Dramatic response to sirolimus in lymphangioleiomyomatosis

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A 41-year-old woman presented with a one-year history of progressive dyspnea, early satiety and abdominal cramping. A clinical diagnosis of lymphangioleiomyomatosis (LAM) was made based on a confirmed right-sided chylous effusion (milky, exudative and triglyceride level of 52.6 mmol/L) and characteristic cystic changes on a high-resolution computed tomography scan of the chest (Figure 1). A large left-sided retroperitoneal lymphangioleiomyoma supported the diagnosis (Figure 2), with no renal angiomyolipoma or clinical evidence of tuberous sclerosis. She was started on sirolimus therapy, which resolved her dyspnea and abdominal symptoms within six months. Her forced expiratory volume in 1 s (FEV₁) (1.6 L) and FVC (2.1 L) had markedly improved during this time interval to 2.6 L and 4.2 L, respectively. Her chest x-ray demonstrated almost complete resolution of the right pleural effusion (Figure 3) and the lymphangioleiomyoma had also decreased in size (Figure 4). The patient remains clinically stable on sirolimus with few side effects and her FEV₁ and FVC on therapy at nine months increased to 3.3 L (96%) and 4.8 L (118%), respectively. The present case is a dramatic example of a chylous pleural effusion and massive intra-abdominal lymphangioleiomyoma responding to sirolimus therapy.

Figure 1) High-resolution computed tomography image demonstrating cystic changes (small arrows) with right-sided chylous effusion (large arrow) before sirolimus therapy

Figure 2) Contrast computed tomography image of retroperitoneal lymphangioleiomyoma (arrows) before therapy. The dimensions of the lymphangioleiomyoma were: craniocaudal 16.5 cm; anteroposterior 6.8 cm; and transverse 8.7 cm. This is a typical computed tomography image of a lymphangioleiomyoma, which are commonly complex lymphatic masses with either thick or thin walls encompassing low-attenuation material consistent with chyle.
KEY LEARNING POINTS

- LAM is a rare progressive lung disease that affects women of child-bearing age.
- It is caused by proliferation of abnormal smooth-muscle-like cells and is characterized by cystic lung destruction, chylous effusions, lymphangioleiomyomas and angiomyolipomas.
- A clinical diagnosis of LAM can be made in patients who present with typical cystic changes on high-resolution computed tomography and angiomyolipomas, tuberous sclerosis or chylous effusion (1).
- Other considerations for multicystic lung disease include the following: emphysema, follicular bronchiolitis with or without lymphocytic interstitial pneumonitis, pulmonary Langerhans cell histiocytosis or Birt-Hogg Dubé syndrome.
- Before 2011, there were no proven treatments available for LAM; however, sirolimus has been shown to stabilize lung function, improve quality of life, reduce chylous effusions and decrease lymphangioleiomyomas (2,3).

REFERENCES