Solitary Fibrous Tumor/Hemangiopericytoma of the Cervical Spine – A Case Report

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Abstract

Introduction: Due to its rarity, the clinical and radiological characteristics of spinal solitary fibrous tumors/hemangiopericytomas remain largely unknown, with only 68 cases documented worldwide between 2000 and 2017.

Case Presentation: We report a case of a 39-year-old female patient who presented to the emergency department with complaints of persisting headaches, a numbing sensation in both arms and a progressive loss of manual dexterity. Physical and neurological examinations were unremarkable. A cervical contrast-enhanced MRI revealed an intradural intramedullary space-occupying lesion at the C3 level, accompanied by extensive edema. The patient underwent a successful cervical laminoplasty under neurophysiological monitoring with en-bloc removal of the mass. She experienced total resolution of her symptoms. Biopsy revealed a grade I solitary fibrous tumor/hemangiopericytoma of the cervical spine, positive for CD34 and STAT6. Ki-67 index was 2%.

Conclusion: This case report combines a rare tumor subtype with a unique presentation, and may further advance our understanding of this clinical entity.

Introduction

First described by Klemperer and Rabin in 1931 [1], solitary fibrous tumors/hemangiopericytomas (SFT/HPC) are an uncommon group of rarely metastasizing tumors of fibroblastic mesenchymal origin. Although commonly perceived as intrathoracic (most often involving the visceral pleura and submesothelial connective tissues), in reality some 60% of these tumors arise outside the thoracic cavity, including the central nervous system, and are now recognized to occur anywhere in the body [1]. These tumors are considered rare – less than 2 percent of all soft tissue tumors in one of the largest retrospective series [2], the most common single site being the pleura, with 900 cases reported worldwide up to 2005 [3]. 16% of SFTs/HPCs occur in the head and neck, including the central nervous system (CNS) [4]. SFTs/HPCs of the central nervous system are rare, usually dural-based, and account for less than 1 percent of all primary CNS neoplasms [4, 5]. These tumors typically occur at younger patients compared with other SFTs/HPCs. No known environmental or genetic risk factors are currently identified for SFTs/HPCs [6]. Of note, since the discovery of a unique and recurrent gene fusion for both SFTs and tumors histologically classed as hemangiopericytomas, the use of the latter term is discouraged in clinical practice, with restructuring of solitary fibrous tumor and hemangiopericytomas as one entity [5, 6].

Grossly, SFTs/HPCs are usually well circumscribed and rarely invade adjacent sites. Microscopically, these tumors are relatively hypocellular and include a thick network of collagen fibers. Mitotic activity is minimal and anaplasia occurs in less than 1% of cases. In contrast, the more aggressive phenotype (previously termed ‘hemangiopericytoma’ by neuropathologists) is highly cellular and tends to recur locally and metastasize [5, 7]. Immunohistochemical testing includes CD34, Bcl2, CD99, vimentin, and the highly specific STAT6 staining. Molecular testing includes the NAB2-STAT6 fusion gene – which appears to be a unique feature of SFTs/HPCs [8, 9].

While SFTs/HPCs are uncommon in the CNS (with only 112 reported cases worldwide until 2012) [10], SFTs/HPCs arising from the spine are very rare, with only 68 cases documented worldwide between 2000 and 2017 [11]. Based on recent reports, these tumors can be divided into 4 subtypes – vertebral, paravertebral, spinal cord and mixed, of which pure spinal cord SFTs/HPCs constitute the minority of cases. Complete surgical resection is the treatment of choice in most cases, as residual tumors may recur in 10–50% of patients and worsen prognosis and overall survival [10, 12, 13].

We hereby present a rare case of an intradural SFT of the cervical spine.

