Coexistence of Spinal Teratoma of the Conus Medullaris and Arteriovenous Malformation in an Adult: A Case Report

Bir Yetişkinde Konus Medullaris Spinal Teratomu ve Arteriyovenöz Malformasyonun Birlikte Bulunması: Bir Olgu Sunumu

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ABSTRACT

The coexistence of spinal teratoma of the conus medullaris and arteriovenous malformation (AVM) is exceptional, which has not been reported previously in the literature. The precise mechanism of the coexistence of these conditions is not known, however, the dysembryonic origin of spinal cord teratoma and AVM seems to play a part in this process. A 34-year-old male patient was admitted with lower back pain, bilateral lower extremity numbness and weakness, and sexual disturbance. Magnetic resonance imaging (MRI) showed an AVM extended cranially from the top of a heterogeneous expansile lesion of the conus medullaris. Surgical exploration and histopathological examination revealed a mature teratoma associated with the AVM. A literature review supported the dysembryonic origin of spinal cord teratomas and AVMs. This unique case may provide insight into the etiopathogenesis of the coexistence of spinal teratoma of the conus medullaris and AVM.

KEYWORDS: Spinal teratoma, AVM, Conus medullaris

ÖZ

Konus medullaris spinal teratomu ve arteriyovenöz malformasyonun (AVM) birlikte bulunması çok nadirdir ve literatürde daha önce bildirilmemiştir. Bu iki durumun birlikte bulunmasının tam mekanizması bilinmemektedir ancak spinal kord teratomu ve AVM’nin disembriyonik kökeninin bu süreçte bir rolü var gibidir. 34 yaşında bir erkek hastalı bel ağrısı, bilateral alt ekstremite uyuşukluğu ve zayıflığı ve cinsel bozuklukla başvurdu. Manyetik rezonans görüntüleme (MRG) konus medullariste heterojen ekspansil bir lezyonun üzerinden kranial olarak uzanan bir AVM gösterdi. Cerrahi eksplorasyon ve histopatolojik inceleme AVM ile ilişkili matür bir teratoma gösterdi. Literatür taraması spinal kord teratomları ve AVM’lerin disembriyonik kökenini destekledi. Bu benzersiz vaka konus medullaris spinal teratomu ve AVM’nin birlikte bulunmasının etiopatogenezi açısından bilgi sağlayabilir.

ANAHTAR SÖZCÜKLER: Spinal teratom, AVM, Konus medullaris

INTRODUCTION

Spinal teratomas are uncommon neoplasms, with an overall incidence constituting 0.2–0.5% of all spinal cord lesions (14). The conus medullaris is the most frequent site of an intramedullary teratoma (1). The majority of spinal teratomas coexist with congenital spinal malformations and tumors, such as split-cord malformations or spinal dysraphisms and carcinoid tumors (5, 20), but rare cases have been reported to coexist with intraspinal vascular malformations. Here, we present a case of the association of spinal teratoma of the conus medullaris with AVM in an adult. A literature review confirmed that this is the first reported coexistance of these conditions. We also consider the pathogenic and neurosurgical issues involved in such cases.

CASE REPORT

History and physical examination

A 34-year-old male patient was admitted with a 2-month history of lower back pain and bilateral lower extremity numbness and weakness. His symptoms were aggravated by exercise, especially walking. In addition, he had experienced sexual disturbance 2 weeks prior to admission. Neurological examination revealed mildly decreased motor function, bilateral grade-IV motor strength in hip flexion, knee extension, dorsiflexion, and plantar flexion, and hypesthesia from the T9–10 to the L1–2 level. Anal wink and sphincter tone were normal. The deep tendon reflexes of the knees and ankle jerks had diminished but were not pathological. No other anomalies, such as hypertrichosis, hemangioma, or dermal sinus tract on the back, were encountered during the physical examination.
**Imaging**

Magnetic resonance imaging (MRI) showed an expansile lesion of the conus medullaris with a heterogeneous appearance indicative of mixed composition at the L1–2 level. Most of the lesion was hyperintense in a T1 fluid attenuation inversion recovery (FLAIR) sequence, and it exhibited mixed signal intensity in T2-weighted images. No areas of the lesion were enhanced following gadolinium injection. These MRI findings suggested a teratoma.

The MRI additionally revealed intramedullary granular-like high signals in the T1-FLAIR sequence and multiple signal voids on T2-weighted images that suggested an AVM. This malformation extended from the top of the mass of the conus medullaris (L1) to the T12 level on the posterior aspect of the spinal cord. The lesion was enhanced after the administration of contrast. No other spinal malformation was detected by MR or radiographic imaging (Figure 1A-F).

