Giant Congenital Cervical Teratoma: Diagnosis, Management and Long-term Follow-up

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SUMMARY

Cervical congenital teratomas are rare conditions associated with abnormal prenatal development.

These tumors are usually benign; however, malignant transformation has been rarely described.

When teratomas are big, they are usually detected prenatally. In these circumstances multidisciplinary management after the delivery is necessary and the prognosis has improved provided that the airway is not obstructed and resection of the tumor is not delayed.

We present two cases of massive neonatal cervical teratoma detected prenatally by ultrasound. In one case, elective caesarean was decided at 40th week. In the other case, the child was delivered at 29th week of gestation, an airway was secured, and total resection of the tumor was performed. None of them have neurological deficit detected on long-term follow-up.

We describe our proposed management with special attention to prenatal diagnosis by ultrasound and review other series in order to provide help in decision-making.

Fetal ultrasound showed a mass tumor with mixed components measuring 90 mm that originated from the right neck without pulsatile components. Oral cavity was not involved. The rest of the structures of the neck were normal.

Prenatal diagnosis of benign tumor arising from the neck (teratoma or cystic hygroma) was made. Follow-up of the fetus was done every week and a cesarean section was performed at 40th week. A female healthy fetus with a huge cystic tumor on the neck was obtained. The baby weighed 3.850 gm and Apgar score was 6/9 at 1 minute. The tumor mass originated in the right neck and had a size of 10 cm. It was not transilluminable and had two major blood vessels surrounding it (Fig. 1).

Endotracheal intubation of the newborn was not necessary and the baby was referred to a neonatal intensive care unit.

Fourth day after birth surgical intervention was considered due to difficult nutrition management related to the tumor mass. At this time, the mass gross appearance had changed showing ulceration over the apex.

During surgical intervention we found that the mass was fixed at the floor of the mouth without any infiltration to the bone structure. The mass was removed and reconstruction of the mouth floor was performed.

Pathological exam revealed a 14 × 12 × 8.5 cm mass with solid components and cystic compartments full of viscous yellow fluid (Fig. 1). Histological examination of the specimen revealed an immature grade II teratoma.

Postoperative course after removal of the tumor was uncomplicated. The baby is now 16-year-old and actually have no neurological sequels neither organ dysfunction.

CASE PRESENTATION

Case Report 1
A 35-year-old multiparous woman was referred at 34th week of gestation for advice about abnormal sac criteria corresponding to her gestational age.
fetus associated with cervical hyperextension. Cystic areas of the tumor were predominant in the right region of the tumor (Fig. 2). The tumor mass approached the esophagus in its proximal superior region but did not involve it. Polyhydramnios was present but there was not impaired swallowing demonstrated on ultrasound. A fluid-filled stomach was present.

Transabdominal cytogenetic amniocenteses showed 46(XX) 9qh+ karyotype. Isolation of trophoblasts showed discrete rise level on basal TSH with decrease of total T3. T4 and TGB-antibodies were normal.

A second ultrasound at 29th week showed polyhydramnios and rapid enlargement of the mass in the fetus neck (9 cm) compromising adjacent neurovascular structures. Then, we decided an elective caesarean in the 32nd week after parents were correctly informed.

A 32nd weeks estimated gestational age female was born with caesarean section. The female baby had respiratory distress with Apgar scores of 5 at 1 minute and clearly showed neck hyperextension. Nasotracheal intubation was further necessary to save baby’s life (Fig. 3).

CT scan showed a non-homogenous tumor mass (12 × 10 × 9) with solid and cystic areas. The esophagus, trachea and larynx were displaced to the patient’s left with flattening of the esophageal lumen. Plain radiographs showed a soft tissue mass with calcium deposits.

Once decision to recommend operative intervention was accepted by the parents surgical intervention was carried out (second day) taking special care to avoid injuries to the thyroid gland and neurovascular structures of the neck (Fig. 4).

The pathological diagnosis was benign cystic mature teratoma. Histology showed neuroepithelium, cystic areas, calcifications, fibroelastic connective tissue and thyroid epithelial cells. The postoperative course was uneventful and at 5-year-follow-up the aesthetic result is excellent. Thyroid
and parathyroid functions are normal and no neurological deficit has been developed.

DISCUSSION

Teratomas are embryonic tumors with variable degrees of differentiation composed of a variety of tissues comprising single germ cells (ectodermal, mesodermal or endodermal) and ectopic embryogenic non-germ cells.1

Despite the ongoing controversy concerning its origin, there are three major accepted theories:2,3

1. Origin during early fetal development (loss of one twin during pregnancy).
2. Pluripotent germ cells.
3. Ectopic embryogenic non-germ cells.

