Introduction
Non-Hodgkin's lymphomas are neoplastic diseases of the lymphatic system, usually involving the lymph nodes, but almost 25-40% of the cases have extranodal onset [1]. Orbital lymphomas represent about 2% of all lymphomas, 5-15% of extranodal lymphomas and approximately 50% of all primary malignant tumors of the orbit [2]. An association between Chlamydia psittaci infection and orbital adnexal lymphoma has been described, and also, thyroid eye disease is considered to be a predisposing factor [3].

Orbital lymphoma can arise from the eyelid, orbit, lacrimal glands or conjunctiva. Clinical signs and symptoms are nonspecific, often delaying the diagnosis. In 25% of the cases the conjunctiva is involved and the patients present with salmon red patches or swollen conjunctiva. In the rest of the patients, presentation is with orbital mass [2]. Common signs at presentation are: palpable mass, pain, exophthalmia, ptosis, dysxia, decreased vision and abnormal ocular movement [4,5]. In approximately 10-17% of cases the lymphoma appears bilaterally, simultaneously in both orbits in 80% of cases and subsequently in 20% of cases. Between 20-40% of the patients have extraorbital lymphoma at onset, especially those with aggressive histology [6,7].

The majority of orbital lymphomas are of low-grade histology (84%), the minority (16%) being aggressive lymphomas [8]. The commonest sub-type of orbital lymphoma is extranodal marginal-zone lymphoma. Other histological types are: follicular, lymphocytic, mantle cell and diffuse large B-cell lymphomas [9,10].

Diagnosis is based on biopsy and staging procedures should include imaging investigations: computed tomography (CT), or magnetic resonance imaging (MRI) to evaluate local extension but also systemic lymphomatous involvement. Differential diagnosis should include other malignant tumors and inflammatory pseudo-tumors of the orbit and thyroid associated orbit disease [1].

The prognosis of ocular lymphoma depends on the histological type, age, localization and stage of disease. Favorable prognostic factors are: low-grade histology, younger age, conjunctival localization and early stage at presentation [11].

Treatment depends on the stage: in localized orbital lymphoma, radiotherapy is highly effective, while in patients with high grade histology or disseminated lymphoma, systemic chemotherapy should be used. In some localized cases, antibiotic therapy against Chlamydia results in complete remission [12,13].

Case report
A case of bilateral orbital lymphoma is presented below, having the approval of the Ethical Committee of the Hospital and the written consent of the patient for publishing his pictures.

A 50 years old male patient referred first to the Ophthalmology Clinic in april 2012 for bilateral exophthalmos and palpebral ptosis. Physical examination revealed bilateral orbital tumors. Biopsy was refused by the patient. In November 2012, he was admitted at the Maxillofacial
Surgery Clinic for bleeding from the periorbital tumors. A biopsy was performed and diffuse large B-cell lymphoma was diagnosed. On admission to the Hematology Clinic the patient presented bilateral periorbital tumors (12 cm in the right and 8 cm in the left side), involving the eyelids and incorporating the eye globes (fig. 1 and 2). The patient also complained of complete loss of vision and constitutional symptoms (night sweats and weight loss).

Laboratory tests revealed leucocytosis (13000/microliter) with 55% atypical lymphocytes in the peripheral blood smear, anemia (hemoglobin=10g/dl), elevated erythrocyte sedimentation rate and decreased level of IgG and IgA. Serological tests for Chlamydia, HIV and hepatitis viruses (B and C) were negative. Bone marrow biopsy showed lymphomatous involvement. Cranio-orbital CT revealed bilateral orbital tumors (14/8/11 cm in the right with 2.7 cm ptosis of the right eye and 7.8/5.6/6.5 cm in the left with 2 cm ptosis of the left eye), without invasion of the eye globes, bones, paranasal sinuses or of the cerebral parenchyma (fig. 3 and 4). Cervical and thoraco-abdominal CT showed enlarged latero-cervical and retroperitoneal lymph nodes.

The patient was diagnosed with stage IVB diffuse large B-cell primary orbital lymphoma and R-CHOP (Rituximab + Cyclophosphamide, Adriblastin, Vincristin and Prednison) chemotherapy was started, which produced a rapid response. After 2 months, the patient presented with relapse. R-ICE (Rituximab + Iphosphamide, Carboplatin and Etoposide) second-line chemotherapy was started and complete remission was obtained after 4 cycles (fig. 5). However, the patient did not regain his vision due to optic nerve atrophy. He did not return for completion of the chemotherapy cycles, nor for follow-up.

Figure 1. Massive right orbital tumor

Figure 2. Left orbital tumor with spontaneous bleeding

Figures 3 and 4. Cranio-orbital CT images before therapy

Figure 5. Resorption of the orbital tumors after chemotherapy
**Discussion**

Primary orbital lymphoma is a rare disease, diagnosis being difficult due to nonspecific symptoms at presentation. Patients usually present with unilateral orbital tumors, only 10-17% of the cases having bilateral orbital involvement at onset. Ocular lymphomas have low-grade histology in the majority of the cases and prognosis is good if diagnosis is made in early stages. However, permanent loss of vision can be seen due to optical nerve atrophy, even in patients with a complete remission of the lymphoma.

Treatment consists of radiotherapy in localized disease but if systemic involvement is present, chemotherapy should be used. The particularity of this case is the presence of massive orbital tumors, due to delayed diagnosis. Long-term prognosis of this patient is unfavorable, due to the advanced stage of the disease at presentation and early relapse after an initial good response to chemotherapy.

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