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<td>Aokage, Keiju; Date, Hiroshi; Okazaki, Megumi; Sano, Yoshifumi; Oto, Takahiro; Kusano, Kengo; Goto, Keiji; Sano, Shunji; Miyoshi, Shinichiro</td>
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Kyoto University
LIVING-DONOR LOBAR LUNG TRANSPLANTATION AND CLOSURE OF ATRIAL SEPTAL DEFECT FOR ADULT EISENMENGER’S SYNDROME

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ABSTRACT

A 38-year-old woman with Eisenmenger’s syndrome underwent bilateral living-donor lobar lung transplantation and simultaneous closure of atrial septal defect. The grafts were a right lower lobe from her husband and a left lower lobe from her brother. Although only two lobes were implanted, the dramatic improvement in pulmonary hemodynamics has been well maintained for more than 5 years. Living-donor lobar lung transplantation and simultaneous cardiac repair may be one of the therapeutic options for adult Eisenmenger’s syndrome with simple congenital heart disease.
INTRODUCTION

Lung or heart-lung transplantation remain the treatment of last resort for adult patients with Eisenmenger’s syndrome who fail medical therapy. Living-donor lobar lung transplantation (LDLLT) was developed to mismatch between supply and demand for those patients awaiting cadaveric lung transplantation. We report the first successful LDLLT and simultaneous closure of atrial septal defect (ASD) for adult Eisenmenger’s syndrome.

CASE REPORT

An adult female patient was diagnosed with ASD in her childhood but did not receive cardiac repair. In 1996, at the age of 31 years old, she delivered a normal baby. Soon after the delivery, symptoms of dyspnea occurred during exercise and she was diagnosed with Eisenmenger’s syndrome. In 2001, at the age of 36, she further developed pulmonary thromboembolism and pulmonary infarction. In 2003, at the age of 38, she became wheel chair bound and transferred to Okayama University Hospital for possible lung transplantation. On admission, she was markedly cyanotic and her arterial blood gas revealed a pH of 7.39, PaO₂ of 35.0 mmHg, and PaCO₂ of 36.2 mmHg, with 3 L/min oxygen administration via a nasal cannula. Chest X-ray showed dilated heart and pulmonary artery (PA) shadow (Figure 1A). The right heart catheterization performed a year ago revealed highly elevated pulmonary arterial pressure (systolic/diastolic/mean: 115/42/72). Two-dimensional echocardiography demonstrated a dilated dyskinetic right ventricle and D-shaped left ventricle. By color Doppler echocardiography, bidirectional shunt through the atrial septal defect with a diameter of 2.8 cm and massive tricuspid regurgitation were demonstrated. Enhanced CT scan showed enlarged pulmonary artery (PA), pulmonary thromboembolism and pulmonary infarction.

The surgical options were discussed with the patient and the family, and they were willing to proceed with LDLLT, which they understood to be the first case for adult
Eisenmenger's syndrome. The case was carefully discussed and approved by the Lung Transplant Evaluation Committee at Okayama University Hospital.

On April 30, 2003, she underwent LDLLT at the age of 38, with a right lower lobe from her husband (36 years old) and a left lower lobe from her brother (34 years old). The height and weight was 146 cm, 40.2 kg for the recipient, 178 cm, 93.0 kg for the husband, and 169 cm, 74.0 kg for the brother. The total force vital capacity (FVC) of the two grafts was estimated to be 2,352 ml, or 89.8% of the recipient’s predicted FVC according to the formula we have proposed.2

Because the patient’s hemodynamics was unstable, cardiopulmonary bypass was instituted with bicaval venous drainage before completing the bilateral hilar dissection. Under cold cardioplegic arrest, the right atrium was opened. We closed the ASD, 25×30 mm, with a pericardial patch. After the aortic clamp was removed, bilateral pneumonectomy was performed. Pulmonary thromboembolism was seen bilaterally distal to the first branch of PA which was removed at pneumonectomy. Two lobes were then implanted in the manner previously described.1 The PA anastomosis was challenging due to marked size discrepancy between the recipient’s main PA and the donor’s lobar PA. The recipient PA was partially closed by placing tacking stitches and then the end-to-end anastomosis was carried out. Cardiopulmonary bypass could be removed without any sign of lung edema after reperfusion.

