CLINICAL STUDY

Routine betamethasone therapy of non-hydrophic fetuses with CPAM – the way to improve perinatal outcome?


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ABSTRACT

Authors discuss novel approach to the management of fetal congenital pulmonary cystic malformation (CPAM) and possible benefit of routine administration of betamethasone, which is currently recommended only for severe cases. The article presents authors’ own experience with antenatally diagnosed CPAM and describes 4 cases of prenatally diagnosed CPAM without hydrops treated by two doses of betamethasone at 21–31 weeks of gestation with the aim of improving the perinatal prognosis by effect on not only mortality but also postnatal morbidity. Article also summarizes current knowledge on all aspects of the prenatal CPAM focusing on its treatment options. Data were obtained from the literature search based on the PubMed and Scopus database with additional search of particular articles from reference list of the selected publications. All 4 patients from the case reports showed regression of the pulmonary mass after maternal administration of betamethasone with 100 % survival rate. Even though it is not possible to say if the CPAM regression was due to the betamethasone treatment, we did not observe any complication in relation to the treatment and all cases survived until discharge. During the literature search, we did not find any data on betamethasone administration in non-hydrophic fetuses with CPAM in relation to the overall perinatal and postnatal morbidity, neither data comparing the outcome between the treated versus observed only fetuses. Routine betamethasone treatment should be discussed in antenatally diagnosed CPAM cases without fetal hydrops in order to reduce the perinatal morbidity associated with CPAM (Tab. 1, Ref. 47). Text in PDF www.elis.sk.

KEY WORDS: betamethasone, CPAM (congenital pulmonary adenomatoid malformation), fetal therapy.

Introduction

Congenital pulmonary cystic malformation (CPAM), previously known as cystic adenomatoid malformation (CCAM), refers to benign multicystic mass of nonfunctioning pulmonary tissue that is usually restricted to one lobe of the lung (1). It represents an abnormality in lung development characterized by the lack of normal alveoli and abnormal proliferation of terminal respiratory bronchioles (2). The exact etiology remains unknown. Histopathology confirms bronchial atresia with absence of normal alveoli due to decreased apoptosis during the embryological development between 7 and 15 weeks of gestation (3). CPAM is usually isolated and sporadic, rarely it can be associated with other anomalies, such as cardiac anomalies, renal agenesis or dysgenesis, gastrointestinal atresia or skeletal anomalies associated particularly with the microcystic form of CPAM (4). Prognosis depends on the CPAM size rather than lesion type. However, natural history of the lesion can differ, as some may come to spontaneous regression, others stagnate or progress to grow (5). Generally, the CPAM is usually highly unpredictable in its growth potential between 18 and 26 weeks of gestation (gw). The fastest growth appears between 20 and 25 weeks with a peak of CVR (CPAM volume ratio) occurring at 25gw (6). There is a plateau in CPAM growth beginning at 25 gw with a decrease in the CVR after 25 weeks gestation reflecting continued fetal growth (7).

CPAM is responsible for 25–30 % of congenital lung malformations and is the most common prenatally diagnosed cystic lung disease (8). Incidence ranges from 1:11 000 to 1:35 000 pregnancies and live births, male fetuses are more likely to be affected (9). The reported perinatal mortality of antenatally diagnosed CPAM has decreasing trend over the world, ranging from 49 % in 1992 to 9 % in 2001 (8, 10).

Mortality results mainly from the pulmonary hypoplasia (11). Postnatally is CPAM associated with moderate respiratory symptoms during childhood, persistence of the lesion can cause recurrent infections and even its evolution into malignant processes cannot be excluded (12, 13).

