Lung Transplantation after Endoscopic Lung Volume Reduction

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Key Words
Lung transplantation · Endoscopic lung volume reduction · Airway colonization · Emphysema

Abstract
Background: Endoscopic lung volume reduction (ELVR) has become an established treatment option in selected patients with end-stage lung emphysema. ELVR, however, does not always prevent disease progression, and patients may inevitably be considered for lung transplantation. Objectives: Currently, limited data exist regarding the impact of preceding ELVR on lung transplantation outcomes. Methods: A retrospective, single-center analysis of lung transplantation (LTx) waiting list candidates, who had previously undergone ELVR for emphysema between 2010 and 2014, was performed. Outcomes were compared to matched (1:2) controls who underwent LTx for emphysema without previous ELVR. The 12-month survival after LTx represented the primary end point. Results: In total 23/693 (3%) patients listed for LTx between January 2010 and May 2014 had undergone ELVR, of whom 20/23 (87%) proceeded to LTx (ELVR group). Forty matched non-ELVR emphysema patients acted as controls. Bronchiectasis on CT prior to LTx was more evident in ELVR patients [11/20 (55%) vs. 12/40 (30%); p = 0.04] as well as airway colonization after LTx [10/20 (50%) vs. 6/40 (15%); p = 0.004]. Among ELVR patients, the most prevalent colonizing organism was Stenotrophomonas maltophilia (4/10 patients, 40%). No significant differences were observed in LTx waiting list time, duration of LTx procedure, ventilatory support, ICU stay after LTx or time to hospital discharge. One ELVR patient (5%) died 189 days after LTx from pneumonia, compared to 1 non-ELVR patient (3%) who died after 269 days (p = 0.61). Conclusions: Previous ELVR treatment was not associated with differing outcomes following LTx. Increased bacterial colonization rates were evident and warrant further investigation.

Introduction
Lung transplantation (LTx) is an established therapy for selected patients with end-stage lung disease [1]. Common conditions requiring LTx are chronic obstructive pulmonary disease (COPD), idiopathic pulmonary fibrosis, cystic fibrosis and pulmonary arterial hypertension [2]. According to current International Society for Heart and Lung Transplantation registry data, COPD represents the leading indication worldwide, accounting for 40% of procedures. In Germany, the pooled prevalence of COPD is 7.6% [3–5] with advanced COPD...
Emphysema is a fundamental component of COPD and is characterized by airspace enlargement, caused by extensive destruction of the alveolar walls, resulting in debilitating hyperinflation. Emphysema is often attributed to smoking, with α1-antitrypsin deficiency being responsible in about 6% of cases [2]. Medical treatment consists primarily of bronchodilators, supplemental oxygen therapy and in advanced cases noninvasive ventilation. Most treatment options improve quality of life but have limited impact on survival.

As an attempt to alleviate hyperinflation, lung volume reduction surgery was proposed as a possible solution some 20 years ago, with acceptable results [6]. This approach employed a buttressed staple excision of hyperinflated, nonfunctioning lung segments, allowing improved function of the remaining lung. Improved exercise ability, quality of life and survival were reported in highly selected candidates (National Emphysema Treatment Trial) [7]. The reported morbidity and an attributable mortality were however considerable at 5%.

In recent years, endoscopic lung volume reduction (ELVR) has emerged in various guises as a less invasive alternative to surgery in these patients [8, 9]. Deployment of one-way valves remains the most popular approach [10–12], with randomized trials reporting benefit in terms of lung function, exercise capacity and potentially survival in highly selected patients with heterogeneous disease and an intact lobar fissure [11–13]. Other ELVR, nonblocking methods include use of sealants [14], thermal therapy and endobronchial coils, which have been the subject of small randomized trials and one multicenter cohort study, which could show improvements in life quality, exercise ability and lung function [15–18]. Endobronchial coaxing attempts to distort segmental airways, restore the tethering effect on small airways, and compress emphysematous lung parenchyma. Another ELVR approach is vapor delivered in targeted areas. Thermal therapy relies upon targeted delivery of hot vapor in an attempt to destroy areas of remodeled lung [18]. No controlled studies regarding this technical approach have been published to date.

