Hirayama’s Disease: Case Report

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Abstract

The Hirayama’s disease affects the neurons of the anterior horn of the spinal cord, usually C7-T1 in young individuals, which causes weakness of the distal muscles of the upper limbs. We report a case of a 14 years old patient presented with significant left hand weakness associated with difficulty gripping objects. After performing a resonance of the cervical spine, there were few hyperintense signals in the anterior horn at the C5-C6 segment. In this situation we can differentiate this case from other conditions, making the correct diagnosis. It is important to report it due to the necessity of better methods that can perform the diagnosis.

Keywords: Lower motor neuron, Monomelicamyotrophy; Hirayama disease; Spinal muscular atrophy

Introduction

Hirayama’s disease (HD), also known as monomelicamyotrophy of distal upper limb, is characterized by pure distal motor atrophy of the upper limbs, affecting young men, in the muscles that are innervated by C7, C8 and T1 segments[1-4]. It is usually sporadic, it has an insidious onset and there is a slow progression followed by stabilization in 2-4 years. The majority of cases are encountered in Asia, mainly in Japan and India. There are few reports of HD in western countries[1]. The disease is relatively rare and predominantly unilateral, but bilateral cases exists in the literature[2]. In same cases the proximal limb can be affected, more than the distal limb[5-9].

Even though the physiopathology still remains unknown, neuropathologic studies reveal a focal lesion in the anterior horn motor cells of the spinal cord. HD is thought to be caused by a compressive myelopathy, which is developed due to the anterior displacement of the dural sac during the neck’s flexion. Its course is generally benign but a few patients remain disabled[10].

Case Report

KRS, a 14 years old man, student, eight months ago realizes difficulty in some activities that demand dexterity and gripping, just like practicing sports (basketball, swimming) due to muscle weakness. Three months ago, he noticed atrophy of his right forearm and hand (Figure 1). His past medical history was unremarkable with no trauma reference. No family members had neuromuscular complaints. Vital signs were normal during Physical Examination. An atrophy of right hand intrinsic muscles was noticed during Neurological Examination as well as a distal paresis with abolition of stilo radial reflex on the same hand. As the same Examination proceeded, no more abnormalities were registered such as no sen-
Hirayama’s disease was reported for the first time in 1959, but the benign course of the disease allowed larger studies in 1982, with the necropsy findings that showed ischemia of the anterior horn of the spinal cord, neuronal loss and gliosis[9].

Our patient is a 14-year-old male presenting an insidious case of distal atrophy of the upper limb, a typical case of Hirayama’s Disease. The diagnoses of the Benign Monomelic Amyotrophy (HD) was elaborated due to the classic history of paresis and unilateral amyotrophy (one member only). Based on diagnostic criteria, the beginning age associated with the absence of bulbar involvement and/or the pyramidal tract and miotomes of other members allows us to consider an Amyotrophic Lateral Sclerosis (ALS)[9].

The progressive spinal amyotrophies, because they are symmetric, bilateral and commonly with positive family history, were excluded too. Structural lesions like, for an example, disc protrusions, syringomyelia, cancer and arteriovenous malformations would show sensibility symptoms and also typical images seen at cervical MRI[9].

There are various theories for the disease’s physiopathology, but the mechanism is still uncertain. The most accepted theory is that a fleeting neck biomechanism, acting in both the dural sac and cervical spine, lead us to a circulation disturb and causing lesions at the anterior horn’s cells. This mechanism has been elegantly demonstrated by the study of somatosensory evoked potentials with neck flexion and neutral position. Patients with Hirayama disease have reduced amplitude of the N13 potential during flexion[7]. It’s known that image exams can show parenchyma changes in the low cervical region. Fleeting neck, the findings become more evident with a large reduction of the dural space[6-13]. At the MRI, findings are more evident, with an important reduction of the dural space. Using electrophysiology, we can see a normal sensorial neuro conduction and the motor conduction amplitude may be reduced, specifically the ulnar and median nerves. Besides this studies changes that match with denervation, it is possible to see fibrillation, positives waves and polyphasic potentials[9].

It’s not known why, but HD affects Japanese men. Some authors have related the association with allergy cases, with serum IgE elevation, considering the hypotheses that this disease may be an allergic myelitis. There are few reports of IgA deficiency in patients diagnosed with Hirayama’s disease, but in our patient this dosage was normal[10,14].

Patients treated with cervical collar, specially those who maintain a flexed neck for long periods of time, have improved. Treatment may still include physical therapy and de compressions might also contribute to clinical improvement[11].

Conclusion

We should be aware that the Hirayama’s Disease or Syndrome mainly affects young male patients who presents distal atrophy of the upper limbs. The patients frequently show improvement with the use of the neck brace, hence it should be introduced as soon as the diagnosis is confirmed.

References