Extralobar lung sequestration associated with fatal neonatal respiratory distress

A. J. BLIEK and D. J. MULHOLLAND

The Departments of Radiology and Pathology, Royal Alexandra Hospital, Edmonton, Alberta

The radiological and pathological findings of lung sequestration in a premature infant are described. There was a left-sided variant of the extralobar type with double systemic arterial blood supply, single systemic venous drainage, and an incomplete bronchial tree for the remainder of the left lung.

A white female infant was born at 34 weeks' gestation by breech extraction with Piper forceps on the aftercoming head. The spontaneous labour had lasted four-and-a-half hours with rupture of the membranes at the time of delivery. Polyhydramnios was noted but not measured: the placenta was unremarkable.

The mother was a healthy 25-year-old gravida 4, para 3, whose antenatal course had been marked by nausea and vomiting throughout with spotty bleeding of short duration at three-and-a-half months. No drugs were administered during the pregnancy other than a single tetanus injection after a foot injury. She had gained 42 lb (19 kg). One live sibling had Down's syndrome.

The infant's birth weight was 6 lb 3 oz (2,800 g), and her total length was 20 in (50 cm). At birth the infant was cyanosed, grey, and gasping with gross indrawing. After suction and intubation she was placed in a respirator, when the Apgar rating improved from 5 to 6. In spite of vigorous treatment of the ensuing hypoxia and acidosis (pH below 6.8), the infant died two hours and six minutes after birth.

![Image](http://example.com/image.jpg)

**FIG. 1.** Antemortem chest radiograph showing dense left hemithorax and faintly visualized trachea and right main bronchus.
A chest radiograph (Fig. 1) showed the presence of air in the oesophagus. The right lung was densely consolidated with a well-developed air-bronchogram. The left hemithorax was opaque without a recognizable air-bronchogram, suggesting occlusion or atresia of the left main-stem bronchus.

Necropsy findings

Obvious anomalies were seen when the thorax was opened, and before evisceration a small polyethylene catheter was introduced into the aorta via the left external iliac artery. Three millilitres of 60% meglumine iothalamate were injected and radiographs were obtained. Two systemic arteries originating from the thoracic aorta were seen supplying a large mass in the left hemithorax (Fig. 2). The ductus arteriosus was patent. Subsequently 2 ml of contrast medium were instilled into the trachea and bronchial tree and a radiograph (Fig. 3) showed a normal right but incomplete left bronchial tree. No connexion with the left intrathoracic mass was evident.

**FIG. 2.** Postmortem aortogram showing two arteries supplying the sequestered lung tissue.

**FIG. 3.** A normal right but incomplete left bronchial tree are demonstrated. No connexion with the mass is evident.

**FIG. 4.** In the opened thoracic cavity the mass is seen displacing the heart and mediastinum.
FIG. 5. The narrow pedicle connecting the mass with the pulmonary ligament contains two arteries and one vein. No diaphragmatic hernia is evident.

FIG. 6. The right and left lungs show a moderate degree of collapse. Alveolar development is present and patchy hyaline membrane formation is noted (×125).
In the thorax there was a left-sided pleural effusion of 20-30 ml of straw-coloured fluid and a small left lung weighing 4 g. Most of the left pleural cavity was occupied by a smooth-surfaced, pink-grey mass of solid tissue, measuring $5 \times 3.5 \times 3$ cm and weighing 35.5 g. The mediastinal structures were shifted to the right compressing the contralateral lung (Figs 4 and 5). Two small systemic arteries located immediately above the diaphragm led directly from the aorta to the mass. A single vein provided drainage to the inferior vena cava. All three vessels passed through a small fibrous stalk at the infero-postero-medial aspect of the mass to the pulmonary ligament. The diaphragm was normal. Approximately 50 ml of bloody fluid associated with a subcapsular haematoma on the anterior surface of the liver was found in the abdominal cavity. Except for the presence of a Meckel's diverticulum the remainder of the gross examination was unremarkable.

Microscopical Examination  Tissue sections of the right lung revealed an advanced degree of alveolar development with slightly thickened but well-vascularized alveolar septa. Patchy hyaline membrane formation was present with epithelial debris and fluid accumulation in some of the airspaces. An uneven degree of dilatation of the alveolar airspaces was probably due to the short period of artificial ventilation (Fig. 6).

The small left lung was similar histologically to the right lung. The left-sided pulmonary mass had a separate pleural covering which in several areas showed much subpleural lymphatic dilatation (Fig. 7). The parenchyma was composed of a multitude of 'terminal bronchiole-like' spaces lined by a single layer of low cuboidal epithelium. Only a few scattered areas showed a pattern compatible with early alveolar development. The septa were thick, poorly vascularized, and composed of a loose cellular stroma. Neither epithelial cells nor hyaline membranes were found within the respiratory spaces (Figs 8 and 9).

Elastic stains of the aberrant arteries supplying the sequestrated pulmonary tissue revealed them to be primarily of a systemic type with negligible medial hypertrophy and abundant elastic tissue throughout the intima, media, and adventitia (Fig. 10).
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**FIGS 8 and 9.** The parenchyma of the sequestrated lobe is composed of 'terminal bronchiole-like' spaces with a single layer of low cuboidal epithelium. Scattered areas show early alveolar development. (× 125 and 300).
Pulmonary sequestration has been documented extensively in the recent and past literature and various theories regarding its embryogenesis have been presented and reviewed (Pryce, 1946; Smith, 1956; Berman, 1958; Swierenga, 1959; Nielsen, 1964; Köhler, 1969). Only a few examples of the extralobar type in the perinatal group have been reported (Potter, 1961; Horowitz, 1965).

The present case illustrates the embryogenesis of the lesion, in particular the pulmonary origin. The extralobar variant differs from the intralobar one in its extrapulmonary location, with its own pleural covering and its lack of connexion with the bronchial tree. It differs from the accessory lung due to the absence of an accessory bronchus or other connexion with the foregut (Spencer, 1962; Blesovsky, 1967).

The lack of respiratory epithelium in the mass, its systemic blood supply, and histological features confirm the common pattern of this lesion (Wagenvoort, Heath and Edwards, 1964; Kissane and Smith, 1967). The spectrum of anomalies may well justify the name of congenital bronchopulmonary-foregut malformation, as suggested by Gerle, Jaretzki, Ashley, and Berne (1968). The dilated subpleural lymphatics have not been reported previously in this anomaly and may be associated with poor lymphatic drainage and the formation of the large pleural effusion.

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