Acquired ichthyosis and false-positive hepatitis A serology as paraneoplastic phenomena in anaplastic lymphoma kinase-positive anaplastic large-cell lymphoma

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DESCRIPTION

A 54-year-old autoclave technician was referred to the ambulatory care department of a UK district general hospital via his general practitioner (GP) with a 5-week history of a ‘profuse, papery’ rash (figure 1A), left groin lymphadenopathy, malaise, myalgia and fevers. He had a medical history of hypertension. Initial blood tests showed an elevated C-reactive protein (188 mg/L), alkaline phosphatase (510 U/L) and alanine aminotransferase (195 U/L) as well as a neutrophilia (9.2×10^9/L), monocytosis (1.2×10^9/L) and thrombocytosis (652×10^9/L). His GP had requested an ultrasound of the left groin; the sonographic appearances favoured reactive lymph nodes, measuring up to 2 cm each in short axis, that were attributed to his extensive active skin condition.

A liver aetiology screen including abdominal ultrasound revealed positive hepatitis A virus (HAV) IgM, negative HAV IgG and a reactive para-aortic lymph node with normal liver echotexture. He was diagnosed with active HAV infection and reactive acquired ichthyosis, which were managed supportively with intravenous fluids and emollients. A skin biopsy from the left flank showed mild epidermal orthokeratosis and spongiosis. The underlying dermis was oedematous with a mild perivascular lymphocytic infiltrate (figure 1B).

The patient developed supraclavicular lymphadenopathy during his inpatient stay, and a CT scan of the neck, thorax, abdomen and pelvis revealed extensive, mostly left-sided lymphadenopathy (figure 2). Lymph node biopsy revealed nodular deposits of lymphoid and plasma cells with clusters and sheets of large atypical multinucleated cells, which were CD3+, mostly CD20−, CD30+, MUM1+, CD4+, CD5+ and anaplastic lymphoma kinase (ALK) positive. HAV RNA was subsequently found to be negative, as was HAV IgM when repeated 3 weeks after the initial sample.

The patient was diagnosed with stage III anaplastic large-cell lymphoma (ALCL) and was commenced on treatment. His ichthyosis and hepatic dysfunction resolved rapidly within weeks, and he is now in complete remission following one cycle of cyclophosphamide, doxorubicin, vincristine and prednisolone (CHOP) and five cycles of CHOP plus etoposide chemotherapy.

Figure 2 Left groin lymphadenopathy demonstrated on CT scan (arrow).

Acquired ichthyosis, derived from the Greek word for ‘fish’ due to its scaly appearance, ichthyosis describes a disorder of cutaneous keratinisation that causes the skin to become dry and cracked. It may be inherited or
acquired, with the latter term referring to those cases caused by an underlying disease process, medication or nutritional deficiency.\textsuperscript{1} Acquired ichthyosis is an established paraneoplastic feature of Hodgkin’s lymphoma and has also been reported in a small number of cases of ALCL. This is the first case of ichthyosis feature of Hodgkin’s lymphoma and has also been reported in a

An important confounding feature in this case was a misleading positive HAV IgM, tested as part of a liver dysfunction screen. While the patient did not have any relevant recent foreign travel, his work as an autoclave technician led us to believe that he may have been exposed to the virus. HAV IgM usually becomes undetectable 6 months after infection,\textsuperscript{3} and as the patient reported no other recent illness nor vaccination for HAV, we presume that this was a false-positive result likely related to his diagnosis of ALCL. False-positive serology has been reported previously in T-cell lymphomas including ALCL for various diseases including Lyme disease,\textsuperscript{4} HIV\textsuperscript{5} and coeliac disease,\textsuperscript{6} with resolution of positive antibody tests after treatment of the lymphoma. There are no previous published cases of false-positive serology for HAV as a paraneoplastic feature of lymphoma.

**Patient’s perspective**

After being sent for an ultrasound by my general practitioner, it was suggested that the lymph node swelling was a result of a skin issue. Later on, I made a number of visits to ambulatory care whereby I was submitting blood tests. During this time, I also had a skin and lymph node biopsy. I got progressively worse and as a result of their findings was placed into quarantine. There was a room but no bed (they then sourced one). I was later released and returned to a ward a week later by my haematology consultant after cancer diagnosis. It took the Christmas period to ascertain which type of lymphoma I had. My all-round care including outpatient care was generally outstanding, and I am particularly thankful to my haematology consultant for her quick reaction and support; she was incredible. I must also mention the junior doctor for her caring support and the various consultants who helped me to get through this. The chemotherapy unit women and outpatient staff were also very professional and caring and kept me smiling. Many thanks to all.

**Learning points**

- Lymphoma should be considered as a differential diagnosis in all patients presenting with acquired ichthyosis.
- Positive serological tests for infectious and autoimmune diseases should be interpreted with caution in patients presenting with signs and symptoms consistent with lymphoma.

**REFERENCES**