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Lung transplantation after ex vivo lung perfusion in two Scandinavian centres

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Objectives: We reviewed our combined clinical outcome in patients who underwent lung transplantation after ex vivo lung perfusion (EVLP) and compared it to the contemporary control group.

Methods: At 2 Scandinavian centres, lungs from brain-dead donors, not accepted for donation but with potential for improvement, were subjected to EVLP (n = 61) and were transplanted if predefined criteria were met. Transplantation outcome was compared with that of the contemporary control group consisting of patients (n = 271) who were transplanted with conventional donor lungs.

Abstract

Is the outcome in patients who receive lung transplants after EVLP different from that in those receiving conventionally transplanted lungs?

Lung function at 1 year and medium-term survival did not differ between groups; however, short-term outcome did.

At our 2 centres, patients with lungs transplanted after EVLP did not have significantly different medium-term outcomes compared to those with conventionally transplanted lungs.

Key question

Key finding(s)

Take-home message

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RESULTS: Fifty-four recipients from the regular waiting list underwent transplantation with lungs subjected to EVLP (1 bilateral lobar, 7 single and 46 double). In the EVLP and control groups, arterial oxygen tension/inspired oxygen fraction ratio at arrival in the intensive care unit (ICU) was 30 ± 14 kPa compared to 36 ± 14 (P = 0.005); median time to extubation was 18 h (range 2–912) compared to 7 (range 0–2280) (P = 0.002); median ICU length of stay was 4 days (range 2–65) compared to 3 days (range 1–156) (P = 0.002); Percentage of expected forced expiratory volume at 1 s (FEV₁) at 1 year was 75 ± 29 compared to 81 ± 26 (P = 0.18); and the 1-year survival rate was 87% [confidence interval (CI) 82–92%] compared to 83% (CI 81–85%), respectively. Follow-up to a maximum of 5 years did not show any significant difference in survival between groups (log rank, P = 0.63).

CONCLUSIONS: Patients transplanted with lungs after EVLP showed outcomes comparable to patients who received conventional organs at medium-term follow-up. Although early outcome immediately after transplantation showed worse lung function in the EVLP group, no differences were observed at a later stage, and we consider EVLP to be a safe method for increasing the number of transplantable organs.

Keywords: Ex vivo lung perfusion • Lung transplantation • Lung reconditioning

INTRODUCTION

Lung transplantation (LTx) is an established treatment option for patients with end-stage pulmonary disease. The number of lung transplant procedures performed annually has consistently been increasing with encouraging and improving long-term results [1, 2].

The published rate of accepted organs from multiorgan donors varies from 15% to 50% [3], implying that in some regions or centres, up to 85% of available lungs are declined. Extended donor criteria and a more aggressive approach to optimize potential donor lung cations may increase organ availability in centres with a low acceptance rate.

Ex vivo lung perfusion (EVLP) has proven its potential by successfully differentiating between reversible and non-reversible lung pathology prior to transplantation [4–12]. Uncertainty over donor lung quality is often the reason for rejecting the organs. Some of these organs may not meet strict transplantation criteria at first evaluation, but in reality they may have the function that allows transplantation. In our setting, the evaluation of marginal organs ex vivo by the transplant team has increased the yield of organs available for transplantation. In the future, EVLP may also be a means of organ improvement by using different treatment strategies during EVLP.

The first successful LTx after EVLP was reported by Steen et al. [13] in 2001. EVLP has since gained increasing interest in transplant centres around the world. In 2008, Cypel et al. [14] in Toronto published an article on the extended EVLP assessment of lung function using a novel strategy, which was the starting point for the ‘Toronto protocol’. In 2011, the group published their first clinical study [15]. This was followed by short- to mid-term results in smaller series and case reports [4, 16–20], while publications on long-term results are still scarce [6, 21].

In our 2 Scandinavian centres, EVLP programmes were introduced in clinical practice in 2011 and 2012 [10, 22, 23]. Herein, we review the outcomes for patients who were transplanted with EVLP lungs and compare results to a control group of recipients who were transplanted with conventional lungs during the same period.

MATERIALS AND METHODS

Study design

The ethics committees of the University of Gothenburg and the University of Copenhagen approved this study. All patients were informed and consented to the possibility of receiving organs that had undergone EVLP when they were listed for transplantation. The organs were otherwise matched according to standard criteria.

