Thanatophoric dysplasia, an enigmatic dilemma: a case report

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ABSTRACT

Thanatophoric dysplasia is a rare, fatal form of skeletal dysplasia that affects fetus in utero. It is characterized by marked underdevelopment of fetal skeleton and short limbs. This disorder can be diagnosed antenatally and the couple should be counselled for termination of pregnancy. 20 years old female came in her second trimester carrying an ultrasound report which showed features of thanatophoric dwarfism. She was advised to terminate her pregnancy but she decided to continue with her pregnancy. She was induced after 40 weeks and she delivered stillborn baby with dwarf like features, a condition identified as thanatophoric dwarfism. Early diagnosis can be done by ultrasound as early as 13 weeks of gestation and since babies born with thanatophoric dysplasia have a very poor prognosis, couples should be counselled regarding early termination of pregnancy. 3D ultrasound scan and molecular analysis can also help identify this disorder.

Keywords: Dysplasia, Skeleton, Pregnancy

INTRODUCTION

Thanatophoric dysplasia (TD) is a form of skeletal dysplasia with very few isolated cases all over the world. It is the second most lethal form of skeletal dysplasia after osteogenesis imperfecta type 2. The word thanatophoric is Greek for “death bearing” or “death bringing”. It is rare with an estimated incidence of 0.28 to 0.60 per 1000 births.1 Very few cases of thanatophoric dysplasia have been reported from India so far. This condition is characterized by marked underdevelopment of the skeleton and short limb dwarfism.2 Phenotypically, these infants have macrocephaly, platyspondyly or flattened vertebral bones, narrow thoracic cage with hypoplastic lung, normal trunk length and rhizomelia or severe shortening of limbs. This condition usually leads to death in neonatal period from respiratory insufficiency. We report one such case of thanatophoric dysplasia here.

CASE REPORT

20 years old patient came to our OPD as primigravida at 16 weeks of pregnancy citing some abnormality in the fetus in her ultrasound report. Her level 2 ultrasound report showed femur length of 10 mm suggestive of skeletal dysplasia. She was counselled regarding termination of pregnancy owing to poor fetal prognosis but the couple wished to continue her pregnancy. There was no history of similar or any other type of malformations in fetus in any of the family members. An MRI was done which showed a large fetal skull with indentations on both sides giving appearance of clover leaf shaped skull (BPD- 96 cm ~ 39 weeks 2 days) with normal brain parenchyma and ventricular system. Fetal neck appeared short with hyper-extended fetal head giving star gazing appearance. Thoracic cavity was small and narrow. Bilateral humerus and femori were extremely short ~ 22 mm and 19 mm, corresponding to 5 weeks 6 days, respectively with metaphyseal flaring. Other bones
like radius, ulna, tibia and fibula were also very short. Pelvis was narrow with small, squared iliac wings. Diffuse skin and subcutaneous soft tissue was thickened. Liquor was found to be increased. Fetal spine was normal.

Since patient was not willing for termination, she was kept in follow-up for the rest of her pregnancy. Her labour was induced at 40 weeks 3 days and had she vaginal breech delivery of a stillborn male fetus, weighing 2.5 kg, who died during the course of labor (Figure 1). The couple did not agree for the autopsy of the fetus. However, an infantogram of the fetus could not be taken post-delivery (Figure 2).

**DISCUSSION**

TD is an autosomal dominant disorder caused by missense mutation in Fibroblast growth factor receptor 3 or FGFR 3. All cases of TD are due to new mutations and due to its sporadic nature, incidence is less than 3%. This condition was first brought to notice by Montreaux et al in 1967. The gene is present on the short arm of chromosome 4 (4p16.3). FGFR3 belongs to tyrosine kinase receptor family which acts as negative regulator of bone growth. Point mutation occurs in FGFR 3 gene in cases of thanatophoric dwarfism which prevents chondrocyte differentiation leading to disorganised endochondral ossification at bone growth plate. Risk of recurrence is less than 5%, possibly caused by gonadal mosaicism. Of the two types of TD, type 1 constitute fetuses with short curved limbs giving telephone-receiver

**Figure 1:** The picture of the male baby shows a large head with bulging forehead, flattened nasal bridge, short neck, long trunk with very short limbs with redundant skin folds.

**Figure 2:** An infantogram was done which showed a large calvarium with narrow thoracic cavity and shortened ribs with anterior splaying, flattened vertebral bodies (platyspondyly), squared iliac wings or which mimic "elephant ears" and short bowed femurs giving "telephone receiver" appearance.
appearance with macrocephaly but normal shaped skull, as seen in our case while type 2 is associated with clover leaf shaped skull and straight femur bone.

Diagnosis is made by ultrasound done in second or third trimester which include the following radiological features:

- Macrocephaly or clover leaf shaped skull or "kleblattschaedel"
- Rhizomelia viz. short, bowed bones esp. of femur
- Narrow rib cage with short, splayed, cupped ribs
- Platyspondyly or flattened 'U- shaped' vertebral bones
- Redundant soft tissues of limbs
- Squared iliac bones
- Normal trunk length.

**Polyhydramnios is commonly associated**

On 3-D ultrasound, features like skin thickening and cranio-facial abnormalities are seen.

MRI can help visualize narrow foramen magnum and platyspondyly. Differential diagnosis includes osteogenesis imperfecta types 2 and 3, achondroplasia, achondrogenesis.

Prognosis is poor as death occurs mostly due to respiratory failure owing to hypoplastic lungs or from brainstem compression from narrowed foramen magnum. However hence, if the condition is diagnosed pre-natally couples should receive genetic counseling and advised for termination of pregnancy. Without proper counseling, families who had a previous child affected with thanatophoric dysplasia would be anxious to think of a future pregnancy. If the diagnosis is made after the birth of the baby, management is solely supportive, as death occurs from respiratory insufficiency within hours to days.

Thanatophoric dysplasia, although rare, is a lethal disorder of fetus and can be detected by ultrasound done in second trimester. Early diagnosis helps in prognostication and termination can be offered to the patients.

Stress should be laid upon proper genetic counseling as couples tend to get anxious about recurrence of this disorder in future pregnancies. Couples should be assured about chances of a normal pregnancy because of the sporadic nature and low risk of recurrence of this disorder. An ultrasound scan can rule out any form of skeletal disorder or any other fetal abnormality in early pregnancy. It is possible to recognize short limbs in fetuses beginning as early as 13 weeks of gestation when femur length can be routinely assessed. 3-D anatomy scan and amniocentesis for molecular analysis can detect the disorder in the second trimester and in case of this abnormality, termination should be advised.

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