Case Presentation

A 39-year-old previously healthy female patient presented to the Rambam Healthcare Campus Emergency Department with complaints of persisting, bilateral temporal headaches for the past 7 days. The patient was of Russian descent, and her familial history was negative for hereditary diseases. The patient reported similar headaches, albeit to a lesser degree, for the past 10 years. During the last 6 months, the headaches had gotten progressively worse and were accompanied by episodes of vomiting. She was also complaining of frequent numbing and tingling sensations in both her arms for the past month, and had increasing difficulty in
performing gentle maneuvers, such as knitting or playing the piano. The patient had denied complaints of fever, blurred vision, weight loss or urinary problems, and was not prescribed any medications for the past year. Her physical and neurological examinations were unremarkable aside from slightly decreased pinprick sensation on the lateral aspects of both her arms. Routine laboratory tests were normal. Her brain MRI was unremarkable, while her cervical scan revealed an intradural, seemingly intramedullary space-occupying lesion at the C3 level, measuring 1.4X1.5X1.6cm. The lesion appeared isointense on T1 and hypointense on T2-weighted images, and was homogeneously enhancing following contrast material injection. A noticeable spinal cord edema was demonstrated above and below the lesion, from C1 through C6 levels. There was no clear visualization of a dural tail (Fig. 1). The patient was offered an elective surgery to relieve the ensuing mass effect and establish a tissue diagnosis. The following week, she underwent a cervical laminoplasty at C2 and C3 levels under electrophysiological monitoring. Following careful incision of the dura, a well-defined white intradural mass was detected and carefully removed en-bloc. Macroscopically, the tumor had both an intra and an extramedullary component, and was oval and viscous in consistency, with scarce vascularity. No major complications were documented throughout the procedure. On the following day, the patient was freely mobilizing. She experienced typical paraspinal muscular pains relieved with acetaminophen and reported a noticeable improvement of her neurological symptoms. Biopsy revealed a grade I solitary fibrous tumor/hemangiopericytoma of the cervical spine, positive for CD34 and STAT6 (Fig. 2–4). Ki-67 index was 2%.

Discussion And Conclusions

Due to its rarity, the clinical and radiological characteristics of spinal SFTs/HPCs remain largely unknown [13]. These tumors pose an immense presurgical challenge, since the vast majority of radiologically based diagnoses are inaccurate, judging by the literature we reviewed, with presurgical MRI sensitivity rates as low as 14% [13, 14]. In this case, the presurgical differential diagnosis was either meningioma or hemangioblastoma, and it was difficult to decide whether the lesion was intramedullary or not. The extensive edema surrounding the lesion made the possibility of a typical meningioma less likely, while hemangioblastoma usually exhibits extensive enhancement associated with a mural nodule, unlike SFT/HPC.

As mentioned, spinal SFTs/HPCs may arise in different locations within the spinal column, and determining the spinal anatomic structure responsible for producing these tumors is often difficult [15]. Table 1 summarizes the three largest case series on spinal SFTs/HPCs to date. Of note, the second series also includes vertebral and mixed vertebral-spinal cord cases, with only 3/11 cases representing ‘pure’ spinal cord involvement, like in our case [13]. The median age averaged from all three series was approximately 45 years, which is somewhat older than our patient. While back pain and localized neck pain were common complaints, headaches and vomiting were not listed in the series, and seem to represent a rare initial presentation of spinal SFTs/HPCs. Upper limb involvement in the form of paresthesias, radiculopathy and myelopathy, as reported in our patient, seem to be more predominant in the Marieniello et al series [10]. Anatomical location of the intraspinal lesion also varies significantly between different publications. Put together, cervical and lumbar lesions constitute the minority of spinal SFTs/HPCs (16/62 and 11/62, respectively). Similar to our case, all lesions were iso to hypointense on MRI T1-weighted imaging. T2-weighted images varied between series, with the majority being hypointense, as in our case. Gross total resection was achieved in the vast majority of patients (52/62 patients), with the posterior approach being the most prevalent. Pathological data, including mitotic activity and immunohistochemical stains were also similar [10, 12, 13].
## Table 1

A Review of the Three Largest Case Series on Spinal Solitary Fibrous Tumors/Hemangiopericytomas