At the outset, inclusion criteria for EVLP evaluation of rejected donor lungs were an arterial oxygen tension/inspired oxygen fraction (PaO₂/FiO₂) ratio <40 kPa and/or X-ray findings that were consistent with pulmonary oedema. Later, criteria were expanded to also include donor lungs for which it was not possible to properly evaluate in the donor (patient on veno-arterial extracorporeal membrane oxygenation (va-ECMO)), or ones with suspected lung injury (donors with pulmonary embolism or severe trauma as cause of death), or donor history, radiological or macroscopic findings suggesting severely impaired lung function that prevents the use of the organs. The decision to proceed to EVLP with lungs that were rejected for direct use was made after discussion between at least 2 transplant surgeons.

The procurement of the donor lungs was performed according to the local standard protocol. During transport, the organs were stored cold on ice. EVLP was performed at the recipient hospital.

Patients from our 2 centres were prospectively and consecutively included in this study between January 2011 and December 2015, and they were followed up until the end of December 2016. Lungs from brain-dead donors that were primarily rejected for transplantation were considered for EVLP. Lungs that achieved acceptable lung function during EVLP were transplanted. In both centres, acceptable lung function was defined as follows: (i) PaO₂/FiO₂ ratio >40 kPa; (ii) pulmonary vascular resistance (PVR) and pulmonary compliance deemed normal under EVLP conditions (350–650 dyn·s/cm² and >50 ml/cm H₂O, respectively) and not deteriorating during EVLP; and (iii) macroscopic appearance and manual inspection without major pathology. Outcomes in recipients who were transplanted with lungs after EVLP or without prior EVLP were compared.

Ex vivo lung perfusion

EVLP at our 2 centres was performed based on a modified version of the procedure described by Steen et al. [24]. Equivalent protocols were applied at both institutions using the Vivoline LS1 device, in which lungs were perfused with Steen solution mixed with red blood cells to a haematocrit of 10–15%, 10 000 U of heparin and 100 mg of meropenem. Evaluation was performed at full perfusate flow. Acceptance criteria differed between our 2 centres with regard to Pao₂/FiO₂ (see below).

Gothenburg. The EVLP procedure has been described in detail previously [25]. Perfusion of the lungs was restricted to 70 ml/min/kg donor weight. The pressure limit of the pulmonary artery...
space fraction (calculated as PaCO₂–EtCO₂/PaCO₂), static lung blood in the PA never exceeded 7 kPa. A physiological dead
simultaneous samples from the PA. The pO₂ of the deoxygenated
gas analysis were drawn from the left atrium and compared to
els under visual inspection was performed. Repeated samples for
manoeuvre with increasing positive end-expiratory pressure lev-
tail previously [23]. Lung protective ventilation was initiated at
Recipient characteristics are reported in Table 1. Surgery was per-
Lung transplantation and outcome analyses

Recipient variables | EVLP (n = 54) | Conventional (n = 271)
--- | --- | ---
Age (years), mean ± SD | 52 ± 12 | 51 ± 13
Diagnosis (%) | | |
IPF | 24 | 25
PAH | 2 | 6
COPD | 33 | 28
Alpha-1-antitrypsin deficiency | 6 | 13
CF | 20 | 12
Other | 15 | 16
Patients on preoperative mecha-
nical ventilation, n (%) | 5 (9.3) | 12 (4.4)
ECMO, n (%) | 1 (1.9) | 16 (5.9)

CF: cystic fibrosis; COPD: chronic obstructive pulmonary disease; ECMO: extracorporeal membrane oxygenation; EVLP: ex vivo lung perfusion; IPF: interstitial pulmonary fibrosis; PAH: pulmonary artery hypertension; SD: standard deviation.

(PA) was increased to 20 mmHg during the evaluation phase. PA
flow and thereby pressure was gradually increased. Mechanical
ventilation with a positive end-expiratory pressure of 5 cm H₂O
and a tidal volume of 6–8 ml/kg were initiated at 32°C. At this
stage, a bronchoscopy was performed to clear any secretions
and to inspect the bronchial tree. At 36°C, a lung recruitment
manoeuvre with increasing positive end-expiratory pressure lev-
levels under visual inspection was performed. Repeated samples for
gas analysis were drawn from the left atrium and compared to
Simultaneous samples from the PA. The pO₂ of the deoxygenated
blood in the PA never exceeded 7 kPa. A physiological dead
space fraction (calculated as PaCO₂–EtCO₂/PaCO₂), static lung
compliance and PVR were continuously monitored.

