Localized Orbital Non-Hodgkin’s Lymphoma

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Abstract

Localized orbital non-Hodgkin’s lymphoma is a rare form of extranodal lymphoma, which has not been reported much in the literature. The orbit differs from the rest of the body, excluding the central nervous system, as concerns lymph drainage. This may be possible to explain some of the peculiar features in lymphoid orbital lesions. A localized orbital non-Hodgkin’s lymphoma in an elderly man is hereby presented for its entry.

Key words

Non-Hodgkin’s Lymphoma, Orbit, Chemotherapy

Introduction

Non-Hodgkin’s lymphoma’s are a group of neoplasms characterized by proliferation of malignant lymphocytes. Localized orbital non-Hodgkin’s lymphoma is a rare form of extranodal lymphoma. The presentation in these cases usually subtle; there are no overt inflammatory clinical signs such as erythema, chemosis of the conjunctiva, or orbital or ocular pain (1-3). The superior orbit is affected most frequently. Grossly, lymphoid tissue as a creamy, yellow, fish-flesh or reddish appearance, and, because it lacks the fibrous stroma characteristic of inflammatory pseudotumor, it tends to be friable. The extraocular muscles can be involved on very rare occasions (4). When this occurs, the superior rectus-elevator complex is most frequently infiltrated and is enlarged to a much greater extent than what one typically sees in the idiopathic myositis or in the Grave’s disease. Unlike idiopathic inflammatory pseudotumor, which presents as multifocal lesions, lymphoid tumors are almost always unifocal.

Case Report

A 65 year old male was referred from a peripheral health centre with a diagnosis of encysted mucocele. The patient had a 2 year history of swelling near the medial canthus of right eye. The swelling was painless, progressive in size, not associated with erythema or ulceration of the overlying skin (Fig. 1.). There was a history of watering from the right eye of almost same duration, which probably was the reason for the wrong diagnosis of the disease at the peripheral health centre. However, the eye was mildly proptosed and displaced slightly laterally. A FNAC of the swelling revealed a poorly differentiated lymphocytic non-Hodgkin’s lymphoma. Computed Tomography (CT) scan of the orbit and brain was done to reveal the extent of orbital involvement and any brain involvement. However, CT orbit revealed a mass occupying mainly the superonasal aspect of the orbit and also extending down below the inferior orbital margin. The medial rectus muscle was free and there was no bony involvement as well. To confirm the FNAC diagnosis, an excision biopsy of the swelling was done. Surgery revealed a well-defined, firm mass extending both superiorly and inferiorly from the nasal aspect of the orbit. Histopathology of the mass confirmed the FNAC finding of a poorly differentiated
It is important to determine the histological features and clinical stages in order to choose the appropriate treatment methods. Rappaport classification divides non-Hodgkin lymphoma’s into sub-groups according to whether the malignant cell is well differentiated, poorly differentiated or histiocytic. The Working Formulation subdivides non-Hodgkin’s lymphoma’s into low-grade, intermediate-grade and high-grade subgroups, based on their 5 year survival statistics. The low-grade group has 5 year survival rates ranging from 50%-70%; intermediate-grade 35%-45% and high grade 23%-32%.

Traditionally, poorly differentiated low grade lymphomas have not been treated until they produce symptoms. This is because, single-agent chemotherapy or combination of agents do not induce complete remissions and more importantly, they do not change the overall survival of patients. These regimens include the use of single alkylating agents like Cyclophosphamide or Chlorambucil or combinations like CVP (Cyclophosphamide, Vincristine and Prednisolone) or CHOP (CVP plus Adriamycin).

Discussion

The etiology and pathogenesis of non-Hodgkin’s lymphoma remain obscure. It has been suggested that lymphomas arise where there is an immunoregulatory defect in the face of a persistent stimulus to polyclonal lymphocyte stimulation. Supporting this hypothesis is the observation that persons with certain autoimmune diseases, congenital immunodeficiency syndromes and those who have received allografts have an unusually high incidence of non-Hodgkin’s lymphoma (5-7).

CT Scan has been a useful tool in the diagnosis of orbital lymphomas. Lymphoid tumors of the orbit present a distinctive molding pattern on CT; they mold themselves to the globe and orbital bones and rarely cause bone destruction (8).

References