Despite the popularity of ELVR, the number of studies remains small. Further evaluation of the value in treating severe hyperinflated emphysema is necessary and the subject of larger ongoing studies. Limited data relating to the potential usefulness of previous ELVR in LTx recipients, with analysis focusing exclusively on the pre-LTx course, exist [19].

**Methods**

A single-center retrospective analysis was performed. Hannover Medical School has a high-volume lung transplant program, routinely performing more than 130 procedures per year. Currently, more than 850 patients actively participate in outpatient LTx follow-up.

Emphysema patients referred for LTx evaluation between January 2010 and May 2014 were reviewed, with those proceeding to transplantation, having previously undergone ELVR, being included. From the remaining emphysema LTx recipients who had not undergone ELVR, a control group matched 1:2 for gender, age and α1-antitrypsin status was generated.

All potential LTx candidates referred to our center undergo multidisciplinary assessment involving surgeons, pulmonologists and psychologists. Prerequisite investigations include extensive laboratory testing to exclude other end-organ disease, sputum samples to assess colonization, a CT of the thorax and vascular imaging to plan perioperative management. Once listed for LTx, candidates were closely monitored at our outpatient clinic. Since December 2011 the lung allocation score (LAS) has been implemented in Germany [20, 21], with relevant data collection having previously been commenced at our center in January 2010. Final LAS values and ΔLAS (LAS last before transplantation – LAS upon listing) were calculated.

Patient and graft survival, duration of transplant procedure, length of ICU and overall hospital stay, airway colonization, quality of life, 6-min walk distance and infection episodes after LTx were analyzed [22]. Quality of life was measured by the 5-dimension Euroqol score [23], from which the visual analog scale was assessed.

After LTx infections requiring hospitalization, infection episodes (detection of a new pathogen followed by anti-infective treatment) and rejection episodes were analyzed.

Our study adhered to ethical guidelines described in the 1975 Declaration of Helsinki. All patients provided informed consent prior to transplantation, using a standardized consent form approved by the local ethics committee, which allowed use of their data for scientific purposes in retrospective analysis.

**Respiratory Sampling**

Bronchoscopy was not routinely performed as part of transplant evaluation. Sputum samples (if possible) were analyzed in each listed candidate before transplantation, and cultures were routinely taken from the recipient bronchi at the time of native lung removal. Following transplantation all patients participated in our long-term surveillance program, which included regular bronchoalveolar lavage and transbronchial biopsies. Colonization was defined as any repeated or intermittent isolation of typical respiratory pathogens [24].

**CT Analysis**

Three lung sections were analyzed representing the lower, middle and upper lung, identified by taking the 75th, 50th and 25th percentile images from the entire number of images in the lung sequence, with the 25th percentile taken to represent the upper lobe. On these reference images, evidence of atelectasis or bronchiectasis was assessed, with the latter being graded using a ‘none’ (0), ‘mild’ (1) and ‘severe’ (2) scale for each individual lung. Reviewers were allowed to examine adjacent CT slices to verify changes. Bronchiectasis was defined as a total score (max. 12) of 3 and more.
Histopathology
All lungs underwent routine histopathological workup following explantation. The formalin-fixed, paraffin-embedded samples were retrieved from the archives of the Institute of Pathology (Hannover Medical School) and re-evaluated by an experienced pulmonary pathologist.

Statistics
Statistical analysis was performed with SPSS 22.0 (SPSS, Chicago, Ill., USA): continuous variables are presented as median and interquartiles. Categorical variables were analyzed by the $\chi^2$ test. Medians were compared with Fisher’s exact test, nonparametric variables with the Mann-Whitney U test. Kaplan-Meier statistics were used to estimate survival rates with statistical significance assessed by the log rank test. For all analyses, $p$ values less than 0.05 were considered statistically significant.

