Difficult Diagnosis and Differentiated Treatment in Synchronous Thoracic Hemangiomas

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Abstract
Thoracic hemangiomas are rare vascular neoplasia among adults. Even with advanced imaging, diagnosis is sometimes impossible, thereby mandating histopathological examination. The differential diagnosis of thoracic hemangiomas can include arteriovenous malformations, thoracic sarcomas (angiosarcoma, paravertebral mesothelioma), neurinoma, and neurofibroma. We herein present a case which was initially suggestive of a malignancy that ultimately proved to be a capillary cavernous paravertebral and vertebral hemangioma. Given the unique topography of this lesion, a correct preoperative diagnosis was not possible and an interdisciplinary surgical approach was required.

Keywords: tumor, capillary-cavernous hemangioma, vertebral and paravertebral, surgical treatment

Introduction
Considered a relatively rare disease in adult pathology, hemangiomas represent a controversial entity in terms of their pathogenic, clinical and therapeutic characteristics. The nosologic framework of hemangioma continues to be debated, having been defined in the past as either congenital vascular malformations (hamartomas) or benign vascular malignancies with a locally destructive nature. In an adult population, hemangiomas, especially the cavernous type, are located in bones, especially the vertebrae, thorax, brain, liver, and skeletal muscle. Since the clinical and imaging apperance of hemangioma can mimic a malignant tumor, histopathological examination is frequently required.

Because there is no universally accepted standard treatment of hemangiomas, several therapeutic alternatives are generally considered: surgery, embolization, direct injection of ethanol (alcohol), chemotherapy, radiotherapy, corticosteroids, immunotherapy depending on age, topography, histopathological type, complications, and local destructive nature. In about 10% of cases, hemangiomas are synchronous. In these situations, the differential diagnosis is even more difficult; it is imperative to rule out malignancy. Ultimately, treatment varies depending on topography and the evolutionary stage of the lesion.

Since hemangiomas are relatively rare entities, with especially misleading clinical and imaging characteristics, their management can be challenging. The case we present herein both illustrates these difficulties as well as demonstrates the value of interdisciplinary collaboration.

Case Presentation
This 45-year-old male patient, without a significant antecedent history of disease, presented with a several year history of progressively severe chest pain that responded poorly to a range of pain medications. Despite his severe chest pain, physical examination was grossly normal. However, chest computed tomography (CT) revealed a solid tumor located in the left paravertebral muscles, adhering to the parietal pleura. Imaging was suggestive for a paravertebral mesothelioma or muscle sarcoma.
Within the posterior mediastinum, a contrast magnetic resonance imaging (MRI) revealed a left paravertebral tumor of 5.5 cm, located in the V-VI intercostal space. The lesion was extra-pleural in location, polilobar in shape and both hyperintense on T2-weighted and hypointense on T1-weighted MRI. Furthermore, imaging demonstrated the mass to be inhomogeneous, well-vascularized, and without invasion or destruction of the adjacent ribs or neural foramen (Figure 2A, 2B).

The appearance was most consistent with an intercostal nerve sheath tumor. Meanwhile, a second lesion was seen on MRI adjacent to and contacting the T6 vertebral body, but without violating the integrity of cortical bone (Figure 3A, 3B).
FIGURE 3: Aspect of magnetic resonance imaging with angiographic study: vertebral hemangioma T6 with cortical bone limits (A,B)
A. Sagital image B. Transversal image a level T6

FIGURE 4: Aspect of magnetic resonance imaging with angiographic study: paravertebral tumor with vertebral T6 contact.

Given a presumptive diagnosis of an intercostal neurinoma, but also considering the possibility of a paravertebral mesothelioma, the patient underwent surgery by the Thoracic Surgery Department of the St. John’s Hospital. A left axillary thoracotomy with sectioning of the latissimus dorsi and serratus muscles was performed. A purple, partially encapsulated polilobar lesion was found intraoperatively located in the
intercostal space, which was attached to a thoracic spine vascular/nervous pedicle. The neural foramen was closed and the tumor was found to have destroyed the adjacent intercostal muscle (Figure 5).

We performed a en-block resection of the lesion, including the parietal pleura of the thoracic spine and the remaining intercostal muscles. Because of serious intraoperative hemorrhage, it proved necessary to dissect, ligate and section the posterior intercostal artery (Figure 6). Parietal plural and paravertebral hemostasis was achieved with TachoSil (Figure 7).

FIGURE 5: Intraoperative appearance of the vertebral capillary cavernous hemangioma with intra-thoracic extension.

FIGURE 6: Intraoperative aspect after paravertebral tumor and parietal
pleura excision

Intraoperative aspect after paravertebral tumor and parietal pleura excision, with intercostal muscles and artery excision, after dissection, ligature and section anterior vertebral artery and dissection paravertebral simpatic lymph nodes at T5-T6 level. Foramen ovale is closed.

FIGURE 7: Intraoperative aspect. Hemostasis with TachoSil (paravertebral application)

Histopathological intraoperative examination revealed a cavernous capillary hemangioma, the findings of which were consistent with final paraffin-embedded microscopy (Figure 8).
FIGURE 8: Capillary cavernous paravertebral hemangioma. Microscopic study (20xHE).

The lesion was positive for CD34 immunohistochemical markers (Figure 9) and actine (Figure 10) within vessel walls, confirming its benign nature and vascular origin, with limited proliferative tendency.

FIGURE 9: Immunohistochemical study. CD 34 marker pozitiv on vessels wall. (20xCD34)
Postoperatively, there was a good outcome with healing per primam and no evidence of tumor recurrence after two years. Postoperative MRI confirmed a significant resection of the lesion which over the ensuing two years remained stable in size and continues to respect the osseous boundaries of the adjacent vertebra (Figure 11A, 11B).
FIGURE 11: IRM aspect two years ago at postoperative care.

A. Paravertebral region after surgery without recidives B. Vertebral body T6 hemangioma with cortical bone limits.

Discussion

Hemangiomas are a relatively rare disease in adult pathology, estimated to account for about 0.5% of all mediastinal tumors [1]. The most frequently thoracic localizations are in the vertebrae, in the intercostal muscles, in subcutaneous tissue, skin, lung parenchyma, in the ribs, and in the posterior mediastinum, and in 25-30% of cases, they may be multiple [2]. Mediastinal localizations are very rare. One hundred cases have been communicated in the literature with the favored location being in the posterior and anterior-superior mediastinum [3].

From a nosological point of view, it has been asserted that hemangiomas are in a class with congenital arteriovenous malformation despite a tumor-like appearance (hamartoma) [4]. However, the modern perspective classifies hemangioma as benign neoplasia of a vascular origin that include neoformation vessels and have the potential to be locally destructive [3, 5]. Furthermore, recently reported chromosomal alterations, “mass” effect with compression of neighboring structures, and a macroscopic appearance of solid or mixed tumors also support the status of a “true neoplasia” [4].

Histopathologically, two main types of hemangiomas are found among thoracic lesions: 1) a cavernous type, frequently with significant expansion of ducts that store considerable quantities of blood and which contains vascular lakes, and a pseudo-capsule without vascular elements as well as fibrosis, post-thrombotic drainage and smooth muscle cells. Such lesions can have either a nodular or diffuse appearance and harbor a tendency for spontaneous regression [4], and 2) a capillary type with smaller neoformation vessels, fibroblasts, and few mitoses within endothelial cells, and which, in the pediatric age group, have a proclivity for spontaneous regression [4].

The pathogenic defect among hemangioblastoma is believed to center on a dysfunction of angiogenesis occurring during periods of blood vessel formation within primitive vascular networks: in the plexiform stage, when there is a network of capillaries that communicate, may occur capillary hemangiomas or in the retiform stage when exist big tubular vessels, with a tendency to coalescence, it forms cavernous
Thoracic hemangiomas are rare malignancies with vascular origin, which have misleading clinical-evolutionary aspects, requiring a difficult differential diagnosis. In terms of imaging, MRI with three-dimensional angiography, and sometimes arteriography, represent methods of choice for diagnosis. The differential diagnosis should include the arteriovenous malformations, malignant cancers, and, in the particular case of synchronous lesions with paravertebral topography, ‘dumbbell shaped’ tumor type, because the treatment is different. The main treatment is surgical, based on the complete excision of the lesion, but there are complementary or alternative therapies, such as embolization, alcohol injection with spinal reconstruction, chemotherapy, corticosteroids, and anti-tumor immunotherapy. There is no standard treatment, but treatment options tailored to topography and evolutionary stage, especially in synchronous forms. The presented case is particular due to topography, histopathological structure, locally destructive nature, and ‘special type’ synchronization, which requires a ‘particular’ differential diagnosis and may represent a model for an interdisciplinary treatment approaches.

Conclusions

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