Diplopia as the presenting feature of acute lymphoblastic leukemia

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ABSTRACT

Purpose: To report a rare case of acute lymphoblastic leukemia presenting with diplopia to an ophthalmologist.

Observations: A 29-year-old male patient presented to ophthalmology department with sudden onset of binocular diplopia in left gaze. Magnetic resonance imaging of brain and orbits revealed a thickened left medial rectus, with enhancement of right sixth nerve, bilateral third and fifth nerves. Bone marrow biopsy revealed acute lymphoblastic leukemia (ALL) with a Burkitt-type chromosomal translocation—t(8; 14) and the patient was started on chemotherapy.

Conclusion and importance: This was a case of incomitant esotropia worse with left gaze due to left medial rectus infiltration mimicking a left sixth cranial nerve paresis. Diplopia can be the only presenting symptom of ALL and it can involve either an extraocular muscle or a cranial nerve.

1. Introduction

Acute lymphoblastic leukemia (ALL) is a hematological malignancy, with less than 10% of adults having central nervous system (CNS) involvement. As cranial nerve palsies occurring due to ALL is rare, a correct and timely diagnosis presents a challenge. A high degree of suspicion combined with neuroimaging, cerebro spinal fluid (CSF) cytology and bone marrow biopsy confirms the final diagnosis. Here, we present a rare case of ALL presenting first to the ophthalmology department with diplopia.

1.1. Case report

A 28 year old male presented with acute onset binocular diplopia in left gaze of 3 days duration. There was no history of trauma. Clinical examination showed orthophoria per corneal light reflex in the primary gaze and esophoria on alternating cover-uncover test. Prism bar cover test showed 16 prism diopter of esotropia on fixing with the right eye. In right gaze, it showed orthophoria and 35 prism diopter esotropia in left gaze. On levoversion there was under action of left lateral rectus. On forced productions, abduction was restricted in left eye which was –3 i.e. inability to abduct the eye >22.5° past midline and hence the movement of left eye was approximately 25%. Diplopia chart showed maximum separation of uncrossed images in levoversion. Hess chart showed under action of left lateral rectus with overaction of right medial and also of left medial rectus suggesting of a possible left sixth nerve paresis. Forced duction test was not done. His best corrected visual acuity was 6/6 in both eyes and intraocular pressure on applanation tonometry was 13 mm Hg bilaterally. Dilated fundoscopy was normal with no papilledema.

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deposition disease (Fig. 2). These findings suggested a possible tubercular or inflammatory etiology deposits or immune complex deposition disease. Thus there was left eye abduction deficit with left medial rectus infiltration/enlargement on diffusion-weighted imaging, and enhancement of right sixth cranial nerve. CSF examination revealed elevated protein (150.4 mg/dl) decreased glucose (59 mg/dl) and elevated WBC count (50 cells/mm\(^3\)) with lymphocyte predominance (90%). With these findings tuberculous meningitis was suspected, as it is endemic in India. However real-time polymerase chain reaction (RT-PCR) for tuberculosis was negative.

Peripheral smear and bone marrow aspiration study was done which showed leukoerythroblastosis, atypical lymphoid cells with round large nuclei, coarse nuclear chromatin and moderate amount of basophilic cytoplasm with multiple cytoplasmic vacuoles (Fig. 3), suggestive of ALL L3 blasts. On flow cytometry, leukemic cells revealed expression of mature B cell markers like CD19, D20 and CD79a. Dim expression of clonality marker kappa-chain was noted. Findings confirmed mature B cell neoplasm. Fluorescence in situ hybridization test was positive for c-myc (8q24.21) rearrangement and karyotype confirmed the translocation t(8:14) (Fig. 4). Patient was treated with chemotherapy and steroids and his ocular symptoms improved after 2 cycles.