The acceptance of organs for transplantation was based on the
following criteria: (i) a PaO₂/FiO₂ ratio >40 kPa during the evalua-
tion phase; (ii) stable haemodynamic and respiratory variables
(PVR, peak airway pressures and lung compliance) during EVLP.
No absolute cut-off levels for these variables were used. A negative
trend with deterioration of physiological variables during EVLP was
considered a relative contraindication for transplantation; (iii) the
absence of macroscopic signs of pneumonic infiltrates or lung
infarctions; and (iv) a normal collapse test. Accepted lungs were
surface cooled in the EVLP system awaiting transplantation.

Copenhagen. The EVLP procedure has been described in de-
tail previously [23]. Lung protective ventilation was initiated at
32°C. At 36°C, blood gases were drawn to assess whether the evalua-
tion phase could be entered. Evaluation was performed at
36°C, following lung recruitment manoeuvres and bronchoscopy.
Lungs were approved if pCO₂ < 6 kPa and if pO₂ > 50 kPa at FiO₂ =
1.0 or pO₂ > 13 kPa at FiO₂ = 0.21. The collapse test was per-
formed to evaluate pulmonary oedema. Accepted lungs were
surface cooled in the EVLP system awaiting transplantation.

Lung transplantation and outcome analyses

Recipient characteristics are reported in Table 1. Surgery was per-
formed as to local preference and routine, either via bilateral se-
quential thoracotomy or sternotomy, either with or without
extracorporeal circulation. After the operation, all patients
received care and treatment according to standard protocols. The
ventilator time was defined as time to extubation in hours.
The time in intensive care unit (ICU) was defined as the number
of days from ICU arrival to general ward discharge. If the recipient
was re-intubated during the index procedure hospitalization, the
ventilator time was defined as the total ventilator treatment time.

Statistical analyses

Continuous data are presented as mean and standard deviation
or median and range. Categorical data are presented as fre-
quency and/or percentage. Differences between groups were
evaluated with the Mann–Whitney U-test or the Student's t-test.
A P-value of <0.05 was considered as statistically significant.
Kaplan–Meier curves were used for survival plots and the log-
rank test for comparison of proportional survival between the
groups. The Shapiro–Wilk test was used for testing normality.
Cox proportional hazard regression was used to examine the
relative risk of death between the studied groups. A test of the
proportional hazards, which was a required assumption of Cox
regression, was performed using a formal significance test based
on the unscaled and scaled Schoenfeld residuals.

RESULTS

During the 4-year study period, from January 2011 to December
2015, with Gothenburg initiating its clinical EVLP programme in
January 2011 and Copenhagen in May 2012, lungs from 1013
donors were offered to our 2 centres (Fig. 1). This number
includes all brain-dead donors (donation after circulatory death
(DCD) donation was not performed in our 2 countries during the
study period), irrespective of donor age. Patients who underwent
all other contemporary LTx (n = 271) procedures during the same
time interval, with organs accepted according to standard selec-
tion criteria not requiring EVLP, were included as the control
group. Retransplantations during the study period were excluded.

Ex vivo lung perfusion

In our combined cohort, 61 patients with donor lungs underwent
EVLP. The mean donor PaO2/FiO2 ratio was 30.6±12 kPa (exclud-
ing 2 donors on VA-ECMO). In cases where donor PaO2/FiO2
met standard acceptance criteria, reasons for EVLP were as fol-
lows: atelectasis unresponsive to ventilator lung recruitment
manoeuvres (n = 4), infiltrates on pulmonary X-ray (n = 4), and
donor on VA-ECMO (n = 2). No EVLP was performed for logistical
reasons.

After EVLP, 47 pairs of lungs were deemed transplantable. In
one of these cases, the lung pair was split and transplanted in 2
different recipients. In 1 case, bilateral bilobar transplantation
was performed. In 5 cases, one of the lungs was used for single
LTx, and the other discarded after EVLP. The conversion rate
expressed as the total number of lungs transplanted to the total
number of lungs placed on EVLP (99/122) was 81%. Another way
of expressing conversion rate would be that in 85% of EVLP runs
either one or both lungs were transplanted. EVLP data during
evaluation are presented in Table 2.

Transplantation

Forty-six bilateral, 1 bilobar and 7 single LTxs were performed
after EVLP, compared to 246 bilateral and 37 single LTxs in the
control group (Table 3). The use of intraoperative extracorporeal
circulation or extracorporeal membrane oxygenation was similar
in the 2 groups.

