Paraganglioma of the cauda equina – Case Report and Literature Review

Paraganglioma de cauda equina – Relato de caso e Revisão da Literatura

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Abstract
Paragangliomas of the cauda equina are tumors of rare incidence, with ~220 cases described in the world literature. They are benign lesions, grade I by the World Health Organization (WHO), whose definitive diagnosis can only be made by immunohistochemical analysis. Its neuroendocrine nature is evidenced by the presence of chromogranin. The relevance of reporting this case is because paragangliomas of the cauda equina should be included among the differential diagnoses of intradural and extramedullary tumors, and especially because they can cause perioperative and intraoperative hypertensive crises by adrenergic discharge.

The present study presents the case of a 36-year-old male patient diagnosed with a lumbar spine tumor located in the central spinal canal that presented as cauda equina syndrome involving 4 months of bilateral sciatica, paraparesis, urinary and fecal retention. The diagnosis of paraganglioma was confirmed by immunohistochemical positivity for chromogranin after microsurgical resection of the tumor.

Keywords
► paraganglioma
► cauda equina
► spinal tumors

Palavras-chave
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**Introduction**

Paragangliomas of the cauda equina region are a rare incidence tumor, with ~220 total cases described in the world literature, and they correspond to ~3.5% of the tumors in this region.\(^1\)\(^,\)\(^2\) The first case was described in 1970 initially as a secretory ependymoma.\(^3\) From that date on, between 4 and 8 cases are usually described annually, with greater evidence since the eighties due to the development of more accurate immunohistochemistry techniques and the use of magnetic resonance imaging (MRI).\(^4\)

Paragangliomas are neuroendocrine tumors classified as grade I by the World Health Organization (WHO) grading system, have a slow-growing behavior, and arise from the paraganglia cells (group of neuroepithelial cells). There are two groups of paraganglia cells: one composed by adrenal medulla cells and the other composed by the extra-adrenal paraganglia cells.\(^5\)

The term pheochromocytoma is used for tumors that arise from the adrenal medulla, while paraganglioma is used for tumors from the extra-adrenal paraganglia. Paragangliomas may occur in any part of the body, but tumors of the carotid body and of the glomus jugulare constitute >90% of the reported paragangliomas.\(^2\)

An accurate preoperative diagnosis of paraganglioma is very difficult to achieve. A definitive diagnosis depends basically on the pathological examination and, especially, of an immunohistochemical study in order to differentiate it from other intradural extramedullary lesions, which should always be considered for differential diagnosis, that is, ependymomas, schwannomas, meningiomas, metastatic tumors, hemangioblastomas, and lipomas.\(^6\)

The goals of the present study are to report a case of a patient with a paraganglioma of the cauda equina region and to perform an extensive literature review on the topic.

**Case Report**

A 36-year-old male was admitted at the hospital with complaints of bilateral low back pain of 4 months of evolution associated with progressive paraparesis, muscle strength grade 4 on the lower limbs, proximal and distal. The patient also had urinary and fecal retention that were progressively worsening. An MRI of the lumbar spine was performed and evidenced an intradural expansive lesion at the level of the L2 vertebrae with an estimated dimension of 3.0 × 2.0 cm (►Fig. 1).

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**Fig. 1** Magnetic resonance imaging of the lumbar spine showing an intradural mass at L2. (A) T1 weighted with contrast sagittal view. (B) T2 weighted without contrast sagittal view. (C) T1 weighted with contrast axial view.
The patient underwent complete microsurgical resection of the tumor which arises on the filum terminale, under general anesthesia. The lumbar and sacral nerve roots were preserved. Intraoperative images are exhibited on Fig. 2. The completely excised tumor is exhibited on Fig. 3. Neuro-monitoring was not available.

An anatomopathological macroscopic examination showed a well-defined encapsulated red lesion measuring 3.0 × 1.5 × 1.5 cm, with a grayish smooth surface, elastic consistency, and small cystic cavities observed after slicing. The microscopic examination was suggestive of myxopapillary ependymoma under hematoxylin and eosin.

However, immunohistochemistry evidenced a paraganglioma with positivity for chromogranin and negativity for glial fibrillary acidic protein (GFAP), S100, and AE1/AE3 (Fig. 4).

The patient was discharged at the 4th postoperative day with complete improvement of motor function and partial improvement in sphincter disorders. There are no signs of recurrence in 2 years of follow-up (Fig. 5).

**Discussion**

The low incidence of this type of tumor makes this report more important in the literature. Although rare, these tumors may cause systemic manifestations because they have the biochemical feature of producing and storing biogenic amines, as well as the potential to produce peptide hormones (adrenaline, noradrenaline, and dopamine).

Even though they are unusual, there are reported cases of hormonal activity with abnormal urine noradrenaline concentration and features of vasomotor amine syndrome. The
limited occurrence of paroxysmal hyperadrenergic states in spinal paragangliomas may have two reasons: first, the incapacity of the tumor cells to secrete stored substances, or second, the incapacity of these substances to cause a clinical syndrome.7

The main symptoms caused by this type of tumors are low back pain, which may be present in 90% of the cases, and sciatica, described in at least 72%. Motor or sensory deficits are detected in 35% of the cases while sphincter and erectile dysfunctions were reported, respectively, in 14 and 1.2% of the patients.8,9

The group of patients at risk described on the literature are within the age range between 12 and 77 years old, with a peak incidence on the 4th and 5th decades of life, with a male predominance. The male to female ratio is 1.54.8,10

The step-by-step of the surgery performed on the aforementioned case is in line with the literature: prone position on surgical table, posterior lower back midline incision, laminectomy of the L2 and of the lower half of the L1, and opening of the dura mater followed by visualization of the nerve roots of the cauda equina. After careful arachnoid microdissection, the nerve roots were separated and a bright red mass was identified on the body of the filum terminale, with a large tortuous exuberant vessel coursing longitudinally on its wall. Bipolar coagulation of the referred vessel was performed, followed by coagulation of the proximal and distal portion of the filum terminale. The filum was proximally and distally sectioned and the tumor removed en bloc to prevent extensive bleeding and subarachnoid dissemination of the tumor.16

In cases in which only subtotal removal was achieved, recurrence will happen in ~ 10% in a period of 1 year. Recurrence does not necessarily mean malignancy, but a sign of suboptimal resection and regrowth. Encapsulated tumors are favorable to complete removal and have greater chance of cure. The options in cases of recurrence include reoperation and radiation therapy, and each case should be judged individually.8,17

Limitations

No immunohistochemical differentiation between chromogranin A or B was made in the current case. Synaptophysin was not available.

Conclusion

Paragangliomas of the cauda equina are rare tumors whose diagnosis is only confirmed by immunohistochemical analysis. They must be included among the differential diagnoses of intradural and extramedullary spinal tumors. Case reports become more important because, in addition to these tumors being in an atypical location, there are studies in the literature describing hypertensive crises during resection of the lesion due to adrenergic discharge.

Conflict of Interests

The authors have no conflict of interests to declare.

References