Prenatal Ultrasonography and Early Surgery for Congenital Cystic Disease of the Lung

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With the recent advent of prenatal ultrasound as a routine screening procedure, diagnosis of congenital cystic lung disease has been made in utero, raising the possibility of elective surgery for these lesions early in infancy before the patient develops respiratory distress or potentially life-threatening infection. From 1979 to 1989 six cases of congenital lung cyst were diagnosed in utero by prenatal ultrasound and followed during pregnancy. Two of the six were not confirmed after birth because the mothers preferred an abortion. The remaining four cases were studied periodically during gestation by ultrasonography. At birth, the first infant developed respiratory distress and underwent urgent left upper lobectomy and left lower segmentectomy at age 18 hours. The other three underwent elective lobectomy at age 10 days, 3 months, and 7 months, respectively. The fourth infant had a normal chest x-ray and ultrasound at birth, and the congenital cysts were confirmed by computed tomography scan. The pathological diagnosis in all four cases was cystic adenomatoid malformation. In two cases, intraoperative measurement of pulmonary function demonstrated significant improvement after resection of the affected lobe. We conclude that congenital lung cysts can be accurately diagnosed by prenatal ultrasound "screening" as early as 18 to 24 weeks' gestation. Advantages of early diagnosis include the option of moving the mother and unborn child to a high-risk obstetrical center for urgent operation on the newborn infant if necessary. Otherwise, once the diagnosis is confirmed, surgical correction can be performed electively and safely before respiratory distress or pulmonary infection complicates the infant's growth and development.

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CONGENITAL cystic disease of the lung includes four classic lesions that may present as abnor mal cystic areas within the pleural cavity in early life: congenital adenomatoid malformation (CAM), pulmonary sequestration, congenital lobar overinflation, and bronchogenic cyst. Although histologically distinct, they share common clinical and embryological characteristics, are often difficult to diagnose, and always require surgical therapy. Diagnosis may be during the newborn period, in cases in which the space-occupying cystic lesion causes respiratory distress, or delayed by months to years until the cyst becomes infected, presenting as recurrent pneumon ia or lung abscess. In the past, late diagnosis and treatment was accompanied by significant morbidity and mortality. During the last 10 years, pediatric surgeons have found that early diagnosis and prompt surgical treatment are both safe and effective.

With the recent advent of prenatal ultrasound as a routine screening procedure, diagnosis of congenital cystic lung disease has been made in utero, raising the possibility of elective surgery for these lesions early in infancy before the patient develops respiratory distress or potentially life-threatening infection. The present study reviews four infants diagnosed with congenital cystic lung disease by prenatal ultrasound with three operated on electively before becoming symptomatic.

MATERIALS AND METHODS

From 1979 to 1989, six cases of congenital lung cyst were diagnosed in utero by prenatal ultrasound and followed during pregnancy in the Department of Pediatric Surgery, University of Milan. Two of the six were not confirmed by us after birth because the mothers preferred an abortion. The remaining four cases were studied periodically during gestation and operated on electively after birth.

Segmentectomy or lobectomy was carried out at operation, and in all four cases a type I CAM was diagnosed.

In two of the four cases (see cases 2 and 4), respiratory function was measured immediately before and after the operation.

A Servoventilator 900 C (Siemens) and a computerized FLEISCH pneumotocograph Type 00 (PN 0079)-Pulmonary evaluation and diagnostic system (PEDS) with two pressure-transducers (for endotracheal and endoesophageal pressure, respectively) were used to study the pulmonary function in cases 2 and 4 just before (time 0 = T0) and 30 minutes after the operation under anesthesia (time 30 = T30).

The following parameters were evaluated: respiratory frequency, tidal volume, pulmonary compliance, transpulmonary pressure, total resistance, and total work. Upon integrating these parameters, pressure-volume loops were obtained in both cases.

The prenatal evolution of the lung cyst was followed periodically in our department by means of real-time ultrasonography with cine tape recording until the time of delivery.

Three of four women delivered vaginally at term. The fourth underwent cesarean section, not because of fetal pathology, but electively because of a previous delivery by cesarean section.

Case Reports

Case 1. The prenatal diagnosis of congenital lung cyst was made at the 24th week of gestation by routine fetal ultrasound (Fig...
The large left-sided cyst displaced the heart and mediastinum, but neither hydrothorax nor anasarca occurred during pregnancy. Cesarean section was performed at the 40th week with delivery of a 2.6-kg female infant. Progressive respiratory distress developed, and a plain chest x-ray confirmed the prenatal diagnosis of congenital lung disease. At age 18 hours, a left upper lobectomy and a double segmentectomy (apical and anteromedial segments) of the left lower lobe was performed with removal of multiple large cysts. Respiratory function dramatically improved, and the baby was extubated 24 hours after surgery. CAM was diagnosed by pathological examination. Follow-up over 4 years showed an active, well-developed girl with apparent normal pulmonary function.

