Congenital lobar emphysema is a rare disease, which affects newborns and infants. It is characterized by air trapping of lung. The incidence is approximately 1 in 70000 to 1 in 90000 live births. Although in 50% of the cases, etiology is miscellaneous; congenital deficiency of the bronchial cartilage, external compression by vascular abnormalities, congenital Cytomegalovirus (CMV) infection, bronchial stenosis, and bronchial mucosal flaps may be responsible for the etiology. Congenital lobar emphysema may be presented with tachypnea, tachycardia, and retraction. Although patients usually admit to hospital with respiratory symptoms; hypertension is rarely reported in the literature. This case of CLE, which represented atypically with respiratory distress accompanied with hypertension is discussed.

Case Report. A 3600 g male infant with a gestational age of 39 weeks was born by spontaneous vaginal delivery without any postnatal complications. When he was 20-days-old, he admitted to hospital with tachypnea (respiratory rate >60 /minutes) and respiratory distress. He was diagnosed as bronchiolitis, received intravenous teicoplanin, cefotaxime, and clarithromycin treatment. Chest x-ray showed hyperinflation of upper left lobe. He was referred to our clinic due to increased respiratory distress symptoms. Physical examination revealed respiratory rate of 65/min, heart rate of 185 beats/min, blood pressure of 130/70 mm Hg (normal range 95/55 mm Hg - 75/37 mm Hg), body temperature of 36.1 C°, and pulse oxygen saturation of 85% at room air. In addition, subcostal and intercostal retractions, decreased air flow entry in both lungs, bronchospasm and 1/6 sistolic ejection murmur at the left second intercostal space were detected. There

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weren't any sign of cyanosis, jaundice, or heart failure. Other systemic examination was normal. Laboratory analysis revealed a hemoglobin level of 12.1 g/dl, white blood cell (WBC) count of 10300/mm³, platelet count of 626000/mm³, serum level of C-reactive protein <1 mg/l, and erythrocyte sedimentation rate of 2 mm/h. Liver function tests, blood urea nitrogen, and serum electrolyte levels were normal. Carbon dioxide retention was detected on the blood gas analysis. Increased aeration of left upper lobe and significant mediastinal shift to the right were observed on the chest x-ray (Figure 1). Echocardiographic examination was normal. He received inhaled salbutamol, intravenous methylprednisolone, intravenous cefoperazone sulbactam, oral amlodipine, and inhaled oxygen treatment with hood. Despite antihypertensive treatment, blood pressure levels remained high. On the computed thorax tomography, emphysematous change was detected in the upper left lobe, which was considered as CLE (Figure 2). Bronchoscopic examination showed focal hyperinflation in the left upper lobe and lingula, mild pneumothorax at right lung, and bilateral atelectasis at posterior portion of lung. Alpha-1 antitrypsin and Cytomegalovirus (CMV) Ig M levels that are used for identifying the etiology of emphysematous changes were negative. In addition, no etiological agents were detected by polimerase chain reaction analysis of respiratory tract secretions. Hypertension accompanied respiratory distress at the same time. The upper left lobectomy was successfully performed by pediatric surgeons. Pathological examination of the tissue revealed emphysematous changes. On follow-up no sign of respiratory distress occurred. Analgesic treatment for pain relief and amlodipine for hypertension were given. Intravenous cefoperazone sulbactam treatment was ceased at the postoperative sixth day. During the postoperative period, the patient's blood pressure levels remained normal. Control chest x-ray demonstrated air trapping of upper portion of the left lung decreased, and right shift of mediastinum regressed (Figure 3). Despite discontinuation of antihypertensive therapy, patient was normotensive. He was discharged on the postoperative seventeenth day. Informed consent is obtained from parents of the cases described.

**Discussion.** Congenital lobar emphysema is a rare congenital abnormality characterised by excessive aeration of the lungs. Abnormal interactions between embryonic endodermal and mesodermal components can cause progressive pulmonary hyperinflation. Massive over distension of the affected lobe and subsequent compression of surrounding structures usually result with respiratory distress. Although any lobe can be

![Figure 1 - Preoperative chest x-ray showed increased aeration in the left lobe and mediastinal shift to the right](image1)

![Figure 2 - Emphysematous changes were seen on the thorax CT](image2)

![Figure 3 - Postoperative chest x-ray demonstrated healing of right mediastinal shift and midline located trachea.](image3)
Congenital lobar emphysema is a rare disease that may cause a wide spectrum of clinical signs, such as tachypnea, tachycardia, and respiratory distress. In addition, it may cause hypertension. As a result; clinicians should consider CLE at the differential diagnosis in the presence of simultaneous occurrence of respiratory distress with hypertension.

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