Bilateral non-tender parotidomegaly: a clue for underlying HIV infection and lymphocytic interstitial pneumonia

Deepanjan Bhattacharya, Pandiarajan Vignesh, Nameirakpam Johnson, Pratap Patra

DESCRIPTION

A 2-year-old boy presented with progressive distension of the abdomen, along with poor weight gain for 3 months. On examination, he had bilateral non-tender parotidomegaly (figure 1A,B), grade II clubbing (figure 2A), generalised lymphadenopathy, severe wasting and stunting, along with firm hepatomegaly. Examination of the respiratory, cardiovascular and central nervous system was unremarkable. He was found to be seropositive for HIV by ELISA. His CD4 count was 1194/mm³. Both mother and father were seropositive for HIV. A chest X-ray revealed bilateral reticular infiltrates (figure 2B). Workup for tuberculosis, cytomegalovirus and Pneumocystis jiroveci was negative. Screening for other opportunistic infections was also negative.

CT of chest revealed areas of ground glass opacities and multiple reticulonodular infiltrates, along with hilar, paratracheal and subcarinal lymphadenopathy (figure 2C).

Fine-needle aspiration from the lymph nodes was suggestive of reactive lymphoid hyperplasia. Echocardiography was not suggestive of pulmonary arterial hypertension. Serology for Epstein–Barr virus viral capsid antigen was positive. The PCR was not done.

A diagnosis of HIV infection with lymphocytic interstitial pneumonia was made, and the child was started on zidovudine, lamivudine and efavirenz. On follow-up for 3 months, there was a significant reduction in the size of the parotid gland.

Bilateral painless parotidomegaly occurs in about 10% of patients with HIV, and is usually secondary to CD8 lymphocyte infiltration and lymphoid proliferation, causing ductal obstruction and cyst formation. Soft tissue imaging is usually suggestive of gland enlargement with multiple cystic lesions, and aspiration cytology reveals foamy macrophages, lymphocytes and atypical squamous cells in a proteinaceous background. Management is usually conservative as most regress with antiretroviral therapy. However, close monitoring is advocated since malignant transformation needs to be ruled out.

Lymphocytic interstitial pneumonia is the second most common disease of the respiratory system in HIV infection affecting 30%–50% of children, after pneumocystis pneumonia, and is recognised as an AIDS-defining condition in the paediatric population. It is characterised by lymphoid hyperplasia of the bronchus-associated lymphoid tissue and infiltration of the pulmonary interstitium with CD8⁺ T lymphocytes and is usually seen with preserved CD4 counts. Children may also have generalised lymphadenopathy, hepatosplenomegaly or painless parotid enlargement secondary to the abnormal lymphoproliferative response, which may be a subtle marker of the respiratory involvement. It is usually a benign disease with 5-year survival more than 60%, however, the malignant transformation may be seen in less than 5% of patients.

Contributors DB, NJ and PP: patient management, literature review and preparation of the initial draft of the manuscript. PV: clinician in-charge, critical review of the manuscript for important intellectual content and final approval of the version to be published.
Learning points

- Bilateral non-tender parotidomegaly is a soft pointer towards underlying HIV infection in children.
- Lymphocytic interstitial pneumonia should be considered in children with HIV infection, clubbing, parotidomegaly and well preserved CD4 lymphocyte count.
- The pathogenesis is usually attributed to the abnormal proliferation of lymphoid tissue due to HIV or Epstein–Barr virus.

Funding
The authors have not declared a specific grant for this research from any funding agency in the public, commercial or not-for-profit sectors.

Competing interests
None declared.

Patient consent for publication
Parental/guardian consent obtained.

Provenance and peer review
Not commissioned; externally peer reviewed.

REFERENCES