Proptosis as Initial Presentation of Acute Lymphoblastic Leukemia in a Child with no associated symptoms: A Case Report

Kourosh Goudarzipour MD¹, Ahmad Mohammadi MD¹, Reza Taherian MD, MPH², Mehran Arab Ahmadi MD, MPH^{3,*}, Behdad Behnam MD⁴, Niloofar Ayoobi Yazdi MD³

1. Pediatric Congenital Hematologic Disorders Research Center, Shahid Beheshti University of Medical Science, Tehran, Iran.

2. Department of Radiology, Faculty of Medicine, Hamadan University of Medical Sciences, Hamadan, Iran.

3. Department of Radiology, Imam Khomeini Hospital Complex, Tehran University of Medical Sciences, Tehran, Iran.

4. Department of Internal Medicine, Firoozgar Hospital, Iran University of Medical Sciences, Tehran, Iran.

*Corresponding author: Dr Mehran Arab Ahmadi, Department of Radiology, Imam Khomeini Hospital Complex, Tehran University of Medical Sciences, Tehran, Iran. Email: mehran_arabahmadi@yahoo.com. ORCID: 0000-0001-9695-1705.

Received: 21 December 2017 Accepted: 05 October 2019

Abstract

Acute lymphocytic leukemia (ALL) is one of the frequent malignancies in pediatrics and involves bone marrow and extramedullary sites. Proptosis as extramedullary involvement of leukemia usually present in acute and chronic myeloid leukemia. It is extremely rare for ALL to present initially as proptosis.Here, a-21-month-old boy was presented with proptosis without any associated symptoms except lymphadenopathy. He was referred with the impression of malignancy from an ophthalmologist. After bone marrow biopsy which showed 33% blast cells, all positive for CD10, CD19, and CD79, the diagnosis of pre-B cell ALL was finally made. His symptoms were improved completely 16 days after starting standard protocol for ALL.Afterone-year follow-up, he was free of any symptoms.According to this initial presentation of ALL and no typical associated symptoms, it is important to make rapid diagnosis and start the treatment in the childhood.

Key words: Acute lymphoblastic leukemia, Proptosis, Childhood

Introduction

Acute lymphoblastic leukemia (ALL) is the most common malignancy of the childhood and it accounts for about onequarter of all childhood cancers (1, 2). Typically, the child is presented with nonspecific symptoms, such as fever, bleeding, bone infection, pain, or lymphadenopathy (3). The most common extramedullary site of ALL after the testicles meninges and is ocular involvement which may be seen on careful ophthalmological investigation in up to one-third of the children with newly diagnosed ALL (4). However, orbital infiltration is more common in other leukemic type hematopoietic cancers and is accounted as a rare finding in ALL (5). It presents with proptosis, lid edema, chemosis, and pain (6, 7) and is usually associated with other non-specific symptoms of ALL (8, 9). This condition might be confused with other diseases involving orbit, such as myeloblastic leukemia, hypereosinophilic syndrome, myelofibrosis with myeloid metaplasia, and polycythemia vera (10). Obviously, the precise diagnosis has an important role in prompt intervention and prognosis of the disease.

Case report

The patient was a 21-month-old boy referred to hematology and oncology department of Mofid Children's Hospital in Tehran (capital of Iran) in 2016 complaining of left eye proptosis without erythema, tenderness, and tearing. His problem was started acutely and he had no other symptoms like bone pain or fever according to his parents. He was presented to an ophthalmologist and he was treated with eye drops but the symptoms were not resolved and with the manifestations of malignancy, he was referred to this center. His past medical history was negative for developmental delay, metabolic disorders, or other medical illnessess. No history of medical illness or malignancy was reported in his parents or other family members. On admission, his vital signs were in normal limits. General physical examination revealed a 1.5 cm left eye proptosis with no erythema or tenderness (Figure 1). He had bilateral submandibular lymphadenopathy sized 2*1.5 cm. He had no hepatosplenomegaly and all other examinations were normal. On cranial nerve examination, his pupil was reactive and other cranial nerves examination was in normal limits. His first complete blood cell count showed white blood cells of 28000, hemoglobin of 9.1, and platelet of 512000. The patient PT and PTT were 11.0 and 27.0, respectively.

Spiral brain computerized tomography scan with contrast demonstrated a 28*16*22 mm homogenous mass with mild enhancement in superolateral aspect of left orbit, and left lateral rectus was also mildly swollen. Mild preseptal thickening is noted in left side. Optic nerve, sheet, and bony structures were intact (Figure 2).

