Antenatal and postnatal management of congenital cystic adenomatoid lung malformation diagnosed by ultrasound and Magnetic Resonance Imaging (MRI)

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ABSTRACT

Purpose: Antenatal diagnosis of congenital cystic adenomatoid lung malformation (CCAM) is vital for disease surveillance and postnatal care. Ultrasonography (US) has been the imaging gold standard for antenatal CCAM assessment. However, one of the limitations of US is the “vanishing phenomenon” caused by isoechogenicity of CCAM tissue and adjacent normal lung parenchyma. Methods: Antenatal serial US were concurrently used with magnetic resonance imaging (MRI) to monitor macro- and microcystic lesions. Results: In both pregnant women, antenatal US and MRI confirmed the presence, in the fetus, of cystic lesions and predicted disease regression/progression as well as the need for postnatal surgical intervention. Several advantages were detected by using both—serial US and MRI (over serial US alone)—including improved signal intensity, exact volume size measurements, precise CCAM location in particular for patients with adverse ultrasound conditions. Both neonates underwent surgical resection and had an uneventful post-operative course. Conclusions: Antenatal use of MRI as well as serial US improved information regarding tissue resolution and delineation of CCAM. The information from two imaging modalities was complementary. Our literature review confirmed the emerging role of prenatal MRI for postnatal monitoring and management of CCAM.

Keywords: Congenital Cystic Adenomatoid Lung Malformation; Antenatal Serial Ultrasound Studies; Antenatal MRI; Imaging Surveillance

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1. INTRODUCTION

Congenital cystic adenomatoid lung malformation (CCAM) is characterized by various cystic intrathoracic masses caused by proliferative terminal respiratory bronchioles and by a reduction in the number of normal alveoli. It usually affects a single pulmonary lobe; multi-lobular or bilateral involvement is rare [1]. CCAM was first reported in 1949 by Ch’in and Tang [2]. The estimated incidence of CCAM ranges from 1:10,000 to 1:35,000 live births [3-5], with male and female neonates equally affected [4]. The possible influences of genes, such as HOXB5, Fdf7, and PDGF-BB, have been described, but, as of now, are not thought to be of any diagnostic or prognostic value [5,6]. The histopathologic classification by Stocker et al. (1977) describes 3 forms of CCAM (type I-III). Type I is the most common form, detected in up to 65% of patients with CCAM: the single or multiple cysts are more than 20 mm in diameter and are lined by ciliated columnar or pseudostratified epithelium, mucus-secreting cells, and, perhaps prominently, cartilage. Type II is the second most common form, detected in up to 20% to 35% of patients with CCAM: the multiple cysts are less than 20 mm in diameter and are lined by cylindrical or cuboidal epithelium, with prominent smooth or striated muscle. Type III is the least common form, detected in up to 10% of patients with CCAM: the solid microcysts are rarely larger than 2 mm in diameter and are lined by cuboidal epithelium, often intricately folded [1]. CCAM ranges in severity from intrauterine regression to progressive nonimmune hydrops and/or mediastinal shifting. Treatment options include prenatal intrauterine laser therapy, postnatal thoracoscopic local excision, or thoracotomy with appropriate surgical resec-
tion. Without treatment, macrocystic lesions, in particular, tend to increase the risk of pulmonary infections, of mediastinal shifting with lung compression, and of malignancy late in life [7-10].

The earliest reported diagnosis of CCAM with ultrasonography (US) was made in 1975 [11]. The sonographic appearance of CCAM was classified by Adzick et al., who distinguished between subgroups of macrocystic (≥5 mm in diameter) and microcystic (<5 mm in diameter) CCAM [12]. Serial antenatal US has been considered the imaging gold standard for antenatal monitoring and management. The disadvantages of US in particular regarding unfavorable ultrasound conditions (i.e. elevated body mass [BMI] indices), however, include imprecise measurement of exact volume size, reduction in signal intensity, and need for a high number of serial imaging studies [13-16].

In contrast, the advantages of magnetic resonance imaging (MRI) especially in cases of attenuate ultrasound signals include exact measurement of the lesion(s)-to-lung volume ratio, superior tissue resolution, with determination of disease progression or regression and precise determination of CCAM location, and need for a low number of studies [17-19].

US and MRI are complementary. Therefore, we used both modalities, in combination, in our two patients; then, we compared our observations with the current literature. Ethical clearance was obtained from the relevant office and experiments conducted in accordance with the Helsinki declaration of 1982 regarding use of human subjects.

