Collet-Sicard syndrome: Case report

Isadora Afiune Thomé de Oliveira¹, Bruna Morais Cordeiro¹, Rafaela Dias Coelho¹, Arthur Fidelis de Sousa¹, Ygor Costa Barros¹, Kim-Ir-Sen Santos Teixeira²


Collet-Sicard syndrome is a unique condition involving lesions of the skull base, which affect both the jugular foramen and the hypoglossal canal, affecting the lower cranial nerves IX, X, XI and XII. Symptoms of this pathology include loss of taste in the posterior third of the tongue (IX nerve), vocal cord paralysis and dysphagia (X nerve), weakness in the sternocleidomastoid and trapezius (XI nerve) muscles and atrophy and paresis of the tongue muscles (XII nerve). The present report aims to describe the case in question, referring to a syndrome rare and little described in the literature. In the case of the patient reported, the conduct adopted was to perform radiotherapy sessions in order to prevent tumour progression since the surgery was contraindicated.

Key Words: Collet-Sicard syndrome; Schwannoma; Radiotherapy

INTRODUCTION

The first mention of the Collet-Sicard syndrome (CSS) occurred in 1915 by Frederic Collet, in reference to a soldier wounded by bullets during The First World War. Subsequently, Jean A. Sicard made new descriptions, culminating in the name of the syndrome [1].

It is described as a rare condition involving lesions of the base of the skull affecting both the jugular foramen and the hypoglossal canal. It is also known as condylar jugular syndrome, resulting from a unilateral and combined affection of the lower cranial nerves IX, X, XI and XII [2].

The possibilities of cause are numerous (Table 1) and among them it is possible to emphasize: Cranial base metastases, primary intracranial tumours (Example: Schwannoma), extra cranial tumours, trauma, vascular lesions, iatrogenic complications, or inflammatory processes [1].

TABLE 1 Causes of Collet-Sicard Syndrome.

<table>
<thead>
<tr>
<th>Causes of Collet-Sicard Syndrome</th>
<th>No. of cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Skull basemetastasis</td>
<td>12</td>
</tr>
<tr>
<td>Prostate adenocarcinoma</td>
<td>6</td>
</tr>
<tr>
<td>Breast adenocarcinoma</td>
<td>2</td>
</tr>
<tr>
<td>Carcinoma of the uterine cervix</td>
<td>1</td>
</tr>
<tr>
<td>Clear cell renal carcinoma</td>
<td>1</td>
</tr>
<tr>
<td>Disseminated adenocarcinoma</td>
<td>1</td>
</tr>
<tr>
<td>Colon carcinoma</td>
<td>1</td>
</tr>
<tr>
<td>Primary intracranial tumours</td>
<td>3</td>
</tr>
<tr>
<td>Jugular glomus tumour</td>
<td>2</td>
</tr>
<tr>
<td>Neurinoma hypoglossus</td>
<td>1</td>
</tr>
<tr>
<td>Extracranial tumours</td>
<td>5</td>
</tr>
<tr>
<td>Multiple myeloma</td>
<td>2</td>
</tr>
<tr>
<td>Hemangiopericytoma</td>
<td>2</td>
</tr>
<tr>
<td>Neck fibrosarcoma</td>
<td>1</td>
</tr>
</tbody>
</table>

In view of this scenario, the present report aims to describe the case in question, referring to a syndrome rare and little described in the literature, comparing clinical findings with current literature.

CASE REPORT

Patient I.L.B, male, 61-years-old, report complaints of dysarthria, dysphagia and reduced tongue mobility. After an otorhinolaryngological evaluation, a CT scan of the skull was requested (Figures 1A and 1B, Figures 2A and 2B) which localized expansive, solid, slow-growing, hypovascular formation located in the jugular foramen and extending inferiorly to the right carotid space of probable neoplastic etiology.

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Figure 1) In A, axial CT of the base of the skull without contrast. The arrow indicates the hypoattenuating lesion in the jugular foramen. In B, axial CT of the skull base, with iodized contrast, the lesion, indicated again by the arrow, does not emphasize.

Figure 2) In A, sagittal CT of the skull with contrast, in which the key (1) indicates the hypodense lesion in its longitudinal aspect. In B, coronal CT, with contrast, demonstrating, at the base of the arrows, the lesion in its frontal aspect eroding the jugular gulf.

Figure 3) In A and B, coronal T2-MRI, demonstrating lesion with hypersignal in the jugular foramen. In C, coronal MRI, T1 after contrast (gadolinium). The tumour enhances intensely. The base of the arrows represents the tumour of the neural sheath (schwannoma).

Figure 4) In A, axial MRI CISS, demonstrating the heterogeneous lesion in the jugular foramen (arrow). In B, T1 axial MRI after contrast, intense tumour enhancement (arrow base).

DISCUSSION

The symptomatology of Collet-Sicard syndrome described in the literature [3] involve loss of taste in the posterior third of the tongue (IX nerve), paralysis of the vocal cords and dysphagias (X nerve), weakness in the sternocleidomastoid and trapezius muscles (XI nerve) and atrophy and paresis of the tongue muscles (nerve XII). Symptoms manifested by I.L.B.

Most intracranial primary tumours affect adults, especially from 45 to 50 years, and among the causes, the most frequent are vascular [4]. The cause of CSS in the patient in question was the intracranial tumour in the neural sheath (schwannoma). Schwannomas are benign tumours, which develop mainly in the sensory part of the nerves (VIII and X) and rarely affect the nerve XII [5]. They mainly affect the female sex, and may be related to hereditary syndromes and exposure to ionizing radiation. By affecting important cranial nerves the surgery is controversial [6]. In this sense, the most recommended treatment is the neuroradiological follow-up associated with curative radiotherapy because it is a slowly evolving tumour. This therapy may be associated with microsurgical resection [7].

In the case of the patient reported, the conduct adopted was the accomplishment of radiotherapy sessions. Surgery was contraindicated due to the risk of affecting the sensitivity and tasting of the posterior third of the tongue, laryngeal and palate sensitivity, speech, sternocleidomastoid and hypoglossal muscle movements and tongue on the injured side if there were recurrences. However, tongue movement, swallowing and elevation of the right shoulder were affected and there was no improvement after treatment.

CONCLUSION AND FINAL CONSIDERATIONS

The present approach on Collet-Sicard syndrome is of extreme relevance for the medical community because it is a rare occurrence with few cases reported. In view of this it is important to question whether the reported syndrome is an underdiagnosed disease or whether it constitutes a really rare condition. Therefore, investments in researches and studies in this area are extremely relevant.

The symptoms initially reported by the patient and the adoption of radiotherapeutic therapy converge with data found in the literature in which the radiotherapeutic alternative is predominant. In addition, there is no vast database of the rare syndrome, therefore, studies on comorbidity are still necessary in order to improve the associated symptomatology. Therefore, it is of fundamental importance to continue research on Collet-Sicard syndrome, including its symptoms, diagnosis, treatment and clinical evolution, since it is a rare condition and still little described in the literature.
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REFERENCES