The original classification of CPAM according to Stocker et al (14), dividing CPAM lesions into three groups, has been recently extended to five types (0–IV) based on the histopathological image and localization of the malformations at various levels of the airway and lung (15–17). Type 0 CPAM is characterized by bronchial-type airways separated only by abundant mesenchymal...
tissue. Types 1 (the bronchial/bronchiolar type) and 2 CPAM (the bronchiolar type) are characterized by cysts > 2 cm in diameter and multiple small cysts, respectively. In type 3 CPAM (the bronchiolar/alveolar type) lesion is solid and not cystic, while type 4 CPAM (the peripheral type) is characterized by peripheral thin-wall, often multiloculated cysts (15, 18). Since the classification is based on the histopathology, other classification of Adzick (19) has become widespread in clinical practice as it is based on ultrasound image and gross anatomy. Adzick et al (19) divides CPAM into two major categories according to the predominant component of the lesion. The macrocystic form of CPAM includes cyst(s) > 5 mm in diameter while the microcystic form consists of pulmonary mass, which is echogenic and more solid with cysts < 5 mm in diameter (19). The volume of fetal CPAM lesion is measured by sonography by assessing the pulmonary mass in three dimensions and using the formula length*height*width*0.52 (20).

Despite numerous case reports on CPAM, there is still lack of evidence on reliable antenatal as well as postnatal management of CPAM mainly because of unknown nature and unpredictable prognosis of this pathology.

Prenatal management includes expectancy with observation, minimally invasive procedures such as single thoracocentesis of the cyst or thoracoamniotic shunt in macrocystic lesions. Large microcystic forms may be treated by invasive open fetal surgery or by non-invasive maternal administration of steroids (21). The exact mechanism of steroid therapy is unknown, it is supposed that it accelerates the lung maturation, which is, in case of CPAM, defected during early intrauterine period.

Even though there is no consensus on CPAM treatment, it is mostly recommended to treat patients at high risk of perinatal death defined by the risk factors such as fetal hydrops and value of CVR. The corticosteroid treatment of asymptomatic CPAM is despite the non-invasiveness highly questionable. It is currently not supported considering the relatively good prognosis of CPAM, lack of experience with treatment of these cases (including lack of case reports in the literature) and concerns about the different response to corticosteroid administration. Therefore, we present four cases of prenatally diagnosed non-hydropic fetal CPAM of various severity. With respect to the risk of perinatal morbidity and patients’ request for active treatment, all patients have been administered corticosteroids as soon as the diagnosis of CPAM was set. The authors also present current knowledge and literature review on the management of antenatally diagnosed CPAM and discuss different approaches to the antenatal management – routine maternal administration of corticosteroids in order to minimize the damage of the lung tissue, size of the lung lesions in postnatal period and to prevent the potential evolution of fetal hydrops with possible brain damage.

Patient 1

32-year-old patient was referred to our center for left intra-thoracic echogenic mass apically (20x11x22 mm) diagnosed by ultrasound at 20 weeks of gestation (gw). Repeated sonography confirmed fetal intrathoracic mass with a mediastinal shift to the right, type 2 CPAM (microcystic form) was diagnosed. The CPAM volume ratio (CVR) at the time of initial diagnosis was 1.5. Ultrasound did not confirm any other congenital anomalies. After a thorough informed consent was obtained, two doses of maternal Betamethasone 12 mg were administrated intramuscularly 24 hours apart at 22 gw. Subsequent serial ultrasound examinations were performed to observe the following course of pregnancy with onward regression of CPAM. At 37 gw male fetus was born by caesarean delivery due to two previous caesarean sections, birth weight 2750g, Apgar score 10/10. Newborn was hospitalized at the neonatal intensive care unit for mild dyspnoe and mild desaturation, required distensive respiration without support of oxygen therapy. CT scan of a newborn’ chest showed pulmonary lesion 47x17x30 mm in size. In the 7th month of life, the child underwent thoracoscopic resection of the affected lobe of the lung, histology confirmed the diagnosis of CPAM type II according to the Stoecker’s classification.

