Parkinsonian Syndrome as a Neurological Manifestation of Behcet’s Disease

D. Bogdanova, I. Milanov and D. Georgiev

ABSTRACT: Background: The central nervous system is often involved in Behcet’s disease. Most common are meningoencephalitic and brain stem syndromes. Although basal ganglia involvement is not an uncommon finding on necropsy, there are only single reports on extrapyramidal syndromes-dyskinesia, chorea and Parkinsonism in patients with Behcet’s disease. Case study: We report a patient fulfilling the criteria of the International Study Group for Behcet’s disease. He had recurrent oral ulcerations, bilateral posterior uveitis and retinal vasculitis, skin papules and pustules, and recurrent monoarthritis. Neurologic examination revealed pseudobulbar palsy, slight and asymmetric bilateral pyramidal syndrome, muscle rigidity involving the four limbs, bradykinesia, masked face, and impaired postural reflexes. There was postural tremor in the extremities and myoclonic jerks involving the tongue and face muscles. Magnetic resonance imaging demonstrated small bilateral multifocal hyperintense lesions, with right predilection, involving the periventricular white matter, brain stem and basal ganglia. Conclusions: The Parkinsonian syndrome found in our patient might be due to involvement of both substantia nigra and basal ganglia. This case further emphasizes the wide spectrum of the neurological manifestations of Behcet’s disease.

RÉSUMÉ: Syndrome parkinsonien comme manifestation neurologique de la maladie de Behcet – À propos d’un cas. Introduction: Le système nerveux central est souvent atteint dans la maladie de Behcet les manifestations les plus fréquentes étaing un syndrome méningoencéphalitique et une atteinte du tronc cérébral. Bien qu’il ne soit pas exceptionnel de trouver une atteinte du noyau lenticulaire, du noyau caudé, de l’anté-mur et du noyau amygdalien à l’autopsie, il y a peu de cas rapportés de syndromes extrapyramidaux-dyskinésies, chorée et parkinsonisme chez les patients atteints de la maladie de Behcet. Étude de cas: Nous rapportons le cas d’un patient qui remplissait les critères du Groupe international d’étude sur la maladie de Behcet. Il avait des ulcères buccaux récurrents, une uvéite postérieure bilatérale et une vasculite rétinienne, des papules et des pustules cutanées et une monoarthrite récurrente. L’examen neurologique a montré une paralysie pseudobulbaire, un léger syndrome pyramidal asymétrique bilatéral, une rigidité musculaire impliquant les quatre membres, une bradykinésie, un faciès parkinsonien et des réflexes posturaux perturbés. Il avait en outre un tremblement postural des extrémités et des secousses myocloniques de la langue et des muscles du visage. La résonance magnétique a montré de petites lésions hyperintenses multifocales et bilatérales, surtout à droite, impliquant la substance blanche pérventriculaire, le tronc cérébral et les noyaux lenticulaire, caudé et amygdalien ainsi que l’avant-mur. Conclusions: Le syndrome parkinsonien observé chez notre patient pourrait être dû à l’atteinte de la substance noire et des noyaux lenticulaire, caudé et amygdalien ainsi que de l’avant-mur. Ce cas démontre le spectre étendu des manifestations neurologiques de la maladie de Behcet.

Behcet’s disease is a world-wide disease, more common in the Middle-East and Eastern Mediterranean.1 It was first described by Behcet in 1937, as a triad of aphthous stomatitis, genital ulceration and uveitis.1 It is a chronic recurrent multisystem disease, now considered as a systemic vasculitis of unknown etiology.23 It affects the skin, joints, eyes, gastrointestinal tract, blood vessels and nervous system.4

Neurological involvement is found in 5.3% to 28% of the patients.56 Male prevalence is predominant in the more severe neuro-Behcet.67 It may occur mainly in two ways: primary (parenchymal) - neuro-Behcet, or secondary to major vascular events-vasculo-Behcet.5,8,9 The neurologic symptoms usually have an abrupt onset, tendency to remit and relapse and a poor prognosis.7

The neurological manifestations are variable, including septic meningitis, brainstem and diencephalon syndromes, pyramidal quadriparesis with pseudobulbar palsy, intracranial hypertension and medullary syndrome.210 Although basal ganglia involvement is not an uncommon finding on necropsy,11 there are only single reports on extrapyramidal syndromes such as dyskinesia,12 chorea13 and undefined “extrapyramidal lesion”14 in patients with Behcet’s disease. The existence of Parkinsonian-like syndromes has been only mentioned.11,14

CASE REPORT:

A 20-year-old man was admitted to the clinic in March 1996 due to acute attack of dysarthria, dysphagia, hands, legs and tongue tremor,
delay in initiation of movements, difficulty in walking and arising from sitting to standing, masked face with neither smiling nor blinking, loss of normal arm swinging when walking, a simian-like posture with flexion of all joints (ankles, knees, hips, back and neck) increased salivation, gait impairment with short and shuffling steps, apathy and inversion of sleep patterns (insomnia and daytime somnolence). He was in good health until October 1995 when he started complaining of blurred vision in the right eye, diagnosed by an ophthalmologist as serosanguineous posterior uveitis and exudative ablation of the retina. In November 1995 he noticed impairment of the speech; CT brain scan exam was normal. One month later in December 1995, blurred vision in the uninvolved eye appeared and a posterior uveitis and ablation of the retina in the left eye was confirmed by an ophthalmologist.

