ABSTRACT

Congenital diaphragmatic hernia is a relatively rare birth defect with unknown etiology. Its association with other anomalies and distinct clinical patterns suggest that several causes may be involved. Congenital diaphragmatic hernia occurs in 1 in 2500 live births. In 85% of cases the defect is left-sided [1]. Most cases of congenital diaphragmatic hernia are sporadic and familial congenital diaphragmatic hernias rare, comprising only 2% of congenital diaphragmatic hernias [2]. This congenital anomaly can almost always be recognized with prenatal ultrasound screening. There is a high degree of variability in both treatment and outcomes. Bilateral congenital diaphragmatic hernia is a rare birth defect, with grim prognosis. We describe a case of bilateral congenital diaphragmatic hernia discovered while repartitioning right sided congenital diaphragmatic hernia. The diaphragmatic defect was repaired and a prolene mesh was placed on the abdominal wound to avoid abdominal compartment syndrome. The patient nonetheless died post operatively due to severe pulmonary hypertension. Bilateral congenital diaphragmatic hernia, priorly identified through a limited number of case reports, is extremely rare. The care of congenital diaphragmatic hernia patients is very difficult for neonatologists and surgeons. Our report particularly the management and outcome of patients with bilateral congenital diaphragmatic hernia.

Keywords: Bilateral congenital diaphragmatic hernia, Pulmonary hypertension.

Bilateral Konjenital Diyafragmatik Herni: Nadir Bir Olgu

ÖZET


Anahtar Kelimeler: Bilateral konjenital diyagrafma hernisi, Pulmoner hipertansiyon.
INTRODUCTION

Congenital diaphragmatic hernia (CDH) occurs in every 2500-3000 live births [1-3]. Bilateral CDH is extremely rare, comprising less than 1% of all CDH cases [4]. Most of them die in utero while less than 35% survive [1-5]. The administration of bilateral CDH is a challenge for pediatric surgeons and the neonatologists. We report a case of bilateral CDH with sac on both sides and discuss the literature.

CASE REPORT

The patient was admitted to our clinic with a 15th gestational week ultrasound revealed the stomach in the thorax next to the fetal heart (Figure 1). The family was informed. Advanced ultrasound was performed and demonstrated bilateral diaphragmatic hernia. At the 29th gestational week, the patient was reinspected and the fetal heart was deviated to the right. Growth retardation was present in the fetus. At the 36th week, the heart of the fetus was severely deviated to the right. The left lobe of the liver was viewed in the thorax (Figure 2). The fetus had severe growth retardation. The patient was called for weekly visits. The risks were explained to the family on week 39 and cesarean section was performed. The baby had birth asphyxia and was resuscitated with bag and mask ventilation. He was a 2600 grams male fetus with APGAR scores 5 and 4 at the first and fifth minutes, respectively. The patient was kept in the intensive care unit under oxygen and feeds were started. On admission his respiratory rate was 70/min, $\text{SpO}_2$ was 87% while receiving oxygen. Chest X-ray observed in both lungs collapsed (Figure 3).

The fetus under went urgent surgical operation. The thorax was entered via left subcostal incision. Spleen, stomach, small intestine, left lobe of the liver and the leftcolon were observed in the thorax. The organs were with drawn from the abdominal cavity and a bilateral wide diaphragmatic hernia was observed. A chest

Figure 1: 15th pregnancy week diaphragmatic hernia image

Figure 2: 36th pregnancy week diaphragmatic hernia image

Figure 3: Preoperative chest radiograph

Figure 4: Postoperative chest radiograph
tube was applied to the left thorax. Underwater drainage was performed. Half of the primary and lateral half of the defect was closed using prolene mesh. The newborn had intestinal malrotation which was closed in accordance with the anatomy of the abdomen. Postoperatively the baby was electrolytically ventilated at rates of 40 per minute and FiO\(_2\) of 85%. The patient, however, deteriorated and developed advanced severe pulmonary hypertension. Although the lung was moderately expanded on a postoperative chest radiograph (Fig. 4), the child went into refractory shock and died 56 hours after surgery.

**DISCUSSION**

The prognosis of CDH is mainly determined by the degree of pulmonary hypoplasia. CDH including the neonatal period is mostly of the Bochdalek type. The defect is usually unilateral and involves the left diaphragm in 75% of cases [6]. Conversely, bilateral CDH is rare and usually fatal. Right-sided defects with delayed diagnoses may cause a wide spectrum of clinical signs [4]. Delayed presentation of right-sided CDH in association with neonatal group B streptococcal pneumonia is a well-recognized entity [4-5]. Antenatal diagnosis of CDH is relatively easy and the sonographic signs of unilateral CDH are mediastinal shift, presence of intestine in the thorax, small abdominal circumference and polyhydramnios sufficiency [1-10]. Mediastinal shift is often the first abnormality observed. In our case the diagnosis was made during week 15th. Song et al [2] recorded antenatal finding of bilateral CDH in a fetus and concluded that bilateral diaphragmatic hernia should be suspected when the degree of mediastinal shift is less than that guessed for a unilateral hernia in existence of other features of CDH.

An identifiable cause is present in about 15–20% of individuals with diaphragm defects [6]. A chromosome abnormality is present in about 10% of all individuals with CDH. The most common abnormalities are trisomy 18 and tetrasomy 12p (Pallister–Killian syndrome). Many additional abnormalities have been reported, such as trisomy 21, trisomy 13, tetrasomy 21, trisomy 8 and other structural abnormalities. Small rearrangements found in unrelated individuals have suggested that one or more genes significant for normal diaphragm development may reside in critical regions such as 15q26.2 [6-10]. In a study made by Furuta et al [3] reported the 11th recorded case of bilateral CDH. The patient was lost in the postoperative period after repair of right CDH while the left side CDH was identified at autopsy. Our case was detected at week 15 since the patient came for her first visit at this time. The intraterine diagnosis was confirmed during operation of the newborn. We favored an abdominal approach since part of the liver had herniated through a large defect in the diaphragm. Another advantage of the abdominal approach is that a prolene mesh can be used. In a study made by Kufjei and Crabbe reported familial CDH in two siblings with a similar type of bilateral CDH in 1999 [7]. Both cases could not be diagnosed antenatally and their outcomes were severe. In a study made by 2003 Neville et al retrospectively reviewed cases of CDH treated in 83 different hospitals in USA.

They found that in bilateral CDH mortality rate was 65% compared with 33% of patients with unilateral CDH [8]. In contrast, reports of patients with favorable outcome have also been published in literature. In 2007 Zaupa et al recorded a case of bilateral CDH with gastroschisis with good outcome [9]. They suggested that a low intrathoracic pressure due to gastroschisis may have been allowed for good lung development and better patient outcome. In our case although the preoperative stabilization was well, there was severe pulmonary hypertension that led to the mortal outcome. In conclusion, although early diagnosis and early intervention was under taken in our patient, the bilateral hernia remained mortal.

**CONCLUSION**

The management of infants with bilateral congenital diaphragmatic hernia remains a difficult problem with significant mortality. Compared to unilateral CDH, bilateral congenital diaphragmatic hernia is more frequently associated with other major anomalies and should immediate further evaluation.
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