Abstract
Hirayama Disease is a rare benign lower motor neuron disorder which is primarily affecting young males. It is characterized by the progressive weakness of the distal upper extremities followed by spontaneous stabilization of the symptoms. In this paper, we describe a 58 year-old female patient with a complaint of weakness in the right hand and forearm. Magnetic resonance imaging of the cervical spine established the final diagnosis of Hirayama disease. Hirayama disease when detected and intervened at an early stage of the disease process, can have a good prognosis. Clinicians and radiologists should be aware of the clinical features, as well as suspicious findings on neutral-position MR imaging and an additional neck-flexion MR imaging study should be arranged to confirm the diagnosis.

Keywords
Hirayama Disease; Motor Neuron Disorder; Myelopathy; Magnetic Resonance Imaging
Introduction

Hirayama disease (HD) is a lower motor neuron disease characterized by unilateral weakness and atrophy of the distal muscles of the upper extremity in the distribution of C7-T1 myotomes. Bilateral involvement is rare. It is a rare non-progressive spinal muscular atrophy seen predominantly in young men in late adolescence and third decade. The onset is insidious, the symptoms progress from three to five years, then the disease progresses spontaneously [1,2].

In this case report, a 58-year-old woman with gradually increasing weakness in the forearm was presented with MRI findings.

Case Report

A 58-year-old female patient; was admitted to the neurology clinic for 25 days with complaints of the increased weakness and occasional numbness in the right forearm. Physical examination of the patient was normal. Neurological examination revealed numbness in the distal right arm and muscle strength was 2/5. No pathology was found in the other neurological examination. Blood investigations including complete blood count, sedimentation rate, renal, liver, and thyroid function tests, creatine kinase, and vitamin B12 and vitamin D3 level were within normal range.

Electromyography (EMG) examination showed that the right radial and ulnar nerve combined muscle action potential (CMAP) amplitude was lower than the left. Other messages were in the normal range. In the right radial and ulnar nerve innervated muscles, there was a slight increase in polyphyletic, motor unit potential (MUP) times. No denervation potential was observed. The other muscles examined were normal.

In the cervical magnetic resonance radiography (MRI), sagittal plane T1 and T2 weighted sequences, axial gradient echo sequence for the discs and axial and sagittal postcontrast T1 weighted sequences were applied at a neutral position. MRI revealed flattening in cervical lordosis, spinal cord atrophy under C5-C6 intervertebral disc in T2-weighted sagittal sections, and increased signal in the posterior spinal cord at C2-C5 levels (Figure 1). In the differential diagnosis of the patient, the cervical MRI examination was refreshed with flexion in view of the Hirayama disease. Flexion MRI, in addition to neutral MRI findings, showed posterior dural displacement between C5-T1 levels, forward displacement in the spinal cord, enlargement in the posterior epidural space, and homogeneous contrast enhancement in the epidural distance in the postcontrast series (Figure 2). The patient was diagnosed with HD by MRI findings and was offered cervical neck and physical therapy exercises.

Hirayama disease was first described by Hirayama et al. in 1959 [1]. The exact pathogenesis of the disease is still unclear. The generally accepted hypothesis is cervical myelopathy induced by flexion. The underlying mechanism is the disproportion with the dura mater and the growth of the vertebral column. While the length of the spinal canal increases with flexion, the dura mater is stretched and separated from the vertebral canal wall. Anterior spinal artery irrigations may occur in the lower cervical spinal cord with anterior spinal cord compression and anterior displacement. Recurrent flexion may lead to chronic circulatory disorder, gliosis, and localized cord atrophy [3].

HD is more common in young males, especially among 15-25 years. Our patient is one of the few cases sampled in the literature in term of age and sex. For HD, self-limiting weakness and insidious onset in the hand and forearm, which are mostly unilateral, are typical. In our case, there were atrophy and numbness symptoms in unilateral forearm and hand muscles and there was no sensory or pyramidal path involvement, typical for HD. In HD, cervical radiographs revealed no specific finding except for flattening in lordosis or scoliosis. Specific features in neutral and flexion MRI have been defined in the diagnosis of HD in the literature. These are mainly spinal cord atrophy, asymmetric cord flattening, intramedullary signal changes in the cervical cord, abnormal cervical axis in neutral position, and the loss of the connection between the posterior dural sac and the lamina, forward displacement of posterior dura, the expansion in the posterior epidural space and contrast enhancement in this region with/without signal void areas suggesting dilated epidural venous plexus in flexion [4].

In our case, findings consistent with the literature were defined in the cervical MRIs in neutral-flexion positions and a diagnosis of Hirayama disease was made.

The differential diagnosis of HD includes spinal muscular atrophy, amyotrophic lateral sclerosis, post-polio syndrome, multifocal motor neuropathy, toxic neuropathy, and structural lesions of the cervical cord such as syringomyelia. These clinical entities are characterized by their characteristic clinical, radiological, and electrophysiological characteristics [1]. In degenerative diseases, disc-osteophyte complex, which may lead to secondary cord compression, can be seen in T1A sequences. In demyelinating diseases, T1A hypointense, T2A hyperintense, focal or diffuse cord lesion with variable diffusion characteristics can be seen in the spinal cord. In spinal cord infarction, T1A hypointense, T2A hyperintense, diffusion-restricting spinal cord lesion is seen in arterial irrigation area [5].

Dynamic cervical MRI, which has an important role in diagnosis in HD, is used in increasing frequency in patients who are investigated in terms of positional spinal cord compression, cervical degenerative disease, cervical myelopathy, rheumatoid arthritis, cervical trauma [6,7].
Conclusion

Limitation of neck flexion can be extremely beneficial if the myelopathy symptoms and signs of Hirayama occur early, even if they are nonprogressive. Clinicians and radiologists should be familiar with imaging findings of the disease, and the key role of flexion cervical MRI should be kept in mind when considering the differential diagnosis.

Scientific Responsibility Statement

The authors declare that they are responsible for the article's scientific content including study design, data collection, analysis and interpretation, writing, some of the main line, or all of the preparation and scientific review of the contents and approval of the final version of the article.

Animal and human rights statement

All procedures performed in this study were in accordance with the ethical standards of the institutional and/or national research committee and with the 1964 Helsinki declaration and its later amendments or comparable ethical standards. No animal or human studies were carried out by the authors for this article.

Conflict of interest

None of the authors received any type of financial support that could be considered potential conflict of interest regarding the manuscript or its submission.

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