Patient 2

36-year-old patient was referred to our tertiary center at 25 gw for a finding of left basal intrathoracic mass. Diagnosis of CPAM was verified by ultrasound and subsequently by MRI. Sonography demonstrated no mediastinal shift, no pericardial effusion, ascites nor polyhydramnios. Other fetal anomalies were not detected by ultrasound. The diagnosis of CPAM mixed type has been set based on the performed examinations. The initial measurements of the mass were 34x23x30 mm. The CVR at that time was 0.6. After thorough consultation of the case, patient agreed with expectation policy. However, subsequent serial ultrasound examination showed enlarging of the mass, the CVR at 30 gw was 0.85, measurements 47x28x30 mm, fetus without hydrops or mediastinal shift. With respect to the growth of the mass and potential complications, we agreed on the betamethasone administration at 31 gw. During subsequent serial examinations, the mass showed slow regression and CVR decreased to 0.57 at 36 gw. At 37 gw labor was induced after premature spontaneous rupture of membranes and male infant was delivered vaginally. Neonate birth weight was 2800 g, Apgar score 10/10. Next course during the hospitalization was without any complications. In the third month of life, a contrast chest CT revealed a residual pulmonary lesion and the patient has recently undergone thoracoscopic surgical treatment.

Patient 3

30-year-old patient was referred to our center for a finding of intrathoracic mass which involved cystoid lesions in right hemithorax at 26 gw. The CVR at referral was 0.24. MRI image pointed to mixed form of CCAM, no other malformations of the fetus were found. Two doses of betamethasone 12mg was administrated 24 hours apart at 26 gw. The CVR was 0.20 at 30 gw and stagnation of the growth of the mass was observed. The patient was observed without any other treatment. Pregnancy was terminated at 38 gw by elective caesarean delivery due to placenta encroaching. Male infant with a birth weight of 2240 g and Apgar score 10/10 was
born. During hospitalization there were no signs of RDS or other respiratory symptoms. In the 6th week of life, the contrast CT of the chest was performed and revealed a pulmonary lesion 29x26x18.5 mm. In the 15th month of life, the child underwent thoracoscopic resection of the affected lobe.

**Patient 4**

28-year-old primigravida at 21 gw was diagnosed with microcystic form of CPAM of the fetus. On ultrasound a hyperechogenic pulmonary mass 22x26x31 mm was described in the left hemithorax, CVR 0.53. Beside the mass, ultrasound examination revealed also mediastinal shift, however there were no signs of fetal hydrops neither cardiac decompensation. Patient’s history did not include any remarkable data. Patient agreed with betamethasone treatment, 2 doses of 12 mg 24 hours apart were administrated at 21 gw. Subsequent ultrasound at 25 gw showed regression in size – 20x20x23 mm, CVR was 0.27.

At 39 gw labor was induced and female infant was vaginally delivered, birth weight 3230 g, Apgar score 9/10. During the postnatal adaptation neonate did not require any breathing support. In 3rd month of life, contrast CT was performed and CPAM type III according to Stoecker was diagnosed. The patient is currently being prepared for thoracoscopic treatment.

**Discussion**

CPAM has become an important study subject due to its different prognosis, clinical manifestation and no clear management guidelines despite the increasing number of prenatally diagnosed cases. The prognosis includes as good course as spontaneous intrauterine resolution of the mass without clinical abnormalities as well as lethal pulmonary hypoplasia, moderate respiratory symptoms postnatally or even progression of the persistent lesion to malignant process (12, 22). An extensive review of childhood lung neoplasms revealed that 8.6 % of malignancies were associated with previously documented cystic malformations (12, 22). Even though prenatal ultrasound screening and postnatal CT scan represent currently gold standard diagnostic tests, repeated ultrasound examination and fetal magnetic resonance imaging (MRI) allow better and more accurate diagnosis (23).

Differential diagnosis of a fetal lung mass is represented by CPAM, bronchopulmonary sequestration (BPS), congenital high airway obstruction syndrome (CHAOS), congenital lobe emphysema (CLE), congenital diaphragmatic hernia (CDH) and mediastinal tumors (16). A similar picture as a microcystic CPAM may have BPS, which appears in the ultrasonographic image as homogeneous echogenic mass. Doppler’s investigation can reveal systemic supply of the mass from the thoracic or abdominal part of the aorta, which is typical of BPS (17). Recent studies show that the distinction between these two entities is not so clear with the discovery of an increasing number of “hybrid” lesions (10).