Results
Waiting List and Referrals
During the study period, 1,562 patients were referred for LTx. Of these, 637 had emphysema as underlying diagnosis (40.8%) with 109 having proven $\alpha_1$-antitrypsin deficiency (17.1%). The number of referrals with previous ELVR has been steadily increasing in recent years: in 2010 $n = 4/203$ (2%), in 2011 $n = 13/135$ (10%), in 2012 $n = 15/105$ (15%), in 2013 $n = 17/102$ (17%) and in 2014 $n = 18/92$ (20%). Overall 23/67 patients (34% of referrals) with previous ELVR were listed for LTx.

Since 2010, 23 (4%) patients died awaiting LTx. None of these patients had previously undergone ELVR.

Transplantations
Between 2010 and 2014, 651 LTx were performed at our center, including 20/23 (87%) of the ELVR patients listed (fig. 1). Patient characteristics are displayed in table 1. In all patients, ELVR had been performed at their local referring hospital.

All patients included in the analysis underwent double LTx. Of the 20 ELVR patients transplanted, 18 had received ELVR with valves (right lower lobe: 3, right upper lobe: 6, left upper lobe: 12, right lower lobe: 2 and bilateral upper lobes: 3). In 1 of these valve patients (1/18) thermal therapy (right lower lobe) followed after removal of prior valves. A single patient received coils (combined upper lobes), and 1 further patient received polymer sealant (right lower lobe). The median interval between ELVR and LTx was 2 years. No difference in the last LAS score or exacerbation episodes before transplantation were noted between groups (table 1).

CT Analysis
Of the 18 patients who underwent ELVR with valves, these were visible on CT in 16 cases (89%). In 6/16 (38%) of these, atelectasis of the target lobe was evident. Bronchiectasis was more prevalent in the ELVR group (11/20, 55%) compared to the control group (12/40, 30%; $p = 0.04$; table 1). Although bronchiectasis was identified in nontreated lobes in the majority of patients (13/18, 82%), 5/18 (28%) patients exhibited bronchiectasis in ELVR-treated lobes (fig. 2).
Outcome
No significant differences were observed between ELVR patients and their matched controls with regard to the principal clinical outcome parameters, e.g. transplant duration, days on ventilator support, ICU stay, length of hospital stay or 12-month graft survival (table 2). In both groups, no patients underwent re-do transplantation during follow-up. In the ELVR group, 1 patient died after virus pneumonia and graft dysfunction 189 days after LTx. In the control group, 1 patient died after 279 days due to restrictive allograft dysfunction. There was no survival difference between groups (table 2). There was no difference in terms of 6-meter walk distance, quality of life, infection or rejection episodes during the 12-month follow-up.

Histopathology
All lung explants were re-evaluated, and findings were correlated with the radiological results lined out above. All specimens showed severe emphysema with prominent bronchiectasis in some, correlating with CT findings.
Microbiology Results

Sputum cultures were available in 42/60 (70%) patients (ELVR 16/20). Intraoperative swabs taken from the recipient bronchus were available for all patients. No differences existed between groups regarding airway colonization rates (p = 0.13) prior to transplantation (table 3). After LTx, airway colonization rates were higher among ELVR patients compared to controls [10/20 (50%) vs. 6/40 (15%); p = 0.004]. The most common organism was Stenotrophomonas maltophilia, being identified in 4/10 (40%) ELVR patients. Other organisms isolated are displayed in table 3.

The single patient who received coils was colonized with Acinetobacter baumannii after LTx. The other non-valve patients (polymer sealant and thermal therapy) demonstrated no colonization before or after LTx.

Bronchiectasis (before LTx) was evident in 5/11 (45%) of ELVR patients with known airway colonization (after LTx). These patients were colonized with: 3× S. maltophilia, 1× oxacillin-resistant Staphylococcus aureus, 1× A. baumannii. None of the control group with evidence of bronchiectasis (12/40, 30%; table 1) was colonized (p = 0.008; table 3).

