MOVEMENT DISORDERS SECONDARY TO LONG-TERM TREATMENT WITH CYCLOSPORINE A

Renato P. Munhoz¹, Helio A.G. Teive¹, Francisco M.B. Germiniani¹, Júlio C. Gerytch Jr¹, Daniel S. Sá¹, Marco A. Bittencourt², Ricardo Pasquini², Carlos H.F. Camargo¹, Lineu César Werneck¹

ABSTRACT - Objective: To analyze the prevalence, severity and functional interference of movement disorders (MD) secondary to chronic use of cyclosporine A (CsA). Method: We conducted a cross-sectional study of 60 patients (58.3% male) with mean age 23.1 (3-75) years, followed at the Bone Marrow Transplantation Service of the Hospital de Clínicas of the Federal University of Paraná, Brazil, taking CsA for at least six months. Our protocol included clinical data, assessment of functional interference of symptoms and neurological examination including observation and grading of MD. Results: Eight (13.3%) subjects reported the presence of tremor at the moment of interview and 29 (48.3%) recalled this symptom at some point during treatment. Neurological examination identified 14 (23.3%) subjects with MD: upper limb symmetric action tremor in 13 (21.6%) and parkinsonism (rigidity and bradykinesia) in 1 (1.7%). No other MD was detected. The mean scores indicated mild clinical signs in all cases. Symptoms were considered subjectively mild with no functional interference. Conclusion: Almost one quarter of patients using CsA chronically presented MD, almost always mild and transitory action tremor, with minimal interference on daily living activities, not requiring any form of intervention in the majority of cases.

KEY WORDS: cyclosporine A, tremor, movement disorders, bone marrow transplant.

Cyclosporine A (CsA) is an immunosuppressive agent used broadly in the prevention of rejection of solid organ transplantation due to its effect as a reversible inhibitor of T4 lymphocytes activity and the subsequent release of lymphokines. Although not a myelosuppressant, its use is limited by several side effects including nephrotoxicity and hepatotoxicity, hypertension, gingival hyperplasia, hypertrichosis and opportunistic infections¹. With the increasing experience with the use of CsA in several centers, neurological complications, once rarely described, are now reported in up to 40% of patients, requiring dosage adjustment in one forth of all cases².

The majority of these side effects are related to the central nervous system, but cases of peripheral neuropathy were also occasionally described. The most commonly described complications inclu-
de confusion, disorientation, a wide variety of changes in behavioral and in the level of conscience, seizures, cortical blindness, aphasia, pyramidal signs and movement disorders (MD)\textsuperscript{3}. Among this particular group, tremor is the best recognized\textsuperscript{4}, but case reports of other forms of MD exist in the literature including ataxia\textsuperscript{5}, parkinsonism\textsuperscript{6,7}, myoclonus\textsuperscript{8} and chorea\textsuperscript{9}.

The objectives of our study were to analyze the prevalence of MD among individuals using CsA chronically after bone marrow transplantation (BMT) and to assess the severity and functional interference of the possible findings.

**METHOD**

We performed an observational study (cross-sectional) assessing 60 patients [35 (58.3%) male and 25 (41.7%) female] with mean age 23.1 (3-75) years, followed by the Service of BMT of the Hospital de Clínicas of the Federal University of Paraná (HC-UFPR). Time for diagnosis of the underlying disorder varied from 0.1 to 6 years (mean 1.6). Previously to BMT, all were submitted to other forms of treatment including chemotherapy, transfusions or both.

Indications for BMT were severe aplastic anemia in 41 (68.3%) cases, chronic myeloid leukemia in 11 (18.3%) cases, acute myeloid leukemia in 4 (6.7%), myelodysplasia in 3 (5%) and acute lymphocytic leukemia in 1 (1.7%). All had documented history of CsA use continuously for at least of six months, preceding the evaluation and following BMT in a target maintenance dosage of 5 mg/kg/day (mean 4.35 mg/kg/day) as part of a protocol established by the service of BMT [associated with methotrexate in 50 (83.3%) and/or corticosteroids in 25 (41.6%)]. According to the same protocol, all received prophylactic sulfamethoxazole, trimetoprine, acyclovir and norfloxacin. Those who were clinically decompensated, had clinical or paraclinical signs of current infection or with a history of drugs that can induce MD such as neuroleptics, tricyclic antidepressants and serotonin reuptake inhibitors, calcium channel blockers, aminophyllin and antiepileptics were excluded.