Generally, the outcome of the fetus with CPAM depends on the complications related to the lesion itself. Lethal pulmonary hypoplasia comes from the compression of the effected ipsilateral lung, polyhydramnios, that may lead to preterm labor, comes from the compression of the esophagus by CPAM mass with inability to swallow, hydrops is related to deterioration of the cardiac venous return due to mediastinal shift (13). Moreover, anemia and cardiac failure with hypotension, which often occur in hydrops, may lead to brain perfusion failure and hypoxic-ischemic changes in fetal brain (24).

Therapeutic options include expectancy with observation or, in case complications occur, invasive techniques such as thoracocentesis, pleuro-atrio ductal shunt placement, percutaneous ultrasound-guided sclerotherapy, or radiofrequency/laser ablation, for microcystic forms of invasive open fetal surgery or noninvasive maternal administration of steroids, none of which are evidence based (17, 25–30).

According to the current literature data, management decision should be based on some of the defined risk factors and active management should be reserved for the mortality high-risk subgroups. The strongest independent prognostic factor which is considered indication for CPAM treatment has been defined to be hydrops (31). A survival rate of more than 95 % of CPAM cases without hydrops has been reported, whereas death occurred before or after birth in 95 % of CPAM cases with hydrops managed expectantly (32). Crombleholme et al (6) introduced a useful sonographic parameter for assessment of the hydrops – the CPAM volume ratio (CVR), which is CPAM mass volume divided by fetal head circumference. CVR then calculates the ratio of the CPAM volume and fetal head circumference ((length (cm) x height (cm) x width (cm) x 0.52) / head circumference (cm)) (7). CPAM lesions with CVR greater than 1.6 were shown to be at high-risk for fetal hydrops and thus at higher risk for fetal or neonatal death, depending on the gestational age at onset of the hydrops (7). Microcystic form is supposed to be associated with poorer outcome than macrocystic type (19). According to the recent studies, microcystic lesions respond to the administration of maternal betamethasone, resulting in maturation of the fetal lung with good effect on large and growing lesions (33). Steroids decrease the production of lung fluid and increase its reabsorption within the thoracic malformation, thus mimicking the physiological third trimester changes (34). They are indicated for microcystic lesions, while it is unclear whether macrocystic forms respond to this treatment (35, 36).

David et al (37) presents an algorithm for prenatal CPAM treatment where he recommends administration of maternal steroid in fetuses less than 32 gw with CVR > 1.6 or fetuses with hydrops regardless of CVR. Even though the predictive value of CVR in terms of the development of fetal hydrops is generally accepted, no threshold was set for the prediction of fetal/perinatal outcomes. Recent systematic review of the studies reporting on the predictive utility of the CVR suggests that a threshold much lower than 1.6 cm2 is likely to be of greater utility for outcomes of perinatal relevance. For neonatal outcomes a CVR on the initial ultrasound scan ranging from 0.5 to 1.0 cm2 appears to have the greatest predictive value (38). Up to now several cases of the microcystic CPAM lesions treated only conservatively with betamethasone have been reported, all of them had hydrops and/or CVR > 1.6. Tsao et al (2) first published the experience with
prenatal steroids in microcystic CPAM in three fetuses who presented with hydrops, with 100 % survival rate. Peranteau et al (39) reported 100 % (11/11) survival after administering betamethasone, 80 % (4/5) hydrops resolution. Curran et al (40) reported following results after betamethasone administration: decrease of CVR in 61.5 % (8/13) and 77.8 % (7/9) hydrops resolution rate, 100 % fetuses survived to delivery and 84.6 % survived to neonatal discharge. Two fetuses whose hydrops did not resolve did not survive to discharge. Loh et al (36) reported 77 % (10/13) survival rate after administration of betamethasone while Derderian et al (41) published 63 % (5/8) survival after repeated administration of betamethasone. Yamashita et al (42), published 100 % (3/3) survival after single dose of betamethasone and 100 % hydrops resolving. Peranteau et al (43) monitored the effect of betamethasone in single dose and multiple dosing, with result of 86 % survival in the multiple administration of betamethasone vs 93 % of a single dose of betamethasone. Review of the published data on standard single dose betamethasone treatment of CPAM can be seen in Table 1. For comparison, Adzick (44) describes 24 cases of open surgery with 11 cases of fetal death. Regarding above data, there is strong evidence that prenatal steroid administration represents first choice treatment in fetuses with high-risk CPAM. It is still unclear if it is also beneficial for patients with non-hydropic microcystic/other CPAM lesions. It is mainly due to a lack of experience with the effect of prenatal steroids on non-hydropic CPAM fetuses and concerns about different response and effects on the child’s future health. On the other hand, even though CPAM is suggested to be disease with good prognosis in case it does not perform with hydrops (11, 21, 45), there is still significant morbidity during postnatal period described in direct association with CPAM (12, 22, 24). This is in accordance with the mentioned finding of systematic review suggesting that threshold of CVR for perinatal outcomes is different and lower than threshold for fetal hydrops (38). Moreover, Gallardo (11) states the prenatal resolution of the lesion should not be accepted as evidence of a true disappearance of the pathology as 45 % of the lesions considered absent in the Nicolaides series (10) were subsequently subjected to resection due to the existence of persistent lesions on CT.

