Ethmoid mucocele in a 5-year-old patient without cystic fibrosis

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Abstract
Paranasal sinus mucoceles are extremely rare in children. We report a case of a 5-year-old boy with no evidence of cystic fibrosis who underwent endoscopic marsupialisation of an ethmoid mucocele.

Case report
A previously well 5-year-old boy presented to our department with a 3-week history of a swollen, teary left eye. He did not have any systemic complaints, and no specific ear, nose or throat symptoms.

On examination he was afebrile and appeared systemically well. Examination of his eyes revealed left-sided proptosis, but his visual acuity and eye movements were normal.

His oral cavity and oropharynx appeared normal and anterior rhinoscopy revealed no abnormality. Endoscopic nasal examination revealed a smooth-surfaced mass protruding from the upper part of the middle meatus on the left side. The rest of the examination was non-contributory.

Laboratory investigations including a full blood count and electrolytes were normal and he had a normal chest X-ray. A sweat test performed was negative for cystic fibrosis.

Computed tomography (CT) (Figs. 1 - 3) of his orbits and sinuses was performed and revealed a mixed-density soft-tissue expansile mass arising in the left ethmoid sinus, causing thinning and bowing of the medial wall of the left orbit. There was minimal rim enhancement of the mass. The mass caused minimal lateral displacement of the left optic nerve and mild left proptosis, but resulted in obstruction of the left maxillary antrum.

The patient underwent endoscopic marsupialisation of the mucocele. Biopsies of the cyst wall showed strips of mucoperiosteum with fragments of reactive bone. The mucosa was lined by respiratory epithelium with subepithelial chronic inflammatory cell infiltrate. There was no evidence of malignancy.

A repeat CT scan was done 4 weeks post surgery and showed no residual mass. At 6-month follow-up, endoscopic nasal examination showed no recurrence.
CASE REPORT

Discussion

A mucocele is an epithelial-lined mucus-containing sac that fills a paranasal sinus and is capable of expansion by bone resorption and new bone formation. They are extremely rare in children and their presence warrants thorough investigation to rule out aetiological entities such as cystic fibrosis. Paediatric mucoceles were previously thought to occur predominantly in patients with cystic fibrosis but some authors have recently reported mucoceles occurring in patients who do not have cystic fibrosis.

Ethmoid sinus mucoceles often appear with painless orbital swellings. They may also produce frontal headaches and proptosis. Differential diagnoses of ethmoid sinus mucoceles in the paediatric population include meningocoele, rhabdomyosarcoma, haemangiomata, neuroblastoma and bony lesions such as fibrous dysplasia and osteomas.

Mucoceles of the frontal sinus can often be diagnosed on plain X-ray films. Because of overlapping bone structures, mucoceles of the maxillary, sphenoid and ethmoid sinuses may not be apparent on X-ray. For this reason CT scan is the primary imaging method of choice.

On CT, a mucocele appears as a non-enhancing, low-density expansile mass filling the sinus, with remodelling and/or thinning of the bony sinus walls. Mucoceles have variable densities on CT depending on protein content, inspissation and superinfection. Following intravenous contrast administration the lining of the mucocele may be enhanced. Magnetic resonance imaging (MRI) is only indicated if uncertainty persists following a CT scan. MRI can provide information about tissue composition, therefore enabling differentiation between different lesions which may be of similar density on CT.

Mucoceles are managed surgically and most authors advocate marsupialisation rather than complete excision. Endoscopic management of paediatric ethmoid mucoceles is recommended as this enables exceptional visualisation of the surrounding anatomy and identification of vital structures which allows for a safe and minimally invasive treatment option. The endoscopic approach avoids an incision and subsequent facial scarring.

In conclusion, ethmoid mucoceles in the paediatric population are rare but can occur in the absence of cystic fibrosis. CT scan is the primary imaging method of choice and endoscopic marsupialisation is recommended.