CASE REPORT

Right Side Congenital Diaphragmatic Hernia and Hydrops: May Not Be Lethal

Maha Y. Bamehriz, MD, SSC-Ped
Department of Pediatric, Faculty of Medicine, King Abdulaziz University
Jeddah, Saudi Arabia

Correspondence
Dr. Maha Yaslem Bamehriz
Department of Pediatric
Faculty of Medicine, King Abdulaziz University
P.O. Box 80215, Jeddah 21589
Saudi Arabia
e.M: mbamehrez@kau.edu.sa

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Abstract
Right sided congenital diaphragmatic hernia associated with hydrops is a rare event and carries a poor prognosis. We report here a case of right side congenital diaphragmatic hernia associated with hydrops which was diagnosed antenatally and survived to discharge with good outcome.

Keywords
Congenital diaphragmatic hernia; Right-side congenital diaphragmatic hernia; Non-immune hydrops fetalis

Introduction
Congenital diaphragmatic hernia (CDH) carries a high risk for neonatal mortality and morbidities[1].

Right sided congenital diaphragmatic hernia (RT-CDH) with presence of the liver protruded into the chest cavity occurs less frequently than left side[2,3], and holds a worse outcome especially when associated with fetal thoracic anomalies or hydrops. Hydrops is defined as the presence of skin and scalp edema or fluid in two or more body compartments.

Case Report
A 34-year-old, G8 P3+4 mother, was referred at 29 weeks to King Abdulaziz University Hospital for detailed ultrasound evaluation and management. Fetal ultrasound was done in the fetomaternal unit and showed a large right side diaphragmatic hernia with a large portion of the liver protruding into the chest cavity and the heart was pushed to the left side. There was a significant hydrops with ascites, pleural effusion and hydrocele. Other organs were normal (Fig. 1). Amniotic fluid reduction was done at that time. Repeated ultrasound after three weeks showed progressive re-accumulation of fluid. Fetal echocardiogram and fetal Doppler showed normal fetal cardiac anatomy with pericardial effusion. Toxoplasma, syphilis, rubella cytomegalovirus, herpes (TORCH) serology was negative.

A male infant of 3800 gm was delivered by cesarean section at 35+5 weeks’ gestation due to preterm labor. Apgar score was 1,5,7 at 1,5,10 minutes, respectively. On
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Figure 1. Fetal ultrasound showed pleural effusion, ascites and liver in the chest.

Figure 2. Chest X-ray showing right side diaphragmatic hernia.

Figure 3. Chest X-ray showing postoperative right-side lung inflation.
examination, there was generalized body edema, and no dysmorphic features suggestive of any syndromes. The initial cord gas showed pH 7.19, pCO2 68 mm Hg, and HCO3 17.8 mmol/L. He was intubated immediately after birth and transferred to neonatal intensive care unit (NICU) for further management.

In the NICU the patient required escalation of mechanical ventilation from conventional to high frequency oscillation ventilation to manage oxygenation failure. Chest X-ray confirmed the antenatal findings (Fig. 2). Echocardiogram was performed and revealed severe pulmonary hypertension, with moderately dilated right ventricle and normal function. The patient received inotropic support for systemic hypotension and Milrinone infusion of 0.5 mcg/kg/hr that was increased to 0.75 mcg/kg/hr, followed by Sildenafil orally for severe pulmonary hypertension. Inotropes were weaned gradually and stopped by day 3 of life. Milrinone was weaned off and patient was continued on sildenafil. Skin edema improved gradually. At age 24 days of life patient underwent corrective surgery where the surgeon replaced the liver back into the abdominal cavity and closed the defect with prolene with no patch used. Intra-operative the antenatal diagnosis was confirmed. The operation was tolerated very well by the patient (Fig. 3). Three days post-operatively, feeding was started, and full feeding was reached on day 50 of life. The patient was extubated to non-invasive ventilation at age 39 days. The patient was discharged home at age 63 days of life on room air and full oral feeding.

Discussion

Congenital diaphragmatic hernia (CDH) is a defect in the diaphragm that allows the movements of abdominal organs into the thoracic cavity. The reported incidence of CDH is 1 in 2500-4000 live birth[1]. Right side diaphragmatic hernia (RT-CDH) occurs less frequently than Left side diaphragmatic hernia (LT-CDH) with an incidence ranging between 14%-25% of all CDH[2,4]. Prenatal diagnosis is less common in RT-CDH compared to LT-CDH due to similar echogenicity of both liver and lung[2-5] and late presentation.

Hydrops associated with CDH is a rare event. It appears that it is more common with RT-CDH and carries a poor prognosis, as presence of the liver in the thorax may obstruct the inferior vena cava and/or other venous inflow. Additionally, the presence of the liver in the thoracic cavity may lead to impairment in the lymphatic flow leading to hydrothorax and lethal pulmonary hypoplasia[6,7]. Sydorak et al.[8] speculated the cause of the hydrops is secondary to venous compression by herniated liver. Another possible cause reported by Narvey and Soni[9] is the direct hepatic compression on the right and left atrium causing decrease in both systemic and pulmonary venous return and causing non-immune hydrops.

Burgos from the Congenital Diaphragmatic Study Group included 3754 infants reported no difference in gender rate, birth weight, gestational age at birth, delivery mood or the rates of associated major cardiac or chromosomal anomalies between RT-CDH and LT–CDH[2,4,10].

In a retrospective chart review of 330 CDH cases, it was found that in comparison to LT-CDH, RT-CDH has stronger association with pulmonary morbidities secondary to severe pulmonary hypoplasia. Such morbidities include longer duration of nitric oxide therapy, increased requirement of tracheostomy, need for oxygen therapy at discharge, and increased chronic pulmonary hypertension with requirement for long-term sildenafil therapy all of which delays repair of the hernia[11].

Mortality rate reported to be either higher in RT sided defect[2], or similar[3,4] to the mortality rate in LT sided defect.

Our patient was diagnosed antenatally at 29 weeks of gestation to have RT-CDH associated with hydrops, born as a preterm, had severe pulmonary hypertension which delayed his operation to repair the diaphragmatic defect until day 24 of life which is consistent with another study[3], oral feeding was started within three days post-operative which is significantly shorter than has been reported previously by Partridge et al[3]. Patient was discharged home at age 63 days, which is consistent with the other report, on full oral feed and on no respiratory support.

Conclusion

In cases of RT-CDH associated with hydrops, prenatal diagnosis combined with optimum respiratory support postnatally in the absence of major congenital anomalies is considered a major contributing factor for survival.
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M.Y. Bamehriz

Conflict of Interest
The author has no conflict of interest.

Disclosure
The author did not receive any type of commercial support either in forms of compensation or financial for this report. The author has no financial interest in any of the products or devices, or drugs mentioned in this article.

Ethical Approval
Obtained.

References
وفق الحجاب الحاجز الخلقي للجهة اليمنى المصاحب للاستسقاء: قد لا يسبب الوفاة

مها يسالم أحمد بامهرز
قسم طب الأطفال، كلية الطب
جامعة الملك عبد العزيز
جدة - المملكة العربية السعودية

المستعرض. فثق الحجاب الحاجز الخلقي للجهة اليمنى المصاحب للاستسقاء هي حالة نادرة لها تبعات غير حميدة. وها نحن نقوم بتسجيل الحالة والتي تم تشخيصها خلال فترة الحمل وقد خرج المريض من المستشفى في حالة جيدة.