Angiocentric immunoproliferative lesion and angiocentric lymphoma of lymph node in children. A report of two cases

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SHORT REPORT

Aim: To report two examples of an angiocentric immunoproliferative lesion (AIL) and angiocentric angiodestructive lymphoma (AL) presenting in lymph nodes in children. Most commonly involving extranodal sites, AIL/AL rarely presents in the spleen and lymph nodes.

Methods/Results: Case 1 presented as a cervical lymphadenopathy in a 3 year old girl being treated for pre-B cell acute lymphoblastic leukaemia. Histological and immunohistochemistry studies revealed an Epstein-Barr virus positive (EBV+), large B cell (CD20 and CD30+) AIL with large areas of necrosis, the whole resembling lymphomatoid granulomatosis. Case 2 presented as a large supraclavicular lymphadenopathy in a 13 year old boy. Histology and immunohistochemistry revealed an EBV–, large T cell (CD45RO, CD56, and CD30+) AL, presenting the features of so called angiocentric T cell/natural killer cell lymphoma, nasal type.

Conclusions: The term AIL/AL refers to a heterogeneous group of conditions not unique to a particular type of lymphoid cell. These lesions are easily recognised by the histopathologist because of their extremely unusual angiocentric pattern. Although rare, AIL/AL may present as nodal lesions in children ab initio.

Angiocentric immunoproliferative lesions (AIL) and angiocentric angiodestructive lymphomas (AL) encompass a heterogeneous group of conditions that have an unusual tendency to present and involve extranodal sites such as lung, skin, sinonasal areas, intestine, and brain. The main proliferative cells may be of the B cell, T cell, or T cell/natural killer (NK) cell phenotype. Occasionally, this peculiar main proliferative cells may be of the B cell, T cell, or T cell/NK cell phenotype. Such as lung, skin, sinonasal areas, intestine, and brain. The unusual tendency to present and involve extranodal sites commonly involving extranodal sites, AIL/AL rarely presents in the spleen and lymph nodes.

CASE 1

A 2 year, 10 month old girl presented in May 2001 with weakness, fever, and hepatosplenomegaly. The peripheral white blood cell count was 100 x 10^9/litre, with 89% of blasts. Flow cytometry revealed 99% CD45+ cells, 97% CD19+ cells, 86% CD20+ cells, 96% CD22+ cells, 92% CD34+ cells, 86% CD10+ cells, 92% HLA-DR+ cells, and 81% terminal deoxynucleotidyl transferase positive cells. The case was interpreted as a pre-B cell acute lymphoblastic leukaemia (ALL), stage III, FAB L1. The patient received steroids and cranial radiotherapy (12 Gy). After treatment, the patient’s white blood cell counts returned to normal. On November 2002, the girl presented with an enlarged, 4 x 4.5 cm lymphadenopathy on the left side of the neck. Human immunodeficiency virus and Epstein-Barr virus (EBV) serology were negative. After the diagnosis rendered with the lymph node biopsy the patient continued her treatment for the leukaemia. Currently (August 2004), she is alive and well.

The material was received as a consultation case. Histological sections showed a lymph node almost completely effaced by irregularly shaped foci of necrosis related to angiocentric and angiodestructive polymorphic cellular infiltrates, which occasionally obliterated the vessels. The foci of necrosis were limited by histiocytes and fibroblasts, focally arranged in a pattern suggesting palisade granulomas, although clearcut granulomas were not present. Among the vascular related cellular infiltrates large lymphoid cells with clear cytoplasm and a prominent lobulated or rounded nucleus with a prominent nucleolus were easily noticed. Some were arranged in concentric rings around the vessels. These cells also extended to the interfollicular areas in the more preserved zones of the lymph node. Immunohistochemistry revealed that the large cell component was CD20+ (fig 1) and CD30+. These lymphomatous cells were associated with a large number of CD45RO+ T cell polymorphic lymphocytes, which were prominent in the angiocentric lesions (fig 2). In situ hybridisation for EBV encoded RNA was positive in the large lymphomatous cells (fig 3). Both the histological picture and the immunohistochemical findings were reminiscent of that of lymphomatoid granulomatosis.