Postoperative immunosuppression was a triple drug therapy consisting of tacrolimus, mycophenolate mofetil and prednisone. Two episodes of acute rejection were successfully treated by steroid pulse. The patient was completely weaned from the respirator on day 22 and discharged from the hospital on day 48 without oxygen inhalation.

The dramatic improvement in pulmonary hemodynamics was seen one month after LDLLT and has been well maintained for more than 5 years (Table I). Her
chest X-ray demonstrated well-expanded grafts without cardiomegaly (Figure 1B). Although she developed right unilateral bronchiolitis obliterans syndrome at 4 years, she is able to carry out normal activities for more than 6 years since she received the LDLLT.

Postoperative course was uneventful in both donors and they have returned to their previous lifestyles during the 6-year observation period.

**DISCUSSION**

There are three transplant options for Eisenmenger’s syndrome: single lung transplantation (SLT), bilateral lung transplantation (BLT), and heart-lung transplantation (HLT). Cardiac repair with BLT appears to have a better post-operative course than SLT, but carries potential morbidity associated with longer cardiopulmonary bypass. HLT is a simpler operation than BLT with repair of congenital heart disease, but requires allocation of two organs from the scarce donor organ supply, and adds risk of cardiac graft coronary vasculopathy and is less tolerant of ischemic time.

LDLLT was pioneered by the University of Southern California group to deal with the shortage of cadaveric donors. They originally applied this procedure almost exclusively to cystic fibrosis patients and then expanded the indications to other diagnoses including pediatric patient with pulmonary arterial hypertension (PAH). Since 2000, we have applied this procedure to both pediatric and adult patients with PAH.

It has been suggested that adults with Eisenmenger’s syndrome have a more favorable hemodynamic profile and prognosis than adults with idiopathic PAH. In contrast, the 5-year survival after BLT or HLT remains to be approximately 50% according to the registry of ISHLT. Our patient with Eisenmenger’s syndrome was critically ill due to associated pulmonary thromboembolism. The realistic option for this patient was only to receive LDLLT in Japan, where obtaining brain death
donors is extremely difficult. The ethical concerns of LDLLT present an obvious dilemma. This procedure subjects two healthy donors to a right or left lower lobectomy associated with an expected risk of death between 0.5% to 1%, a complication rate of 10-20%, and the inevitable 15% reduction in pulmonary function. While there have been no deaths in the donor cohort, these disadvantages are to be carefully explained to the potential donors during the process of obtaining informed consent. Because of these ethical concerns, we accept only very sick patients as LDLLT recipients.

Because limited amount of lung tissue is implanted in LDLLT, size matching is an important issue. Matsuda and his colleagues reported successful LDLLT and simultaneous closure of ASD for an 11-year-old boy with irreversible pulmonary hypertension. The present case was the first successful LDLLT for adult Eisenmenger’s syndrome. Luckily, our patient had two male donors who were taller than her and the total FVC of the two grafts was estimated to be 89.9% of the recipient’s predicted FVC. Although only two lobes were implanted, the dramatic improvement in pulmonary hemodynamics has been well maintained for more than 5 years. Although her FEV1 decreased at 5 years due to right unilateral bronchiolitis obliterans syndrome, she is maintaining a good quality of life because her unaffected left graft is functioning normally.

LDLLT and simultaneous cardiac repair may be one of the therapeutic options for adult Eisenmenger’s syndrome with simple congenital heart disease.

Acknowledgement

The authors have nothing to disclose on this article.
REFERENCES


**Figure Legend**

**Figure 1.** Chest X-ray of a 38-year-old female with Eisenmenger’s syndrome. (A) Before transplantation. (B) One year after receiving bilateral living-donor lobar lung transplantation.
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PAP, pulmonary arterial pressure; PCWP, pulmonary capillary wedge pressure
CVP, central venous pressure; PaO₂, arterial oxygen tension; PaCO₂, arterial carbon dioxide tension
VC, vital capacity; FEV₁, forced expiratory volume in one second