His past medical history included recurrent oral aphthous stomatitis, skin papules and pustules, and recurrent monoarthritis in right knee joint.

Physical examination showed aphthous stomatitis on the buccal mucosa. The lingual mucosa was covered with yellowish patches. He had inflamed pustular lesions on his face, neck and back and monoarthritis involving the right knee joint. The lungs were clear, cardiovascular exam was unremarkable, there was no hepatosplenomegaly and renal function was normal.

Ophthalmological examination disclosed bilateral posterior uveitis and retinal vasculitis. Visual activity was down to light perception in both eyes.

Neurologic examination revealed pseudobulbar palsy with dysarthria, dysphonia, positive snout, suck and palpmemtal reflexes and pathologically brisk jaw jerk. Slight and asymmetric bilateral pyramidal syndrome with brisk tendon reflexes predominantly in the left limbs, and left Babinski sign were evident. Muscle strength was preserved and there was no spastically increased muscle tone with clasp-knife phenomenon. Muscle lead pipe rigidity involving the four limbs with cogwheel phenomenon in the wrists, bradykinesia, masked face with loss of facial expression and impaired postural reflexes were found. There was loss of normal arm-swinging when walking, and when turning, balance was uncertain. There was gradual reduction in the size and legibility of his handwriting (micrographia). There was postural tremor in the extremities and myoclonic jerks involving the tongue and face muscles. The patient was in an apathetic state.

Laboratory examinations revealed normal haemoglobin, red and white blood cells, platelet and differential count, erythrocyte sedimentation rate, liver and kidney function tests and serum electrolytes. Blood glucose, prothrombin time, partial thromboplastin time, fibrinogen, serum protein and C-reactive protein were within normal limits. Serum immunoglobulins IgG, IgA, IgM, C3, C4, C9 were within normal limits. The titre of cryoprecipitates was increased, the ratio T4/T8 was normal, circulating immune complexes were within normal limits, antibody reactions against human mucosal tissue and antinuclear antibodies to ssDNA, dsDNA were positive, while reactions to RNP were negative. Cardiolipin antibodies examined by ELISA were positive (IgG-24 GPU, IgM-80 MPU). Serologic tests for syphilis, toxoplasmosis and neuroborreliosis were negative. Anti-HIV antibodies were not present.

Cerebro-spinal fluid (CSF) characteristics were: normal pressure, proteins 75 mg/dl, lymphocytes 112.10⁶/l. Electrophoretic examination was normal. The pathergy test was non-reactive at 48 hours. On haplotype analysis the patient expressed HLA B5 and HLA B27.

EMG findings confirmed the presence of myoclonic jerks of about 3 Hz frequency. The EMG examination also revealed postural tremor with 12 Hz frequency and synchronous pattern in antagonistic muscles, EEG showed increased slow waves and disorganization of the background activity.

Magnetic resonance imaging (MRI) T2/ISE and TSE/DUAL demonstrated small bilateral multifocal hyperintense lesions with predilection in right in periventricular white matter, brain stem and basal ganglia (Figures 1 and 2). The patient was treated by pulsed megadose steroids: 1000 mg of intravenous methylprednisolone and 1000 mg intravenous cyclophosphamide, both once a month and oral colchicine 1.5 mg daily. Anti-parkinsonian therapy with Sinemet® 750 mg daily was applied. No clinical improvement was achieved.

**DISCUSSION**

There are no specific laboratory or neuroimaging abnormalities in Behçet’s disease and the diagnosis is made essentially on clinical grounds. The diagnosis in our case was made on the criteria of the International Study Group for Behçet’s Disease:¹⁵ there were recurrent oral ulcerations, appearing three times in a six month period, observed by the patient and the physician; eye lesions (bilateral posterior uveitis and retinal vasculitis) observed by ophthalmologist and skin lesions (papulo-pustular) observed by physician before the corticosteroid treatment. The symptomatic triad is often not complete.¹¹ Our patient had no recurrent genital ulcerations, which may be absent in nearly half of the patients.¹¹ The pathergy test was negative, but in spite of this the diagnostic criteria were fulfilled.