The assessments were performed after a mean 223.1 (192-367) days post-BMT and included demographic and treatment data, and a directed questionnaire inquiring specifically on the subjective presence of MD described in lay terms. If the answer to this questionnaire was positive, its interference on activities of daily living (ADLs) was assessed using practical examples. One of the authors with experience in MD (DSS) completed a standardized protocol that included body weight measurement, application of the Bain et al.\textsuperscript{10} scale for tremors and the Unified Parkinson’s Disease Rating Scale\textsuperscript{11} part III (UPDRS-III), as well as the observation and rating of other possible MD (ataxia, dystonias, chorea and myoclonus).

The HC-UFPR ethics committee approved the study and all included subjects gave their informed consent before the beginning of the evaluation process.

**RESULTS**

Eight (13.3%) out of 60 interviewed subjects reported the subjective finding of MD at the time of assessment, in all cases tremor. This number increased to 29 (48.3%) when the question embraced the retrospective finding of these symptoms at any point of CsA treatment, 21 (72.4%) of these were observed during the first three months of treatment, not requiring any form of intervention. The objective assessment performed by the examiner found 14 (23.3%) patients with MD: 13 (21.6%)
with action tremor (postural or kinetic) and one (1.7%) with parkinsonism (Figure). No other form of abnormal movement was reported by the subjects identified by the examiner.

All the eight cases of tremor reported subjectively were confirmed objectively by physical assessment. Five additional subjects were identified with this signal did not mentioned it during interview; the same observation was valid for the one individual with parkinsonism.

In all 13 cases of action tremor, the phenomenological description was that of a symmetric upper extremities movement. One (1.67%) of these described as purely postural. In the remaining 10 (16.6%) cases, tremor had a concomitance of a kinetic component of the same or lower intensity. Two (3.3%) presented tremor that was more noticeable during movement with no evident dysmetria or intentional tremor (Figure). One individual presented concomitance of symmetric tremor in the lower limbs.

Mean upper extremity Bain et al.10 tremor scale score was 3.8 (2-7) of possible 20 points, indicative of mild tremor. The subject with parkinsonism had a total UPDRS-III score of 6 out of possible 56 points, with mild scores on items related to rigidity and bradykinesia, with no resting tremor or postural instability. In accordance with these scores, symptoms were considered subjectively mild with no ADL functional interference, except for one case. Duration of CsA treatment in the individuals with tremor did not differ significantly in comparison with those who presented no signs or symptoms.

**DISCUSSION**

The results of our study showed that almost one forth of all subjects receiving CsA chronically presented with MD, almost exclusively action tremor. This finding, though, is mild as demonstrated by the observation the examiner and the patient, been frequently transitory and not requiring any form of intervention in the majority of cases. Our study differed from others that evaluated adverse effects secondary to CsA treatment because it included exclusively patients taking this drug for at least 6 months, excluding symptoms reactive to introduction and dosage adjustment. In accordance with this observation, almost half of the interviewed subjects retrospectively reported the presence of transient tremors that did not required any form of change in therapeutic regimen as well as any other form of intervention. Previous studies assessing the presence of neurological complications emphasized part of our findings: Pirsch et al.3, for example, found postural tremor prospectively in 33.8% of subjects taking CsA for one year, the majority during the first 3 months of treatment. Accordingly, two studies evaluating side effects in patients taking CsA found an incidence of tremor varying from 12 to 21%.8,12 concluding that this symptom is generally mild and does not require dosage adjustment or any form of intervention. These studies demonstrated that tremor usually occurs during initial attempts to increase daily dosage, but is also described in individuals receiving stable maintenance dosages within therapeutic blood levels, as in the cases presented here. The only study that analyzed late neurological complications of CsA was the one published by Trocha et al.13 including patients taking CsA for at least 5 months, with findings very similar to ours: 28% presented tremor with minimal interference on ADL. On the other hand, one form complex neurotoxic syndrome that invariably includes postural tremor and confusion was described in the series published by Wijdicks et al.8 and Menegaux et al.14 almost always related to elevated serum levels and/or intravenous CsA infusion.

