DEPRESSION IN PARKINSON’S DISEASE

Study of 60 cases

Roberto César Pereira do Prado¹, Egberto Reis Barbosa²

ABSTRACT - Depression is very frequent in Parkinson’s disease (PD) and largely unrecognized by neurologists, emphasizing the need of an approach to psychiatric symptoms by non psychiatrists in order to ensure an early diagnosis of depression in PD; clinical characteristics and the prevalence rate of depression in PD were evaluated and the relationship of depression in PD with other variables were determined. Sixty PD subjects, who fulfilled the clinical criteria for primary PD, 56.6% males, age range from 44 to 85 years old, in different stages of the disease were investigated. All subjects were submitted to the UPDRS-III, V and VI, Clinical Interview Schedule and the Hamilton depression scale. A significant correlation was found between depression and UPDRS-III, V and VI, anxiety and irritability. The frequency of depression in PD in this study was nearly 40% possessing specific features. Structured interviews and evaluation scales are essential for an accurate diagnosis and proper treatment of depression in PD.

KEY WORDS: depression, Parkinson’s disease, neuropsychiatric disorders, diagnosis, comorbidity.

Depressão na doença de Parkinson: estudo de 60 casos

RESUMO - A depressão é manifestação freqüente na evolução da doença de Parkinson (DP), gerando a necessidade de nova abordagem neuropsiquiátrica por parte dos médicos não psiquiatras, visando o reconhecimento precoce do quadro depressivo na DP. Foram estudadas as características clínicas e frequência da depressão na DP e correlacionadas com outras variáveis. Sesenta pacientes que preenchiam os critérios clínicos atuais para DP, sendo 56.6% do sexo masculino (44 a 85 anos), em diferentes estágios clínicos da doença, foram submetidos às escalas de avaliação para DP (UPDRS-III, V e VI), para transtornos neuropsiquiátricos - Entrevista Clínica Estruturada e Escala de Hamilton. Houve associação estatisticamente significante entre depressão e UPDRS-III, V e VI, ansiedade e irritabilidade. A frequência de depressão, situou-se em torno de 40% apresentando características próprias. Entrevistas estruturadas e escalas de avaliação são essenciais para o diagnóstico preciso e tratamento adequado do fenômeno depressivo na DP.

PALAVRAS-CHAVE: depressão, doença de Parkinson, manifestações neuropsicossomáticas, diagnóstico, comorbidade.

The nonmotor symptoms (e.g., dementia, autonomic dysfunction, and mood disturbances) are rather frequent in Parkinson’s disease (PD), being as common as the motor symptoms that are highly characteristic of this disease¹². These manifestations can be the most disturbing and disabling aspects of the disease. Depressive symptoms are particular a significant cause of functional impairment for these patients. Depressive symptoms have also been described in other movement disorders such as Huntington disease, basal ganglia idiopathic calcification, Wilson’s disease, vascular parkinsonism, progressive supranuclear palsy, neuroanionicytosis, dystonia, spino-cerebellar degeneration¹. The assessment of comorbid psychiatric symptoms in PD subjects should take into account if symptoms are 1) anti-parkinson medication-induced; 2) secondary to PD disease phenomena, like depression or anxiety secondary to motor fluctuations; 3) combination of disease factors and medication side-effects. The presence of psychiatric symptoms will demand different actions as anti-parkinson medication reduction or addition of a psychoactive drugs. Among several psychiatric symptoms, depression stands out because of its unique features and high prevalence rate in PD.

Depression is by far the most common psychiatric finding in PD. Prevalence rates span from 4 to

¹Departamento de Medicina, Universidade Federal de Sergipe, Aracaju SE, Brasil: Professor Adjunto; ²Departamento de Neurologia, Universidade de São Paulo, São Paulo, SP, Brasil: Livre Docente

Received 1 November 2004, received in final form 6 May 2005. Accepted 10 June 2005.

Dr. Roberto César P. do Prado - Rua Celso Oliva 321/301 - 49020-090 Aracaju SE - Brasil. E-mail: rcprado@infonet.com.br
70% depending on the methodology employed. Data from studies employing the Beck Depression Inventory (BDI) or the Hamilton Rating Depression Scale (HRDS) showed a prevalence rate of 40%, an incidence rate of 1.86% a year and a cumulative risk of 8.60%4. Depressed PD patients typically present lack of motivation, reduced self-esteem, high level of anxiety, pessimism about the future, irritability, suicide ideation and rarely suicide attempt and hallucinations and delusions5. The diagnosis of depression in PD as well as in other hypokinetic movement disorders is particularly difficult because, perhaps more than any other medical disorder, the symptoms of PD overlap with the symptoms of depression1,6,7. Most patients with PD appear depressed to the passive observer. The “masked facies”, psychomotor slowing and fatigue of PD give the appearance of depression in euthymic patients5. On the other hand, depression in PD is often not recognized by neurologists.

