long follow up of these patients, 47% were diagnosed with dementia\(^16\). Cognitive deterioration in PD may not always be parallel with motor performance. The tendency for negative association between UPDRS and MMSE scores in one study shows that cognitive dysfunction increases as motor disability increases\(^17\). In our study, a negative association was found between H&Y staging and MMSE while a positive association was found between Blessed test.

| Table 2: Correlation between cognitive test scores and SCOPA-AUT score |
|-----------------|-----------------|-----------------|-----------------|-----------------|
| SCOPA-AUT       | MMSE | Blessed score | FAB  | DSTf | DSTr | p value |
| 23.6            | 15.8 | 10.8           | 5.1  | 2.0  | p>0.05 |

| Table 3: Correlation between cognitive test scores and Heahn-Yahr stage |
|-----------------|-----------------|-----------------|-----------------|
| Heahn-Yahr      | MMSE | Blessed score | FAB  | DSTf | DSTr |
| 23.6            | 15.8 | 10.8           | 5.1  | 2.0  |
| p value         | p=0.044 | p < 0.01         | p=0.040 | p=0.005 | p=0.025 |
| r value         | r: -0.333     | r = 0.41         | r: -0.340        | r: -0.451        | r: -0.367 |

Various studies have reported autonomous symptoms to occur at every stage of Parkinson's disease, in different severity and between 70-93\(^%\)\(^18\). Clinical presence and severity of autonomous involvement in PD is assessed effectively and correctly by using SCOPA-AUT questionnaire\(^8\). SCOPA-AUT study showed association between the degree of motor exposure in PD and autonomous symptoms. Our study and the study of Oh et al. 
did not find any association between SCOPA-AUT score and H&Y staging\textsuperscript{19}. In their study, Magerkurth et al. reported that one or more types of autonomous dysfunction could be seen and autonomous dysfunction was not associated with the duration and severity of the disease\textsuperscript{20}. Orthostatic hypotension is one of the most common symptoms of PD and it is seen with a rate of between 20-50\% and it is more common in the advanced stages of the disease. In another study, no association was found between the disease severity and orthostatic hypotension\textsuperscript{21}. While there are studies which have not found an association between autonomic dysfunction and cognitive exposure\textsuperscript{19,22}, there are also studies which show an association between the two\textsuperscript{23}. Prospective studies are needed to show the association between autonomic dysfunction and cognitive exposure.

There are limited numbers of studies about the effects of autonomous dysfunction on cognitive exposure. Unlike other studies, we assessed the clinical autonomous dysfunction in PD cases with detailed cognitive tests (Mini Mental State Examination test, Blessed scoring, Frontal Evaluation Test and number sequence test). In parallel with some studies in literature, no association was found between cognitive tests and SCOPA-AUT scores. However, while there was a negative association between cognitive test scores and H&Y staging, there was a positive association between Blessed score and H&Y staging. Autonomic dysfunction, which can easily be determined clinically with SCOPA-AUT test, was found to occur in almost every stage of our cases. There are studies which show parallelism or non-parallelism between SCOPA-AUT test and H&Y staging. Our study did not find a parallelism between the two.

When the results of studies were assessed together, it can be concluded that autonomic dysfunction can be seen in every stage of PD and it can cause cognitive exposure in some cases.

Since our study used a test which was proved to be efficient and reliable in finding out autonomous dysfunction, orthostatic hypotension and postural hypotension were not checked.

In addition to the cognitive tests we conducted, we believe that a study which assesses cognition with watch drawing and Montreal Cognitive Assessment: MoCA test, a study which assesses orthostatic hypotension and postural hypotension with holter through 24 hours of record in addition to SCOPA-AUT test, a study with many cases and in which cases are classified according to the drugs used and the duration of drug use will make valuable and important contributions to literature.

**Declaration of Conflicting Interests:** The authors declare that they have no conflict of interest.

**Financial Disclosure:** No financial support was received.

**REFERENCES**