Early postoperative results

One patient in the EVLP group and 4 in the conventional group
died within the first 48 h after transplantation (Table 3). Death in
none of these cases was attributable to insufficient lung function.
PaO2/FiO2 at arrival in ICU was 30±14 kPa in the EVLP group
compared to 36±14 in the control group (P = 0.005). When com-
paring the EVLP group to the control group, median time to
extubation was 18 h (range 2–912) vs 7 h (range 0–2280 h)
(P = 0.002), median ICU stay was 4 days (range 2–65) vs 3 days
(range 1–156) (P = 0.002) and time to discharge to home or a re-
habilitation facility was 30 days (range 17–112) vs 28 days (range
12–268), respectively (P = 0.35).

Follow-up

One-year retransplantation-free survival was 87% [confidence
interval (CI) 82%–92%] in the EVLP group and 83% (CI 81%–85%)
in the conventional group. Cumulative retransplantation-free sur-
vival did not differ significantly between the groups (log rank,
P = 0.63) (Fig. 2) during the entire period in our combined
cohorts. Causes of death in the EVLP group are reported in
Table 4. Cox proportional hazards regression was performed. The haz-
ard ratio for EVLP was 1.14 (95% CI 0.67–1.93), P = 0.62. The as-
sumption of proportional hazards was tested as described in the
statistics section and fulfilled. Cox regression did not show a sig-
ificant difference in survival between groups. Data indicated a
14% higher relative risk for death or retransplantation in the
EVLP group with a wide CI that did not reach significance.

Pulmonary function test

FEV1.0% was 75±29% and 81±26% at 1 year in the EVLP and
control groups, respectively (P = 0.18).

DISCUSSION

The main finding of the present study in 2 Scandinavian centres
was that the cumulative retransplantation-free survival for up to
5 years in patients transplanted with EVLP-evaluated lungs was comparable to a control group of all contemporary patients transplanted with non-EVLP organs. \(P_{O2}/FiO2\) at arrival in ICU was lower, and time to extubation and time in ICU were significantly longer in the EVLP group. There was no difference in time to discharge from hospital, lung function at 1 year, or mortality or need for retransplantation. This indicates that the selection of donor organs for EVLP, as well as the selection of lungs for transplantation after EVLP, was adequate. This study is based exclusively on organs transplanted from brain-dead donors.

During the study period, the acceptance rate at our 2 institutions for conventional lungs without prior EVLP was 27%, a number higher than what many international centres report [1, 2]. Of the organs not fulfilling standard criteria, 61 were selected for EVLP evaluation. In contrast to others, while applying wider indications for EVLP [6] we have only accepted lungs with poor gas exchange or other clear contraindications for transplantation for EVLP. Fifty-four patients were transplanted with EVLP lungs, in which the conversion rate for EVLP-evaluated lungs was 81–85%. The selection for EVLP is still mainly based on clinical judgement, but the criteria for the selection of donor lungs suitable for EVLP will need to be explored further in future studies.

Implementing an EVLP programme can significantly increase the number of organs available for transplantation, possibly even utilizing up to as many as 50% of donor lungs. Centres with an already high acceptance rate can expect to have a significant addition of available organs by the introduction of EVLP.

EVLP strategies at our 2 institutions have been described previously [22, 23] and are based on the protocol initially developed by Steen et al. [13], using a cellular perfusate with banked blood added to a haematocrit of 10–15%, an open left atrium and evaluation of the lungs at relatively high pressure and flow. Since the introduction of EVLP, this protocol has been implemented at our institutions after minor modifications. No study has so far presented a validated algorithm for which lungs to accept for transplantation following EVLP. P/F-ratio and macroscopic appearance are still the main determinants of whether to proceed to transplantation, while other variables such as compliance, PVR dynamics and dead space fraction may provide supporting evidence.

The optimal yield of transplantable organs after EVLP depends on several factors, including selection criteria for considering EVLP, the experience of the transplant team and thresholds for proceeding to transplantation. At our two centres, the decision to use initially non-acceptable marginal donor lungs was initially made by junior retrieval surgeons in the donor hospital, followed by more experienced surgeons and anaesthesiologists after EVLP.

