In this case report we describe a 27-year-old man who presented with progressive enophthalmos for 5 months without any other associated ocular symptoms such as pain, diplopia, or visual disturbance. Computed tomography showed that his progressive enophthalmos originated from a frontoethmoidal mucocele and this caused destruction of the lamina papyracea and shrinkage of the ethmoidal air cell. Finally the enlarged orbital space caused an inward deviation of the eyeball. Endoscopic marsupialization was successfully performed by an otolaryngologist and did not result in any ophthalmologic sequelae. Although frontoethmoidal sinus mucoceles mostly frequently originates from orbital mucoceles, enophthalmic manifestations are very rare. Enophthalmic conditions are not as responsive to surgical interventions as exophthalmic conditions.

Key Words: Enophthalmos, Sinus mucocele
T1 high signal intensity, which may have been due to protein content and a chronic state of sinusitis, suggesting that it was most likely a frontoethmoidal mucocele (Fig. 1). We concluded that this lesion was a chronic mucocele and we referred the patient for otolaryngologic consultation.

The otolaryngologist performed endoscopic sinus drainage surgery during which a bulging mucocele that originated from the right frontal sinus and extended toward the anterior ethmoidal sinus; this mucocele was filled with a white to yellowish mucous discharge. The lamina papyracea had a nearly total bony defect, but the orbital muscle was not exposed. All of the mucous material was removed by curved suction and part of the middle turbinate was resected using straight cutting and the mucosal cavity was enlarged by marsupialization.

Eight months after surgery, the lesion was completely resolved, but the patient still complained of an enophthalmic condition (Fig. 2). At that time, the Hertel examination with the same base (96 mm) showed 11.0 mm in his right eye and 14.0 mm in his left eye, including that a 3.0 mm enophthalmic condition remained. We discussed the need for orbital reconstruction with the otolaryngologist, but we collectively decided not to perform additional ocular reconstruction surgery because we were concerned that the ocular reconstructive material might result in a recurrent mucocele [6,7]. Institutional review board approval was obtained in Seoul St. Mary’s Hospital.

**Discussion**

Frontoethmoidal mucoceles are cystic, slowly expanding lesions that usually originate from the frontal or ethmoidal sinuses. They may also involve the orbital space and cause a downward and lateral globe displacement with proptosis [5]. These mucoceles may be accompanied by ophthalmic manifestations such as displacement of the eyeball, proptosis, restricted ocular movement, diplopia, lid swelling, chemosis, optic neuropathy, decreased vision and increased
intraocular pressure [8-10]. More rarely, a frontoethmoidal mucocele may be accompanied by “dynamic proptosis” which presented spontaneous and dynamic proptosis in both eyes, pulsatile in nature [11]. To our knowledge, however, this is the first case of frontoethmoidal mucocele presenting with an enophthalmic condition that gradually worsened over 5 months, which was not recovered from a successful endoscopic surgery.

Following endoscopic or external surgical removal of most of the frontoethmoidal mucoceles that has orbital involvement, most patients show a good recovery and improvements of their ophthalmic symptoms such as proptosis, limitations in eye-movement, and increased intraocular pressure [6,8-10]. Patients presenting with an enophthalmic condition, however, do not show symptom improvement after successful sinus surgery [5,9,10]. Ophthalmic enophthalmic conditions are usually caused by orbital blow out fractures, which result in enlargement of the orbital space [1]. In our patient, the bulging frontal sinus mucocele grew toward the ethmoidal sinus, destroying the lamina papyracea, and the ethmoidal air cells became hypoventilated. The frontal sinus mucocele finally resulted in ethmoidal atelectasis. During this time, the enlarging the orbital space and the deviated medial wall bony component made the eyeball deviate inward, similar to the effect of a right orbital medial wall fracture. These ethmoidal bony defects and the shrinkage of ethmoidal air cells were confirmed by the otolaryngologist who performed the functional endoscopic surgery.

Enophthalmos of a sinus origin is associated with silent sinus syndrome and this is an uncommon condition. It usually presents with progressive enophthalmos and hypoglobus due to the gradual collapse of the orbital floor and with opacification of the maxillary sinus that is then followed by subclinical maxillary sinusitis [2]. Patients with ethmoid silent sinus syndrome have recently been described in other studies; in these patients, enophthalmos resulted from the negative pressure in the anterior ethmoidal air cells on the defective side, and this increased orbital volume and markedly reduced the sinus volume [3,4]. The cosmetic problems of patients with enophthalmos may be corrected by orbital reconstructive surgery. In our patient, this would have consisted of reduction of the intraorbital contents and insertion of alloplastic material into the right medial wall, which had been almost totally destroyed by the melting effect of the sinus mucocele. But the orbital reconstructive material can cause recurrent sinus mucoceles by blocking the outflow of sinus mucous material [6,7]. Thus, after consultation with the patient’s otolaryngologist, we concluded that an additional operation should await improvements of the symptoms.

Most patients with ocular symptoms associated with sinus mucocele show good recovery after correction of the sinus lesion [9,12]. We encountered a patient with enophthalmos associated with a frontoethmoidal mucocele. However, the patient’s symptoms were not recovered after successful surgical removal of the mucocele.

Conflict of Interest

No potential conflict of interest relevant to this article was reported.

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