Congenital Lobar Emphysema: Perioperative Evaluation and Management

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ABSTRACT

Background: Congenital lobar emphysema (CLE) is a lung malformation characterized by overdistension and air trapping in the affected lobe. It is one of the causes of neonatal and infantile respiratory distress. This study aimed to evaluate our experience regarding perioperative and surgical management in children with CLE.

Methods: A retrospective observational study was done for all CLE patients who underwent surgery at Menoufia University Hospital. Perioperative data collected included demographic, clinical, and radiological findings, as well as operative and postoperative data.

Results: We included 30 neonates and infants who suffered from CLE between January 2013 and December 2020; the mean age was 111.43 ± 65.19 days, and 21 were males. All cases presented with respiratory distress; 19 had cyanosis, and 15 had recurrent pneumonia and fever. Plain chest x-ray and computed tomography (CT) revealed emphysema in all cases. Lobectomy was done in all cases; the mean age at surgery was 147.58 ± 81.49 days. Postoperative complications occurred in 147.58 ± 81.49 days. Postoperative complications occurred in 15 patients, and 2 of them needed mechanical ventilation. The follow-up duration ranged from 3 months to 1 year (except 1 case lost to follow-up after 3 months), and all patients were doing well.

Conclusion: CLE is a rare bronchopulmonary malformation that requires a high index of clinical suspicion, especially in persistent and recurrent infantile respiratory distress. Chest CT is the most useful diagnostic modality. Early management of CLE improves outcome and prevents life-threatening complications. Surgical management is the treatment of choice in our center, without recorded mortality.

INTRODUCTION

Congenital lobar emphysema (CLE) is a rare developmental pulmonary anomaly that usually occurs due to aberrations in the development of lung parenchyma during the third week of gestation [Kerstine 2010]. CLE is defined as overinflation of ≥1 lung lobes due to partial obstruction of the bronchus, leading to pressure manifestations on the mediastinal structures [Olutoye 2000]. Most patients’ symptoms appear during the first 6 months of life; half of patients have no clinical manifestations at birth. Ventilation and perfusion are usually impaired owing to hyperinflation of the diseased lobe, and with progressive hyperinflation, compression on the other lobe and mediastinal structures occurs [Mendeloff 2004; Paramalingam 2010].

The management of patients with CLE depends on the severity of clinical manifestations. Half of patients have manifestations in the first months of life. Although a few asymptomatic patients have been described in previous studies, most cases are diagnosed in the first 6 months of life [Karnak 1999; Markowitz 1989]. The management of patients with CLE depends on the severity of clinical manifestations. Half of patients have manifestations in the first months of life. Although a few asymptomatic patients have been described in previous studies, most cases are diagnosed in the first 6 months of life [Karnak 1999; Thakral 2001]. Conservative management is recommended in moderate cases, whereas severe cases are usually treated by lobectomy. Controversy regarding management is still present, especially if the manifestations of CLE are mild or moderate, for which some surgeons prefer conservative management and others prefer surgical intervention. But if there is severe respiratory distress, lobectomy is the treatment of choice [Mei-Zahav 2006; Ulku 2008].

We present our experience in the surgical management and follow-up of 30 cases admitted to our hospital.

METHODS

This is a retrospective observational study of 30 patients with CLE who were admitted for surgical management at the pediatric and cardiothoracic surgery departments, Menoufia University Hospital, Egypt, between 2013 and 2020. Their clinical presentation, diagnosis, treatment, and 6-month follow-up data were studied. The diagnosis of CLE was confirmed by clinical and radiological findings, in addition to histopathological examination. The parameters analyzed included age, clinical manifestations, chest x-rays, and chest CT. All patients underwent lobectomy. Elective lobectomy was done in 24 cases, and emergency lobectomy in 6, because of respiratory distress. Postoperative data included need for mechanical ventilation, timing of oral feeding, clinical
picture, analysis of radiological findings, and 6-month follow-up examination. Follow-up was done in outpatient clinics for all patients. The study was approved by Menoufa Faculty of Medicine Hospital ethics committee.

