Thanatoporic dwarfism

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Thanatoporic dwarfism, a lethal form of neonatal dwarfism is a congenital malformation, the cause of which is still not certain. In the past, many cases of thanatoporic dwarfism were confused with achondroplasia and achondrogenesis. Prenatal diagnosis of thanatoporic dwarfism has been achieved by ultrasonography and radiography in the second or third trimester of pregnancy.

A case of thanatoporic dwarfism diagnosed at the early beginning of the third trimester is presented. Early prenatal diagnosis and elective abortion may avoid problems in obstetric management which may occur later in pregnancy.

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รายงานผู้ป่วยทั้งหมด 253 ราย ประกอบด้วย 219 รายเป็นชาย และ 34 รายเป็นหญิง ปรากฏการณ์ที่มีความเกี่ยวข้องกับภาวะการเติบโตเร็วขึ้น ได้แก่ ภาวะการเติบโตที่ผิดปกติ metaphysis ซึ่งมีเนื้อเยื่อตันที่มีการเจริญเติบโตที่ผิดปกติ และการเจริญเติบโตของกระดูกที่ผิดปกติ achondroplasia หรือ achondrogenesis เป็นต้น กระบวนการเจริญเติบโตของกระดูกที่ผิดปกติในระยะยาวไม่สามารถคงอยู่ได้ การให้การรักษาภาวะการเติบโตที่ผิดปกติจะช่วยให้การเจริญเติบโตของกระดูกที่ผิดปกติ สามารถคงอยู่ได้ เมื่ออาการหรืออาการที่เกี่ยวข้องถึงระดับหนึ่ง ให้ทำการรักษาด้วยการใช้ผลิตภัณฑ์ที่มีประสิทธิภาพในการดูแลรักษา
Thanatophoric dwarfism is one of the severe types of congenital malformation which occurs in the fetal skeletal system at the metaphysis\(^1\). This malformation was first reported by Maroteaux et al in 1967\(^2\) using a term derived from Greek which means “death bearing”, because neonates afflicted with this type of malformation died a short time after birth\(^3\). In the past, this condition had often been misdiagnosed as achondroplasia since the two conditions share very similar features but recently more studies have classified the differences between the two conditions.

Case report
A 26 year old primigravida who lived in Bangkok first attended the antenatal clinic at a Bangkok Metropolitan Administration’s Health Centre when she was approximately 11 weeks pregnant but uncertain of the date of the last menstrual period. Physical examination and results of routine laboratory were within normal limits. She had no family history of congenital malformation nor had her husband who was a 36 year old taxi driver.

Subsequent 4 visits to the antenatal clinic showed slightly less than average weight gain of 1.5 kg. in 4 weeks but normal increase of uterine size until 28 weeks gestation when uterine fundus was at the level of 3/4 above the umbilicus, larger than gestational age with inaudible fetal heart sound.

She was referred to the antenatal clinic at Chulalongkorn Hospital where her fundus was found to be at the level of 3/4 above the umbilicus with a live fetus as confirmed by fetal heart sound auscultation. Her BP was 120/80 and she had no glucose or albumin in her urine. Since she was uncertain of the date of her last menstrual period, ultrasonographic examination was performed to confirm gestational age. A live fetus was found in cephalic presentation with Enlargement of the cerebral ventricle (Figure 1,2), short arms and legs, polyhydramnios and a posterior placenta. An initial diagnosis of hydrocephalus with short limbs and polyhydramnios was made.

![Figure 1. Normal fetal head from ultrasound scanning.](image-url)
Figure 2. Dilatation of cerebral ventricle in this fetus.

A plain abdomen X-ray 2 weeks later revealed a fetus with short arms and legs and small spinal column (Figure 3). A definitive diagnosis of thanatophoric dwarfism was made. Two weeks later, the patient presented at the labour room with a two hour history of ruptured membrane and one hour later labour occurred spontaneously. Labour progressed uneventfully and she had a normal delivery, of a live female neonate weighing 1,500 G., 8 hours later. The neonate was 34 cm. long with abnormally large head, short arms and legs (Figure 4), a rather small thoracic cage, and survived for only 35 minutes. The normal looking placenta weighed 440 G and the cord was 35 cm. long, unfortunately the placenta and cord were not sent for histopathology. The patient’s stay in hospital was uncomplicated and she was discharged 3 days after delivery.

Figure 3. X-ray of fetal spine.
Comment

Thanatophoric dwarfism is a type of dwarfism which is incompatible with life and is the most common cause of death in dwarf neonates. Its aetiology is uncertain and most occurrences are sporadic with male to female sex ratio of 2:1. There have, however, been reports of autosomal recessive type of inheritance in 2 offsprings with thanatophoric dwarfism, from consanguineous parents.

Characteristics of thanatophoric dwarfism include micromelia, shortened body, averaging 40 cm at term. The skull is enlarged with widened fontanelles and small foramen magnum. The skull base is narrow with small face and flattened nasal ridge. The thoracic cage is narrow with short ribs and curved arms and legs. Microscopic examination of the bones reveal that chondrocytes and bony trabeculae alignment are not in an orderly fashion, especially around the central epiphyseal—metaphysis area. The vertebrae are flattened and there are wide intervertebral gaps.

Prenatal diagnosis can be made using ultrasonography, X-ray, or fetography where some of the above characteristics may be apparent. Occasional difficulties may be encountered when differentiating this condition from other types of congenital dwarfism such as achondroplasia, achondrogenesis or osteogenesis imperfecta and the final diagnosis will depend on detailed history and above mentioned investigations. In the case of homozygous achondroplasia, both parents will have features of the condition. If one parent is normal, achondroplasia can then be excluded. In the case of heterozygous achondroplasia, the thoracic cage is not as small and narrow as that of thanatophoric dwarfism whose body is of normal length but with short arms and legs. In the case of achondrogenesis, the body, arms, legs are all short with less than normal ossification in the spinal column, the pelvis and the skull. In the case of osteogenesis imperfecta, ultrasonography or X-ray will reveal similarly curved arms and legs but there are areas of demineralization of the skull, arms and legs as well as several fracture sites.

Compared with ultrasonography, X-ray has the advantage of more definite imaging of various bones, and ossification centers are better visualized.

Thanatophoric dwarfism is mostly diagnosed during the 3rd trimester because the uterine size is larger than average for gestational age, as in this case,
due to the fact that thanatophoric dwarfism is frequently associated with polyhydramnios. Other complications may be found such as abnormal presentation, preterm labour, and cephalopelvic disproportion.

Diagnosis during early pregnancy will mean easy and safe termination of pregnancy by the vaginal route with avoidance of unnecessary abdominal delivery. In a case of near term pregnancy with a breech presentation, the short and rigid neck of thanatophoric dwarfism has limited movement and will lead to difficult delivery of the after coming head with subsequent birth canal injuries if vaginal delivery is attempted.

Prenatal diagnosis of congenital anomalies is therefore advisable especially among those with risk factors such as history of congenital anomalies in previous pregnancies and elderly gravidae, in order to avoid dangerous complications of pregnancy and delivery as well as psychic trauma to the mothers.

**Summary**

Thanatophoric dwarfism is a type of congenital malformation which is not compatible with life. Its aetiology is still uncertain, therefore prenatal diagnosis especially using ultrasonography to detect abnormality of fetal limbs, vertebrae and skull in early pregnancy will result in safe and efficient management of these unfortunate gravidae.

**References**