Acute onset of paraganglioma of filum terminale: A case report and surgical treatment

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ARTICLE INFO

Article history:
Received 20 February 2017
Received in revised form 6 May 2017
Accepted 7 May 2017
Available online 19 May 2017

Keywords:
Paraganglioma
Filum terminale
Cauda equina
Spinal tumor
Case report

ABSTRACT

INTRODUCTION: Paragangliomas of filum terminale are rare benign tumors, arising from the adrenal medulla or extra-adrenal paraganglia. These lesions usually present with chronic back pain and radiculopathy and only two cases of acute neurological deficit have been reported in literature.

PRESENTATION OF CASE: A case with an acute paraplegia and cauda equina syndrome due to an hemorrhagic paraganglioma of the filum terminale is described. Magnetic resonance imaging showed an intradural tumor extending from L1 to L2 compressing the cauda equina, with an intradural and intradural bleed. An emergent laminectomy with total removal of the tumor was performed allowing a post-operative partial sensory recovery. Histopathological examination diagnosed paraganglioma.

DISCUSSION: Paragangliomas are solid, slow growing tumors arising from specialized neural crest cells, mostly occurring in the head and neck and rarely in cauda equina or filum terminale. MRI is gold standard radiological for diagnosis and follow-up of these lesions. They have no pathognomonic radiological and clinical features and are frequently misdiagnosed as other spinal lesions. No significant correlation was observed between the duration of symptoms and tumor dimension. Acute presentation is unusual and emergent surgical treatment is fundamental. The outcome is very good after complete excision and radiotherapeutical treatment is recommended after an incomplete resection. Conclusion: Early radiological assessment and timely surgery are mandatory to avoid progressive neurological deficits in case of acute clinical manifestation of paraganglioma of filum terminale.

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1. Introduction

Paragangliomas of filum terminale are rare benign neuroendocrine tumors, arising from the adrenal medulla or extra-adrenal paraganglia, representing approximately 3% of cauda equina tumors [1]. The mean age of presentation is approximately 40–60 years with a slight male predominance [2]. Clinically and radiologically these lesions can be misdiagnosed as schwannomas or ependymomas and they often present insidiously with chronic back pain and radiculopathy. Reviewing the pertinent literature, only two cases of acute manifestation are described. We describe a case with an acute paraplegia and cauda equina syndrome because of an hemorrhagic paraganglioma of the filum terminale. We analyse the clinical, histopathological and radiological findings of these tumors, and discuss surgical treatment in line with the SCARE criteria [3].

2. Presentation of case

A 56-year-old man with a 10 years history of progressive low back pain and bilateral radicular leg pain without evidence of bowel or bladder incontinence or myelopathy, presented with a 4-day history of the acute-onset of a flaccid paraplegia with urinary retention, accompanied by complete sensory loss below L1. Emergent MRI (Magnetic Resonance Imaging) demonstrated a large intradural well encapsulated tumor extending from L1 to L2 compressing the cauda equina, measured about 4 cm in crano-caudal diameter, heterogeneously enhancing after gadolinium injection with serpentine flow-voids in the subarachnoid space cranial to the tumor and with FSE (Fast Relaxation Fast Spin Echo) MR images showing uniform hyperintensity suggestive of an intraslesional and intradural bleed (Figs. 1 and 2a–c). An emergent D12-L3 laminectomy was performed. Intraoperatively, prior to dural opening, the dural sac was noted to be dark blue in color and tense, because of underlying hematoma. The dura was initially opened above the tumor to prevent downward herniation of the mass, observing the egress of bloody cerebrospinal fluid under pressure and a large, well-encapsulated, dark red in color, mass pushing the dorsal nerve roots out of the dural opening. Caudally extending the midline duro-
Fig 1. Sagittal (B) T1-weighted (A), T2-weighted FrFSE (B) and T2-weighted FrFSE fat sat (C) magnetic resonance images revealing a large homogeneously hyperintense intradural lesion (yellow arrow) extending from L1 to L2, with enhancement after gadolinium and flow voids cranial to the mass indicative of venous congestion or high vascularity of the tumor.

Fig 2. Axial T1 (A) and T2-weighted (B) and coronal T1-weighted (C) images showing a large lesion (yellow arrow) taking up the vast majority of the cross-sectional area at L1-L2 level, obscuring the visibility of the cauda equina. Photograph of the hemorrhagic paraganglioma (4 × 2 cm) after excision en bloc (D). One year post-operative, sagittal T1-weighted image after gadolinium (E) demonstrating no residual or recurrent contrast enhancing tumor.