Kinetic tremor, isolated or as a part of a cerebellar syndrome, seems to be less frequent than postural tremor15. Other MD are rare and described only in case reports. The series of Wijdicks et al.8, for instance, describes 3 cases of myoclonus with speech disturbances responsive to reduction of CsA dosage. Two cases of chorea (one of these possibly pseudoathetosis secondary to peripheral neuropathy) related to CsA have been published in the medical literature, the first described in a series of 46 patients post-BMT16 and the second described after liver transplantation (LT) in Wilson’s disease9. Cerebellar ataxia was described in up to 5% of patients starting treatment, specially following renal transplantation (RT) in children17,18. Belli et al.5 described one case of cerebellar syndrome detected 6 months after LT, symptoms improved after CsA withdrawal. Although used for prevention of rejection of neural xeno and allogenic transplantation in Parkinson’s disease (PD) and possibly providing improvement in motor aspects of animal models of this disorder19-21, CsA has been considered the cause of parkinsonism in a few case reports. Wasserman and Honig7 reported two patients with resting tremor and bradykinesia taking CsA post-BMT, one responded to levodopa while the one
that did not tolerate levodopa improved after immunosuppressant dosage was reduced. Another case report describes a patient with symmetric resting tremor, rigidity and bradykinesia non-responsive to levodopa and trihexyphenidyl but improved when CsA was withdrawn after RT. The case of parkinsonism observed in our study did not require specific treatment because symptoms did not interfere with the ADL. It was not possible to establish a cause and effect correlation among CsA and parkinsonian signs since the individual was in the age range for idiopathic PD.

Possible risk factors for neurological complications of CsA include intensive pre-transplantation chemotherapy, radiation, drug interactions, metabolic disturbances, tissue rejection, fluid retention and hypertension related to corticosteroids, hypercholesterolemia, hypomagnesemia and aluminum overload in RT. Other drugs that interfere with CsA metabolism may increase blood levels to potentially toxic levels. Although anatomicopathological studies did not find a histological substrate for tremor and other MD in patients taking CsA, the mechanism for neurotoxicity seems to be related primarily to the fact of CsA being highly lipophilic, crossing easily the blood brain barrier. From this point on potentially reversible interactions may happen interfering with central neuronal circuits including direct changes in neurotransmission and receptor function, phosphorylation pathways and regulation of transcriptional factors. Feinstein et al. demonstrated that CsA modulates dopaminergic transmission through its stimulating effect on phosphorylation of dopamine and adenosine dependent phosphoprotein 3′5′ monophosphate (DARPP-32) on medium spined striatal neurons. DARPP-32 has a fundamental effect on dopaminergic neurotransmission and CsA increases its levels up to 17-fold. Accordingly, this CsA dopaminergic modulation seems to have an inducer effect of rodent hyperactivity. On the other hand, CsA potentiates domperidone-induced antidopaminergic effect. Amongst such evidences, the reasons why certain patients manifest hyper or hypokinetic syndromes, as well as the fact that such toxicity being manifested most frequently as postural tremor remains obscure.

Our study has important limitations: although been indicated in several situations, CsA evaluated here exclusively post-BMT, hence our findings should not be extensive universally even if other studies in different population have provided similar results. It would be important to correlate symptoms not only with the dosage taken but also with serum levels. This correlation was not performed initially because it would only be valid if the exact moment when the MD would be detected clinically could be determined for each subject and, at that moment, serum levels measured. This can only be performed prospectively, using a methodological process that escapes from the one proposed for our study. We could not speculate any hypothesis linking the presence of the MD with treatment timing or its improvement with CsA dosage reduction since these side effects are usually mild, transient and also because in the vast majority of cases the drug was been used on its lowest effective daily dosage. Another limitation was the lack of a control group. This problem was assumed considering that a healthy control group would bring significant biases for being an entirely distinct population in respect to potential neuro-pathological substrates (underlying condition, pre-BMT treatment, transplantation, etc) that may predispose our patients to MD. On the other hand, once all patients received the same drug regimen, we cannot count on the possibility of a comparison with a group that is not receiving this form of treatment. The only form of comparison that would avoid such bias would be between groups receiving different forms immunosuppressive regimens, such as tacrolimus versus CsA, OKT3 versus CsA. Once again this possibility was not feasible by the fact that all studied subjects were following the same protocol established by the Service of BMT of the HC-UFPR.

We conclude that action tremor is the most common MD in patients treated with CsA, it is relatively more frequent in the beginning of treatment, being reported retrospectively in almost half of the individuals on our study. The assessment after at least 6 months demonstrated the transient character of this complication that has minimal if any impact on ADL and almost never requires specific treatment. Therefore, the initial approach to the patient taking CsA who presents this form of side effect should be reassuring about its benign and transient character. In the rare cases when there is significant functional interference, the interventions described in the literature can be resumed to CsA dosage adjustment or symptomatic treatment directed to the specific MD.
REFERENCES