Discussion

To our knowledge, this is the largest study assessing the impact of previous ELVR in emphysema patients proceeding to LTx. Previous ELVR treatment was not associated with significant differences in outcome, but increased rates of airway colonization were observed.

The St. Louis group has previously reported on LTx following lung volume reduction surgery [25]. In the largest series involving 36 patients, Backhus et al. [26] identified both a longer transplantation duration and greater length of hospital stay. Thirty-day mortality and major morbidity rates were unaffected.

EVLR after LTx has been reported by Perch et al. [27], who inserted endobronchial valves in hyperinflated na-
Table 3. Airway colonization

<table>
<thead>
<tr>
<th></th>
<th>ELVR group (n = 20)</th>
<th>Control group (n = 40)</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Any colonization before LTx</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Stenotrophomonas maltophilia</em></td>
<td>5 (25)</td>
<td>4 (10)</td>
<td>0.13</td>
</tr>
<tr>
<td>ORSA</td>
<td>1</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>MSSA</td>
<td>1</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td><em>Escherichia coli</em></td>
<td>1</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td><em>Proteus mirabilis</em></td>
<td>1</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td><em>Enterobacter aerogenes</em></td>
<td>1</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Any colonization after LTx</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Stenotrophomonas maltophilia</em></td>
<td>10 (50)</td>
<td>6 (15)</td>
<td>0.004</td>
</tr>
<tr>
<td><em>Pseudomonas aeruginosa</em></td>
<td>4</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>ORSA</td>
<td>1</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>MSSA</td>
<td>1</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td><em>Klebsiella pneumoniae</em></td>
<td>1</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td><em>Escherichia coli</em></td>
<td>1</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td><em>Acinetobacter baumannii</em></td>
<td>1</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td><em>Proteus mirabilis</em></td>
<td>0</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Different organism colonization</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(compared to pre-LTx n = 5)</td>
<td>7 (35)</td>
<td>5 (13)</td>
<td>0.04</td>
</tr>
<tr>
<td>Bronchiectasis before LTx</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(CT score &gt;2)</td>
<td>11</td>
<td>12</td>
<td></td>
</tr>
<tr>
<td>Any colonization before LTx</td>
<td>3/11 (27)</td>
<td>1/12 (8)</td>
<td>0.23</td>
</tr>
<tr>
<td>Any colonization after LTx</td>
<td>5/11 (45)</td>
<td>0/12 (0)</td>
<td>0.008</td>
</tr>
</tbody>
</table>

ORSA = Oxacillin-resistant *Staphylococcus aureus*; MSSA = methicillin-sensitive *Staphylococcus aureus*. Values are numbers with percentages in parentheses.

The majority of ELVR patients in this series received endobronchial valves (n = 18), of which approximately one third were evident on CT, and atelectasis of the target lobe could be identified. Atelectasis after valve implementation is considered a predictor for clinical benefit from ELVR. Atelectasis itself does not appear to influence subsequent LTx outcome [12]. The low rate of atelectasis indicates a minority of ELVR nonresponders.

In this series, only single patients underwent coiling, polymer sealant or water vapor treatments, which limits the transferability of all techniques. At least treatment with endobronchial coils has recently been shown to be a promising treatment in emphysema patients in a not yet well-defined subgroup of patients [16]. Further evaluation of this treatment strategy and its impact on LTx needs to be evaluated.

Previous studies have shown that de novo airway colonization of the lung allograft with Gram-negative bacteria is associated with subsequent chronic lung allograft dysfunction [24, 28, 29]. Other studies have demonstrated that donor colonization impairs LTx outcomes [30], with the impact of donor-to-host transmission being previously described [31]. In our study, airway colonization was evident in the bronchoalveolar lavage fluid of 50% of EVLR patients after LTx, much higher than would be expected or indeed observed among non-ELVR patients. Gram-negative bacteria predominated, with *S. maltophilia* attracting particular attention (table 3). The majority of ELVR patients had received valves, which potentially increase the risk of colonization, as airway foreign bodies may result in biofilm formation, impeding ciliary clearance.

Potential confounders accounting for airway colonization aside from ELVR may include previous antibiotic treatment in both the recipient and donor, as well as donor transmission. Nevertheless, bronchiectasis was more frequently seen in ELVR patients, although no difference in the incidence of airway colonization existed prior to transplantation between patient groups. CT scans prior to ELVR treatment were unavailable for retrospective evaluation, so the true prevalence of bronchiectasis prior to ELVR or even the possibility of ELVR inducing bronchiectasis remains unclear. Reports associating ELVR with an increased incidence of infective exacerbations as a potential risk factor for pathological colonization do exist [12]. Fruchter et al. [32] have recently published a case series of bacterial colonization in ELVR patients that may potentially explain the higher incidence of bronchiectasis in these. Noncystic fibrosis bronchiectasis is a further, separate risk factor for Gram-negative bacteria colonization.
tion [33] and may possibly overlap with a clinical diagnosis of COPD. Important differences exist in the ability to screen for colonization before and after LTx, with reliance on differing sources needing to be considered (sputum and bronchus swab vs. bronchoalveolar lavage after LTx).

This work has several limitations. The sample size is small and reflects a single-center experience. All reported data have been evaluated retrospectively, and minimal data exists regarding nonvalve ELVR. Patient selection for LTx after ELVR creates inherent negative bias towards ELVR nonresponders within our cohort. Long-term data, in particular the development of chronic lung allograft dysfunction and the potentially negative impact of higher colonization rates, are lacking. The safety profile of all ELVR techniques and their impact on LTx outcome require further studies.

In conclusion, ELVR treatment in LTx candidates does not appear to influence long-term outcomes. ELVR appears potentially associated with increased rates of airway colonization and bronchiectasis before LTx, which may subsequently impact negatively on long-term survival after LTx. Further research into long-term outcomes is therefore required.

Acknowledgment

The authors thank Paul Borchert for his technical support.

Financial Disclosure and Conflicts of Interest

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References


Table 5 in the paper by Confalonieri M et al. entitled 'Opening of a respiratory intermediate care unit in a general hospital: impact on mortality and other outcomes' [Respiration 2015;90:235–242] should read:

Table 5. Management attitude and treatment timing in a sample of matched patients with ARF admitted in different hospital setting

<table>
<thead>
<tr>
<th></th>
<th>RICU</th>
<th>Emergency unit</th>
<th>Internal medicine wards</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Median time to second blood gas check, h</td>
<td>1.56 (0.4)</td>
<td>4.26 (3.4)</td>
<td>17.1 (10.9)</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Mean time to antibiotics initiation, h</td>
<td>0.84 (0.3)</td>
<td>1.63 (1.6)</td>
<td>2.2 (2.12)</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Median time to mechanical ventilation, days</td>
<td>0.3 (0.6)</td>
<td>0.7 (0.7)</td>
<td>4.8 (3.2)</td>
<td>0.0001</td>
</tr>
<tr>
<td>Use of NIV, n (%)</td>
<td>42 (70.0)</td>
<td>27 (46.5)</td>
<td>12 (19.3)</td>
<td>0.0001</td>
</tr>
<tr>
<td>Use of corticosteroids, n (%)</td>
<td>58 (96.6)</td>
<td>46 (79.3)</td>
<td>39 (62.9)</td>
<td>0.0012</td>
</tr>
<tr>
<td>Use of chest physiotherapy, n (%)</td>
<td>43 (71.6)</td>
<td>11 (18.9)</td>
<td>6 (9.6)</td>
<td>0.0001</td>
</tr>
</tbody>
</table>

Data in parentheses are interquartile ranges or standard deviations unless otherwise indicated.