Acquired diaphragmatic hernia in pediatrics after living donor liver transplantation

Three cases report and review of literature

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Abstract
Rationale: Diaphragmatic hernia (DH) in pediatrics following living donor liver transplantation (LDLT) has been seldom reported in the past.

Patient concerns: We report successful diagnosis and treatment of three pediatric cases with DH secondary to LDLT, discuss the possible etiology, and review the relevant literature.

Diagnoses: The primary disease was biliary atresia and DH was diagnosed by computed tomography scan or x-ray of chest.

Interventions: Laparotomy was performed successfully to repair the DH.

Outcomes: The respiratory and digestive function was gradually recovered in 1 to 2 weeks after repair operation. In 2 to 8 months follow-up, patients were asymptomatic without any respiratory or digestive complications.

Lessons: DH post-LDLT should be recognized as a possible complication when a left lateral segment graft is used. Careful clinical examination and prompt surgery could minimize complications.

Abbreviations: BA = biliary atresia, CT = computed tomography, DH = diaphragmatic hernia, ESLD = end-stage liver diseases, GRWR = graft weight/recipient body weight ratio, LDLT = living donor liver transplantation, LE = laparoscopic exploration, LLS = left lateral segment, LT = liver transplantation, MP = methylprednisolone, Tac = tacrolimus.

Keywords: diaphragmatic hernia, living donor liver transplantation, pediatric

1. Introduction
Since donor shortage became more severe for patients with end-stage liver diseases (ESLD), liver transplantation (LT) with partial grafts, including living donor liver transplantation (LDLT) and split LT, is being paid more attention, due to its advantage of significantly shortening waiting time.[1] Diaphragmatic hernia (DH) is one of the rare complications occurred in pediatric LT, accompanying with multiple patho-physiological factors.[2] DH is usually urgent and requires speedy intervention by surgical treatment. From August 2000 to April 2017, 680 pediatric patients have received LT in our center, and more than half received LDLT. Among which, 3 patients suffered from DH. We, here, report successful diagnosis and treatment of the 3 pediatric cases with DH secondary to LDLT and review the literatures to investigate and analyze the possible etiology.

2. Cases report
In total 680 pediatric patients, 471 patients received partial graft LT, in which 407 patients received LDLT and 64 patients received split LT. In pediatric LDLT, 3 cases were diagnosed of DH. The primary diagnosis in these children were biliary atresia (BA) and all of them were performed LDLT at the age of <1 (Table 1). Two of the 3 patients received Kasai operations pre-LDLT. Case 1, 2, and 3 were 9-month-old male, 6-month-old male, and 2-year-old male, who received LDLT from father, mother, and father at 6-month-old, 3-month-old, and 7-month-old, respectively. All LDLT procedures and the postoperative courses were uneventful.

All subjects signed the Informed Consent Form. The study protocol was approved by the Ethics Committee of the hospital and the study was also conducted in accordance with the principles delineated in the Declaration of Helsinki.

The authors have no conflicts of interest to disclose.

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mosed to grafts’ portal vein end to end. Roux-en-Y hepaticojejunostomy was performed, no matter whether the recipients had received Kasai operation pre-LDLT or not.

Immunosuppression regimen was tacrolimus (Tac) combined with methylprednisolone (MP) in the first 3 months post-LDLT, then followed by Tac mono-therapy. No complications were found prior to DH.

Three pediatric recipients were diagnosed with DH post-LDLT (Table 2). Case 1 and case 2 were diagnosed with DH in <3 months, whereas DH was found in case 3 at 16 months post-LDLT. The symptoms of DH in case 1 were mainly respiratory distress with emergent cyanosis. In the early stage of DH, the degree of blood oxygen saturation was down to 70%, and could be relieved by oxygen inhalation. However, respiratory distress and cyanosis symptom reappeared without oxygen therapy. Right DH was confirmed by urgent chest computed tomography (CT) scan. Other symptom in case 1 included mild dyspepsia for >2 weeks. The symptom of DH in case 2 was early stage mild dyspepsia. The patient was diagnosed as reversed stomach by chest x-ray at the beginning and conservative therapy was adopted for several months. Twelve months later, gastrointestinal symptoms deteriorated, accompanied by anepithymia and vomiting, left DH was then diagnosed in the operation. Case 3 recovered successfully in the following year post-LDLT but presented symptoms of digestive obstruction, such as high fever, frequent vomiting, grievous abdominal distention, and pain. When patient was admitted to hospital, CT scan showed right DH.

