Case Report

Hydrops fetalis secondary to supradiaphragmatic extrapulmonary sequestration with congenital cystic adenomatoid malformation

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ABSTRACT

Congenital cystic adenomatoid malformation (CCAM) is a hamartomatous lesion. A 30 year old woman, G2P1L1, in the 29th week of gestation presented with pain abdomen, chest pain, cough with expectoration, fever and inability to appreciate fetal movements of 2 days duration. Clinically, she had pneumonia. An ultrasound revealed a single fetus in breech presentation with features of hydrops fetalis, hypoplastic left lung, mediastinal shift to the left and poor diastolic and systolic flow in the umbilical artery.

A still born male baby delivered subsequently was found at autopsy to have hydrops fetalis, supradiaphragmatic right sided extralobar sequestration (ELS) with associated congenital cystic adenomatoid malformation (CCAM). The right lung also showed CCAM. There were no other associated anomalies. We present a rare case of ELS with CCAM.

INTRODUCTION

Congenital cystic adenomatoid malformation (CCAM) is a hamartomatous lesion characterized by abnormal bronchial and bronchiolar-like structures of varying sizes and distribution. It is associated with 34.4% stillbirths. Non-immune hydrops fetalis has been described in association with numerous conditions of the fetus in-utero. Thoracic pathology commonly associated with hydrops include congenital cystic adenomatoid malformation of the lung (CCAM) (30%), Chondrodysplasia (25%), right sided diaphragmatic hernia (15%), pulmonary sequestration (8%), etc. Extra-lobar pulmonary sequestration (ELS) is defined as the presence of a mass of abnormal pulmonary tissue outside the visceral pleura, that does not communicate with the tracheobronchial tree and is supplied by an anomalous systemic artery. Congenital cystic adenomatoid malformation is a hamartomatous lesion characterized by the presence of abnormal bronchial and bronchial-like structures of varying sizes and distribution. In a series of 15 cases reviewed by Stocker and Kagan-Hallet, one case contained CCAM with extra-lobar pulmonary sequestration. Subsequently, Zangwill and Stocker described a case of CCAM with an ELS and stated that association occurred.
in approximately 25% of cases accessioned to the Armed Forces Institute of Pathology. Conran and Stocker et al studied a series of 50 cases of ELS and found associated CCAM in 50%. We present one such case in a 30 year old lady who presented with intrauterine fetal death.

CASE REPORT

A 30 year old lady G2P1L1 in the 29th week of gestation, presented with pain abdomen, cough with expectoration of 2 weeks and fever with inability to appreciate fetal movements of 2 days. Chest X-ray and laboratory investigations confirmed lobar pneumonia. USG abdomen revealed a single fetus in breech position with features of hydrops fetalis and hypoplastic left lung. There was no evidence of blood group incompatibility. She delivered a still born, male baby weighing 1900gm with features of hydrops fetalis. An autopsy was performed.

On examination of the thoracic cavity revealed a hypoplastic right lung, large supra-diaphragmatic extra-pulmonary mass attached by a thin stalk to the mediastinal tissue, mediastinal shift to the left. Cut section of both the lung and extrapulmonary mass was homogenous, pale white with numerous randomly distributed cysts 0.1 to 0.5cm in diameter.(fig. 1and 2). The histopathology of the extra pulmonary mass showed a fibrous capsule with underlying dilated lymphatics and peripheral lung parenchyma with dilated acini and interspersed bronchioles. The rest of the parenchyma showed multiple cystically dilated spaces and bronchiolar-like structures arranged in a back to back configuration. These structures were lined by cuboidal to columnar epithelium and separated by thin fibrovascular septae. The right lung showed compressed lung parenchyma and similar cysts as described above. The left lung and other viscera were normal. (fig. 3A and 3B)

Based on these features the diagnosis of nonimmune hydrops fetalis with ELS with associated congenital cystic adenomatoid malformation, type – 2 and right pulmonary hypoplasia was rendered.

DISCUSSION

Congenital cystic adenomatoid malformation of the lung is a rare condition occurring in 1 in 25,000-35,000 pregnancies. At the time of prenatal USG, there may already be evidence of considerable fetal distress, including non-immune hydrops fetalis (NIHF) in these cases (12.43%). Congenital cystic adenomatoid malformation is further subdivided into 5 pathologic types. Type 0 or acinar dysplasia is characterized by firm and small lungs. It is incompatible with life and is seen in association with cardiovascular anomalies and dermal hypoplasia. Type 1 has multiple large cysts or a
single dominant cyst with surrounding smaller cysts. The lining of the cysts is ciliated columnar to pseudostratified ciliated columnar epithelium. Type 2 is composed of multiple evenly distributed cysts that rarely exceed 1.2cm in diameter lined by ciliated cuboidal to columnar epithelium. Type 2 lesion is seen in association with extralobar sequestration in 40% of the cases. Type 3 lesions present with mediastinal shift and large bulky masses with small cysts. Type 4 CCAM known as the peripheral cyst type is distal acinar in origin. The common clinical presentations of CCAM are stillbirth or neonatal death associated with anasarca, prematurity and maternal polyhydraminos, acute respiratory distress in the newborn or an indolent course with recurrent pulmonary infections. Congenital cystic adenomatoid malformation can cause considerable morbidity and mortality secondary to pulmonary hypoplasia (PH) because of lung compression and NIHF due to compression of the heart and great vessels. The presence of hydrops has emerged as the single most important poor prognostic indicator in fetuses with congenital cystic adenomatoid malformations. The co-existence of CCAM in ELS has been rarely reported. Extralobar sequestration cases containing back-to-back cystic structures of varying sizes lined by bronchiolar type epithelium are classified as CCAM arising in an extralobar sequestration. The age at diagnosis ranges from prenatal period to 73 years. There is an equal gender distribution with 74% being left sided lesions. Whereas ELS can be readily distinguished from intralobar sequestration by the complete separation of ELS from the normal lung, the presence of CCAM within on ELS requires careful microscopic examination to identify the features mentioned above. CCAM in association with ELS is invariably of the type 2 morphology. Thus, ELS, a developmental anomaly of the lung seen in the thoracic spaces, anterior and posterior mediastinum, and within the diaphragm, may contain a second anomaly, CCAM usually type 2, in up to 50% cases.

REFERENCES


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