Figure 1  Polymorphic angiocentric proliferation with CD20+ large cells arranged around the vessel. The lower left hand corner depicts an area of necrosis. Case 1.

Abbreviations: AIL, angiocentric immunoproliferative lesion; AL, angiocentric lymphoma; ALL, acute lymphoblastic leukaemia; EBV, Epstein-Barr virus; NK, natural killer
CASE 2
This 13 year old boy was being followed up for a left sided supravacular lymphadenopathy that enlarged progressively over six months. Surgical resection produced a 5 × 4 × 4 cm lymph node with increased consistency, showing pink to whitish patches of the sectioned surface. Histology of the lymph node revealed features similar to those of case 1. However, in this specimen, the lymphomatous infiltrates were somewhat more monomorphous and there were clearcut remnants of lymph node histology at several points at the periphery. Less involved areas of the lymph nodes contained numerous eosinophils. The areas of necrosis showed geographical contours partially delimited by fibroblast spindle cells (fig 4). Densely collagenised areas were also seen. The whole picture suggested a progression from angiocentric and angiodestructive lesions leading to infaracts, which were then organised and transformed into scarred areas. Immuno-histochemistry showed the lymphomatous cells to be CD45RO+ (fig 5), CD30+ (fig 6), CD56+, and CD20−. In situ hybridisation for EBV encoded RNA was negative. The histological and immunohistochemical findings were similar to so called angiocentric T/NK cell lymphoma, nasal type.

The staging investigation revealed probable involvement of the mediastinal lymph nodes. Nose and paranasal sinuses were unremarkable.

DISCUSSION
AIL lesions involving lymph nodes are rare. Notably, a recent series from Japan include 14 cases grouped under the heading of AIL type II (four cases) and type III (10 cases), following an already reported classification. Not all were of the same immunophenotype, and the series encompassed B cell, T cell, and NK cell cases, sometimes associated with EBV. None presented in children.

AIL can occur in children but is seen in extranodal sites. To our knowledge, there are no reports of AIL or AL initially affecting lymph nodes in children.

Immunosuppressed patients, especially those undergoing treatment for leukaemia, can present EBV induced lymphoproliferative lesions, either nodal or extranodal, which usually respond well after the patient recovers an adequate immune status. However, we are not aware of descriptions of lesions with an angiocentric pattern under these circumstances. Lymphomatoid granulomatosis has been described in patients with Wiskott-Aldrich syndrome, a type of primary immunodeficiency, and in other conditions in which there is an underlying immune defect. It seems to be agreed that some cases originally defined as lymphomatoid granulomatosis are indeed examples of B cell lymphomas associated
with EBV, and that high grade AIL (indistinct from angiocentric lymphomas) can be shown to express EBV genomes by in situ hybridisation. Therefore, grade III lymphomatoid granulomatosis/AIL is regarded as a subtype of diffuse large B cell lymphoma. Notably, lymph nodes and spleen seem to be very rarely involved in lymphomatoid granulomatosis.

That is why we entertained the diagnosis of lymphoma in case 1, a child undergoing treatment for ALL for 18 months, who presented with an EBV+ B cell AIL in a cervical lymph node. EBV associated B cell lymphoproliferation may occur in children undergoing treatment for ALL. Reported cases have been in extranodal sites, have represented fatal acute EBV infection with haemophagocytic syndrome, or the patients have recovered after discontinuation of immunosuppressive drugs. Although speculative, we hypothesised that the uneventful recovery of our present patient may have resulted from the effect of the chemotherapy given for ALL acting on the EBV positive B cell angiocentric lymphoma.

The microscopic findings of case 2 correspond with those found in angiocentric T/NK cell lymphoma, nasal type, extranasal.

Molecular studies are beyond our scope and were not performed in these two cases.

The angiocentric distribution of the lymphomatous cells in these lesions is so unusual that it readily suggests the diagnosis.

In summary, we report two examples of the unusual involvement of lymph nodes by AIL/AL, which only very rarely presents in children.

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The patients gave their permission for these cases to be published.

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