Case 2. The prenatal diagnosis of congenital lung cyst was made by routine fetal ultrasound at the 20th week of gestation (Fig 2). A large cyst involved the left lower lung. Neither hydrothorax nor anasarca were observed during the remaining gestation. Term vaginal delivery produced a 3.2-kg girl with no evidence of respiratory distress. Plain x-ray and computed tomography (CT) scan confirmed the prenatal diagnosis suggesting CAM. A left lower lobectomy was successfully performed on the 10th day of life. Pulmonary function measured at the time of surgery improved immediately and significantly after lobectomy, and the postoperative course was uneventful. Pathological examination confirmed CAM. Follow-up at 11 months showed a healthy infant with apparent normal pulmonary function.

Case 3. The prenatal diagnosis of congenital lung cyst was made at the 18th week of gestation and followed by periodic ultrasonography until birth (Fig 3). The differential diagnosis initially included pulmonary sequestration, but the typical image of cysts at the 19th week strongly supported the diagnosis of CAM. The cysts did not increase in size during gestation, and no fetal complications were observed. Term vaginal delivery produced a 3.9-kg boy with no evidence of respiratory distress. X-ray at birth was equivocal for right lower lung cysts, but these were demonstrated later by ultrasound and CT scan. The child remained free of symptoms and underwent surgery at 3 months of age. At surgery the right lower lobe was distorted by one large cyst and two smaller cysts with the clinical appearance of CAM. A right lower lobectomy was performed. Due to technical difficulties with measurement, the improvement of the pulmonary compliance, although appreciated clinically, could not be reliably recorded. Histological examination confirmed the diagnosis of CAM. The postoperative course was uneventful, and the infant was doing very well with no respiratory symptoms at the 3-month follow-up visit.

Case 4. The prenatal diagnosis of congenital lung cyst was made at the 20th week of gestation (Fig 4). Follow-up ultrasonography showed normal fetal growth without hydrothorax or anasarca. At term, a 3.4-kg girl was born by spontaneous vaginal delivery and had no respiratory symptoms. The cysts could not be demonstrated after birth either by plain chest x-ray or by ultrasonography. However, CT scan confirmed the prenatal diagnosis, and the infant underwent surgery at age 7 months. A left lower lobectomy was performed, and intraoperative measurement of pulmonary function was otherwise uneventful, and spontaneous vaginal delivery of a 3.2-kg girl occurred at the 39th week. The female infant was free of respiratory symptoms. Plain x-ray and computed tomography (CT) scan confirmed the prenatal diagnosis suggesting CAM. A left lower lobectomy was successfully performed on the 10th day of life. Pulmonary function measured at the time of surgery improved immediately and significantly after lobectomy, and the postoperative course was uneventful. Pathological examination confirmed CAM. Follow-up at 11 months showed a healthy infant with apparent normal pulmonary function.
RESULTS

The prenatal diagnosis of congenital cystic lung disease was confirmed at birth, and the diagnostic impression of CAM was confirmed by histological examination at time of surgery in all four cases. In each of the four infants, the cysts did not increase in size during fetal life. In two cases we observed the evolution of the cystic abnormality from the 18th and 20th week onward, respectively. Although no further in utero complications were observed in patient 1, she was shown to have multiple large cysts displacing the mediastinum at 24 weeks, and progressive respiratory distress during the first 18 hours of life led to urgent surgery. In the 24 hours following operation, there was dramatic clinical improvement in respiratory function, as previously described. In cases 2 and 4, intraoperative measurement of pulmonary function showed less dramatic but significant improvement immediately after removing the affected lobes. In both cases, a significant improvement in both tidal volume and compliance after lobectomy at T30 was seen (Table 1). This is shown by the “sigmoid shape” slope in the inspiratory half of the curve, which reflects a larger volume at the same pressure after the operation (Fig 5). Because of technical measurement difficulties during the third case, this demonstration was not possible.

Lobectomy in three cases and combined lobectomy-segmentectomy in one case were more readily accomplished because of the excellent condition of the lung parenchyma adjacent to the cysts; namely, neither atelectasis nor inflammatory adhesions of the nearby segments or lobes were observed.

Surgery was easily performed without significant blood loss. In all cases, the remaining lung tissue reexpanded to fill the hemithorax within a few hours.

