Introduction

Teratomas are rare congenital tumors composed of tissues derived from the embryonic germ cell layers.\(^1\) They may occur in the gonads and extragonadal locations. The incidence of teratoma is 1:4000 live births with the most common site being the sacrococcygeal area.\(^2\) Head and neck teratomas are very rare accounting for <5% of all teratomas with nasopharyngeal ones being extremely rare.\(^2,3\) Teratoma of the nasopharynx may arise from the skull base or the posterior pharyngeal wall. Nasopharyngeal teratomas can cause tracheoesophageal obstruction,\(^2\) and we present a case report of a neonate whose nasopharyngeal teratoma presented with airway obstruction.

Case Report

A term male neonate delivered by spontaneous vaginal delivery weighing 3.5 kg following an uneventful pregnancy and was admitted shortly after birth to the nursery with grunting respiration and cyanosis (saturation of 70% in room air), Apgar scores were 5\(^1\) then 8\(^2\). His mother attended antenatal clinic; she had normal booking blood and her antenatal ultrasound scans (weeks 28 and 36) were unremarkable. Clinical examination was unremarkable except for weak cry and slight resistance during the insertion of a nasogastric tube through the left nostril. A presumptive diagnosis of choanal stenosis and possible recurrent laryngeal nerve palsy were made.

The infant was placed on the continuous positive airway pressure (CPAP) in which he was comfortable on FiO\(_2\) of 21%. Chest and postnasal space X-ray were obtained and were normal. An ear, nose, and throat surgeon were consulted, and a planned examination under anesthesia (EUA) was organized. The initial EUA revealed nothing; infant continued to need CPAP.

Abstract

Teratoma in the nasopharynx is one of the rare causes for nasal symptoms in the newborn. The present report was on a term “neonate” who had respiratory distress from a hidden teratoma. Cheaper and readily available investigation modalities including postnasal space X-ray and examination under anesthesia revealed nothing. A postnasal space computed tomography also revealed nothing. Magnetic resonance imaging revealed a mass which was resected endoscopically and histology revealed an immature teratoma. Herein, we present the case, management challenges and literature review to emphasis that negative investigations with persistent nasal symptoms should prompt further evaluation and teratoma should be considered when encountering newborn with nasal symptoms without obvious cause.

Keywords: Airway, Immature, Teratoma

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Head and neck computed tomography (CT) was ordered which revealed nothing [Figure 1]. Magnetic resonance imaging (MRI) which was not readily available was ordered, and it revealed a large heterogeneous enhancing lobulated mass in the left naso- and oral-pharyngeal cavity measuring 1.7 cm × 2.1 cm × 2.4 cm [Figure 2].

Excision of the mass was performed endoscopically [Figures 3 and 4] and the histology revealed immature teratoma. The infant was followed up for recurrence with a repeat EUA conducted at 3 months of age, and the site looked clear. Alfa-fetoprotein level was also done at 3 months and was 47.8 IU/ml which was normal for age.

**Discussion**

Teratoma and neuroblastoma are the most common tumors in the fetus and the newborn with the prevalence of congenital teratoma estimated at 1:4000 live births.[2] The most common site for teratomas is the sacral region[3] while nasopharyngeal teratomas are very rare.[3] Many theories have attempted to explain histogenesis of teratomas with the most popular theory suggesting displaced totipotent cells during migration from the York sack to the gonadal ridge.[1] Nasopharyngeal teratomas tend to arise from the skull base or lateral nasopharyngeal walls. Our patient’s teratoma was arising from the skull base.

Four basic histologic classification of teratomas are recognized i.e., dermoid cysts (epithelium-lined with skin elements, composed of ectodermal, and mesodermal layer), teratoid cyst (composed of all the three germ elements but poorly differentiated), true teratoma (three germ elements differentiated into specific tissue or organs), and epignathi (tumor with developmental fetal organs and limbs).

Teratomas may be identified in utero if they are very large or if it causes polyhydramnios by impeding the fetus from swallowing liquor. The mother to our patient had an uneventful pregnancy with normal obstetric ultrasound scans on two occasions. At birth, the tumor may cause respiratory embarrassment as it did in our patient in which case, the main-stay of care is prompt assessment and management of airway. Our patient was managed on CPAP to keep the airway patency.

![Figure 1: Computed tomography scan at the level of nasopharynx](image1)

![Figure 2: Magnetic resonance imaging showing mass in the left nasal space](image2)

![Figure 3: Mass seen in the nasal pharynx](image3)

![Figure 4: Nasopharynx after excision of mass](image4)
Nasopharyngeal teratomas tend to be associated with other abnormalities, and reported ones include cleft palate, cardiac abnormalities and microcephaly. It is speculated that these abnormal tissues inhibit fusion of normal tissues. The case we present had no associated abnormalities. Although nasopharyngeal teratomas may protrude through the oral or nasal cavity and associated morbidity and mortality are related to the location and size of the tumor; from our case, we learn that lack of an obvious protruding mass does not preclude the presence of teratoma.

The differential diagnosis for nasopharyngeal teratoma includes meningoencephalocele, encephalocele, glioma, neurofibroma, rhabdomyosarcoma, or hemangioma.

Imaging methods including CT scan and MRI are important; CT scan will characterize bone changes and detect intralesional calcification but the investigation modality of choice for teratoma would be an MRI which demonstrates the soft tissue characteristic of the tumor, and it aids in determining the relation of the mass to other vital structures such as blood vessels. In our case, the X-ray and CT did not identify the mass, but it was on MRI. Immature teratomas such as, we found in our patient, do not have calcified structures and may be missed on X-ray and CT scan. Where resources are constrained, this can become a diagnostic dilemma. It is important that extension of the mass into the brain is ruled out before surgery and in our case, there was no intracranial extension seen on MRI.

The majority of congenital teratomas are benign although malignant ones have been reported. Teratomas that are histologically immature and those containing York sack tumor have a higher tendency for malignant transformation. The treatment of choice is complete excision coupled with close follow-up with tumor marker assay (alpha fetal protein) and imaging. Recurrence has been reported in some cases including those with incomplete resection, tumor rupture, nongonadal teratoma, especially sacral, immaturity, and raised alpha fetal protein level. Although at 3 months, our patient did not have makers of recurrence, close follow-up is necessary.

Conclusion

- A differential diagnosis of nasopharyngeal teratoma should be considered in any newborn with persistent nasal symptoms
- Immature teratomas can be challenging to identify where resources are limited, but persistent symptoms with negative investigations should prompt further evaluation. MRI is the investigation modality of choice in identifying teratomas.

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Conflicts of interest

There are no conflicts of interest.

References