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Case report - Thoracic oncologic

Mediastinal Castleman’s disease mimicking thoracic paravertebral schwannoma

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Abstract

A 51-year-old female underwent resection of a solid lesion in the posterior mediastinum, preoperatively interpreted at imaging as thoracic schwannoma, requiring double sequential surgical procedure to be resected. The histologic examination of the resected mass diagnosed a hyaline-vascular Castleman’s disease.

1. Introduction

Castleman’s disease (CD) is a rare lymph proliferative disorder of unknown aetiology that often involves the thorax [1]. Since Castleman’s first description, three sub-types have been described with characteristic histological features, localisation, clinical presentation and prognosis: the hyaline-vascular subtype is generally localised, benign, asymptomatic, resectable with good prognosis and low recurrence rate; the plasma cell type and mixed type are more aggressive diseases with multifocal involvement of nodal stations and organs. This disorder is often misdiagnosed or undiagnosed without histological confirmation, even though an incorrect surgical approach can lead to intraoperative complications, such as massive bleeding [2]. We report a case of hyaline-vascular symptomatic CD of the posterior mediastinum, with the radiological appearance of a thoracic paravertebral schwannoma.

2. Clinical summary

A 51-year-old female presented for a physical examination for persistent cough. A chest radiograph revealed a left paravertebral nodular opacity. Computed tomography (CT) of the thorax performed with intravenous administration of iodine contrast revealed a multilobulated solid lesion in the posterior mediastinum, partly involving the lateral foramen between T-6 and T-7 (Fig. 1). The images were suggestive of thoracic schwannoma and the patient underwent gross-total resection of the tumour through a combined anterior and left trans-thoracic approach. The postoperative course was free of major complications, but the patient experienced important thoracotomy-related pain, not responsive to NSAID. At histological examination, the nodal tissue showed follicular hyperplasia, focally associated with atrophic germinal centres penetrated by hyalinised capillaries, resulting in the so-called ‘lollipop follicles’. Some foci of lamination of the mantle cell layers could also be seen. According to these microscopic findings and the clinical picture, a diagnosis of hyaline-vascular CD was then rendered (Fig. 2). At six-month follow-up patient is alive, painless and without evidence of recurrence of disease.

3. Discussion

CD is a rare lymphoproliferative disorder, first described in 1956 by Castleman et al. [3]. It is usually benign, idiopathic and the median age of presentation ranges from adolescence into the seventh decade [4]. It is rarely associated to autoimmune phenomena or within POEMS syndrome or in conjunction with a non-Hodgkin lymphoma especially in HIV positive patients. Clinically CD presents both as benign localised resectable disease or as an aggressive multicentric variant associated with systemic symptoms and frequent recurrence; in the latter, symptoms tent to mimic a lymphoma (fatigue, fever, night sweats, weight loss). Histologically three forms have been described: the hyaline-vascular type is the more frequent (85%) and often asymptomatic, the other sub-types are less frequent (15%) and more aggressive and symptomatic. About 70% of cases occurs in the chest [5], usually (90%) involving the mediastinum. Radiological evaluation generally shows a mediastinal or hilar mass, homogenously enhanced after contrast,
be considered, including CD. In both cases surgical approach is curative, so every effort must be made for the procedure to be radical.

References

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