**Operative Technique**

General anesthesia via proper-sized single-lumen endotracheal intubation is induced, and excessive ventilatory pressures are avoided until the chest is opened. This precaution is necessary to avoid further expansion of the obstructed lobe and complication of remaining pulmonary function. A lateral position was used in all cases in this study, with a muscle-sparing posterolateral thoracotomy through the fourth intercostal space. The skin incision typically starts at the anterior axillary line just below the nipple level and extends posteriorly below the tip of the scapula. The incision then proceeds in a cranial direction halfway between the vertebral border of the scapula and the spinous processes of the vertebrae. The latissimus dorsi is divided, and the serratus anterior is retracted. Typically, at the fourth interspace the intercostal muscles are divided using electrocautery above the fifth rib. A rib spreader is placed into the thoracic cavity and minimally opened. Cautery can then be used to perform an internal thoracotomy by continuing the division of the intercostal muscles more anteriorly (up to the level of the internal mammary artery) and posteriorly (up to the level of the paraspinal tendons). The internal thoracotomy will prevent rib fracture during subsequent spreading of the retractor.

At thoracotomy, the emphysematous lobe is pale, spongy, and frequently prolapsed through the wound. It does not collapse and feels like a thickened sponge. The bronchus is soft, and the bronchial stump is closed by suture. Extrinsic causes, such as bronchial obstruction by an anomalous pulmonary artery, prehilar bronchogenic cyst, aneurysmal ductus arteriosus, or enlarged mediastinal nodes, should be excluded by careful mediastinal exploration before lobectomy. Before closure, it is necessary to test for air leak by saline. Usually, the anesthetist is asked to start manual ventilation to help inflate the residual lobe to fill the thoracic cavity. Two chest tubes are inserted, and the chest is closed in layers. Postoperative follow-up for 6 months in all cases is our practice to detect recurrence of manifestations or appearance of complications.

**Statistical Analysis**

Data were coded, entered, and analyzed using Microsoft Excel software. Data were then analyzed in Statistical Package for Social Sciences (SPSS version 20) software. Qualitative data are presented as number and percentage, and quantitative continuous groups are presented as mean ± standard deviation (SD).

**RESULTS**

Of 30 patients with CLE, 21 were males and 9 were females. The mean age of onset of symptoms was 32 ± 33.7 days. Presenting features varied: tachypnea was present in all cases, dyspnea in 18 cases, cyanosis in 6, fever in 15, failure to thrive in 3, and irritability in 9. Tachypnea, cyanosis, wheezing, and failure to thrive were evenly scattered in all age groups, whereas cough, fever, and recurrent respiratory tract infection were significantly more common in older patients than younger ones. In contrast, dyspnea was more common in younger patients and correlated positively with bad general condition.

Fifteen cases presented with bad general condition (irritability, apneic spells, and neonatal reflex depression). Six cases needed assisted mechanical ventilation, and 12 needed mask oxygen; the remaining 12 cases were mildly symptomatic and able to maintain good oxygen saturation and blood gases on room air without oxygen supplementation. Arterial blood gas analyses revealed hypoxia in 18 cases and hypercapnia in 12 cases.
A plain chest radiograph was done in all patients, with the following characteristic findings: increased translucency of affected side in 30, mediastinal shift in 24, herniation of overexpanded lung into contralateral side in 15, and atelectasis in 15. CT of the thorax confirmed the findings of chest radiography and was beneficial in follow-up (Figures 1 and 2).

The left upper lobe (LUL) was affected in 24 cases, right middle lobe (RML) in 3, and right upper lobe (RUL) in 3. All patients underwent total lobectomy. Lobectomy was elective in 24 cases and emergency in 6 cases, due to severe respiratory distress. No postoperative oxygen supplement was required in 24 cases, whereas 3 cases needed assisted ventilation. Oral feeding was started on postoperative day 2 in 15 patients, day 3 in 9, day 4 in 3, and day 5 in 3. Follow-up after 6 months revealed no complications in 24 cases, recurrent attacks of coughing in 3, and recurrent respiratory tract infection in 3.

**DISCUSSION**

The etiology of CLE is unknown in half of cases, and hyperplasia, dysplasia, or absent bronchial cartilage are
indicated in a quarter of cases [Karnak 1999]. Acquired causes of CLE included meconium aspiration, hypertrophic mucous membranes, and foreign body aspiration [Michelson 1977; Aslan 2005]. Bronchogenic cysts and mediastinal tumors are considered rare extrinsic causes of CLE [Khemiri 2008; Kumar 2008].

CLE is a rare disease more common in males than females, at a ratio of 3:1 [Demir 2019], which was consistent with our study, in which the male:female ratio was 2.34:1, and a study by Cataneo et al [2013]. In our study, the age of patients ranged from 2 days to 5 months. The majority (70%) presented within the first 6 weeks of life, as supported by Thakral et al [2001], in whose study 81% of the cases presented within the first 6 weeks of life, as well as Kunisaki et al [2019]. Delayed respiratory distress in children 5 years of age was reported by Man et al [1983]. Cataneo et al [2013] reported that delayed diagnosis of CLE was due to combined recurrent respiratory tract infections and respiratory distress in some patients in their study.

