Persistent tachypnoea in an infant with cystic lung lucencies on CT scan

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A 1 year and 8 months old boy, with mild motor developmental delay and right hemiparesis following a perinatal brain injury, was brought with a history of fast breathing and cough for 2 months and low-grade fever of 10 days duration. He was second born to non-consanguninously wed parents by spontaneous vaginal delivery with delayed cry at birth and neonatal intensive care unit stay for 10 days. There was no exposure to cigarette smoke. On examination there was mild central cyanosis with pan digital clubbing; pulse rate was 102 per minute; respiratory rate was 44 per minute; oxygen saturation of 89% which improved to 95% with 2 L/min of oxygen. On systemic examination, cardiovascular status was found to be normal; respiratory examination revealed coarse crepitations bilaterally in the entire lung fields; abdominal examination revealed normal sized liver and a palpable spleen tip; neurological examination revealed mild right hemiparesis. His investigations revealed normal blood counts with mild microcytic hypochromic anaemia; sterile blood culture; chest radiograph revealed bilateral streaky interstitial opacities with honey comb pattern; work-up for TB and immunodeficiency states returned negative results. He was treated with ceftriaxone injection for 7 days without any improvement. A high-resolution CT scan of the chest revealed multiple cystic lesions of irregular size involving the parenchyma of entire

Figure 1 High-resolution CT scan of the thorax showing multiple cystic lesions of irregular size involving the entire lung fields.

Figure 2 The cellular infiltrate composed of Langerhans cells (red arrow) admixed with eosinophils (black arrow) (H&E ×100). Left upper inset—Langerhans cells with convoluted grooved nuclei (red arrow ×1000). Lower inset—Langerhans cells showing CD1a, S100 and CD68 positivity (DAB ×400).
Lung biopsy revealed nodular infiltrates of Langerhans cells, alveolar macrophages with immunohistochemical staining positive for CD1a, CD68, S100 (figure 2). Bone marrow examination, liver biopsy, bone scan and echocardiogram were normal. MRI of the brain was uninformative except for gliosis in left frontoparietal lobe. He was managed as isolated pulmonary langerhan cell histiocytosis (PLCH) with pulmonary dysfunction and treated with 6 weeks course of oral prednisolone and weekly vinblastine therapy. On follow-up his tachypnoea had decreased gradually and considerably with improved effort tolerance.

Langerhans cell histiocytosis (LCH) is a rare multisystem disease of unknown aetiology characterised by clonal proliferation and accumulation of dendritic cells in many organ systems resulting in a heterogeneous presentation.1 Lung involvement is a well-known complication of multisystem LCH, both in children and in adults. Isolated PLCH, on the other hand, has been reported largely in adult smokers, but very rarely in children.2 The classical radiological changes described in pulmonary LCH include bilateral nodular and reticulonodular infiltrates predominantly involving the upper and mid-lung zones (in adults) in the early stages and becoming predominantly cystic at later stages.3 In children, the subpleural parenchyma and costophrenic recess are more often involved than in adults. The pathological changes in PLCH consist of granuloma (rich in Langerhans cells, eosinophils, lymphocytes and macrophages) formation and peribronchiolectasia. Intervening lung parenchyma may appear normal.4 Langerhans cells can be identified based on the presence of an eosinophilic cytoplasm, elongated nuclei which have clefts and folds. Immunohistochemical staining will identify S100, CD1a antigen and CD207 (Langerin) which are expressed by histiocytes. For patients with progressive decline in lung function, corticosteroid therapy is indicated. If unresponsive, treatment with vinblastine or methotrexate is initiated.1

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