Three pediatric patients were performed DH repair operations followed by chest CT scan or x-ray diagnosis (Fig. 1). All surgical repairs were performed by laparotomy along the incision of LT. A 2cm × 2cm and a 2cm × 3cm post-medial defect were found in the right diaphragm in case 1 and case 3, respectively. In case 2, a 2cm × 2cm front medial defect was found in the left diaphragm. Defect in case 1 was extended laterally to return the herniated content back into abdomen, while the herniated contents in case 2 and case 3 were easily returned without extending the defect. Herniated contents included partial intestines among all 3 pediatric patients and partial colon in case 1, proximal stomach in case 2, distal stomach and intestinal loop in case 3, respectively. In case 1 and case 2, the bowel looked viable, however, the herniated intestinal loop in case 3 showed obvious sign of ischemic damage, and Roux-en-Y hepaticojejunostomy was reperformed. Defects in all cases were closed by primary closure without mattresses. The 2 right DH patients were each inserted with a chest tube, whereas no tube was placed in the left DH patient.

Postoperative courses were uneventful, and respiratory and digestive function was gradually recovered in 1 to 2 weeks post repair operation for all 3 cases. At 2 to 8 months follow-up, patients were asymptomatic, without any respiratory or digestive complications.

### 3. Discussion

The most common surgical complications after pediatric LDLT are biliary stricture, biliary leakage, portal vein complications, and hepatic artery thrombosis.[13] DH post-LT is very rare, however, was mostly found in pediatric recipient. Close to 30 cases (27 cases) of DH post-LT in pediatrics have been reported in recent 10 years (Table 3).[3,4-11] All reported patients received grafts on the left side, including left lateral segment (92.6%) and left lobe (7.4%). As to the site of DH, it was more frequent to be on the right side (25 cases, 88%), compared with only 1 case on the left side and 1 case on both sides. In our hospital, the morbidity of pediatric DH post-LDLT was 0.74% (3/407), which is similar to that in other transplant centers. Although the incidence of DH is lower than other complications, the effect of DH can be dramatic and should be paid more attention and be disposed in emergency.

The potential risk factors of DH post-pediatric LDLT include surgical trauma during hepatectomy, condition of left lateral grafts, increased pressure of intra-abdomen, thin musculature of the diaphragm, malnutrition, and immunosuppressants administration.[12] Receiving partial grafts is common, especially for LLS in pediatric LDLT. Majority of reported DH have been left lobe

### Table 1

Demographic data of patients in DH pre-, intra-, and post-LDLT.

| Case | Sex | Age at LDLT | Diagnosis | Previous surgery | BMI | PELD score | Graft type | Operation types of LDLT | Total operation time | Abdominal closure | Immunosuppressant |
|------|-----|-------------|-----------|------------------|-----|------------|------------|------------------------|---------------------|-----------------|-----------------|
| 1    | M   | 6 mo       | BA        | None             | 19.08 kg/m² | 24 | LLS 2.7%   | Piggy-back | Formal Tac + MP        | 520 min            | Formal Tac + MP  | Tac + MP        |
| 2    | F   | 5 mo       | BA        | LE               | 14.07 kg/m² | 11 | LLS 2.5%   | Piggy-back | Formal Tac + MP#       | 470 min            | Formal Tac + MP# | Tac + MP#       |
| 3    | M   | 7 mo       | BA        | LE               | 19.40 kg/m² | 23 | LLS 3.0%   | Piggy-back | Formal Tac + MP#       | 535 min            | Formal Tac + MP# | Tac + MP#       |