The objective of this study was to assess neuropsychiatric symptoms, specially depression, in a selected group of PD patients.

**METHOD**

Sixty consecutive PD subjects were included in this study. There were 34 males (56.6%) and 26 females (43.3%) and average age was 64.83 ± 11.64 years (age range from 44 to 85 years old). The London Brainbank diagnosis criteria was employed for the clinical diagnosis of PD in all patients8. All patients were examined at the University Hospital - Federal University of Sergipe - Unity of Movement Disorders and were seen from November 1997 through May 2001.

Unified Parkinson’s Disease Rating Scale (UPDRS) part III (motor signs), part V (Hoehn and Yahr PD staging scale), part VI (Schwab and England daily activities scale) and the Mini-Mental State test - MMS - were applied to all patients to assess respectively, the degree of clinical parkinsonian symptoms and cognitive status of the studied group9.

A three-level stratification was necessary for the MMS scores because the studied population included several patients with minimal educational background or illiterate11,12. Three patient groups were created according to educational level: illiterate group, minimal educational level (< 8 years of school education) and elevated educational level (> 8 years of school education). Three distinct MMS scores corresponding to the patient grouping described above were fixed: 18 points for the illiterate group, 21 points minimal educational level and 24 points for elevated educational level as suggested by some authors12.

The Portuguese version of the Clinical Interview Schedule (CIS-R) was applied to each individual patient by the first author to determine the presence of psychiatric symptoms13. Specific CIS-R questions concerning symptoms primarily related to the Parkinson’s disease resembling somatic symptoms, body functions, tiredness were excluded as recommended by Botega et al.13. Clinical suspicion for depression was achieved in all subjects who scored two or more points in the depression or depressive thoughts.

The Portuguese version of the HAM-D was subsequently applied to all patients bearing a clinical suspicion of depression14. The authors adopted the 17-item score range 0 to 50 points HAM-D scale instead of the full 21-item scale. The remaining 4-item list was not included because it assesses findings as diurnal mood variation, delusions, paranoid and obsessive symptoms which not only are uncommon in PD but it does not significantly contribute to the overall estimative of the affective disorder. HAM-D scores up to seven points were considered normal, a score between 7 through 17 points was indicative of mild depression, 18 up to 24 points was considered moderate and above 25 points was considered depression of a severe degree14.

A brain CT scan with contrast was obtained for all patients. Those presenting CT scan abnormalities other than volumetric reduction of the brain were excluded from the study.

**Statistical analysis** – The clinical characteristics and the prevalence rate of depression as well as age, gender, age at disease onset, duration of motor dysfunction, motor dysfunction laterality, PD stage, functional disability, cognitive status were analysed. The Student’s t test was employed to compare the average of the quantitative variables between the depressed and non-depressed subjects. The chi-square or Fischer-s tests were applied to the quantitative variables and the odds-ratio between the depressed and non-depressed patients to evaluate more important features. Person’s and Spearman’s correlation measures were employed to assess the continuous variables. An alpha of 5% was considered statistically significant for all tests. Descriptive indexes lower than p=.05 were statistically significant. Stata 7.0 version software was employed for the analysis.

**RESULTS**

Thirty-four males (56.6%) and 26 females (43.3%) were included. Age range was 64.83 (44 to 85 years), disease onset age was 61.31 (41 to 82) and disease average duration was 4.42 anos (0.16 to 13). The relative frequency of neuropsychiatric manifestations is depicted in Figure 1.

Mild depression was recorded in 82.60% whereas moderate and severe depression rates were 8.69% as shown in Figure 2.

There was a statistically significant correlation between depression and motor symptoms evaluated by UPDRS part III (chi-square = 6.7685, p = 0.009
- Table 1); disease staging scored by UPDRS part V (chi-square = 6.3316, p = 0.012 - Table 2) and for performance at daily activities, assessed by UPDRS part VI (chi-square = 16.4076, p = < 0.001 - Table 3).

A statistically significant association was found for depression and irritability (chi-square = 8.8187, p = 0.032 - Table 4), worrying thoughts (chi-square = 19.1640, p = 0.001 - Table 5) and anxiety (chi-square = 19.6005, p = 0.001).

**DISCUSSION**

The frequency of depression (38.33%) recorded in the present investigation is in keeping with the current literature data. Specific clinical features of the depression in the present investigation were sleep complaints, anxiety, irritability and lack of concentration and interest. However, symptoms like guilty feelings, self-destruction, suicidal thoughts were not recorded in this sample as they usually are in a primary depression population.