3. Discussion

Paragangliomas are solid, well-encapsulated, highly vascular, slow-growing neuroendocrine tumors arising from specialized neural crest cells [4]. Paraganglionic cells and the neural crest have a common origin, and during embryogenesis, they migrate along the neural tube. Paragangliomass result from dysfunction of embryonic paraganglia cell migration or non-regression. They can be found in adrenal and extra-adrenal tissues, and the extra-adrenal paragangliomas can be divided into sympathetic and parasympathetic types [1]. The sympathetic paragangliomas are usually secretory and produce catecholamines while parasympathetic paragangliomas tend to be non-secretory [5]. In the central
Table 1

Review of acute onset of paraganglioma of filum terminale/cauda equina.

| Author          | Age (years) | Sex | Symptoms                          | Location | Surgery (Emergent) | Post-operative outcome                                      |
|-----------------|-------------|-----|-----------------------------------|----------|--------------------|-------------------------------------------------------------|
| Nagarjun et al. | 36          | F   | Paraplegia, Cauda equina Syndrome | T12-L2   | GTR                | Partial motor-sensitive recovery, partial bladder incontinence |
| Ma et al.       | 51          | M   | Paraparesis, Cauda equina Syndrome| L1-L5    | GTR                | Partial motor-sensitive recovery                             |
| Present case    | 56          | M   | Paraplegia, Cauda equina Syndrome | L1-L2    | GTR                | Partial sensory recovery                                     |

Legend: GTR: Gross Total Resection.

The nervous system nearly 80–90% paragangliomas (predominantly parasympathetic) occur in the head and neck and typically arise in the carotid body or the glomus jugulare but other sites may be sella turcica, cavernous sinus, pineal gland, pituitary gland, cerebellopontine angle, and petrous ridge. At level of cauda equina or filum terminale they are very rare accounting for 2.5–3.8% of cases [6]. The first documented description of a paraganglioma in the cauda equina region was performed by Lernan in 1972 [4]. The diffusion in the central nervous system and distant metastases is rare [7]. These lesions are commonly encountered in the fifth and fourth decades of life with male predominance as reported by Gutenberg [2], and they are sporadic neoplasms, but approximately 1% of cases are autosomal dominant [5]. MRI is the gold standard radiological exam for the diagnosis and follow-up of paragangliomas or cauda equina or filum terminale, that may appear isointense on T1-weighted images and hyperintense on post-contrast T2-weighted sequences with homogeneous or heterogeneous enhancement [8]. However, these spinal tumors have no pathognomonic features and are frequently misdiagnosed as schwannomas, ependymomas, meningioma, teratoma, and hemangioma [9]. A serpiginous flow void from vessels associated with the upper pole of the tumor and a T2-weighted images revealing a hypointense tumor rim (cap sign) due to presence of hemosiderin, caused by a prior hemorrhage, can be present. Optional spinal angiography can reveal the highly vascularized pedicle [10]. Yang et al. reported intratumoral hemorrhagic cyst fluid [11], Miliaras et al. concluded chronic hemorrhage can occur in these tumors [12], whereas Li et al. reported the first case of spinal paraganglioma exhibiting subarachnoid hemorrhage [13]. Histologically paragangliomas are slow-growing benign tumors (WHO Grade I), comprised of two cell types, chief cells and spindle shaped sustentacular cells, which are classically described as “Zellballen” or nesting pattern. S100 staining is positive in sustentacular cells and can also be positive in tumor cells of ependymomas. Ependymal cells are GFAP positive whereas GFAP staining is negative in neoplastic cells of paragangliomas [1]. Nearly half of paragangliomas of cauda equina contain mature ganglion cells, but origin of this variation remains unclear [14]. The most common symptom is the low back pain with or without radiculopathy (50%), motor or sensory deficits can be present in less than 10% of patients and bowel or bladder incontinence is quite rare (3%) though some authors reported a higher frequency of sphincter and genital disturbance, compared to other tumors of cauda [12,14]. Paraplegia and sympathetic secretory symptoms related to catecholamine are uncommon [15]. No significant correlation was observed between the duration of symptoms and tumor dimension. In literature, only two cases with an acute flaccid paraparesis/cauda equina syndrome attributed to an intratumoral hemorrhagic paraganglioma, (Table 1) have previously documented [15,16]. Because the relatively benign natural history, observation can be a reasonable option in asymptomatic cases. Jansen et al. estimated the growth rate of head and neck paragangliomas with doubling time of >10 years with an average of 4.2 years [17]. Intra-operatively paragangliomas are most often located in the intradural-extradural compartment and they appear well encapsulated, purple, friable and hemorrhagic. The outcome is excellent after complete excision of the mass and the main technical problem is dense adhesion to nerve roots [18]. In the case of a total removal, however some authors recommend a long-term follow-up due to a possibility of recurrence, which is rare about 1–4% [7,11]. With subtotal resection, 10% paraganglioma of cauda equina recurred within one year following surgery, though Landi et al. reported a relapse 30 years after incomplete resection [19]. Radiotherapy is recommended after a subtotal resection [20].

4. Conclusion

Paragangliomas of the filum terminale are solid, well-encapsulated, highly vascular, rare benign lesions, which very infrequently present with acute manifestations. We report a case of acute flaccid paraplegia and cauda equine syndrome from an hemorrhagic paraganglioma, treated via emergent surgery with partial neurological recovery. Due to the lack of pathognomonic radiological findings, histopathological examination is the gold standard for diagnosis of these lesions. In case of acute presentation early recognition and timely surgical treatment are mandatory to avoid progressive neurological deficits while adjuvant radiotherapeutical treatment is important if an incomplete resection is performed.

Conflicts of interest

No conflict of interest.

Funding sources

No funding has been used for this research.

Ethical approval

No ethical approval has been applied for this case report study, only the written and oral consent by the patient.

Consent

A written consent has been obtained from the patient for publication of this case report and accompanying images and is available for review on request.

Author contribution

All the authors has contributed equally to the paper.

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