Sight-threatening optic neuropathy is associated with paranasal lymphoma

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Abstract: Malignant lymphoma around the orbit is very rare. We present a rare case of optic neuropathy caused by lymphoma. A 61-year-old Japanese woman was referred to our hospital for evaluation of idiopathic optic neuropathy affecting her right eye. The patient was treated with steroid pulse therapy (methylprednisolone 1 g daily for 3 days) with a presumed diagnosis of idiopathic optic neuritis. After she had been switched to oral steroid therapy, endoscopic sinus surgery had been performed, which revealed diffuse large B cell lymphoma of the ethmoidal sinus. Although R-CHOP therapy was immediately started, prolonged optic nerve compression resulted in irreversible blindness. Accordingly, patients with suspected idiopathic optic neuritis should be carefully assessed when they show a poor response, and imaging of the orbits and brain should always be done for initial diagnosis because they may have compression by a tumor.

Keywords: optic neuropathy, malignant lymphoma, paranasal lymphoma, rhinogenic optic neuropathy

Introduction
Paranasal mass lesions can potentially cause visual loss by compressing the optic nerve, but only a few such cases have been reported in the literature.1–3

Here, we present a rare case of optic neuropathy caused by paranasal lymphoma.

This tumor generally has a poor prognosis, and early diagnosis is essential for effective treatment. Imaging modalities such as computed tomography (CT) and magnetic resonance imaging (MRI) are important for making a correct differential diagnosis.

Case report
A 61-year-old Japanese woman was referred to the Department of Ophthalmology at Yokosuka Kyosai Hospital for evaluation of optic neuropathy affecting her right eye. Her past history included surgery on the paranasal sinuses 50 years earlier, and uterine myomectomy 20 years earlier. There was no relevant family history.

The best corrected visual acuity was counting fingers on the right and 20/30 on the left.

Optic disk and retinal findings were normal. A left relative afferent pupillary defect was present, but the findings on neuro-ophthalmological examination were otherwise unremarkable. There were no abnormal findings on slit-lamp examination, or on general physical examination. The critical flicker frequency was undetectable in the right eye and was 40 Hz in the left eye. Goldman perimetry showed centrocecal scotoma, indicating an abnormality of the optic nerve. However, fluorescein angiography did
not reveal any abnormal optic disc findings, such as disc hyperfluorescence or a wedge-shaped filling defect. CT and MRI of the brain and orbits demonstrated a mass occupying much of the ethmoidal sinus (Figure 1A, B). The mass extended into the extraconal right orbit. Since the mass did not seem to be causing compression, the patient was treated with steroid pulse therapy (methyl-prednisolone 1 g daily for 3 days). This was followed by oral prednisolone, which was started at a dose of 30 mg/day and then tapered. After she was switched to oral steroid therapy, endoscopic sinus surgery was performed under a diagnosis of sinus mucocele by an otolaryngologist. In addition, sinus biopsy was performed.

Despite this treatment, marked improvement of her visual acuity on the right was not obtained and the acuity was 20/2000.

On the day after sinus surgery, the patient developed tender proptosis. The best corrected visual acuity was counting fingers on the right, while ptosis, dilation of the pupil, and loss of eye movements indicated oculomotor nerve palsy. It was impossible to use the Hess chart, because her vision was too poor. MRI showed expansion of the ethmoidal tumor with compression of the right optic nerve and right oculomotor nerve (Figure 1C). On the same day, biopsy revealed diffuse large B cell lymphoma of the ethmoidal sinus (Figure 2A–D). Staging investigations did not show any evidence of disseminated disease, and stage 1E extranodal non-Hodgkin's lymphoma (NHL) was diagnosed. The patient was treated with R-CHOP therapy (rituximab, cyclophosphamide, doxorubicin, vincristine, and prednisolone), followed by loco-regional irradiation at the Hematology Department. The tumor initially responded to treatment and decreased in size. Despite achieving remission, the patient became blind.

Discussion
Malignant lymphoma around the orbit is very rare, but we should remember that it can threaten vision by compressing the eyeball and/or optic nerve. Malignant lymphoma rarely arises in the paranasal sinuses. For NHL of the head and neck region, less than 5% of tumors arise at extranodal sites, while the majority develop in the lymphoid tissue of Waldeyer's ring. Correct diagnosis of paranasal lymphoma is usually delayed. Because tumors at this site cause few early symptoms, the diagnosis is usually made at an advanced stage. Detailed investigation is often postponed at the initial stage and is delayed until the tumor causes clear symptoms. When such a tumor is still small, CT scanning often shows clear paranasal sinuses and leads to a false sense of security.
However, sometimes obstruction of a sinus will create a mucocele or pyocele, which is detected by surgery. Orbital invasion is a common manifestation of malignant sinus tumors, and ocular findings can be useful to assess the expansion and status of a primary sinus lesion when clinical evaluation is difficult. The common presenting signs and symptoms are proptosis, visual loss, lid edema, diplopia, limitation of ocular motility, conjunctival chemosis, blepharoptosis, conjunctival injection, globe displacement, epiphora, anisocoria, eye pain, and photophobia. From 50% to 70% of the patients with orbital and ocular manifestations show orbital involvement at the time of initial radiological evaluation. The abnormalities detected on X-ray films of the sinuses range from a soft tissue mass to orbital bone destruction. The most common sinus to be involved by malignancy is the maxillary sinus and squamous cell carcinoma is the most frequent malignant tumor.

The available reviews include only a few cases of sinonasal lymphoma. The majority of sinonasal lymphomas are classified as clinical stage 1E by the Ann Arbor system, and as large B cell lymphoma by the REAL classification.

Some studies have suggested that radiation alone provides good regional control at an early stage, while additional chemotherapy can be reserved for more extensive disease. However, other studies have shown that combined modality treatment with chemotherapy and loco-regional irradiation improves both disease-free and overall survival. The overall mortality rate is 55%. For effective treatment of these life-threatening malignant tumors, early detection by investigation of orbital symptoms may be important.

Our initial diagnosis was idiopathic optic neuritis because we did not consider optic nerve compression based on the imaging findings. However, repeat imaging after the diagnosis had been made revealed evidence of compression, suggesting that the patient already had optic nerve compression which the initial imaging study failed to detect. We started R-CHOP therapy immediately after we identified malignant lymphoma.
but prolonged optic nerve compression resulted in irreversible blindness. Despite this, we do not think that starting therapy earlier would have led to a good outcome because vision was already reduced to counting fingers (indicating irreversible damage) at her initial presentation. Accordingly, patients with suspected idiopathic optic neuritis should be carefully assessed when they show a poor response, and imaging of the orbits and brain should always be performed for initial diagnosis because they may have compression by a tumor. This case serves to remind ophthalmologists about the pitfalls of diagnosing and treating optic neuritis. If ophthalmologists investigate patients who have headache or a history of chronic sinusitis with bone destruction on X-ray films, they should strongly suspect a malignant tumor of the sinuses.

Disclosures
The authors disclose no conflicts of interest.

References