Approximately 30 percent of cervical teratomas are composed of thyroid epithelium. Sometimes the thyroid gland is occupied or even substituted for teratoma tissue.4,5 This relationship to the thyroid gland has been argued to be so constant that Roediger and associates6 have postulated that all teratomas presenting in the anterior neck region arise from embryonic cells in the primitive analogue of the thyroid gland.7 Other authors do not share this opinion and regard the relationship with the gland as entirely fortuitous, in view of the extreme rarity of intrathyroid origin.

Teratomas can be broadly classified into mature or immature. Most neck teratomas are reported as mature, and malignant immature teratomas have been said to occur in approximately 5 percent of teratomas of the neck.

Mature teratomas usually contain well-differentiated tissues from the ectodermal, mesodermal, and endodermal germ cell layers, and any tissue type may be found among the tumor. Mature teratomas are benign, though some mature and immature teratomas may secrete enzymes or hormones. Immature teratomas also contain tissues from all 3 germ cell layers, but immature tissues, primarily neuroepithelial, are present. Immature teratomas can be graded from I to IV based on the amount of immature tissue found in the tumor specimen. High grade immature teratomas are associated with aggressive behavior, malignant foci and local recurrence.3 There is usually no correlation between tumor grade and patient age.
Malignant teratomas are rare in the neonatal period; however, adults are at high risk for malignancy.\(^7,9\)

Common risk factors related to malignancy include: age (specially adults), high grade immature tumors, and neonatal tumors that have been not removed before two months of age.\(^3\)

The prognosis of neck teratomas varies. Unless surgery is instituted without delay, the prognosis of cervical teratoma can be serious. In the past, a large proportion of patients with cervical teratomas (more than 25% of those reviewed by Silberman and Mendelson)\(^10\) died prior to surgery.

The diagnosis of congenital cervical teratomas may be made in the antenatal, perinatal or postnatal periods. Frequently the diagnosis is made in utero by ultrasound, as is the case for our two patients. However, the diagnosis of cervical teratomas can be difficult before excision. Important improvement in the management of neonates suffering from congenital cervical teratomas and tumors in the neck has occurred during the past decade. This development is partly due to the widespread use of prenatal ultrasonography enabling prenatal diagnosis of these tumors.

Some neoplasm show specific ultrasonographic findings suggesting the differential diagnosis, but others do not. Knowledge of the presence of a neoplasm in the fetus may alter the prenatal management of a pregnancy and the mode of delivery, and facilitates immediate postnatal treatment. Our patients are a good example to show this final assessment. In both cases we decided an elective caesarean in order to avoid accidental tearing of the tumor and damage to the baby.

During pregnancy, the reason most often cited for routine ultrasound is to detect fetal anomalies. Nowadays, ultrasound is extensively used in obstetrics and gynecology. The potential benefits of ultrasound include:

1. Low cost.
2. Easily accessible.
3. Parental counseling.
4. Ultrasonography is the most innocuous and noninvasive procedure.
5. It can detect polyhydramnios (one-third of the patients; secondary to difficulty in swallowing of amniotic fluid).
6. It can detect cervical masses with solid and cystic areas (high suspicious for teratoma).
7. Ability to characterize the composition of teratomas.
8. Prenatal diagnosis permits the careful arrangement of the time, location, and mode of delivery.
9. Ability to characterize relations of the mass to the large vessels of the neck.
10. It can be useful to planned preoperatively the surgical treatment of cervical teratomas.
11. It can be useful as a guidance to localize the tracheal rings during EXIT procedure.
12. It can confirm endotracheal tube placement.
13. It may help us as assistance and guidance in surgical dissection intraoperatively.

The differential diagnoses for prenatally diagnosed neck masses are cystic hygroma, congenital goitre, thyroid masses, neuroblastoma, thyroglossal duct, branchial cleft cyst, cervical hemangioma, cervical lymphangioma, laryngocele, meningocele and macerated twin fetus.\(^2,3\) Actually, further improvements in other imaging modalities such as 3D ultrasound and ultrafast MRI has enabled us to delineate the tumor with high specificity to assure an adequate prenatal and postnatal management and parental counseling.\(^11\)

There are many reports of the features of head and neck teratoma on MRI, but its use to diagnose these lesions has not been fully established. High signal intensity from fat on T1-weighted images may be helpful but is not always present. Multiloculated cystic tumors with high signal on T2-weighted images may be indistinguishable from cystic hygroma.

In an elegant retrospective study aiming to evaluate the role of MR imaging to improve sonographic prenatal diagnosis of congenital anomalies the authors found that fetal MRI were most effective to study retroperitoneal pathology, meningocele and visualizing the airways, brain parenchyma, and congenital brain malformation.\(^11\)

Therefore, ultrafast MR imaging should be used as an adjunct to ultrasound when findings are equivocal or when the ultrasound images are difficult to interpret in the prenatal diagnosis of cervical teratomas. However, the prenatal differential diagnosis of these tumors is often difficult or even impossible (Figs 5 and 6).