With the manifestations of malignancy or solid tumor, he underwent bone marrow aspiration which showed 33% blast cells, all positive for CD10, CD19, and CD79, suggesting pre-B-cell ALL diagnosis. Cerebrospinalfluid examination was negative for malignancy. Standard chemotherapy protocol for ALL was started (11). His proptosis was improved 3 days after treatment initiation and finally disappeared on day 16. He was on maintenance therapy for 12 months. After one-year follow-up, he was free of any symptoms. Written informed consent was obtained from the patient for publication of this case report.



Figure 1. A large left orbital mass with impending vision compromise.



Figure 2. Contrast-enhanced computed tomography on the orbit and the brain. The star depicts a 28*16*22 mm homogenous mass with mild enhancement in superolateral aspect of left orbit. Mild preseptal thickening is noted in left side. Optic nerve, sheet and bony structures are intact.

Discussion

Neoplasms are common etiologies of pediatric proptosis. These neoplasms include retinoblastoma, optic glioma, rhabdomyosarcoma, leukemia, lymphoma, histiocytosis, Ewing sarcoma, and metastasis (12). Leukemic presentation as an orbital mass is usually observed in acute forms ofmyelogenous origin (13). Orbital involvement in patients with ALL is exceedingly rare and can be a manifestation of a primary or relapsed disease. Presentation of proptosis in a patient which is not known for ALL, i.e. primary ALL, will be a diagnostic challenge for physicians.

In this case report, presentation of pre-Bcell ALL as proptosis in a 21-month-old boy is presented. The diagnosis of pre-Bcell ALL in this child was made based on the bone marrow aspiration. In B-lineage ALL, the most important markers for differential diagnosis diagnosis. and subclassification are CD19, CD20, CD22, CD24, and CD79; besides, the presence of CD10 antigen defines the "common" ALL subgroup (14). Bone marrow aspiration of our case showed 33% blast cells, all positive for CD10, CD19, and CD79, suggesting pre-B-cell ALL diagnosis.

Presentation of primary ALL as proptosis was reported to 9 previous case reports (8, 9, 15-21). The age group described in these cases ranged between 6 months and 16 years with no gender dominance. Bilateral involvement of the orbit was observed in only one report (18) and other cases had unilateral involvement similar to the presented case. Leukemic infiltration of orbit possibly originates from marrow cavity of the orbital wall; therefore, simultaneous ocular and orbital infiltration seems unusual. Consistently, only two studies reported simultaneous ocular and orbital involvement (20, 21). In most of these cases, proptosis is associated with other non-specific signs and symptoms of ALL, such as fever or weight loss, which help the physician to think about systemic etiologies. However, in the presented case, initial presentation of proptosis in ALL was not associated with such symptoms, potentially leading to diagnosis such as orbital cellulitis and orbital pseudo-tumor which are common etiologies of proptosis in children (12). Initial presentation of primary ALL as proptosis in a previous healthy child was only reported in two studies (16, 20). In both of these studies, a

progressive proptosis was emerged during two weeks. In the presented case, the patient had a history of proptosis at the same eye about one year ago which made the diagnosis more difficult.

Although presence of orbital involvement in acute myeloid leukemia, known as granulocytic sarcoma, does not significantly alter the survival in these patients (22), the prognostic influence of orbital involvement in ALL is not known. Nevertheless, ocular involvement in ALL is associated with a poor prognosis (23). Among the previous reports who did not have an ocular involvement, all of the cases, except one case (18) survived at the end of follow up; however, length of follow-up was very short in some of these precluding reports. conclusive a interpretation. Anyway, it should be noted that the orbital manifestations of ALL regressed after chemotherapy in all of these cases similar to the presented case, addressing that a right diagnosis in these patients can completely resolve the proptosis.

Conclusion

This patient sensitized us to an unusual presentation of ALL in children. ALL should be kept in mind as a probable diagnosis in every child with acute proptosis even when the child has no past medical history and no associated symptoms in first presntation. Longer follow-up may be promising in determining the effect of orbital involvement on prognosis of the pediatric ALL.

Conflicts of interest

The authors declare no conflicts of interest for this manuscript.

References

1. Greaves MF, Alexander F. An infectious etiology for common acute lymphoblastic leukemia in childhood? Leukemia 1993;7(3):349-360.

2. Wartenberg D, Groves FD, Adelman AS. Acute lymphoblastic leukemia: epidemiology and etiology. Acute Leukemias 2008;77-93.