**Table 2**

Clinical characteristics of pediatric patients related to DH.

| Case | Chief complaints | Period between LDLT and DH | Period between DH and DH repair operations | Site | Operation of DH | Follow-up period post-DH | Prognosis |
|------|-----------------|---------------------------|------------------------------------------|------|-----------------|--------------------------|-----------|
| 1    | RD              | 3 mo                      | In emergency                             | Right posterior medial                  | Laparotomy                 | 10 mo                    | No recurrence |
| 2    | GS              | 1 mo                      | 13 mo later                              | Left front medial                      | Laparotomy                 | 9 mo                     | No recurrence |
| 3    | GS              | 16 mo                     | In emergency                             | Right posterior medial                  | Laparotomy                 | 4 mo                     | No recurrence |

**Note:** DH = diaphragmatic hernia, GS = gastrointestinal symptoms, LDLT = living donor liver transplantation, RD = respiratory distress.
or LLS transplantation which resulted in right diaphragm being unprotected and susceptible to overextension. Severe adhesion between diaphragm and diseased liver formed by previous Kasai operations or injury of bare area caused by dissection might have led to delayed necrosis and perforation.\textsuperscript{[6,13]} Similarly, DH occurred in some cases which were only performed with hepatectomy without LT.\textsuperscript{[14,15]} Furthermore, pediatric recipients were malnourished in general, mostly suffered with hyperbilirubinemia, and were more prone to be injured with delayed healing. It has been confirmed that immunosuppressants, such as steroid or macrolide immunosuppressants, could impair healing post-LT.\textsuperscript{[16]} and Rossetto et al\textsuperscript{[17]} had reported DH post-LT was related with mammalian target of rapamycin. To our experience, surgical trauma in hepatectomy, especially to the bare area, and malnutrition in perioperative period were the main causes of DH.

The most common complaints in DH of pediatric recipients post-LT were respiratory distress, as well as symptoms of gastrointestinal obstruction, including abdominal pain, nausea, and vomiting.\textsuperscript{[2,4–11]} However, DH was not found until respiratory distress or gastrointestinal obstruction. So it was speculated that the trauma of diaphragm happened more earlier than hernia itself in which the hernia hole was enough to afford gastro-intestine through it. In our data, the diameter of hernia hole was 2 cm at least. Earl et al\textsuperscript{[18]} reported an incidence of small bowel obstruction of 3.2\% after orthotopic LT in the pediatric population, with diaphragmatic hernias being responsible for nearly half of the cases and lymphoproliferative disease for the remainder. Besides patients’ statements, diagnosis of DH is usually confirmed by chest x-ray or CT scan. CT scan can also accurately detect any major vasculature and adhesions in abdomen post-LT.\textsuperscript{[9]}

As to the therapy of DH, it has been agreed that once DH is identified, operation should be performed immediately. Delayed treatment could result in disorder of cardiac and pulmonary function, loss of bowel, and/or severe abdominal infection. Laparotomy was the routine therapy. In 2015, Lee et al and Yeung et al had reported 3 cases of DH, and 1 case was performed by thoracoscopy.\textsuperscript{[10,11]} The courses of DH after surgical repairing were mostly uneventful without any surgical complications. Our experience showed that pediatric patients with DH should receive operation in time, even if the symptoms related to DH are mild. As to the approach of operative intervention, we suggest laparotomy to be the primary option, considering possible obstruction of digestive tract and likely adhesions of abdominal tissues post-LDLT.