2. Discussion

There have been cases reported of leukemia presenting with orbital muscle infiltration, but in most cases it is during relapse of the disease,\(^1\),\(^2\) or in children.\(^3\) Recently Alrobaian et al. published a case of ALL presenting with sixth nerve palsy along with retinal hemorrhages, and postcontrast MRI of their case through the brainstem showed left sixth nerve enhancement involving the cisternal segment.\(^4\) However our case showed left medial muscle thickening and enhancement of multiple cranial nerves. In our case, the left sixth nerve paresis was mainly due to the intrinsic mechanical restriction caused by the thickening of the left medial rectus rather than a central cause. Version, duction, saccade amplitudes and forced duction can help distinguish muscle vs. nerve and extrinsic restriction. Apart from left medial rectus thickening, although the MRI did pickup multiple cranial nerves enhancement, the left 6th nerve was normal in its entire course and our patient had no obvious clinical signs or symptoms of the involved cranial nerves.

Workup of a case presenting with diplopia should include a thorough anterior segment examination, documentation of extraocular movements, fundus examination mainly to rule out papilledema, diplopia and Hess charting, routine blood work up (including blood sugar and serum lipid profile) and blood pressure measurement. Trauma should be ruled out. Forced duction test is to be performed if suspicious of restrictive palsy. All cranial nerves should be checked to localize the lesion. MRI brain with orbit and contrast should be performed. In our case diplopia
was primarily due to incomitant esotropia worse with left gaze, due to left medial rectus infiltration and restriction, rather than left cranial nerve 6 paresis causing left eye abduction deficit.

ALL is a hematological malignancy of lymphoid line that originates in the lymphoid precursors of bone marrow, lymph nodes and thymus. Less than 10% of adults with ALL have CNS involvement with significant neurological dysfunction such as cranial nerve palsy.6 ALL with CNS involvement can be rapidly fatal if untreated. Several risk factors are associated with the development of CNS involvement in ALL. Age seems to be a key factor with a higher incidence in young adults.7 Mature B cell subtype is also associated with an increased risk of CNS spread. Leukemia causes neurological dysfunction directly by invading the CNS or indirectly by altering hematological factors; the former occurs by dissemination of leukemic cells in the subarachnoid space and leptomeninges, while latter occurs because of hyper viscosity syndrome secondary to high levels of intravascular leukemic cells. Infiltration of individual nerves with leukemic cells or ischemia induced damage of the vasa nervorum causes cranial nerve manifestations.7

Besides the clinical evaluation of neurological signs and symptoms, three independent techniques are used to diagnose CNS disease in ALL patients: CNS neuroradiology, CSF cytology and flow cytometry.8 When one is considering an infiltrative etiology based on neuroimaging showing enhancement of cranial nerves with heterogeneous extraocular muscle enhancement and leptomeningeal/ependymal involvement, as in our case, CSF studies should be performed looking for malignant cells (flow cytometry included, especially when protein and cell count in the CSF is high), in addition to the routine tests. From endemic countries, more common diagnosis like tubercular meningitis should also be ruled out first. In our case RT-PCR was negative for tuberculosis. Bone marrow biopsy has to be performed if the CSF analysis comes inconclusive.

The concomitant use of neuroradiology, bone marrow biopsy and flow cytometry confirmed the diagnosis of ALL. We believe that a correct diagnosis has been made here without resorting to much intervention, utilizing the unique capabilities of both radiology and histopathology.

3. Conclusion

Thus a case of ALL presenting only with diplopia is reported here. Though the MRI showed multiple cranial nerves affected, the patient had no other symptoms or signs. Diplopia was probably due to the leukemic infiltrate on the medial rectus, again an unusual presentation. So a high degree of suspicion, a combined use of neuroimaging CSF analysis, cytoflowmetry and bone marrow biopsy confirmed the final diagnosis of acute lymphoblastic leukemia.

Patient consent

Consent to publish this case report has been obtained from the patient in writing.

Disclosures

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Authorship

All authors attest that they meet the current ICMJE criteria for Authorship.

Declaration of competing interest

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