| Series                        | Number of cases | Median Age | Clinical Presentation                           | Anatomical Location | Radiologic Appearance | Treatment and approach | Pathological data                                      |
|-------------------------------|-----------------|------------|-----------------------------------------------|--------------------|-----------------------|------------------------|------------------------------------------------------|
| Mariniello G et al (2012)     | 35 'intraspinal' cases (2 cases plus a review of 33 case reports), no bone involvement mentioned | 50 years | Myelopathy (57.1%), Myeloradiculopathy (20%), radiculopathy (22.9%) | Thoracic (54.3%), cervical (34.3%) and lumbar spine (11.4%) | MRI – 35/35 were isointense on T1 and hypointense on T2-weighted images. | All cases were treated surgically with laminotomy or laminectomy. 14/35 were intradural extramedullary, 9/35 were intramedullary and 8/35 were mixed intra and extramedullary lesions. Gross total resection was achieved in 33/35 cases. In all reports, the tumor was of hard consistency, scarcely vascular with a well-defined surface. | 35/35 were positive for CD34. The mean Ki-67 index was below 3%. |
| Yi X et al (2017) [13]        | 11 cases        | 42 years   | Back pain (45%), lower extremity hypoesthesia, paresthesia and urinary dysfunction (18%) | Lumbar (36.4%), thoracic (36.4%) and cervical spine (18.2%, mostly involving C2-C3 level) | 8/11 were well-defined. 5/11 were oval. CT – 11/11 slightly hypodense, MRI – 11/11 tumors were iso to hypointense on T1 and hyperintense on T2-weighted images. 9/11 showed marked heterogeneity on contrast enhanced CT or MRI. The mean long diameter was 6.24 cm. | All patients were treated surgically. 10/11 were operated on using the posterior approach, and only 1 patient was operated on using the anterior approach. Gross total resection was achieved in 10 patients (10/11, 90.9%), and subtotal resection (STR) was achieved in 1 patient. | Osteolytic pattern was found in 7/11 cases. Only 3/11 cases purely involved the spinal canal. 9/11 were positive for CD34, 7/11 had very low mitotic rate (less than 4 mitoses/10 HPF). |
| Series                  | Number of cases | Median Age | Clinical presentation                                                                 | Anatomical location | Radiologic Appearance                                      | Treatment and approach                                                                 | Pathological data |
|------------------------|-----------------|------------|---------------------------------------------------------------------------------------|---------------------|-------------------------------------------------------------|----------------------------------------------------------------------------------------|------------------|
| Wang J et al (2019) [12] | 16 (pure spinal cord involvement, no bone impairment) | 42.6 years | Localized pain (backache and neck pain, 94%), lower extremity weakness (25%), upper limb weakness (only one case) | Thoracic (56.2%), lumbar (18.8%) and cervical spine (25%) | CT – iso to hypodense (7/7), MRI – Intradural, intra or extramedullary lesions, iso to hypointense on T1 and slightly hyperintense on T2-weighted images (16/16). 11/16 were homogeneous and 5/16 were heterogeneous on post contrast MRI. Mean long diameter was 2.1 cm. | 6/16 were intramedullary and 10/16 were extramedullary. All patients were treated surgically with a posterior laminectomy. Gross total resection was achieved in 9/16, while the rest were treated with subtotal resection. No complications were observed. 6/16 underwent postoperative radiotherapy. | 16/16 were positive for CD34 and STAT6. The mean Ki-67 index was 4.6%. |

Spinal SFTs/HPCs are a rare and relatively benign clinical entity, often misdiagnosed preoperatively on imaging studies. Clinical presentation varies and ranges from localized pain to neurological complaints. Most lesions will be treated operatively with extremely high success rates, as gross total resection is the usual end result.

**Declarations**

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- **Availability of data and materials:** not applicable
- **Authors’ contributions:** HS performed the literature review, summarized the case and prepared the final manuscript, GR and AE analyzed the radiological data and provided important clinical insights when drafting this manuscript. GS provided neurosurgical insights, assisted in gaining access to all relevant materials and contributed to the final manuscript. All authors read and approved the final manuscript.
- **Ethics approval and consent to participate:** not applicable
- **Consent for publication:** Written informed consent was obtained from the patient for publication of this case report and any accompanying images. A copy of the written consent is available for review by the Editor-in-Chief of this journal.

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Figures
Figure 1

Sagittal T2- (1.a) and T1-weighted contrast-enhanced (1.b) MRI images through the cervical spine demonstrating a round intradural lesion at C3 level with extensive cord edema. Of note, the lesion is hypointense on T2-weighted images, with no clear evidence of a dural tail.
Figure 2

SFT/HPC, grade I – spindle cell tumor with a typical "patternless pattern", collagen bundles and staghorn blood vessels. Hematoxylin and eosin, x100
Figure 3

SFT/HPC, grade I – Positive staining for CD34, immunoperoxidase, x100
Figure 4

SFT/HPC, grade I – Diffuse positive nuclear stain for STAT6, immunoperoxidase, x100

Supplementary Files

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