We described 4 cases of prenatally diagnosed CPAM at 20–26 gw, 2 microcystic and two mixed forms. Even though none of the cases performed hydrops, one case presented with mediastinal shift and CVR close to the risk value. In the other cases, CPAM lesions continued to grow even after 28 gw. Betamethasone in standard regimen of 2 doses, 12 mg intramuscularly, 24 hours apart was administered in all of the cases. All fetuses responded to the treatment with either stagnation or regression of the size of the CPAM mass. We did not record complete resolution of the mass in any of the patients nor did we observe other complications of the pregnancy that could be related to the steroid treatment. All patients delivered at 37 gw or later after spontaneous onset of labor or premature rupture of membranes without complications, 2 delivered vaginally, 2 by caesarean section due to previous caesarean section and placenta previa. All children survived until discharge and all of them had to undergo thoracoscopic pulmonary resection of the remaining lesion during first year of life. The fact that postnatally diagnosed persistent pulmonary mass diagnosed by CT was larger than that diagnosed prenatally by ultrasound may be explained by the results of Nicolaides series (10).

We currently continue to follow up the presented children and collect data regarding their postnatal development and health state. Even though our case series does not prove the benefit of antenatal administration of betamethasone in non-hydropic fetuses with prenatally diagnosed CPAM, we believe it supports the “safety” of the betamethasone administration in these cases at least during the perinatal period.

The current recommendations to treat only high-risk fetuses with CPAM come from the concerns about the possible effects of the antenatal corticosteroids to the child. However, there are already several longer-term follow-up studies such as the extended original Liggins and Howie randomized cohort (46) that were followed up at age 30 years. Even though the betamethasone in this study was administered in order to manage patients with imminent preterm birth, data show that exposure to 12 mg betamethasone, two doses 24h apart or double this dose did not alter growth parameters or cardio-metabolic risk, the only difference being a higher insulin level after glucose tolerance test, suggesting possible mild insulin resistance. Psychological testing showed no difference in cognitive functioning, working memory and attention, psychiatric morbidity, handedness, or quality of life (47). Considering our data, uncomplicated course of the pregnancy after betamethasone administration, calculating the risk and benefit together with its non-invasive and simple use, we think that all patients with prenatally diagnosed CPAM could be offered this treatment as soon as the diagnosis is set, particularly patients requesting active management. We believe it is more beneficial to start the steroid therapy before the hydrops occurs, as this state may already be associated with brain perfusion failure and ischemic changes (24). Moreover, the treatment might help to avoid other complications such as pulmonary compression of any grade,
References


Received February 22, 2021.
Accepted March 15, 2021.