Cervical Disc Herniation and Central Horner Syndrome

Hérnia de disco cervical e síndrome de Horner central

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Abstract

Central Horner syndrome is a rare condition, comprising a unique pathophysiological phenomenon. It results from vascular lesions, head or thoracic trauma. We describe a case of Horner syndrome associated to cervical disc herniation, and first-order neuron compression. To our knowledge, this is the second case reported to date in the literature.

Keywords

► síndrome de Horner central
► spinal cord lesion
► cervical disc herniation

Introduction

Horner syndrome, or oculosympathetic dysfunction, was described by the Swiss ophthalmologist Friedrich Horner in 1869,1 and represents the interruption of the cervical sympathetic nervous system. It is characterized by a symptom triad: ptosis (upside down ptosis), pupillary miosis and facial anhidrosis, although some authors associate a fourth symptom to the syndrome, facial hyperemia.2 It is an unusual entity that has no relation to age, gender or race. The syndrome prognosis depends on the etiology, and underlying patient condition. This case report refers to a rare cause for this entity, which resulted from the compression of the cervical sympathetic preganglionic nuclei by cervical disc core herniation.3,4

Case Report

A fifty-five years-old patient with complaints of neck and shoulder pain for a week, without reference to a triggering factor, namely exercise or a traumatic event. The symptoms had increased in the 24 hours previous to the hospital observation, without any relief with immobilization or positional correction. Four hours after pain exacerbation, he described motor deficit of right side limbs, without other neurological disorder.
Physical examination showed decreased muscle strength on the right arm and leg, grade 4 on Medical Research Council (MRC) scale, impairment of sensory function below C6-C7 dermatome level, increased deep tendon reflex at upper and lower right limbs, and positive Babinski sign. The cranial nerves exam showed anisocoria, with right pupil miosis and slowed photo-motor response, combined with upper right eyelid ptosis and elevation of the lower eyelid (Fig. 1). The patient presented facial anhidrosis, but he refused starch-iodine test. The ciliospinal reflex was absent on the right side, and no other abnormal finding was detected through cranial nerves exam. Brain computed tomography (CT) was normal, but cervical CT and cervical magnetic resonance imaging (MRI) showed a stenotic cervical spinal canal with disc protrusions at C3-C4 (major) and C5-C6, with evident spinal cord compression. Magnetic resonance imaging (T2 sequence) revealed spinal cord hyperintensity on the described levels; however, it had been conditioned by motion artifact (Fig. 2).

Patient was submitted to anterior cervical discectomy and fusion of C3-C4 and C5-C6, with placement of intervertebral polyetheretherketone (PEEK) cages on both levels (Fig. 3). He presented postoperative motor recovery, but sustained right pupil miosis and upside down ptosis (Fig. 4). On the first month evaluation, patient showed clear recovery, without motor or sensory deficit, and no evidence of oculosympathetic dysfunction (Fig. 5). Due to the progressive improvement of all the deficits, and the absence of identifiable lesions on brain CT, we decided not to conduct postoperative brain MRI.
Horner Syndrome etiology is diverse, from congenital, hereditary to acquired causes, which highlights the trauma. It may arise from brachial plexus lesions, chest, cervical or thoracic spine injury, tumor lesions, such as Pancoast, or carotid dissecting aneurysms. It is an entity repeatedly associated with thoracic, throat and neurological surgical procedures. A concern on anterior cervical disectomy, related to longus colli sympathetic chain retraction injury.

The sympathetic chain is a three-neuron pathway that originates in the hypothalamus. The first-order neuron runs from the posterolateral hypothalamus to the midbrain and pons. It extends to spinal cord intermediate-lateral column, between C8 and T2 (ciliospinal center of Budge). The second-order neuron consists on preganglionic pupil-motor fibers, emerging from T1 and extends to the cervical sympathetic chain, in close relation to the pulmonary apex and subclavian artery. The trunk enters the superior cervical ganglion where it synapses with postganglionic neurons at the carotid bifurcation level (C3–C4). It surrounds the internal carotid artery to the distal structures, covering the cavernous sinus, and follows the VI cranial pair (abducens) and ophthalmic branch of V cranial pair (trigeminal), innervating the pupillary dilator system, Muller muscle and the vascular complex of the face. The rostral ventrolateral medulla contains premotor neurons controlling cardiovascular conditions, whereas rostral medullary raphe regions are a candidate source of sympathetic premotor neurons for thermoregulatory functions.

The network interruption can be central, from the hypothalamus to C8-T2 level, or peripheral, including the cervical sympathetic chain, superior cervical ganglion or the internal carotid artery, with the classification of pre or postganglionic.

The occurrence of central Horner syndrome is relatively uncommon, and can usually be identified by the concomitant presence of hypothalamic, brainstem, or spinal cord signs and symptoms, which help to localize the lesion. This entity is not associated to the so common cervical spondylotic myelopathy. Myelopathy occurs as result of three important pathophysiological factors. Static mechanical factors, dynamic-mechanical factors, and spinal cord ischemia. It damages the posterior columns, spinocerebellar, and corticospinal tracts. Although as the corticospinal tract is first affected during spinal cord compression because it has a watershed arterial supply, the first-order neurons of sympathetic chain localized on Budge center, immediately lateral to the dorsal gray, are preserved by a common anterior and posterior blood supply. The descending sympathetic tract presents a lateral location that preserves it from mechanical trauma or compression.

The most typical presentation occurs as part of the lateral medullary syndrome (Wallenberg syndrome) due to posterior-inferior cerebellar artery or distal vertebral artery stroke. It manifests with the common symptom triad, but anhidrosis usually affects ipsilateral hemibody (not only the face). Patients with preganglionic lesions may have flush. This symptom, also called harlequin effect, occurs with exercise in some patients. Patients with postganglionic lesions may report ipsilateral orbital pain or migraine-like headache.

Differential diagnosis is required in all situations, especially on such unusual clinical setting, which combines a pyramidal pathway lesion together with the sympathetic nervous system impairment. It is necessary to consider the hypotheses of ischemic vascular injury, subarachnoid hemorrhage, epidural hematoma and space occupying lesions. As a complementary diagnostic method, MRI assumed main importance to differential diagnosis in Horner Syndrome, like in the case of carotid artery dissection, spinal cord infarction, and cervical or thoracic injury.

As shown, the present case constitutes a first-order neuron lesion. It resulted from the spinal cord compression by a right lateralized disc herniation and narrowing of the spinal canal, which affected the intermediate-lateral column. Radiographic evaluation excluded major brain lesions but showed spinal cord hyperintensity on T2 Fluid-Attenuated Inversion Recovery (FLAIR), mainly at C3–C4 level, an unusual location to cause Horner syndrome. Even in a trauma situation this would generally be ascribed to lesions at C5-C6 and T1-T2.

Neurosurgical intervention was crucial in this case. Recovery did not happen immediately after surgery, although motor function, pupillary and eyelid symmetry were restored in the following days.

In our research on PubMed and Google Scholar, we found that only the work of Hyunjin Ma and Insoo Kim reports Horner Syndrome due to a large left paramedian disc herniation with cord compression at the C4–5 level. Similarly to our case, it also presented total recovery after decompression with cervical disectomy and arthrodesis.
Conclusion

Horner syndrome is a rare clinical entity, even more unusual when related to cervical disc herniation. This is a case report with fast clinical progression, without previous symptoms, obvious trauma or cervical spine instability signs. In such cases, clinical and image findings are paramount, and timely surgery might dictate the patient’s prognosis.

Conflict of Interest
None of the authors has any potential conflict of interest.

References