In the current study, CLE was diagnosed by clinical findings and confirmed by plain chest x-ray, with atelectasis found in most cases. We confirmed the diagnosis by CT scan, also to discover other vascular anomalies. The diagnosis can be made prenatally by fetal ultrasound, but some cases may be missed and discovered later on during the postnatal period due to progressive overinflation of the emphysematous lobe that leads to pressure manifestations on mediastinal structures [Correa-Pinto 2010]. In our study, the LUL was most often involved, then RUL and RML. This finding is in agreement with the study done by Bush et al [2019]. Thacker et al [2014] also reported that the LUL was the most affected lobe followed by RUL and RML, whereas Berrocal et al [2004] reported that CLE was confirmed in lower lung lobes and bilobal lobes in some patients.

In our study, 100% of cases had hypertranslucency of the affected lobe, mediastinal shift to the opposite side was found in 80%, and 50% had herniation of the affected lobe to the opposite side. Atelectasis was found in 50% of the cases, and no patient had pneumothorax. In a study conducted by Ozcelik et al. [2003], the radiological findings of 30 patients consisted of hyperlucent of affected lobe in 100%, mediastinal shift to the opposite side in 52%, atelectasis in 23%, and pneumothorax in 4%.

Our study included 6 cases (20%) with associated heart anomalies by echocardiography, which must be routinely ordered preoperatively. Our finding was consistent with other studies [Mendeloff 2004; Bush 2019; Ozcelik 2003] that reported 14% to 20% of patients having congenital cardiac anomalies in association with CLE. These anomalies included patent ductus arteriosus, atrial septal defect, ventricular septal defect, and tetralogy of Fallot.

The standard surgical intervention in our study was lobectomy; lobectomy was also the standard management for other studies [Cataneo 2013; Perea 2017]. There is some controversy about whether to manage CLE patients conservatively or surgically, especially in mild clinical presentations; Mei-Zahav et al [2006] and Ceran et al [2010] prefer nonsurgical treatment in mild cases. Fierro et al [2018] discussed the decision in 6 cases with CLE, deciding that lobectomy was beneficial in such infants.

All our cases were operated via standard muscle-sparing posterolateral thoracotomy, which is our preferred approach in this age group and disease. The emphysematous lobe usually expands and occupies the whole thoracic cavity, which is managed with difficulty in thoracoscopy. Some authors who have used thorascopic lobectomy experienced difficulty and converted to thoracotomy [Rahman 2009; Zoeller 2018].

In a study conducted by Lincoln et al [1971], bronchial cartilage deficiency was found in 88% of cases, with no abnormal findings in 12%. In our study deficiency of bronchial cartilage was found in 70% of cases, with no abnormal findings in the other 30%. Approximately 20% of our patients had postoperative complications. Three cases suffered respiratory failure, needed mechanical ventilation, and were successfully weaned 3 to 4 days later; 2 patients had postoperative atelectasis; and 1 patient had pneumothorax.

Thakral et al [2001], in a 6-month follow-up of CLE cases, found 81% of the cases to be free of symptoms; 14% had recurrent attacks of respiratory tract infection, and 5% had recurrent attacks of coughing. In our follow-up at 6 months, no symptoms were found in 80% of the cases, 10% had attacks of cough, and 10% had recurrent respiratory tract infection.

No death occurred after 6 months of follow-up in our study. Ozcelik et al [2003] reported a survival rate of 88.5%; in Lincoln et al [1971], 79%; in Davis et al [1979], 88%; and Nazem et al [2010], 87%. Nazem et al [2010] referred to the number of affected lobes and base deficit at presentation.

Limitations

This study may be limited by the small number of patients and being retrospective in design.

Conclusion

Congenital lobar emphysema (CLE) is an important cause of respiratory distress in neonates and infants and should be put in the scope and mind of surgical practice. The diagnosis of CLE is established by combined evaluation of history, clinical examination, and radiological imaging and especially benefits from the accuracy of CT scan, the gold standard for diagnosis. CLE varies greatly in its presentation. The earlier the clinical presentation, the more severe the lesion and the greater the need for urgent surgical intervention. We emphasize that early surgical intervention (lobectomy) in mild and moderate disease is better than waiting for occurrence of complications. It is the appropriate treatment, and CLE is highly curable with lobectomy.

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