**Operation**

The patient underwent a complete L1–2 and partial T12 decompressive lumbar laminectomy and excision of the spinal mass thought to be a teratoma. During surgery, the dura was opened through a midline incision to reveal a yellowish, irregular, fatty, cystic mass at the conus medullaris. The AVM was observed just superior to the mass, connecting

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**Figure 1:**

A, B) Preoperative T1-FLAIR and T2-weighted images showing expansile lesion of conus medullaris with heterogeneous appearance, associated with multiple signal voids (AVM) superior to the lesion that connect with the upper pole and extend cranially. C, D) T1-FLAIR images showing unenhanced lesion and enhanced AVM after administration of contrast. E, F) MR and radiographic images showing no other spinal malformations.
The adjacent cauda equina was compressed laterally by the cystic mass (Figure 2A). Needle aspiration was first performed to prevent the leakage of cystic fluid into the intradural space. A hairy substance was found in the aspirated fluid. After opening the capsule wall, the entrapped contents were released completely. Because the tumor adhered strongly to the conus medullaris and cauda equina, the spinal cord and the tumor were not separated by a distinct border. Only subtotal tumor resection could be performed under a surgical microscope. Because the upper part of the mass harbored a component of the AVM, they were excised together. No attempt was made to address other components of the AVM. Somatosensory evoked potentials that were used during surgery to avoid further morbidity during the resection of the tumor capsule from the nerve component.

**Postoperative course**

The patient experienced no postoperative neurological complications and his strength soon returned to normal. A postoperative MRI revealed that the teratoma had been removed and the AVM was intact (Figure 3A, B). The patient continued to experience bilateral lower extremity numbness and lower back pain, but these symptoms resolved completely within 2 months after the surgery. The patient’s sexual disturbance did not improve.

**Histopathological examination**

Tissues resected from the lesion on the conus medullaris, including the piece harboring the AVM component, were processed in the conventional manner with formalin fixation followed by paraffin embedding. Histopathological analysis indicated that the lesion was composed of mixed tissues from two germ layers, the ectoderm and mesoderm. Microscopic findings showed that the tumor contained stratified squamous...
epithelium with underlying sebaceous glands and other skin adnexal structures. Mature adipose tissue and blood vessels (primarily from the exophytic component) were also found, indicating a mesodermal origin. Microscopic examination showed an AVM in the teratoma, in which dilated thick-walled arteries were prominent. The histopathological diagnosis was a mature cystic teratoma associated with an AVM (Figure 2B–E).

**DISCUSSION**

Mature spinal teratomas are uncommon lesions, particularly in adults (6). Most reported patients with spinal teratomas presented with associated spina bifida or other dysplastic spinal abnormalities (14). In a recent study of 27 spinal teratomas, 21 patients had associated features of spinal dysraphism (16). Other lesions have been reported to coexist with teratomas, including lipomas, lipomeningoceles, and carcinoid tumors (5, 22). An association with a spinal-cord AVM, however, has not been reported previously in the English-language scientific literature. Spinal-cord AVMs are a relatively rare congenital condition that usually occur alone (4), but they may coexist with other vascular, cutaneous, and skeletal malformations. Some cases of intramedullary AVMs have been reported to coexist with lipomyelomeningoceles (9, 10, 19).

Teratomas are uncommon intraspinal lesions and their pathogenesis remains unclear (18). A dysembryonic or prenatal origin has recently been proposed, due to the dysfunction of several factors involved in gene functions and cellular inductive interactions (8). Ijiri (6) posited that the misplacement of pluripotential embryonic caudal mesenchymal cells during the secondary neurulation period, preceding closure of the neural tube, resulted in the formation of a teratoma without other spinal malformations. The development of this pluripotent mesenchyma may also expand into other tissues, including blood vessels. The combined lesion may thus be explained by the focal maturation of this pluripotent mesenchyma (primarily as a teratoma) in the primitive vascular plexus, followed by the failure of the capillary component of this plexus to develop (21). Thus, the combination of a spinal-cord teratoma with an AVM most likely originates from mesenchymal tissue that did not regress during the embryonic period.

The preoperative clinical diagnosis of a spinal-cord teratoma is difficult (12). MRI is regarded as the gold-standard technique for the preoperative diagnosis of a teratoma, given the differential intensity of the mass components and the presence of fatty tissue (13, 17). In our case, preoperative MRI defined the morphology and localization of the tumor in relation to the conus medullaris, leading us to suspect a teratoma. An AVM was also noted just superior to the tumor, connecting with the upper pole and extending cranially.

Spinal-cord teratomas are treated surgically, and radical removal of the tumor should be the aim of surgical intervention whenever possible (4, 9, 10, 12, 16). However, complete surgical resection is difficult in about 50% of cases because of the intimate adhesion of the tumor to the spinal cord and to the roots of the cauda equina (1, 2, 11, 14–17). After incomplete removal of the tumor, the remaining endodermal or ectodermal elements may resume growth. Recurrences are most common in immature or malignant teratomas, but growth occurs at a slow rate and the overall survival time for patients with mature teratomas is good (15). The management of spinal-cord AVMs is challenging because treatment-related morbidity is relatively high (3, 7). Spontaneous prognosis data are lacking, particularly for asymptomatic AVMs (3, 23).

In our case, only partial resection was possible due to the lack of a distinct border between the spinal cord and the tumor. Because surgical obliteration or excision of intramedullary malformations is difficult at best, and the natural history of asymptomatic unruptured AVMs is unknown (7), only the AVM harbored in the upper teratoma was excised.

Although the coexistence of spinal teratoma of the conus medullaris and AVM is exceptional, neurosurgeons should be aware of their potential association. The present case supports the dysembryonic origin of spinal-cord teratomas and AVMs. Radical resection should be the goal of spinal teratoma treatment, and long-term follow-up is necessary. Long-term observation is the recommended course of action for an associated asymptomatic and unruptured AVM.

**REFERENCES**