2. METHODS AND RESULTS

2.1. Case 1

A 29-year-old woman (primigravida) was referred, at 21 weeks of gestation, to our tertiary center for a comprehensive US study to assess fetal chest masses. Earlier in her pregnancy, echocardiography excluded any associated anomalies; amniocentesis excluded chromosomal pathology (46 XX). We diagnosed CCAM in the right lung of the fetus with transabdominal sonography, using Acuson 128 XP/10 (Acuson, Mountain View, CA, USA) and Sonoline G 60 (Siemens Medical Systems, Inc., Issaquah, WA, USA) systems, both equipped with a probe of 3.5 to 5 MHz. Our initial US study revealed multiple macro- and micro-cystic malformations, 5 to 12 mm in diameter, in the right lung (Figure 1(a)). Adverse ultrasound conditions caused by maternal increased BMI let to fast $T_2$-weighted MRI (acquisition time, about 10 seconds, HASTE sequence) at 21 weeks of gestation; we used a Multiarray GyroScan 1.5-T MRI unit (Philips, Germany). Our MRI study not only confirmed the initial US diagnosis but also provided superior tissue resolution and revealed the exact location of the lesions (lower lobe of the right lung; Figures 1(b) and (c)).

Our US follow-up studies also confirmed the primary location (lower lobe of the right lung) and subsequent CCAM regression. During the course of gestation, our serial prenatal US studies excluded the presence of hydrops, of mediastinal shifting, of polyhydramnios, and of other anomalies. Spontaneous premature delivery occurred at 35 weeks of gestation (the neonate’s birth weight, 3160 gm; Apgar score, 9/10/10; pH value, 7.39). Initially, the neonate had no respiratory symptoms of concern. For postnatal monitoring, the neonate was transferred to the neonatal intensive care unit, given its capability for extracorporeal membrane oxygenation (ECMO) and intensive pediatric surgical care.

![Figure 1. (a) Sonographic feature of the chest 21 weeks of gestation; (b)-(c) MR images of the macro- and micro-cystic chest masses.](image-url)
The postnatal chest computed tomography (CT) scan confirmed the diagnosis of macro- and micro-cystic lung lesions of the lower lobe of the right lung. The initially asymptomatic neonate developed clinical signs of CCAM during the first week of life, including tachypnoe and dyspnea. To prevent the risk of recurrent infection and late malignant transformation, we performed an elective posterolateral thoracotomy, with resection of the lower lobe of the right lung, when the neonate was 11 days old (Figure 2(a)).

Histopathologic examination confirmed the diagnosis of CCAM type II (Figure 2(b)). The neonate’s postoperative course was uneventful.

2.2. Case 2

A 24-year-old woman (gravida II, para I) was referred, at 30 weeks of gestation, to our tertiary center for comprehensive US screening. Earlier in her pregnancy, elective amniocentesis showed no chromosomal pathology (46 XX). Our initial US study showed a single macrocystic lesion, 20 mm in diameter, in the left lung of the fetus (Figure 3(a)).

After detection of CCAM, we performed an extensive sonographic examination, including fetal echocardiography, which excluded fetal cardiac failure and other associated anomalies. Inappropriate terms of ultrasound caused by reduced amniotic fluid volume let to fast T<sub>2</sub>-weighted MRI at 30 weeks of gestation; we used a Multiarray GyroScan 1.5-T MRI unit (Philips). Again, our MRI study not only confirmed the initial US diagnosis but also provided superior tissue resolution and revealed the exact location of the lesion (lower lobe of the left lung; Figure 3(b)); concurrent microcystic lesions in other parts of the lungs were not present. During the course of gestation, our serial prenatal US studies revealed a decrease in the size of the single cystic malformation; at term, it was 16 mm in diameter. Our follow-up US studies confirmed the location of the macrocystic lesion (lower lobe of the left lung) and excluded the presence of hydrops, of mediastinal shifting, and of polyhydramnios. At 40 weeks of gestation, the woman underwent a cesarean section (the neonate’s birth weight, 3510 gm; Apgar score, 8/8/10; pH value 7.31). A postnatal chest CT scan confirmed the diagnosis of a single macrocystic lesion in the lower lobe of the left lung.

The primary asymptomatic condition of the neonate changed within the first three days of life, when clinical signs of infection developed. Five days after delivery, before serious respiratory infection and/or dysfunction.
could occur, the neonate underwent elective segmentectomy of the lower lobe of the left lung. Histopathologic examination confirmed the diagnosis of CCAM type I. The neonate’s postoperative course was uneventful.

3. DISCUSSION

CCAM is the most common (77%) of congenital cystic lung diseases, followed by pulmonary sequestration, bronchogenic cysts, and bronchial atresia [7-9,20]. The disease spectrum of CCAM is quite variable. Lesions with spontaneous regression and, ultimately, complete disappearance during the third trimester have been reported; so have rapidly growing lung masses with non-immune hydrops and in utero demise [7,16,20]. Close perinatal monitoring is essential to optimize outcome [7-9,16,20-22]. Serial antenatal US studies have been the imaging gold standard to characterize the features of CCAM, including the size of lesions, their appearance (macro- vs micro-cystic), and their location (lobe involvement) [13-16,23]. However, US studies, even when done serially, have disadvantages, including less than ideal CCAM resolution and imprecise initial CCAM location, particularly in obese pregnant women [13-16].