Table 1. Preoperative and Postoperative Data from Cases 2 and 4

<table>
<thead>
<tr>
<th>Case 2</th>
<th>Case 4</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Preoperative</td>
</tr>
<tr>
<td>T (min)</td>
<td>0</td>
</tr>
<tr>
<td>Rf (breaths/min)</td>
<td>30</td>
</tr>
<tr>
<td>Vt (mL/kg)</td>
<td>5.1</td>
</tr>
<tr>
<td>Ct (mL/cm H$_2$O/kg)</td>
<td>0.248</td>
</tr>
<tr>
<td>P (cm H$_2$O)</td>
<td>22</td>
</tr>
<tr>
<td>Rt (cm H$_2$O/L/s)</td>
<td>152</td>
</tr>
<tr>
<td>Wt (g/cm/kg)</td>
<td>216</td>
</tr>
</tbody>
</table>

Abbreviations: Rf, respiratory frequency; Vt, tidal volume; Ct, compliance; P, pressure; Rt, transpulmonary resistance; Wt, total work.
of the operation. A silastic chest tube was placed at
the time of surgery in each case and removed 5 days
after the operation.

Follow-up of these four patients at 4 years, 11
months, 3 months, and 3 months, respectively, shows
apparent normal function of the remaining pulmo-
nary tissue. All children are growing and free of
respiratory infection.

DISCUSSION

With the advancing technology and ease of prena-
tal ultrasound, congenital cysts of the lung can be
diagnosed with increasing accuracy between 18 and
24 weeks of gestational age. The advantages of
prenatal diagnosis of these lesions include the option
of moving the mother and unborn child to a high-risk
obstetrical center where delivery can be planned with
expert neonatologists, pediatric anesthesiologists, and
pediatric surgeons available. Once the diagnosis is
confirmed, surgical correction can be carried out
electively before respiratory distress or pulmonary
infection complicate the infant's growth and develop-
ment.1,2,4,5 Postnatal plain chest x-ray is not always
sufficient to diagnose congenital lung cyst, nor is
prenatal ultrasound always sufficient, as we discov-
ered in case 4. Rather, CT scan was required to
demonstrate the congenital lung cysts in our fourth
patient, an important point in confirming the accu-
ry of carefully performed prenatal ultrasonogra-
phy.

We followed all fetuses from the time of prenatal
ultrasound diagnosis of congenital lung cysts to the
birth of the infant. Although the four diagnoses were
made early in pregnancy, the cysts did not increase
during fetal life and neither hydrothorax nor anasarca
occurred.

If the evolution of the congenital cyst remains
uncomplicated during fetal life, the infant can be
delivered at term and elective surgery subsequently
carried out. Since all congenital lung cysts eventually
require surgery, a precise diagnosis as to specific type
(CAM, pulmonary sequestration, congenital lobar
overinflation, or bronchogenic cyst) is not important,
even though we correctly predicted the diagnosis of
CAM in our four cases, as confirmed by pathological
examination.

Early elective operation is preferred before respira-
tory distress or infection complicate the procedure.
Surgery carried out in our four cases was straightfor-
ward and unencumbered by adhesions or destroyed
tissue planes which characteristically affect patients
diagnosed after multiple bouts of pulmonary infec-
tions.1,6,7 Moreover, in two of the three asymptomatic
cases, pulmonary function improved immediately af-
after lobectomy, as documented intraoperatively prior
to thoracotomy and 30 minutes postoperatively under
anesthesia.

We conclude that the advantages of early surgery
for congenital cystic lung disease are manifest by our
experience with the four cases reported. Prenatal
ultrasound is an accurate and important screening
procedure; and, once a prenatal diagnosis of cystic
lung disease has been established, CT scan should be
obtained to confirm the diagnosis whenever a chest
x-ray or repeat ultrasound is equivocal or "normal"
after birth. Definite improvement in pulmonary func-
tion can be expected (as measured in cases 2 and 4)
after early elective surgery for congenital lung cyst,
and the otherwise inevitable complications of respira-
tory distress or recurrent infection can be avoided.

REFERENCES

and management of congenital cystic disease of the lung in

adenomatoid malformation of the lung. Potential diagnostic pitfall.

3. Stocker JT, Madewell JE, Drake RM: Congenital cystic
adenomatoid malformation of the lung. Hum Pathol 8:155-171,
1977

4. Monclaire T, Schistad G: Congenital pulmonary cysts versus a
differential diagnosis in the newborn: Diaphragmatic hernia. J

adenomatoid malformation of the lung. Potential diagnostic pitfall.

6. Lilly JR, Wesenberg RL, Shikes RM: Segmental lung resec-

cystic adenomatoid malformation, pulmonary sequestration, and
bronchogenic cyst in infancy and children: A clinical group. J
Pediatr Surg 9:85-93, 1974