Atypical clinical and radiological presentation of Juvenile Nasopharyngeal Angiofibroma

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ABSTRACT

Juvenile nasopharyngeal angiofibroma (JNA) constitutes less than 0.5% of all head and neck neoplasm. It is a histopathologically benign, yet locally aggressive, vascular tumor that occurs most frequently in males from 5 to 25 years. But it could mimic the clinical course of antrochoanal polyp. The clinical and radiological evaluated with computer tomography and with endoscopic resection of the tumor (an antrochoanal polyp), a histopathological report was angiofibroma.

Background:

Nasopharynx angiofibroma is a combined vascular and fibrous neoplasm arising in the nasopharynx of prepubertal and adolescent males. The tumor exhibits a strong tendency to bleed and, despite being microscopically benign, frequently exhibits destructive and aggressive behaviour.

Histopathology

Histologically, JNA exhibits cells of myofibroblast origin surrounded by a fibrous pseudocapsule. There are multiple vascular channels dispersed within the neoplasm composed of abundant endothelial cells embedded in a collagenous tissue network. An important hallmark is the lack of a true muscular layer, and this absence precludes any form of vasoconstriction and is felt to contribute to the tumor’s high propensity for hemorrhage.
Diagnosis

Key features to support a diagnosis of JNA include the presence of a vascular mass with an epicenter at the posterior nasal cavity near the medial pterygopalatine fossa, the presence of bony modelling—but not destruction—with tumor growth, and the lack of regional or distant metastasis. For atypical extension or unexpected rapid growth, a biopsy should be considered to assess for other neoplasms aside from JNA. The computed tomography (CT) and magnetic resonance imaging (MRI) are the primary diagnostic tests. JNA is commonly associated with known as the Holman-Miller sign. Widening of the sphenopalatine foramen may also be observed on CT and MR scanning offers improved soft tissue delineation. The tumour may display a “salt and pepper” appearance.

Angiography is of diagnostic value with its characteristic tumour blush. It provides information about the major vascular supply and allows for pre-operative embolisation. The major arterial supply to these tumours is typically the ipsilateral internal maxillary artery, with occasional additional vessels from ascending pharyngeal artery, and branches from cavernous ICA or contralateral external carotid system.

Clinical presentation

Clinical presentation is usually with nasal obstruction or recurrent severe epistaxis in an adolescent male. Clinical features vary depending upon the extent of disease, but extensions into the sphenoid and other sinuses, orbit, infratemporal fossa and middle cranial fossa are often silent. Alternately, such extensions may manifest with cheek swelling, proptosis or visual loss.

Treatment

The primary treatment modality is surgical excision and it remains a viable option in cases of intracranial extension. However, the surgical plan needs to consider that a more extensive resection in an attempt to prevent recurrence is inherently associated with a higher degree of morbidity. Areas of most concern include the pterygoid fossa, clivus, basisphenoid, sphenoid diploe, cavernous sinus, and intracranial vault.

Case Report:

A 15-year-old boy was referred to our department from another hospital with difficulty in breathing and swallowing for 6 months. Over time, his complaint had increased. He had been complaining of nasal obstruction for a long time. There was no nasal discharge or allergic tendencies with only one episode of mild epistaxis occurred 2 months ago. The child was in good physical health with hypo-nasal speech. On examination of the oral cavity, there was a large, smooth surfaced, reddish mass hanging behind the soft palate into the hypopharynx. Anterior rhinoscopy was showing smooth surface whitish polyp, on rigid endoscopic examination of the nasal cavities and the nasopharynx, the same polyp.

On examination of the neck, there was no palpable lymph node. Computerized tomography (CT) of the paranasal sinuses demonstrated free sinuses with tumor extends from the right pneumatized pterygoid air cell into nasopharynx and continuing into the oral cavity (Fig. 1 A&B).
Tumor extends from the right pneumatised pterygoid air cell into nasopharynx and continuing into the nasal cavity on the right side (Fig. 2 A&B). The patient was operated under general anesthesia, in the supine position with his head slightly elevated, a pedicle of the polyp on the lateral nasal wall passing through the right choana into the nasopharynx, the pedicle of the polyp at level below the middle turbinate. The tumour removed completely endoscopically (Fig. 3).

A nasal packing merocel® was used. The histopathological examination showed features of a nasopharyngeal angiofibroma with highly fibrous elements. The outcome of surgery was evaluated clinically, endoscopic and by CT scans.

Discussion:
Angiofibromas are histopathologically benign but potentially locally destructive vascular tumors. They are un-encapsulated neoplasms composed of a rich vascular network within a fibrous stroma. Angiofibromas originate predominantly in the posterior-lateral wall of the nasopharynx. Angiofibromas arise typically in the nasopharynx, specifically at the trifurcation of the sphenoidal process of the palatine bone, the horizontal ala of the vomer, and the roof of the pterygoid process.

These vascular tumors expand commonly beyond the nasopharynx into the cranium, nose, and paranasal sinuses. Classically, JAF are confined to boys in adolescence and early adulthood. JNA accounts for less than 0.5% of all benign lesions that originate in the nasopharynx. JNA is an uncommon tumor, with reported incidence between 1 in 5000 and 1 in 60,000 otolaryngology patients, and

In young adults, differential diagnosis of nasopharyngeal masses includes juvenile nasopharyngeal angiofibroma, meningoencephalocele, nasal glioma, hemangioma, grossly enlarged adenoids, and nasopharyngeal malignancies.
Antrochoanal polyp (ACP) is a maxillary sinus polyp that originates in the maxillary sinus, passes through a sinus ostium and extends into the choana. ACPs represent approximately 4-6% of all nasal polyps in the general population. The more common manifestation of Antrochoanal polyp is unilateral nasal obstruction, but it may sometimes be bilateral, depending upon the blockage of the nasopharynx.

The more common manifestation of ACP is unilateral nasal obstruction (especially during the expiratory phase), but may sometimes be (20-25% of cases) bilateral, depending upon the blockage of the nasopharynx. Other clinical manifestations are rhinorrhoea, bleeding, snoring, foreign body sensation, halitosis, headache, post nasal drip and loss of sense of smell. There are occasional reports of cases starting with epistaxis, polyp strangulation, spontaneous amputation, dyspnoea and dysphagia, with extension to the mouth producing dysphagia, speech disorders and obstructive sleep apnoea.

In our case, the presentation was similar to that of antrochoanal polyp.

In a study of 72 patients, Lloyd et al. reported, three findings on CT and MR imaging that should suggest a diagnosis of JNA: (1) a soft tissue mass in the nasopharynx or nasal cavity, (2) a mass in the pterygopalatine fossa, (3) erosion of posterior osseous margin of the sphenopalatine foramen, Holman-Miller sign or Widening of the sphenopalatine foramen.

Conclusion:

A unilateral nasal polyp in childhood should not be taken lightly as an antrochoanal polyp, especially there is a juvenile nasopharyngeal angiofibroma mimicking an antrochoanal polyp.
References:


