Percutaneous Drainage for Giant Pulmonary Interstitial Emphysema in a Tiny Infant with a Birth Weight of 327 g

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Abstract

Giant pulmonary cyst in extremely low birth weight (ELBW) infants has been described as one of severe pulmonary diseases. Any definitive therapy for refractory cases, where conservative methods of treatments are not effective, has not been established as a standard. Herein, we report an ELBW infant with a giant pulmonary cyst cured by percutaneous drainage without any adverse events. A female infant was born with a birth weight of 327 g. Surfactant was administered on days 1 and 2 of life to treat respiratory distress syndrome. Tracheal intubation was performed and synchronized intermittent mandatory ventilation was promptly initiated following birth. On the course, right giant pulmonary cyst developed on day 9 after birth. Although we started conservative therapy, including right lateral decubitus positioning, high-frequency oscillatory ventilation, and systemic corticosteroid administration, the diameter of the cyst had reached 34 mm, and mediastinal displacement was observed on day 28 after birth when she weighed 393 g. She recovered by percutaneous drainage followed by suction with a pressure of –10 cm H2O under mild sedation for 3 days. We believe that percutaneous drainage can be one of the available options for unilateral pulmonary interstitial emphysema.

Keywords
► drainage
► extremely low birth weight infant
► pulmonary interstitial emphysema
► respiratory

Case Presentation

A female infant with a birth weight of 327 g was delivered in the 23rd gestational week by emergency cesarean section with percutaneous drainage without any adverse events. This study was approved by the ethics committee of Nagano Children’s Hospital, and written informed consent was obtained from the patient’s parents for publication.
owing to exacerbation of preeclampsia, so that antenatal steroids were not administered. Her mother, gravida 2, para 1, without premature rupture of membranes was treated for hypertension during pregnancy, and restricted fetal growth had been diagnosed. Apgar scores were 1 at 1 minute and 2 at 5 minutes. No abnormalities were observed during general examination, and there were no evidence of meconium staining nor chorioamnionitis. Placental inflammation was not diagnosed pathologically. Tracheal intubation was performed, and synchronized intermittent mandatory ventilation (SIMV) was promptly initiated following birth. Surfactant was administered on days 1 and 2 of life to treat respiratory distress syndrome. A pulmonary cyst emerged in right lung on day 9 (Fig. 1A), accompanied by poor oxygenation and ventilation (fraction of inspired oxygen [FiO2] 0.4 to keep oxygen saturation [SpO2] > 94% on SIMV at positive end-expiratory pressure 7 cm H2O and peak inspiratory pressure [PIP] 18 cm H2O). Conservative therapy was started, including right lateral decubitus positioning, HFOV at an inspiratory to expiratory ratio of 1:2 to reduce gas leak (from day 14 after birth to avoid high pressure ventilation), and systemic corticosteroid administration: hydrocortisone 5 mg/kg/d for 2 days, tapered to 1 mg/kg/d every 2 days. However, the cyst continued to enlarge. We again changed the ventilation mode from HFOV to SIMV at PIP 22 on day 21 after birth to deal with frequent desaturation.

On day 28 after birth, the infant weighed 393 g. On the course, the setting of PIP was required to be 25 mm Hg, and diameter of the cyst had reached 34 mm and mediastinal displacement was observed, indicating pulmonary interstitial emphysema (PIE).

After informed consent of the parents of the infant was obtained on day 29, we punctured the cyst guided by X-ray under local anesthesia, with xylocaine 1% in addition to midazolam and fentanyl injection. Considering the physique of the infant and suction efficiency, we placed a 20-gauge venous catheter with needle (Terumo Corp., Tokyo, Japan) via the fifth intercostal space on mediastinal axillary line in a dorsal direction (Fig. 1B). After confirming the successful discharge of air by manual suctioning, we started suction drainage with a pressure of −10 cm H2O under mild sedation with the injection of phenobarbital, as needed. On the course of continuous drainage, the cyst gradually diminished to a negligible size, and blood SpO2 improved (FiO2 0.3 on HFOV at mean airway pressure [MAP] 14 cm H2O) at day 3 after the procedures. Consequently, we removed the catheter on the same day (Figs. 1C and 2). The pulmonary cyst did not recur, and no adverse event was observed. This infant was discharged to home without oxygen on 201 days.

**Discussion**

We reported the possibility of percutaneous drainage for PIE in an infant who weighed 393 g when the procedure was performed. The pulmonary cyst of the infant did not recur, and no adverse event was observed.

When physicians treat PIE, conservative therapy is generally considered at first. For example, positioning the infant with the affected side down determined to collapse lung, minimal chest physiotherapy, and endotracheal suctioning, and if possible, decreasing ventilator pressure and inspiratory times. Additionally, HFOV may be effective to avoid higher MAP. If PIE is resistant to conservative therapy, more invasive regimens, such as selective intubation, lobectomy, or pneumonectomy, are reported. Conversely, there are a few case reports on percutaneous drainage for PIE.

The cause of PIE is leaked gas trapped in the interstitium from alveoli. Especially, preterm infants have a higher risk of PIE as their perivascular connective tissues are abundant and relatively more compliant. Additionally, atelectasis in respiratory distress syndrome and plugged small airways in meconium aspiration syndrome can increase the risk of air leak because the ventilation in lung can be uneven. We hypothesized that percutaneous drainage would be also effective for PIE based on its mechanism exhibited by trapped air in thorax like in pneumothorax. Especially, our case was advantageous owing to sufficient size of the cyst enabling drainage and its proximity to thoracic wall on X-ray. In a previously reported case, the cyst was multilobulated, so that physicians languished to perform drainage. As for the device, we could not clarify but 24-gauge peripheral intravenous catheter might have been used.

The infant in our case had considerably smaller weight compared with those previously reported cases of drainage.

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**Fig. 1** X-rays of the patient’s giant pulmonary cyst. (A) Day 9 at the time of emergence of PIE. (B) Day 29 at the time of cyst puncture. (C) Day 32 at the time of catheter removal.
Further research is required to determine the optimum approach for this procedure in terms of adequate patient’s condition, concrete device, and appropriate suction pressure.

**Conclusion**

We believe that percutaneous drainage is a potentially effective treatment for ELBW infants when refractory PIE leading to breathing and circulation failure is diagnosed.

**Conflict of Interest**

None declared.

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**References**


![Fig. 2](image-url) Course of RSS (consisting of mean airway pressure multiplied by FiO2). FiO2, fraction of inspired oxygen; HFOV, high-frequency oscillatory ventilation; PIE, Pulmonary interstitial emphysema; RSS, respiratory severity score; SIMV, synchronized intermittent mandatory ventilation.