Orbital Burkitt Lymphoma in an Immunocompetent Pediatric Patient

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1. INTRODUCTION

Burkitt lymphoma (BL) has been classified by the World Health Organization as a mature B-cell neoplasm, described in 1958 as a small, no cleaved B-cell, no-Hodgkin lymphoma, that was observed in African children in the endemic form, usually involving mandible, maxilla, orbit or adjacent orbital soft tissue structures like the eye1-3. Other two forms of BL are the sporadic and the immunodeficiency-associated. The sporadic form occurs in non-endemic areas and typically begins with an abdominal tumor with bone marrow involvement, rarely with ocular or orbital involvement. The immunodeficiency-associated form has been reported in association with HIV infection, SIDA and Epstein-Barr virus infection4-6.

2. CASE REPORT

A 5-year-old male patient is admitted with 2 weeks’ history of rapid progressive proptosis of his left eye with inferior dystopia, also presents upper gaze and levoversion limitation, with vertical diplopia in all extraocular movements (Fig. 1). He denies trauma history, systemic symptoms, weight loss or lymphadenopathy. His best corrected visual acuity was 20/40 in right eye and 20/200 in left eye, with a normal examination of the right eye, baseline Hertel exophthalmometer reading was 15 mm on the right eye and 19 mm on the left eye.

The following tests were requested: blood count, CT-scan and A scan ultrasound biometry and B scan ocular ultrasound.

The blood count shows lymphocytosis (9500/mm³), monocytopenia (53/mm³) and neutropenia (430/mm³). A 16.5 mm homogeneous ocupative lesion in the orbital roof was showed by B scans ocular ultrasound. The tumor caused indentation of the eye. With the A-scan ultrasound biometry, the tumor had low reflectivity, without flow. The CT-scan showed an isodense well-defined lesion in the orbital roof, with inferior displacement of the globe. There was no bone erosion (Fig. 2A,2B).

Fig. 1. External appearance with proptosis of his left eye with inferior dystopia

An incisional biopsy was performed by a superior eyelid crease approach under general anesthesia. Pathology reported a dense
lymphoid infiltrate disposed in a diffuse pattern, composed of medium size cells, with scarce cytoplasm. Multiple mitosis figures (16 in 10 high power fields), with apoptosis figures. A “starry sky” pattern was described, corresponding with an orbital Burkitt lymphoma (Fig.3). Epstein-Barr Ag test and ELISA for HIV were negative.

3. DISCUSSION

BL was initially described in the sub-Saharan Africa in pediatric population by the high incidence of jaw malignancy, and it was associated with Epstein-Barr infection. This tumor can arise from any location and the typically image observed in histologic exam is the “starry sky” pattern of lymphocytes. After some years a sporadic form was described in North American population. In contrast with the African tumors in jaw, maxilla and orbital bones, this sporadic form involving the abdomen, typically the ileocecal region, and rarely involving orbital or ocular structures.

Originally the BL immunocompromised variant was described in patients with AIDS. The infection by Epstein-Barr virus has been implicated in BL, being positive in 20% of sporadic cases, 40% in cases and as high as 95% in the endemic form. One large study has reported that cases with positive Epstein-Barr antibody had better outcomes.

BL is associated with c-myc translocation from chromosome 8 to 14, and this was observed in 66% of patients with orbital BL in some series. In the literature is reported too that some cases present a translocation of the Ig light chain from chromosome 2 or 22 to the c-myc region in chromosome 8.

Burkitt lymphoma has been described in people as old as 84 years, and this case was an immunocompetent woman which referred diplopia as main symptom, and was found to have a conjunctival mass including the parotid gland, lymph nodes and bone marrow.

Levine et al. conducted an 8-year review of 421 patients from the American Burkitt Cancer Registry, and reported a male to female ratio of 2.65:1 in patients younger than 13 years, and 1.35:1 in older than 13 years.

Knowles and Jakobic in their review of 60 patients with orbital lymphoid neoplasm reported that the most common physical finding was a palpable mass in 90% of cases, and proptosis is in 33%, like the presented case.

Central nervous system disease has been reported by Levine et al. as one of the most common sites for BL, with 5% of involvement and therefore a high suspicious must be maintained when patients presents with orbital signs.

Some cases could present systemic spread in sporadic BL but in the presented case the patient didn’t have it.
Immunophenotypically, all cytological variants express germinal center antigens BCL-6, CD10, as well as superficial IgM. Also, you can find leucocyte antigens CD45, CD43, as well as B-line antigens PA X5, CD79a, CD19, CD20, CD22 and plasmatic cell antigen CD38.

Prognosis remains guarded, with significant mortality within 1 year of presentation. Our patient was treated with systemic chemotherapy with the CODOX-M/IVAC regimen (cyclophosphamide, vincristine, doxorubicin, high-dose methotrexate / ifosfamide, etoposide, high-dose cytarabine), with remission after one year.

4. Conclusions

Sporadic orbital Burkitt lymphoma occurs in immunocompetent individuals with a wide age range. Almost 50% presents with an adjacent paranasal sinus involvement. Concurrent or eventual systemic involvement is common. Prognosis remains guarded, with significant mortality within 1 year of presentation. This is the first Orbital Burkitt Lymphoma in immunocompetent patient reported in Mexico.

References