A CASE OF CONGENITAL HIGH AIRWAY OBSTRUCTION SYNDROME CAUSED BY COMPLETE TRACHEAL OBSTRUCTION WITH ASSOCIATED ANOMALIES

Hyun Jin Cho, MD1, Yeon Mee Kim, MD, PhD2, Hee Young Lee, MD3, Hye-Sung Won, MD, PhD3

Departments of 1Obstetrics and Gynecology, 2Pathology, Haeundae Paik Hospital, University of Inje College of Medicine, Busan; 3Department of Obstetrics and Gynecology, Asan Medical Center, University of Ulsan College of Medicine, Seoul, Korea

Congenital high airway obstruction syndrome (CHAOS) is caused when the upper airway is obstructed or severely narrowed. The prenatal ultrasound findings of CHAOS include large echogenic lungs, inverted diaphragms, dilated airways, and fetal ascites and/or hydrops. Recently, ex utero intrapartum treatment (EXIT) procedure or fetoscopic tracheostomy are being widely used for the treatment of CHAOS. However, CHAOS with early presentation of hydrops confers ominous sign even with EXIT procedure. We report a case of CHAOS with hydrops and associated anomalies that was confirmed by autopsy.

Keywords: Congenital high airway obstruction syndrome; Prenatal ultrasound; Fetal hydrops; Autopsy

Case Report

A 29-year-old woman (gravida 1, para 0) was referred to the Asan Medical Center at 20+5 weeks of gestation for evaluation of a fetal lung mass. Her menstruation had been on a regular cycle of 28–30 days. Her medical and familial medical history was unremarkable. Amniocentesis had already been performed before the referral because of fetal hydrops and a normal fetal chromosome was confirmed. A high resolution ultrasound examination was carried out at the Asan Medical Center. At the ultrasound examination, fetal weight was estimated at 542 g and the amount of amniotic fluid was normal. An ultrasound examination showed bilateral, expanded, hyperechoic lung, inverted diaphragm, anterior displacement of the heart, a visible trachea, massive ascites, pleural and pericardial effusion, and generalized edema (Fig. 1). These findings were strongly suggestive of CHAOS with fetal hydrops. The patient and her husband were counseled about the natural history and prognosis of CHAOS. They decided to terminate the pregnancy. The patient was transferred to Haeundae Paik Hospital near her house three days later. Labor was induced with laminaria and intravenous sulprostone. The patient delivered vaginally a breech-presented fetal body at 6 pm and a fetal head...
disconnected from the body came out immediately thereafter. An autopsy was performed to establish causes of the upper airway obstruction. A postmortem finding of a fetus that weighed 451 g was consistent with CHAOS. The fetus had a total blockage of the trachea due to thickened tracheal cartilage and soft tissue, herniation of brain and spinal cord into the dilated upper trachea, massively enlarged lungs bilaterally, flattened diaphragm, soft tissue edema of the trunk, and a compressed, relatively small heart (Fig. 2). There were micrognathia and a left 3rd toe that over-rode the 2nd toe. On microscopic examination, increased proliferation of fibroblasts in the tracheal cartilage and soft tissue was found (Fig. 3). Two days later, the patient was discharged uneventfully.

**Fig. 1.** Ultrasound images of a fetus with congenital high airway obstruction syndrome at 20 weeks of gestation. (A) Dilated fluid filled trachea and massive ascites. (B) Transverse scan of an enlarged echogenic lung, compressed heart and fetal hydrops.

**Fig. 2.** Postmortem findings show a totally obstructed trachea (arrow).

**Fig. 3.** Microscopic examination of the obstructed portion of the opened trachea shows proliferated fibroblasts around soft tissue (arrow). In this view, a totally obstructed wall opposite the trachea was confirmed (arrowhead) (H&E, ×40).
Discussion

CHAOS is caused by complete obstruction or severe narrowing of the upper airway. Causes of obstruction are laryngeal atresia, sub-glottic stenosis, laryngeal web, a completely occluding laryngeal cyst, or, rarely, a tracheal obstruction. Complete or near complete obstruction of the upper airway in utero causes retention of fetal lung fluid and increases in intratracheal pressure. Massive distension of lung by intratracheal pressure results in cardiac and caval compression and subsequent heart failure, ascites, pleural effusion and hydrops in utero [2]. The true incidence of CHAOS is unknown because this is a very rare and life-threatening condition with high mortality and morbidity. Survival is dependent on early prenatal diagnosis, close prenatal follow up, and fetal or immediate postnatal intervention. If unrecognized before delivery, the reported mortality is 80% to 100%. In 1994, Hedrick et al. [1] first described the constellation of findings of CHAOS: large echogenic lungs, flattened or inverted diaphragms, dilated airways distal to the obstruction, and fetal ascites and/or hydrops. Advancements in ultrasound technology and introduction of fetal magnetic resonance imaging can provide visualization and localization of the level of the obstruction [3].

