Case Report

Prenatal Diagnosis of Congenital High Airway Obstruction Syndrome: Report of Two Cases and Brief Review of the Literature

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Congenital high airway obstruction syndrome (CHAOS) is the obstruction of the fetal upper airways, which may be partial or complete. It is usually incompatible with life. Prenatal recognition of the disease is quite important due to the recently described management options. We report here two cases of CHAOS due to tracheal atresia diagnosed by antenatal ultrasonography and fetal MRI. We also briefly review the relevant literature with the associated management options.

1. Introduction

Congenital high airway obstruction syndrome (CHAOS) is defined as complete or partial obstruction of the fetal upper airways. This clinical condition was brought into notice firstly by Hedrick in the late 1900s [1]. CHAOS is usually caused by atresia or stenosis of the larynx or trachea. The true incidence of CHAOS is unknown. If the syndrome is unrecognized during the prenatal period, it usually results in stillbirth or death shortly after delivery [2]. Fortunately, more cases can be recognized in utero nowadays, as there are significant technical improvements in prenatal imaging. Bilaterally enlarged hypechogenic lungs, dilated airways, and flattened or inverted diaphragm are the typical prenatal sonographic findings. Fetal ascites and non-immune hydrops may also be associated with the clinical condition [3]. Due to the recently described management options, prenatal definition of fetal airway obstruction has come into prominence with the hope of neonatal outcome improvements [4]. We report here two cases of CHAOS due to tracheal atresia diagnosed by antenatal ultrasonography and fetal MRI.

2. Case 1

A 31-year-old woman with a previous live birth was referred for a routine second trimester antenatal ultrasound at 19-week gestational age. Ultrasound examination revealed that the fetus had bilateral large echogenic lungs. The principal bronchi appeared dilated (Figure 1). The diaphragm was inverted. The heart was centrally placed and seemed to be compressed by the enlarged lungs, and there was moderate ascites. However, amniotic fluid was normal. An MRI study was performed which again showed bilateral large and distinctly hyperintense lungs. Significantly, dilated main bronchi could be clearly seen. Inversion of the diaphragm, the typical small compressed fetal heart, and ascites were also confirmed and no additional anomaly could be found (Figure 2).

Based on ultrasound and MRI findings, diagnosis of CHAOS due to tracheal atresia was made. The parents were counselled regarding the relatively poor prognosis of the syndrome, and the pregnancy was terminated according to the choice of the family. The fetus had severe hydrops manifested by ascites and pericardial and pleural effusions. Postmortem examination was compatible with a complete tracheal atresia.
Figure 1: Antenatal sonography of case 1 showing dilated main bronchi (a), large hyperechoic lungs (b), inverted diaphragm (c), and ascites (d).

Figure 2: Antenatal MRI of case 1 showing dilated airways (arrow), hyperexpanded lungs, inverted diaphragm, and ascites.

Figure 3: Postmortem autopsy findings of case 1 compatible with tracheal atresia (arrow): voluminous lungs and dilated upper airways.

Figure 4: Antenatal sonography of case 2 showing large hyperechoic lungs (a), dilated main bronchi (b), large hyperechoic lungs, inverted diaphragm, and ascites.

as the cause of airway obstruction (Figure 3). The lungs of this fetus were very enlarged (Figure 3).

3. Case 2

A 24-year-old primiparous woman presented for a routine second trimester antenatal ultrasound at 18-week gestational age. Ultrasound examination showed that the fetus had bilateral large hyperechoic lungs. The diaphragm was inverted. The heart seemed small and was centrally displaced. There was massive ascites. The tracheobronchial tree appeared dilated (Figure 4). However, amniotic fluid quantity was normal.

Based on ultrasound findings, diagnosis of CHAOS was made. The parents were counselled regarding the relatively poor prognosis, and the EXIT procedure was offered. However, the parents decided to get an elective termination, and the pregnancy was terminated. The fetus had severe hydrops manifested by anasarca and pericardial and pleural effusions. Postmortem examination was compatible with a complete tracheal atresia as the cause of airway obstruction (Figure 5). The lungs of this fetus were significantly enlarged (Figure 5).

4. Discussion

Tracheal atresia is a very rare congenital malformation which takes place by deficient recanalization of the upper airways around the 10th week of gestation resulting in a clinical spectrum defined as congenital high airway obstruction syndrome (CHAOS) [1]. In a healthy fetus, the fluid secreted by fetal lung is absorbed through the tracheobronchial tree. However, in case of any obstruction in the tracheobronchial tree, this fluid cannot be cleared. The accumulation of the fetal lung fluid results in gradual increase of intratracheal pressure leading to enlargement of the lungs. It is the beginning of a chain reaction: the enlarged lungs cause compression of the heart and great veins. Due to the compression, the heart replaces centrally and becomes small and dysfunctional. Decreased venous return and dysfunctional cardiovascular system end in ascites and hydrops. The diaphragm flattens or inverts according to the severity of the process [2]. Besides tracheal atresia, the other rare underlying causes of CHAOS are laryngeal agenesis, subglottic stenosis or atresia, and laryngeal webs or cysts. However, the obstruction is mostly secondary to laryngeal atresia [1, 4].
malformation, cervical teratoma, and vascular rings like double aortic arch [1, 3].

CHAOS is mostly sporadic, and the exact incidence is not known [1, 2, 7]. In utero death of the affected cases; being a part of some genetic syndromes; better detection rate of the anomalies by the means of technical improvement of imaging tools; and data only from isolated cases instead of studies consisting of large series may explain this indefinite incidence. The most common associated genetic disorder with CHAOS is Fraser’s syndrome which is inherited by autosomal recessive form and characterized by urogenital defects, laryngeal atresia, syndactyly, and cryptophthalmos [2]. CHAOS may be also a part of Cri-du-Chat syndrome, short-rib polydactyly syndrome, and velocardiofacial syndrome [2, 8].

The point to take into consideration is the necessity of a detailed evaluation of all CHAOS suspected cases due to possibility of coexistence of any genetic syndrome and significant implications of inheritance for future pregnancies [8].

In the past, CHAOS was thought to be equivalent to a certain fetal death. However, nowadays, especially if CHAOS due to incomplete obstruction is diagnosed in the late 2nd or in the 3rd trimester and if severe hydrops has not occurred yet, the EXIT procedure (ex utero intrapartum treatment) can be offered. The common objective of the procedure is to settle an intact airway for the baby before the fetomaternal circulation is stopped [9].

There are few cases with spontaneous antenatal improvements. This progress may be due to spontaneous perforation or tracheoesophageal fistula with the refining of the obstructed fluid leading to decrease in the pressure of airways and reversal of the process [10].

As a result, CHAOS is a rare and fatal cause of congenital airway obstruction if unrecognized during prenatal period. Antenatal sonographic imaging shows typical findings which can lead to a diagnosis. MRI is superior to sonography in demonstrating the level of obstruction and in assisting in the differential diagnosis by excluding extrinsic causes of obstruction. This is important especially if any fetal intervention is considered.

Conflict of Interests

The authors declare that there is no conflict of interests.

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