Polyhydramnios can complicate 20 to 40 percent of the prenatally diagnosed cases. Over-distended uteri from polyhydramnios have been known to cause maternal complications such as ureteral obstruction.\(^2,8\) Also, the increase in uterine size due to polyhydramnios can precipitate preterm labor and delivery. Serial ultrasound examinations should be performed to monitor amniotic fluid volume. In some cases, an excess amniotic fluid level can be reduced to avoid complications. Premature labor and delivery in cases of giant cervical teratoma are common. Prenatal diagnosis allows for a management approach tailored to meet its anticipated complications.\(^12\) Airway obstruction can be life-threatening and accounts for up to 50 percent of the mortality associated with this tumors. When upper airway obstruction is anticipated, an early elective cesarean delivery should be performed (before spontaneous labor begins) and the baby should be left on fetomaternal circulation until the airway is secured.\(^12\) Tracheostomy should be reserved for airway obstruction, when orotracheal intubation is not possible. Fiberoptic bronchoscopy may obviate the need for tracheostomy in some patients if attempted at initial intubation maneuver. In our experience, one patient required nasotracheal intubation with EXIT procedure while in the other was not necessary to secure the airway. In both cases a multidisciplinary team was present at delivery.
Figs 5A to C: (A) Cervical mass with solid and cystic areas (arrow); high suspicion for teratoma. (B) Cervical teratoma compressing the trachea extrinsically. Tracheal stenosis in the inferior third. Strict follow-up allows for a management approach to avoid airway obstruction at delivery. (C) Ability to characterise relations of the mass to the large vessels of the neck and blood supply of the tumor.

Fig. 6: Magnetic resonance of the tumor mass showed a well surrounded multiloculated cystic tumor.

Two procedures have been used from the past to secure airway and avoid obstruction of the baby at birth: EXIT and OOPS procedure.

The EXIT procedure refers to Ex-utero Intrapartum. It refers to a treatment in which placental support is maintained until the airways are evaluated, surgically corrected and secure. The technique leaves an intact fetoplacental circulation and guarantees a normal fetal oxygenation while fetal airway patency is secured. To perform this procedure the head and at least one hand of the fetus are delivered and the rest of the body along with the umbilical cord and the placenta remain in utero. Only after the baby is intubated is the umbilical cord clamped and the body delivered.12-15

The OOPS procedure (operation on placental support) is a method in which umbilical cord is preserved and only clamped after the airway has been secured by intubation or by tracheotomy. Maximum uterine relaxation is important in both procedures in order to avoid separation of the placenta from the uterus and maintain fetomaternal circulation ensured.16

In addition to the OOPS and EXIT procedures, extracorporeal membrane oxygenation (ECMO),1 and cannulation of the umbilical vessels have been suggested.

Cesarean section, especially for large tumors, is indicated to avoid birth dystocia and teratoma avulsion. In rare cases in which hydrops develops earlier in gestation, open fetal surgery to remove the cervical teratoma may be necessary to save the baby. Early surgical resection of the tumor should be anticipated in cases of large cervical teratomas in order to avoid complications and malignant degeneration. Preoperative evaluation should include a chest X-ray, and CT scan of the head, neck, and chest to provide vital information regarding invasion and extension of tumor. The team approach is of utmost importance in providing an organized and coordinated care plan. An obstetrician and/or perinatologist, neonatologist, pediatric surgeon and if necessary, anesthesiologist and otolaryngologist should all be available in the delivery suite for the birth, resuscitation and possible surgical intervention of the neonate.17 At present, surgical removal offers the best approach for survival and cure. Our two babies (at just two and four days old) underwent surgical intervention to avoid further complications.

Functional and aesthetic results after surgery are usually excellent. Most common postoperative complications include recurrent laryngeal, marginal mandibular and vagus nerve damage.4,8,17 In part this is due to distortion of the neck anatomy secondary to the mass. Transient hypothyroidism and hypoparathyroidism has been described.8 Mental retardation ranging from mild to profound is probably secondary to hypoxia associated with prolonged resuscitation maneuvers.

Alpha-fetoprotein is elevated in cases of fetal neural tube defects, congenital defects, and anomalies such as cervical teratoma. Usually, a preoperative alpha-fetoprotein level is recommended. Obtaining serial beta-hCG and alpha-fetoprotein levels will allow for follow-up of the completeness of tumor excision or recurrence.
In a recent review it was stated that the most important variable affecting long-term survival in literature cases has been successful and complete resection of the tumor. At long-term follow-up (at least 5 years), our patients had no evidence of local or distant recurrence on examination and radiological images.

Even though cervical teratomas are considered benign tumors in infancy, occasional malignant transformation and consequent death have been reported. Until 1998, true malignant teratomas have been reported in only 9 of 220 cases. The diagnosis of malignancy was made secondary to metastasis and/or histological evidence of germinoma, embryonal carcinoma, or choriocarcinoma.

REFERENCES