Other imaging modalities have hardly been used in the prenatal period. Antenatal CT studies are not indicated, because of the amount of irradiation. In contrast, MRI, particularly in the third trimester, appears to be relatively safe and provides excellent tissue resolution [17,18]. Of note, MRI was used in one of our cases as early as 21 weeks into the pregnancy, i.e. at an early stage of gestation. MRI also distinguishes CCAM from other chest lesions, clearly identifies the exact lung location, and demonstrates the presence of compressed normal lung [10,16]. Thus, antenatal MRI enables clear identification of high-risk fetuses and may help optimize postnatal intensive care. In addition, quantification and assessment of pulmonary hypoplasia by antenatal MRI can offer insights about such options as termination of pregnancy, elective delivery, and planned postnatal surgical procedures [16,23].

In our experience, especially in cases of unfavorable terms of US, information provided by the initial MRI study was superior to the initial US study with regard to tissue resolution and precise CCAM location. The advantage of serial US studies lies in the documentation of morphologic changes (e.g., size, appearance) over time. Therefore, we believe that in cases of attenuate ultrasound conditions the combination of an initial MRI study and serial US studies provides the best information regarding changes in CCAM appearance.

To date, despite the use of serial prenatal US studies to monitor the lesions of CCAM, no standard predictors for disease progression or regression have been pinpointed. It remains to be determined whether or not a second MRI study, during or close to the end of the third trimester, will help pinpoint any such predictors, which are essential to optimize outcome [17,18]. As mentioned above, the clinical symptoms of neonates with CCAM vary in severity. Neonates with severe and symptomatic cases, including in utero development of nonimmune hydrops or polyhydramnios require ventilatory support, and sometimes high-frequency ventilation or even ECMO shortly after birth because of hypoplasia of the lungs [24,25]. Initially asymptomatic neonates with isolated CCAM, in the absence of hydrops, may develop, shortly after birth, recurrent respiratory tract infections (10%) [15,23]. In addition, pleuropulmonary blastoma is a risk in childhood, and bronchoalveolar carcinoma is a risk in adolescence [20,21]. In contrast, nonhydrionic fetuses with isolated CCAM have little or no hypoplasia of the lungs and, therefore, a much better prognosis [26-29]. Irrespective of the absence or presence of symptoms, close prenatal surveillance, carefully planned delivery, and rigorous postnatal evaluation at a facility with ECMO capability and pediatric surgical care all contribute to favorable outcome [21-25]. Aggressive in utero management, using both thoracoamniotic shunting (for macrocystic malformations) and intruterine laser therapy or fetal surgery (for microcystic solid malformations), has been proposed in cases with a poor prognosis [7-10]. However, compared to noninterventional obstetric care, open fetal surgery has failed to improve survival rates [7,13]. Antenatal high-risk CCAM fetuses (with either mediastinal shifting or progressive non-immune hydrops) and symptomatic CCAM neonates (with respiratory insufficiency) require immediate postnatal emergency surgery [7-10, 25].

In contrast, treatment of asymptomatic lesions is controversial. Simple monitoring of asymptomatic CCAM has been suggested, but monitoring carries the risk of ongoing and recurrent pulmonary infections [26]. Although early surgical resection (within the first 4 weeks of life) is associated with an increased anesthetic risk, it maximizes compensatory lung growth at this early stage [9,26,28]. A consensus seems to be growing against the simple monitoring “wait and see” approach in cases of asymptomatic CCAM, because postnatal spontaneous resolution may not occur. Elective resection within the first 3 to 6 months of life has also been suggested, given the lifelong risk of recurrent lung infections and occult malignant lung cell proliferation [3,10,25-27]. The youngest described patient with malignant malformation due to CCAM was 13 months old [30]. Elective surgery is usually well tolerated, the risk of infection is lower, and compensatory lung growth with normal long-term respiratory function has been observed [13,20]. Despite current limitations, prognostic information is necessary to provide parental counseling throughout the pregnancy.
and appropriate perinatal management [7,8,27,31]. In hydropic CCAM fetuses, the presence of additional life-threatening anomalies might lead to the decision to terminate the pregnancy. Prognosis is very favorable in fetuses without hydrops and with isolated, unilateral CCAM [13,14,20,25]. After extensive parental counseling, a noninvasive obstetric strategy should be adopted, including serial US studies and an initial antenatal MRI study. The value of a subsequent MRI study in the third trimester needs to be determined [17,18].

4. CONCLUSION

In summary, CCAM of the fetal lung remains a challenge for obstetricians, neonatologists, and pediatric surgeons. Improvements in antenatal monitoring are necessary to improve and better predict outcomes. The combination of prenatal MRI and serial US studies appears to optimize fetal surveillance and postnatal care.

REFERENCES


