Lymphangioleiomyomatose Pulmonaire (IAM) : Typical CT image Thoracic

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Clinical Image

Pulmonary lymphangioleiomyomatosis (LAM) is a rare lung disease characterized by interstitial proliferation of cells resembling dystrophic smooth muscle cells. It can occur sporadically or as part of a genetic disease; tuberous sclerosis of Bourneville.

This condition mainly affects young women of childbearing age, however it can occur after menopause [1].

LAM is characterized by progressive lung cystic destruction, frequently associated with renal angiomyolipomas. Patients with AML generally present with recurrent pneumothorax due to cyst rupture, or progressive dyspnea that progresses to chronic respiratory failure [2].

Imaging, in particular the high-resolution scanner, plays a key role in the diagnosis and follow-up of patients with AML. Typical appearance of LAM on the HR chest CT scan is characterized by the presence of multiple cysts (n >10), rounded, with regular contours, whose size varies between 2 to 6 mm in diameter, with regular thin walls not exceeding 2mm in diameter. thickness [3,4] (Figure 1, 2). Cystic lesions are homogeneously and diffusely distributed within normal lung parenchyma (Figure1, 2).

A very discreet Interstitial Syndrome can be associated. The presence of nodules with well-defined contours is found in patients with tuberous sclerosis [5].

Infracentimetric mediastinal adenomegaly are frequent. The main complications to look for on the chest CT scan: pneumothorax and chylothorax.

Chest CT is also performed as part of screening in patients with TSC looking for radiological signs of AML [6].

Figure 1 and 2: Chest CT; axial and coronal section; showing multiple diffuse cystic lesions in favor of pulmonary lymphangioleiomyomatosis.
AML should be differentiated from other cystic lesions of the lung, mainly histiocytosis X; which affects the smoking subject with a characteristic appearance on the scanner of a cyst of irregular lace shapes, predominant in the middle and upper lobe, but the distinction sometimes remains difficult in the advanced stage of the disease, and also with lymphocytic interstitial pneumonia (LIP) which is characterized by ground glass patches around the cysts, distributes around the bronchovascular sheaths.

Although the chest CT scan may be compatible or characteristic of AML, the definitive diagnosis is based on the combination of clinico-radiological criteria and the histological confirmation obtained by lung biopsy or biopsy of a lymph node or lymphangiomyoma [6].

References