An important point in an investigation pertaining to PD and depression concerns the most adequate instruments to assess the affective symptoms. Shulman et al. from the University of Miami have
assessed the local staff neurologists accuracy rate of the diagnosis of depression and other psychiatric symptoms in PD patients. Clinical impression based on a routine consultation was compared with specific data obtained from specific clinical scales for depression, and other psychiatric symptoms. Clinical assessment accuracy was 35% for depression.

The overlap of motor symptoms as bradykinesia and psychomotor retardation poses as a real challenge for the diagnosis of affective disorders in PD. For this reason, the authors elected the CIS-R structured interview, an easily applied instrument, useful to assess the neuropsychiatric symptoms. Additionally, the CIS-R yields a high detection rate of psychiatric symptoms whereas the HAM-D was meant to measure the severity degree of depressive symptoms in this PD population. The combined use of both research instruments minimized the rate of misdiagnosis of psychiatric symptoms by non-psychiatrists. The newly edit version of the CIS-R can be easily applied by other health-related professionals removing thus the time and economic...
The present study revealed no statistically significant correlation between depression and gender. This was also reported by Ehmann et al.\(^9\). However, Warburton\(^24\) found a female predominance for depressive symptoms in a group of 140 PD patients selected for thalamotomy and 140 controls.

There was no statistically significant correlation between the cognitive changes assessed by the MMS and the depression ratings. One should bear in mind that the MMSE is just a screening test and that mild depression rates predominated in the present series of patients, whereas other investigators as Starkstein et al.\(^25\) showed a remarkable intellectual deterioration in the severely depressed patients. These authors even proposed two distinct forms of PD, one with rapidly progressive cognitive changes and severe depression and the second type with a slowly cognitive decline without depression. The relation between depression and PD, yet controversial, presents two clinically relevant points according to the authors\(^25\). Firstly, the treatment for depression bears the potential to arrest the cognitive decline and secondly, the life expectancy of the PD patients with cognitive decline becomes significantly shorter.

Concerning the correlation between motor dysfunction laterality and depression no statistically significant results were found in this series, as determined likewise by Aarsland et al.\(^15\). Nonetheless, Direnfeld et al.\(^26\) data showed that patients with predominance of motor symptoms on the right (corresponding to left side of the brain dysfunction) presented higher Beck depression inventory scores as opposed to individuals with bilateral motor dysfunction or individuals with left-side predominant motor symptoms. Starkstein et al.\(^27\) findings also indicate a relationship between elevated depression scores and right side motor dysfunction.

The severity degree of the motor dysfunction assessed by the UPDRS part III and V and depression was statistically significant in the present series. In other words, the more severe the motor impairment was, the more elevated the scores for depressive symptoms. However, this finding does not achieve support in investigations published by other authors\(^9,19,22,27\) in which this relation was not observed. Likewise, the correlation between depression and daily activities scale measured with UPDRS - VI was statistically significant. Kostic et al.\(^28\) results are in keeping with the findings mentioned above despite these authors employed a different scale to evalu-
ate daily activities. The authors showed an association between depression and anxiety, as 12 patients from the fifteen that presented a CIS-R score for anxiety ($\geq 2$), also presented significant scores for depression and/or depressive ideas. Menza et al.\textsuperscript{29} reported that 12 from 42 patients of a PD population sample fulfilled the criteria for anxiety and in this subgroup 11 patients were also depressed.

The association between depression and irritability, depression and worrying thoughts was a significant one in the present series. Out of 18 subjects who scored $\geq 2$ in the CIS-R for worrying thoughts, 13 also presented a significant score $\geq 2$ for depression.

This study points at some directions in terms of an ethiopathogenesis for the listed psychiatric changes. The notion of a reactive depression is supported by the correlation between motor changes documented by the UPDRS (parts III, V, VI) and depressive scores. However, the lack of a statistically significant correlation between the subject's age, disease duration and previous history of depression all point in favor towards the hypothesis that the depression in PD is related to the CNS chemical imbalance. We would like to stress that the ethiopathogenesis of psychiatric symptoms in PD is a very complex issue and new studies should be carried out for a better understanding of this topic.

We concluded that: a) the CIS-R (Clinical Interview Schedule - CIS-R) is an useful clinical instrument that can be applied by non-psychiatrists in the determination of neuro-psychiatric changes; b) the depression rate found by the authors equivalent to 38.33% is in keeping with others literature data; c) the depressive syndrome found in this group of PD patients showed somewhat specific characteristics as: sleep complaints, anxiety, lack of concentration, worrying thoughts, irritability, panic symptoms; no guilty feelings, self-punishment, delusions or suicide were observed in the patients studied; d) statistically significant correlations between depression and the blocks III (motor examination), V (Hohne e Yahr modified scale) and VI (Schwab e England scale) of the UPDRS were observed; e) statistically significant correlation between depression and irritability was found\textsuperscript{40}.

**REFERENCES**