Polyminimyoclonus in Hirayama disease

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DESCRIPTION

A 26-year-old man, right-handed, otherwise healthy warehouse worker presented with painless left-hand weakness. He noticed difficulty with fine finger movements and thumb articulation that had been gradually worsening over the past 2 years. He denied any sensory changes or involvement of any other muscles. There was no history of recent trauma, neck pain or bulbar symptoms. He had no regular medications. On examination, mental status and cranial nerve testing were unremarkable. Tone was normal. He had atrophy of the left first dorsal interosseus (FDI) and tremulous movements of the left digits with irregular jerky movements suggestive of polyminimyoclonus (video 1). Weakness in his left hand was measured by Medical Research Council grade: left FDI (4−/5), second palmar interosseus (3/5), abductor digiti minimi (4/5), flexor pollicis longus (FPL, 4−/5) and abductor pollicis brevis (4+/5); with normal strength in all other muscle groups. Right arm and leg strength, reflexes, sensation and gait were normal.

MRI of the cervical spine without contrast revealed left hemicord flattening between C5 and C7 without cord compression, with epidural venous distension on flexion view (figure 1), characteristic of Hirayama disease. Sensory and motor nerve conduction studies were unremarkable, whereas electromyography showed chronic neurogenic changes in the left FPL, flexor digitorum profundus (median) and FDI muscles. He was treated conservatively with soft collar and had no progression of weakness at 6-month follow-up.

Hirayama disease, or monomelic amyotrophy, is an uncommon cause of distal arm weakness, clinically distinct from other motor neuron diseases as it has a benign prognosis. It primarily affects adolescents with male predominance. It is characterised by progressive distal upper extremity weakness and atrophy in the C7-T1 distribution, followed by spontaneous arrest of disease progression. The majority of cases begin unilaterally progressing to asymmetric bilateral involvement. Cold paresis and polyminimyoclonus are widely observed. MRI of the cervical spine may show subtle or no findings, but when performed in neck flexion classically shows anterior displacement of the posterior dura, enlarged epidural spaces with flow voids, flattening of the cervical cord with or without signal change and an enhancing crescentic-shaped mass in the posterior epidural space. Unlike other motor neuron diseases, the majority of cases stabilise within 5–6 years.

Polyminimyoclonus is defined as intermittent, low-amplitude, arrhythmic movements most

Learning points

► Hirayama disease should be suspected in young males presenting with progressive weakness of the upper extremities, especially when polyminimyoclonus is seen.
► Polyminimyoclonus should be distinguished from tremor, and when seen may be a clue toward a diagnosis of Hirayama disease or other conditions associated with anterior horn cell disruption.
► Investigation of Hirayama disease should include MRI in neck flexion, as the expected findings may not be present in neutral position, and establishes the diagnosis.
Images in…

commonly seen in the hands and fingers. The movements are accentuated when the hands are outstretched, fingers are extended or during action. It has been proposed that polyminimyoclonus associated with anterior horn cell disruption, such as in Hirayama disease, is a result of fasciculations occurring so frequently that it gives the appearance of myoclonic movement. This sign is one of the clinical hallmarks of Hirayama disease and has been reported in 61%–80% of cases. It should not be mistaken for tremor and when seen may be a clue towards a diagnosis of Hirayama disease as in this case.

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