Introduction

Cervical teratomas are rare tumors comprising only 3%-5% of all pediatric teratomas, with the incidence of 1 in 20,000–1 in 40,000 live births. They are usually diagnosed on antenatal scan as neck mass or postnatally as cervical mass with or without airway compromise. Although mostly benign in nature, large lesions can cause perinatal mortality because of upper airway obstruction and so require an extensive perinatal planning for upper airway management.

The diagnosis of teratoma can be challenging, especially if it is a purely cystic teratoma as then it becomes difficult to differentiate it from lymphatic malformation.

We present a rare case of cervical cystic immature teratoma in a newborn resembling lymphangioma.

Case Report

This was a 1-day-old, 2.7 kg full-term female child, born with cesarean delivery. The child was referred to us with complaints of large midline cervical mass with stridor. The antenatal history was uneventful. The child had mild respiratory distress and was accepting feeds well. On examination, there was a large neck mass of size about 10 cm × 12 cm, firm in consistency, and noncompressible involving almost whole of the neck [Figure 1].

The clinical findings were suggestive of tumor. However, on ultrasound and MR angiogram [Figure 2], it was a well-defined lesion with multiloculated cystic areas without calcification. There was some internal hemorrhage, and it was a non-enhancing lesion on the arterial and venous phase. The mass was indenting the trachea. A very likely possibility of lymphatic malformation was given. Serum alphafetoprotein (AFP) levels were 35,000 ng/ml.

With a high suspicion of cystic teratoma in view of a firm, well defined, noncompressible, and noninfiltrative lesion, the baby was planned for the surgery. Intraoperatively, it was large thick-walled lobulated mass involving bilateral submandibular triangles and submental triangle [Figure 3]. It was adherent densely posteriorly with the thyroid gland and pretracheal fascia. The mass was completely excised. The postoperative period was uneventful. The child was shifted on ventilator and was extubated after 48 h.

On gross examination, the specimen was of size 10 cm × 8 cm with multiple cystic areas and mucoid contents. Microscopic examination showed neuroglial tissue, cartilage, squamoid epithelium, focal calcification, and small foci of primitive neuroepithelial tissue suggestive of immature teratoma Grade II.

After follow-up of 1 year, child is fine with no recurrence and normal alpha fetoprotein and normal thyroid profile.
Large cervical teratomas may extend into the mediastinum or compress trachea, causing pulmonary hypoplasia and many of these may require an extensive perinatal multidisciplinary planning for EXIT (exutero intrapartum treatment) procedure. If left untreated, cervical teratomas have an estimated mortality rate of 80%–100%.

The diagnosis of the cystic teratoma can be confusing, especially if it is predominantly cystic lesion as then it may resemble lymphatic malformation. Both are located in the anterior and posterior triangle of the neck which further add to the diagnostic challenge. The risk of airway compromise with teratomas is more as they are more firm and less compressible than lymphangioma. When the diagnosis is in question, a high-resolution magnetic resonance imaging (MRI) or computed tomography scan can help in confirming the diagnosis; however, in some cases, it can be challenging. MRI is preferred modality as it is radiation free and provides better soft tissue detail.

On ultrasound, cervical lymphangiomas appear as infiltrating cystic lesion with fine internal septations. While cervical teratomas are usually thick walled, well-defined noninfiltrating lesions with solid and cystic components. The characteristics of thick wall with solid and cystic components and the presence of calcification, which is seen in 50% of cases, are useful in distinguishing a cervical teratoma from lymphangioma. The unique imaging characteristics are helpful in making the diagnosis but sometimes can be confusing as was in our case.

Some studies have suggested that origin of neck teratomas is from embryonic cells in primitive thyroid gland which explains there close relationship with thyroid tissue requiring thyroidectomy in some cases. After surgical removal, cervical teratoma patients should be evaluated for hypothyroidism. In a series of 23 children presented to Great Ormond Street Hospital, London, with a head and neck teratoma, 5 cases of neck teratoma were found to have origin from the thyroid gland. In our case, also the origin of mass was from the thyroid tissue.

Serum AFP level is a tumor marker for making the diagnosis, but its interpretation can be fallacious as it can normally be elevated in the neonatal period and can vary depending upon the age and birth weight.

Prompt surgical excision of a teratoma is the treatment of choice. If needed, stabilization of airway should be done before the surgery. Cervical teratomas in children are almost always benign and rarely can be immature or malignant. However, it has been reported that immaturity of various constituent tissues in childhood teratomas (unlike adult) does not impact on the otherwise favorable prognosis of the teratoma. Chemotherapy is indicated in case of recurrence, disseminated disease, invasive tumor, incomplete resection, and malignancy. The disease and

Discussion

Congenital cervical teratomas are rare in newborns, accounting for 1%–5% of all pediatric teratomas. They predominate in females. Large cervical teratomas may extend into the mediastinum or compress trachea, causing pulmonary hypoplasia and many of these may require an extensive perinatal multidisciplinary planning for EXIT (exutero intrapartum treatment) procedure. If left untreated, cervical teratomas have an estimated mortality rate of 80%–100%.

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recurrence can be monitored by serial AFP measurements at regular intervals.\[3\]

Few prognostic factors have been reviewed by Alexander et al.\[7\] which includes polyhydramnios and the tracheoesophageal index (sum of the lateral and ventral displacements of the tracheoesophageal complex from its normal anatomical location at the ventral aspect of the cervical spine) which have been found to be associated with the increased risk of neonatal respiratory compromise.

Long-term outcome in these patients has been found to be excellent after complete surgical excision.\[4\] Outcome is certainly excellent when compared to lymphatic malformations in terms of quality of life, persistent or recurrent disease.\[10\]

**Conclusion**

A large neck mass in the neonatal period requires a high index of suspicion based on the clinical features and imaging to differentiate a teratoma from a lymphangioma as prompt surgical excision is mandatory in the former. Although usually benign, cervical teratomas carry a significant risk of intrauterine or neonatal death because of polyhydramnios and upper airway obstruction. A good planning and imaging ensures complete excision without any significant morbidity.

**Declaration of patient consent**

The authors certify that they have obtained all appropriate patient consent forms. In the form, the legal guardian has given his consent for images and other clinical information to be reported in the journal. The guardian understands that names and initials will not be published and due efforts will be made to conceal identity, but anonymity cannot be guaranteed.

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Nil.

**Conflicts of interest**

There are no conflicts of interest.

**References**