Angiomatous Meningioma Arising in Tonsil: A Rare Tumor at Rare Site

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Abstract

Angiomatous meningioma (AM) is a rare histological variant of meningioma, accounts for 2.1% of all meningioma, accounts for 2.1% of all meningiomas. It is World Health Organization Grade-I histological subtype of meningioma having predominance of blood vessels over that of tumor cells. We report a case of 45-year-old male presented with extradural AM in very unusual site, i.e., rectal tonsil, the great diagnostic difficulty was noticed due to its unusual site. Histopathology of this tumor showed thin walled vascular channels and cells with bland morphology in the background. The diagnosis was confirmed by immunohistochernistry. Tumor cells showed positivity for epithelial membrane antigen, vimentin, CD3₁, and progesterone. Primary extradural meningioma is a rare entity, extradural AM being rarer still. We present this rare case to highlight the unusual ectopic site of AM and its histopathological features to distinguished it from closely mimicking vascular tumors such as hemangioblastoma and hemangiopericytoma.

Key words: Angiomatous meningioma, Extrudal, Tonsil

INTRODUCTION

Neoplasms of the meninges are derived from mesenchymal elements normally present at this site and the most common tumor is meningioma.¹ This distinctive group of tumor accounts for approximately 35% of all primary central nervous system (CNS) neoplasms, 15% of intracranial tumors, and about 25% of intraspinal tumors, with an estimated incidence rate of 6.29/100,000 person-years.² Angiomatous meningioma (AM) is a rare subgroup of meningiomas, constitute 2.1% of all meningiomas.³ It is World Health Organization (WHO) Grade-I histological subtype of meningioma which has a predominance of blood vessels over that of the tumor cells.⁴ Vast majority of these tumors are histologically and clinically benign.⁵ Primary extradural meningioma is a far more rare entity. They have histologic features comparable to those observed in the CNS, including meningothelial, transitional, psammomatous, fibrous, angiomatous, and chordoid variants.⁵,⁶ They have been reported to occur in a variety of locations including calvarium, orbit, middle ear, paranasal sinuses, nasopharynx, neck, skin, lungs, mediastinum, and retroperitoneum.⁷

CASE REPORT

We present the case of 45-year-old male patient who came to the ear-nose-throat outpatient department with the complaints of difficulty in swallowing, persistent cough, and cold with a slight change in voice. His complaints began 1-year ago with slowly growing soft reddish brown mass over rectal tonsil. The case underwent tonsillectomy on the lesional side.

Pre-operatively the tumor was highly vascular present within the parenchyma of the tonsil. Complete surgical excision of the rectal tonsil done and the specimen was sent for histopathological examination in 10% buffered...
formalin to histopathology laboratory for routine paraffin sectioning. Hematoxylin and eosin (H and E) stained sections on examination revealed numerous small and medium sized thin walled vascular channels exceeding 50% of the area of the tumor. Most of them are small with hyalinized walls with intervening areas showing oval cells with abundant cytoplasm and vesicular nuclei (Figure 1). Some cells showed vacuolated and clear cytoplasm. The mitotic activity was inconspicuous. Mild degenerative nuclear atypia was present in few cells.

The impression of a vascular lesion was made, and the differential diagnosis included were AM, capillary hemangioblastoma and hemangiopericytoma depending on the prominence of vessels and occasionally non-meningothelial appearance of the tumor cells. However, the diagnosis of AM was confirmed by immunohistochemistry. The tumor cells showed positivity for vimentin (Figure 2), epithelial membrane antigen (EMA) and CD3 (Figure 3) and progesterone (Figure 4).

DISCUSSION

Meningiomas are meningothelial cell neoplasms, typically attached to the inner surface of the dura matter, emerges from meningothelial cells. Most meningiomas are benign and correspond to WHO Grade-I. The WHO classification scheme attempts to predict which tumors are most likely to recur, placing lesions with worrisome histopathological features into Grade-II (atypical) or Grade-III (malignant) categories. This grading system was developed primarily through clinicopathologic correlations using intracranial meningiomas. The current WHO classification has not been applied to a large series of extradural meningiomas. Hence, we discussed the pathology and possibility of developing primary extradural AM in an unusual site.

AM is one the rare subgroup of meningiomas, constitutes 2.1% of all meningiomas. It has the histological and clinical features of benign meningioma in which vascular component exceed 50%. Until now clinicopathologic
characteristics of AM have not been systematically analyzed and studied in detail, first of all by Hasselblatt et al. in 2004. They studied 38 cases of AM and confirmed the existence of this rare subgroup of meningioma.

Primary extradural meningiomas account for <2% of all meningiomas and angiomatous extradural meningiomas are rarer than that. Another large series from Thailand reports that 8% of meningiomas are primary extradural lesions. However, the meaning of extradural is quite variable in the published literature, with some authors including primary intracranial tumors that have extended extra cranially by direct extension or by distant metastasis.

Extradural AM have clinical, histological and immunohistochemical features identical to their CNS counterpart; such as the mean age of onset (AM: 51.8 years and meningioma 49.7 years) 5 years recurrence rate (AM 5.3%, meningioma 7%). Male to female ratio (AM 1:1.4, meningioma 1:2).

Histologically AM is composed of meningothelial and vascular elements. The vascular component consisted of a multitude of blood channels which ranged from capillary or sinusoidal type to vessels of relatively large caliber with markedly thickened walls.

The intervascular spaces were occupied by sheets of uniform polygonal cells with poorly defined borders of cytoplasm and centrally situated large oval shaped nuclei.

Because of such extremely vascularized nature of this meningioma, previously (it is a generally held) belief that the term angiomaticus is more a descriptor than a diagnosis and has no prognostic or any other clinical significance. Furthermore, it has been argued that the presence of this variant might be only justified to represent a niche for highly vascularized meningiomas to distinguish them from vascular tumors like hemangiopericytoma and hemangioblastoma. These three entities share similar clinical features, radiological findings, and histopathological features. Immunohistochemistry can be used to distinguish and to reach the final diagnosis. AM shows similar pattern of immunoreactivity as that of typical meningioma. Tumor cells of AM show positivity for EMA, cytokeratin, vimentin, desmoplakin, and focal positive reaction to antibodies to progesterone. Meningiomas also express steroid hormone receptors, more so for progesterone than estrogen. Lack of progesterone receptor has been correlated with a less favorable clinical outcome.

Further confirmation may be done by elector microscopy. Tumor cells on electron microscopy show cytoplasmic extensions occupying intervascular spaces formed varying degrees of inter-digitation.

AM had been divided into two pattern according to the diameter of vessels as macrovascular and microvascular by Hasselblatt et al. in 2004. According to their study macrovascular pattern shows >50% of vessels having larger size than 30 µm in diameter and microvascular pattern in which shows >50% of vessels were smaller than 30 µm in diameter.

It is the microvascular pattern which can be confused with hemangioblastoma.

Hemangioblastoma is slow growing highly vascular tumor of adults, histologically comprised two main components; stromal cells that are large and vacuolated and abundant vascular cells.

Stromal cells of hemangioblastoma lack endothelial cell markers, such as Von Willebrand factor and CD, and do not express endothelium associated adhesion molecules such as CD. Vimentin is the major intermediate filament expressed by stromal cells. Endothelial cells express receptors for platelet derived growth factor and vascular endothelial growth factor.

Hemangiopericytoma is a highly cellular and vascularized mesenchymal tumor composed of irregular cells with ill-defined cytoplasmic margins and staghorn vascular pattern.

Immunohistochemical stains play a vital role in differentiating hemangiopericytoma and hemangioblastoma from AM. Hemangiopericytoma is immunoreactive to vimentin and CD, but stains negatively with EMA, which is positive in AM. Even hemangioblastomas fail to stain with EMA and CD, which gives positivity in AM. The importance of differentiating these tumors needs to be emphasized due to the prognostic differences. Because AM belongs to Grade-I meningiomas, it has a favorable prognosis.

The embryonic development of CNS is a complex process involving migration, induction, specialization, differentiation, and remodeling. This process occurs in a system of co-developmental cooperation in coordination with the entire body. Malformations of the CNS are frequently associated with the malformations of the systemic organs and major heterotropias. Our case of primary extradural AM could be the result of these developmental malformations.

Each and every clinician should also consider ectopias and hamartomas in the differential diagnosis of the tumors arising in tonsils.
CONCLUSION

Although AM is a rare variant of meningioma, it shares similar clinical features and prognosis with benign meningiomas and has some unique clinical, radiological, histopathological, and immunohistochemical features. Primary extradural AM rarely presents with tonsilar involvement hence clinician should consider this entity in the differential diagnosis of tonsilar tumors. It may mimic other vascular neoplasms like hemangioblastoma or hemangiopericytoma creating a diagnostic dilemma which requires immunohistochemistry for final diagnosis.

REFERENCES


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