Figure 1. Chest computed tomography scan or x-ray of the patients pre-, intra-, and post-repair operation of DH (case 1 and case 3 were both right DH and case 2 was left DH). 1. Arrows head the position of DH intra-operation. 2. Words on the top of the columns means the time of images. 3. Words on the left of the rows means the order of patients.
### Table 3
**DH in pediatric recipients post-LT reported in literature.**

| Authors         | Year | Patients (n) | Period between LT and DH | Previous surgery | Graft type | GRWR (%) | Chief complains                                      | Period from DH to repair operation | DH location                                      | Content in hernia                                      | Follow-up (m) |
|-----------------|------|--------------|--------------------------|------------------|------------|----------|-----------------------------------------------------|-----------------------------------|---------------------------------------------------|------------------------------------------------------|---------------|
| Okajima et al[4] | 2007 | 1            | 2 mo                     | No               | LLS        | 1.35     | Abdominal pain, vomiting, respiratory distress      | Emergency (5 mo later)            | Right postero-lateral                             | Convoluted ileum                                   | 15            |
| Kazimi et al[5] | 2010 | 2            | 3 mo, 14 mo              | No               | LLS, left lobe | 0.94, 0.77| Abdominal pain, nausea, vomiting, respiratory distress, high fever | Emergency, –                       | Right-side                                       | Small bowel                                         | 6, 4 y        |
| Shigeta et al[6] | 2012 | 3            | 42 d, 8 d, 3 mo          | 2 Kasai          | LLS, reduced LLS | 3.21, 2.39, 2.57| Respiratory distress, vomiting, bowel obstruction | Emergency, 99 d later               | Right postero-lateral, right postero-medial | Small bowel, transverse colon                     | 15, 1 yr, –   |
| Moon et al[7]   | 2012 | 4            | 28 d, 7 mo, 46 d, 81 d   | 2 Kasai          | LLS        | 1.62, 2.52, 1.55, 3.77 | Abdominal pain, vomiting, respiratory distress, headache, dizziness, anoxia | 1 d, 14 d, emergency                   | Right postero-medial                               | Colon, part of the small bowel and part of ileum | 1 y, 3, 1, –  |
| Lam et al[8]    | 2013 | 1            | 3 m                      | No               | Left lobe | –         | Abdominal pain, anoxia, respiratory distress        | –                                 | Right postero-lateral                               | Small bowel, a portion of the Roux limb            | 2 y           |
| Cortes et al[2] | 2014 | 10           | 3.864 y (57–5501 d)      | –                | LLS        | 3.0 ± 1.22 (1.7–6.0) | Small bowel obstruction                          | Emergency                          | Right postero-lateral, with a left DH             | Small bowel                                         | 3.5 y         |
| Dökümçü et al[9] | 2015 | 1            | 10 mo                    | Laparoscopic cholangiography | LLS | 2.1 | Tachypnea, respiratory distress                      | Emergency                          | Right postero-lateral                              | Ileal segments, ascending colon                     | 2 y           |
| Yeung et al[10] | 2015 | 3            | 15 mo, 9 mo, 15 mo       | 1 Kasai          | Reduced LLS, LLS | 4.4, 2.4, 2.5 | Incidental, repeated vomiting, respiratory distress | 10 mo later, emergency, –             | Right cardiophrenic region, right postero-medial   | Large bowel, mesenteric fat, terminal ileum         | –             |
| Lee et al[11]   | 2015 | 3            | 5 mo, 2 mo, 28 mo        | –                | LLS        | 4.6, 4.8, 3.26 | Vomiting, decreased activity, high fever, respiratory distress | Emergency                          | Left medial, right side                            | Stomach, small bowel                                | 42, 5, 8      |

DH = diaphragmatic hernia, GRWR = graft weight/recipient body weight ratio, LLS = left lateral segment, LT = liver transplantation.
4. Conclusions

DH post-LDLT is unusual, however, should be recognized as a possible complication when a left lateral segment graft is used. In some cases, DH is found acutely and can even be life-threatening. Diagnosis can easily be made with chest x-ray or CT scan. A high index of suspicion and prompt surgery could minimize complications. When unexplained respiratory or gastrointestinal symptoms after LDLT occurred in pediatric patients, DH should be highlighted in differential diagnosis.