The real problem in diagnosis of Behçet’s disease is differentiating from other forms of vasculitis such as systemic lupus erythematoses (SLE), which can produce oral ulcerations and iritis as well central nervous system involvement.¹⁴ Although anti-ds DNA antibodies are typically seen in SLE, low titers...
have also been demonstrated in the sera of patients with other connective tissue diseases.16 Our patient had oral ulcerations, an abnormal titer of antinuclear antibodies and anti-ds DNA antibodies, but had no malar or discoid rash, photosensitivity, polyarthritis, serositis, renal disorder, hematologic disorder, seizures or psychosis, and so did not fulfill the requirement of the American Rheumatism Association for 4 of the 11 criteria for SLE.16

Neurological involvement in Behcet’s disease usually begins with a definite acute attack1 and the differential diagnosis with multiple sclerosis may be difficult on purely clinical grounds.10 Some MRI findings are in favour of Behcet’s disease: involvement of basal ganglia, thalamus, central part of pons, and absence of periventricular predominance in white matter lesions.17 Lesions of the brain stem are consistently associated with severe neurological impairment such as tetraparesis and pseudobulbar palsy.17 The MRI abnormalities are not specific and can be observed in other central nervous system vasculitides; however, the brain stem involvement is rarely found in SLE. Cerebro-spinal fluid examination might be normal,7 or reveal slight lymphocytosis and mild elevation of protein,18 as in our patient. However the electrophoretic examination is usually normal in contrast to multiple sclerosis.19 High levels of intracranial pressure, high protein content and/or pleocytosis usually predict a poor prognosis.7 Although there are no specific laboratory investigations, an elevated sedimentation rate, C-reactive protein and serum IgA, IgG, IgM, and/or C9 levels have been reported.19 Elevated cerebro-spinal fluid IgM, IgG and IgA, and C3 and C4 concentrations have been reported,20 although in our case they were normal. Circulating immune complexes and autoantibodies to mucosal tissue and a decreased proportion of helper/inducer T4 lymphocytes have also been reported.9,19 Our investigations revealed antibodies to human mucosal tissue, antineural antibodies and increased cryoprecipitates, while indices of circulating immune complexes, the third (C3) and the fourth (C4) components of complement and the T helpers/inducers ratio were within normal limits. Anticardiolipin antibodies (ACA) have been found in some patients with Behcet’s disease, and are clearly associated with thrombosis at multiple sites.21,22 ACA antibodies IgG and IgM were positive in our case. It is known that HLA B5 is related to the ocular type, HLA B27 to the arthritic type, and HLA B12 to the mucocutaneous type of Behcet’s disease in some populations.19 The reported patient with ocular, arthritic and mucocutaneous symptoms was positive for HLA B5 and HLA B27.

The patient presented here has all the previously described main features of Behcet’s disease. The neurological involvement was extremely severe, despite the short course of the disease. Brain stem involvement is reported to be the most common and disabling manifestation of neuro-Behcet’s syndrome.6 Bilateral pyramidal lesions and pseudobulbar palsy might mimic parkinsonian syndrome due to muscle weakness and lability in facial expression. Apathy also adds to lack of normal facial expression. However, our patient had a clear lead-pipe rigidity with cogwheel phenomenon, postural disturbances, flexed gait and increased salivation, while the muscle strength was preserved. The impairment of vision in both eyes further increases the patient’s disability. It is noted that uveitis is more common among the patients with neurologic involvement.6,7 Generalized myoclonic jerks attributed to brain stem involvement,6 as well as diencephalic symptoms including sleep disturbances have been reported.23 Dyskinesia in patients with Behcet’s disease due to a discrete lesion in the internal segment of the globus pallidus has been described.12 Chorea in Behcet’s syndrome has also been described.13 Parkinsonian-like syndromes with rigidity and akinesia have been reported in some patients.11,14 In one case, Parkinsonian symptoms were combined with intention tremor.14 We were not able to find a report on patients with typical static Parkinsonian tremor. In our patient there was also no evidence for any static tremor. However, a postural tremor, with the characteristics of enhanced physiological tremor23 was found. Such tremor may sometimes be present in Parkinsonian patients, even without any static tremor.24 In patients with Behcet’s disease, foci of necrosis in the substantia nigra,25 and basal ganglia have been found on necropsy.11 Both gray and white matter are usually involved and microglial infiltration of the gray matter has usually been observed.11 Thus the Parkinsonian syndrome found in our patient might be due to involvement of both substantia nigra, and basal ganglia. However, the lack of improvement after levodopa therapy suggests predominantly basal ganglia involvement. Our patient had no improvement from treatment with high doses of corticosteroids and immunosuppressive agents, probably because of the multiple and severe involvement.

The presented case has a rare combination of all the main features of Behcet’s disease and Parkinsonian syndrome. Bradykinesia, muscle rigidity and impaired postural reflexes were the most predominant features. No typical static tremor presented. We were not able to find any publication on typical Parkinsonian syndrome in patients with Behcet’s disease. This case further emphasizes the wide spectrum of the neurological manifestations of Behcet’s disease.

References


