Neuro-Behçet’s disease: a clinical and radiological dilemma

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DESCRIPTION
Behçet’s disease (BD) is a clinical syndrome characterised by systemic vasculitis commonly involving the central nervous system. However, brain stem involvement is less pronounced. The lack of definitive conclusive tests with prime dependence on clinical criteria led to low applicability and diagnostic ambiguity in challenging cases of isolated organ involvement or atypical presentation.

We report a 44-year-old man who presented to the emergency department with 1-day history of undocumented febrile illness, drowsiness, nausea and headache. In further exploration of his presenting complaint, collateral history of mental and behavioural changes was prompted manifesting as paranoia, social withdrawal and memory disturbances over 2 months. Detailed assessment for proceeding head trauma, exposure history and substance misuse was unremarkable as well as thorough systemic review. Clinically, patient appeared apathic and hypophonic with delayed verbal responses. Meningeal signs were elicited coupled with cranial palsy (bilateral trigeminal, facial and bulbar involvement), subtle cerebellar dysfunction plus generalised hypotonia and brisk reflexes and distal muscle weakness primarily of lower limbs. Head-to-toe examination otherwise was unremarkable including funduscopic assessment.

Initial metabolic (electrolytes, serum osmolality, glucose, thyroid function, cortisol, angiotensin-converting enzyme), infectious (HIV, hepatitis, syphilis and Brucella serologies), serum electrophoresis and autoimmune (antinuclear antibody, rheumatoid factor, anti-neutrophilic cytoplasmic autoantibodies) screens were non-diagnostic. He underwent lumbar puncture twice during his admission, cerebrospinal fluid examination showed leucocytes 157/cumm (normal 0 to 10) with 95% polymorphs, erythrocytes 157/cumm (normal 0), protein 1.62 g/L (normal 0.15 to 0.45) and glucose 5.7 mmol/L (normal 2.2 to 3.9) and unremarkable microbiology, cytology, limbic, paraneoplastic and oligoclonal bands’ screens. A non-contrast CT scan of head did not show any acute insult; however, MRI revealed abnormal signal intensity involving brain stem, internal capsule and bilateral cerebellar peduncle (Figure 1). Review of performed electroencephalogram demonstrated evidence of mild-moderate diffuse encephalopathy with no epileptiform discharges or lateralising signs.

In his journey to recovery, he was initially treated in line of aseptic pyogenic meningoencephalitis. Where his meningism and acute complaints marginally improved, the consolation of other subacute neurological symptomology and deficits did not. After multidisciplinary case review with general medicine, diagnostic radiology, infectious diseases, rheumatology and neurology specialists; patient was started on 5-day course of pulse steroids and intravenous immunoglobulin in view of high suspicion for probable neuro-Behçet’s disease. By day 10 of admission, marked clinical and radiological response (Figure 2) was observed.

On discharge, patient was partially ambulatory with minimal assistance and near normalisation of his overall status. Patient sustained follow-up on outpatient basis with rheumatology and neurology services and currently on maintenance mycophenolate mofetil while tapering prednisolone therapy.

Behçet’s disease is unique model of systemic vasculitis with multiorgan involvement. The diagnosis is ascertained clinically in majority of cases. This case represented diagnostic challenge for a patient with BD’s neurological involvement in the absence systemic disease. The lesion mainly involved the brain stem carrying risk of haemodynamic-respiratory deterioration; thus, it
Images in... was crucial to reach a diagnosis and initiate the treatment in a timely manner.

Learning points

- Behçet's disease could present with isolated and critical neurological manifestations warranting prompt diagnosis and treatment.
- In the presence of undifferentiated heterogeneous complex medical presentation, a multidisciplinary team approach offers diagnostic superiority over fragmented care.

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