We present a case of CHAOS caused by complete tracheal obstruction. We found increased proliferation of fibroblasts in the tracheal cartilage and the soft tissue around the cartilage. The estimated abdominal circumference by ultrasound was 27 cm (which is normal for 31+3 gestational weeks) because of fetal ascites and generalized edema, whereas the estimated fetal biparietal diameter was 5.7 cm (normal for 22 gestational weeks). There was no abnormal finding regarding the fetal head at the prenatal ultrasound examination. Therefore, at the time of vaginal delivery of the breech-presented fetus, brain herniation and separation of the fetal head from the body may have occurred by traction of the fetus for delivery of the fetal abdomen trapped tightly in the uterine cervix.

CHAOS might be associated with other structural abnormalities. In the case presented, the fetus with CHAOS was accompanied by micrognathia and abnormalities of the toes. When CHAOS is diagnosed with the presence of severe associated abnormalities, a fatality is even more likely. A notable example of disease with upper airway obstruction with associated anomalies is Fraser syndrome. Fraser syndrome is characterized by variable defects including laryngeal atresia, cryptophthalmos, renal agenesis, syndactyly and genital abnormalities. Fraser syndrome is an autosomal recessive disorder - the recurrence rate among siblings is 25%. A high maternal serum alphafetoprotein level may increase the suspicion of Fraser syndrome [4,5]. Laryngopharyngeal/pharyngotracheal pinpoint fistula on the obstructed portion or tracheoesophageal fistula may allow communication of lung fluid and release of increased intrathoracic pressure. Patients with pinpoint fistulas often do not have severe hydrops in utero [6]. Isolated airway obstruction without hydrops has a relatively favorable prognosis. CHAOS with early presentation of hydrops is an ominous sign with a high rate of fetal demise and a poor survival rate even with the ex utero intrapartum treatment (EXIT) procedure. Roybal et al. [7] reported the prognosis and severity of twelve cases of CHAOS. According to their study, there was no evidence of fetal hydrops in four of the five cases whose EXIT procedures were successful. The number of fetuses who had fetal hydrops was eight, and only one fetus among the eight survived for more than one year.

The EXIT procedure for a various fetal conditions was first reported by Mychaliska et al. [8] in 1997. The first successful EXIT procedure for a case of CHAOS was reported by DeCou et al. [9] in 1998, and there have been many studies of EXIT for fetal airway obstructions. The EXIT procedure is performed using high doses of inhaled halogenated agents to facilitate uterine relaxation during cesarean section. The fetus remains hemodynamically stable, as the uteroplacental circulation is maintained. A variable strategy is attempted at intubation, including direct laryngoscopy, rigid bronchoscopy, and tracheostomy. And then the uteroplacental circulation is interrupted [8]. In recent years, fetoscopic tracheoscopy has been performed to eliminate the cause of obstruction. It was possible to create a pharyngotracheal fistula for the decompression of the lung. However, further studies are necessary to determine the long term effects and safety of fetoscopic tracheoscopy [2,10].

In the case presented, the fetus with CHAOS not only had severe hydrops, but also had associated anomalies. We could not try the EXIT procedure or tracheoscopy in utero. This procedure requires cooperation of obstetricians, pediatric surgeons, anesthesiologists, and neonatologists. This report is a rare case of CHAOS with severe hydrops and associated anomalies in Korea. We hope this case report will encourage others to try to perform the EXIT procedure and other fetal treatments.

References


다른 기형을 동반한 완전 기도 폐쇄에 의한 선천성 상기도 폐쇄증후군 증례

인제대학교 의과대학 해운대백병원 1산부인과, 2병리과, 3울산대학교 의과대학 서울아산병원 산부인과 조현진1, 김연미2, 이희영3, 원혜성3

선천성 상기도 폐쇄증후군은 상기도가 완전히 폐쇄되거나 심하게 좁아지는 경우에 발생한다. 선천성 상기도 폐쇄증후군은 초음파상 커지면서 고에코성으로 보이는 폐와 역위된 횡경막, 늘어난 기도, 그리고 태아 복수 혹은 수종이 생길 때 산전진단이 가능하다. 최근에는 분만 시 시행하는 자궁외 수술이나 태아경을 이용한 기관절개술이 이 질환의 치료로 이용되고 있다. 그러나 임신초기에 태아수종을 동반하는 경우에는 치료를 하더라도 예후가 매우 나쁘다. 저자들은 다른 기형과 태아수종이 동반된 선천성 상기도 폐쇄를 산전에 진단하고 부검으로 확인하여 이를 보고하고자 한다.

중심단어: 선천성 상기도 폐쇄, 산전 초음파검사, 태아수종, 부검