3. Ries LA, Harkins D, Krapcho M, Mariotto A, Miller BA, Feuer EJ, et al. SEER cancer statistics review, 1975-2003. 2006; 1-10.

4. Schachat AP, Markowitz JA, Guyer DR, Burke PJ, Karp JE, Graham ML. Ophthalmic manifestations of leukemia. Arch Ophthalmol 1989;107(5):697-700.

5. Charif CM, Belmekki M, Hajji Z, Tahiri H, Amrani R, El Bakkali M, et al. [Ophthalmic manifestations of acute leukemia]. J francais d'ophtalmologi 2002; 25(1):62-66.

6. Çavdar AO, Gözdaşoğglu S, Arcasoy A, Demirağ B. Chlorama-like ocular manifestations in Turkish children with acute myelomonocytic leukaemia. Lancet 1971; 297(7701):680-682.

7. Patel AV, Miller JB, Nath R, Shih HA, Yoon MK, Freitag SK. Unilateral eye findings: a rare herald of acute leukemia. Ocul Oncol Pathol 2016;2(3):166-70.

8. Ramamoorthy J, Jain R, Trehan A, Saxena AK, Ahluwalia J. Orbital Mass in a Child With Acute Lymphoblastic Leukemia: A Case Report and Review of the Literature. J Pediatr Hematol Oncol 2016;38(8):646-648

9. Chaudhry SR, Kreis AJ, Underhill HC, Madge SN. Orbital mass secondary to acute lymphoblastic leukaemia in a child: a rare presentation. Orbit 2014;33(6):421-423.

10. Pui MH, Fletcher BD, Langston JW. Granulocytic sarcoma in childhood leukemia: imaging features. Radiology1994;190(3):698-702.

11. Cooper SL, Brown PA. Treatment of pediatric acute lymphoblastic leukemia. Pediatr Clin North Am 2015;62(1):61-73.

12. Rao AA, Naheedy JH, Chen JY-Y, Robbins SL, Ramkumar HL. A clinical update and radiologic review of pediatric orbital and ocular tumors. J Oncol 2013; 2013:1-9.

13. Kincaid MC, Green WR. Ocular and orbital involvement in leukemia. Surv ophthalmol 1983;27(4):211-232.

14. Chiaretti S, Zini G, Bassan R. Diagnosis and subclassification of acute lymphoblastic leukemia. Mediterr J HematolI2014; 6(1):e2014073-e2014080.

15. Rubinfeld RS, Gootenberg JE, Chavis RM, Zimmerman LE. Early onset acute orbital involvement in childhood acute lymphoblastic leukemia. Ophthalmol 1988; 95(1):116-210.

16. Thakker MM, Rubin PA, Chang E. Pre–B-cell Acute Lymphoblastic Leukemia Presenting as an Orbital Mass in an 8-Month-Old. Ophthalmol 2006; 113(2):343-346.

17. Humayun M, Bernstein SL, Gould HB, Chavis RM. Orbital childhood acute lymphoblastic leukemia as the initial presentation. J Pediatr Ophthalmol Strabismus 1992;29(4):252-254.

18. Önder F, Kutluk S, Cosar CB, Kural G. Bilateral orbital involvement as a presenting sign in a child with acute lymphoblastic leukemia. J Pediatr Ophthalmol Strabismus 2000; 37(4):235-237.

19. Yuh WT, Flickinger FW, Willekes CL, Sato Y, Nerad JA. Intraorbital tumor as the initial symptom of childhood acute lympoblastic leukemia magnetic resonance imaging findings. Clin Imaging 1990; 14(2):120-122.

20. Kiratli H, Bilgiç S, Emeç S. Simultaneous conjunctival, uveal, and orbital involvement as the initial sign of acute lymphoblastic leukemia. Jpn J Ophthalmol 2007; 51(2):139-141.

21. McManaway JW, Neely JE. Choroidal and orbital leukemic infiltrate mimicking advanced retinoblastoma. J Pediatr Ophthalmol Strabismus1994; 31(6):394-396.

22. Davis JL, Parke DW, Font RL. Granulocytic sarcoma of the orbit: A clinicopathologic study. Ophthalmol 1985; 92(12):1758-1762.

23. Russo V, Scott I, Querques G, Stella A, Barone A, Noci ND. Orbital and

ocular manifestations of acute childhood leukemia: clinical and statistical analysis of 180 patients. Eur J Ophthalmol 2008; 18(4):619-623