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Author contributions

Wei Gao proposed the study. Kai Wang performed research, analyzed the data, and wrote the first draft. Nan Ma, Xing-Chu Meng, Wei Zhang, Chao Sun, Chong Dong, Bin Wu helped to collect the data. All authors contributed to the performance of LDLT, the repairing operations of DH, the design and interpretation of the study and to further drafts. Wei Gao is the guarantor.

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References

[1] Lauterio A, Di Sandro S, Concone G, et al. Current status and perspectives in split liver transplantation. World J Gastroenterol 2015;21:11003–15.
[2] Cortes M, Tapuria N, Khorsandi SE, et al. Diaphragmatic hernia after liver transplantation in children: case series and review of the literature. Liver Transpl 2014;20:1429–35.
[3] Neto JS, Pugliese R, Fonseca EA, et al. Four hundred thirty consecutive pediatric living donor liver transplants: variables associated with posttransplant patient and graft survival. Liver Transpl 2012;18:577–84.
[4] Okajima H, Hayashida S, Iwasaki H, et al. Bowel obstruction due to diaphragmatic hernia in an elderly child after pediatric liver transplantation. Pediatr Transplant 2007;11:324–6.
[5] Kazimi M, Iibs C, Alper I, et al. Right-sided diaphragmatic hernia after orthotopic liver transplantation: report of two cases. Pediatr Transplant 2010;14:e62–4.
[6] Shigeta T, Sakamoto S, Kanazawa H, et al. Diaphragmatic hernia in infants following living donor liver transplantation: report of three cases and a review of the literature. Pediatr Transplant 2012;16:496–500.
[7] Moon SB, Jung SM, Kwon CH, et al. Posterior mediastinal diaphragmatic hernia following pediatric liver transplantation. Pediatr Transplant 2012;16:E106–9.
[8] Lam HD, Mejia J, Soltys KA, et al. Right diaphragmatic hernia after liver transplant in pediatrics: a case report and review of the literature. Pediatr Transplant 2013;17:E77–80.
[9] Dökücü Z, Divarci E, Erdener A, et al. Acquired right diaphragmatic hernia following pediatric living donor orthotopic liver transplantation. Pediatr Transplant 2015;19:E149–51.
[10] Yeung F, Chung PH, Wong KK, et al. Iatrogenic diaphragmatic hernia in pediatric patients. Pediatr Surg Int 2015;31:389–92.
[11] Lee S, Soo JM, Younes AE, et al. Thoracoscopic approach for repair of diaphragmatic hernia occurring after pediatric liver transplantation. Medicine (Baltimore) 2015;94:e1376.
[12] McCabe AJ, Orr JD, Sharif K, et al. Right-sided diaphragmatic hernia in infants after liver transplantation. J Pediatr Surg 2005;40:1181–4.
[13] Suh Y, Lee JH, Jeon H, et al. Late onset iatrogenic diaphragmatic hernia after laparoscopy-assisted total gastrectomy for gastric cancer. J Gastric Cancer 2012;12:49–52.
[14] Vernadakis S, Paul A, Kykalos S, et al. Incarcerated diaphragmatic hernia after right hepatectomy for living donor liver transplantation: case report of an extremely rare late donor complication. Transplant Proc 2012;44:2770–2.
[15] Soufi M, Meillat H, Le Trait YP. Right diaphragmatic iatrogenic hernia after laparoscopic fenestration of a liver cyst: report of a case and review of the literature. World J Emerg Surg 2013;8:2.
[16] Kahn J, Muller H, Iberer F, et al. Incisional hernia following liver transplantation: incidence and predisposing factors. Clin Transplant 2007;21:423–6.
[17] Rossetto A, Baccafan D, Calliari M, et al. Diaphragmatic rupture in a liver transplant patient under chronic immunosuppressive therapy with sirolimus: rare complication after liver transplantation. Updates Surg 2001;63:51–3.
[18] Earl TM, Wellen JR, Anderson CD, et al. Small bowel obstruction after pediatric liver transplantation: the unusual is the usual. J Am Coll Surg 2011;212:62–7.