![Figure 2](https://academic.oup.com/ejcts/article-abstract/55/4/766/5146506)

**Table 4: Cause of death in EVLP recipients**

| Diagnosis       | Type of transplantation | Time after operation | Causes                                        |
|-----------------|-------------------------|----------------------|-----------------------------------------------|
| PAH             | Bilateral sequential    | 31 months            | Chronic graft failure                         |
| Fibrosis        | Bilateral sequential    | 24 months            | Squamous cell carcinoma, bronchial stenosis after radiotherapy |
| Fibrosis        | Bilateral sequential    | 18 months            | Graft failure secondary to pneumonia          |
| \(\alpha\)AT deficiency | Bilateral sequential | 25 months            | Graft failure secondary to recurrent pulmonary infections |
| COPD            | Bilateral sequential    | 18 months            | Graft failure secondary to pneumonia          |
| CF              | Bilateral sequential    | 5 months             | Graft failure secondary to pneumonia          |
| \(\alpha\)AT deficiency | Bilateral sequential | 13 months            | Chronic graft failure, bronchial stenosis    |
| COPD            | Bilateral sequential    | 36 months            | Respiratory failure secondary to recurrent infections and rejection |
| COPD            | Bilateral sequential    | 5 months             | Respiratory failure secondary to recurrent infections and rejection |
| Fibrosis        | Bilateral sequential    | 1 day                | Bleeding, graft failure, ECMO                 |
| Fibrosis        | Bilateral sequential    | 1 day                | Bleeding                                      |
| Fibrosis        | Bilateral sequential    | 61 months            | Unexpected death during sleep at home         |
| Fibrosis        | Bilateral lobar         | 14 months            | Pulmonary embolism and BOS                   |
| Fibrosis        | Single                  | 18 months            | BOS                                           |

\(\alpha\)AT: alpha 1-antitrypsin; BOS: bronchiolitis obliterans; CF: cystic fibrosis; CNI: calcineurin inhibitor; COPD: chronic obstructive pulmonary disease; ECMO: extracorporeal membrane oxygenation; PRES: posterior reversible encephalopathy syndrome; PAH: pulmonary artery hypertension.
It could be argued that sending more experienced senior surgeons for organ retrievals could further increase the acceptance rate, thereby avoiding the quite substantial costs associated with EVLP evaluations in selected cases.

During the introduction of the EVLP programme at our 2 centres, funding was provided by research grants. However, after being adopted as a clinical routine procedure, it is now financed via the tax-based, general public health care system, in line with other medical procedures.

In this study, the conversion rate was 81%, which is high, but in agreement with most previous studies reporting a conversion rate between 46% and 100% [4, 6, 22]. The DEVELOP-UK study stands out in this aspect, reporting a conversion rate of only 34% [26]. It was an ambitious, multicentre observational study that compared EVLP to conventional lungs, which was terminated early due to slow recruitment and concerns about high levels of the use of ECMO in the EVLP arm. One could speculate that one reason for the low conversion rate may be that several of the participating centres had little experience in EVLP and consequently adopted a more conservative approach in accepting organs.

It could be argued that donor lungs that other centres would have considered for conventional transplantation were exposed to unnecessary EVLP in our centre, i.e. selecting very good lungs for EVLP. However, as our acceptance rate is already comparatively high, in combination with the fact that lungs subjected to EVLP in this study were first declined by multiple centres for conventional transplantation, we do not believe that other centres would have used any of these lungs for conventional transplantation. The high conversion rate may indicate that even more lungs should undergo EVLP, hopefully resulting in more lungs for transplantation. Martens et al. [27] published a retrospective database analysis of unused lung donors, identifying a large potential for EVLP to further increase the donor pool in transplant centres, even when the majority were already extended criteria donor lungs.

Different strategies for EVLP have been suggested and reported [14]. This study indicates that good long-term results can be achieved with more than 1 EVLP protocol. In a recently published study, where 2 EVLP strategies were compared in a porcine experimental setup, we could not show any significant difference in lung performance [28].

Our results indicate a significantly longer time on ventilator and a longer length of stay in ICU in patients receiving EVLP-treated lungs. This is not a surprise, because functionally less optimal organs were selected for EVLP and later transplanted. The time on EVLP is relatively short with a median of 175 min (range 76–577 min). It has been suggested that longer EVLP time could reduce oedema. Our clinical data show that lungs might lose or gain weight, i.e. fluid, during EVLP, and EVLP does not necessarily decrease oedema content in a specific pair of lungs. This has further been evaluated by our group in a recent publication [29]. Hemofiltration during EVLP, to increase perfusate oncotic pressure and thereby optimize oedema reduction, may be a way forward [25].

Our data show a comparable outcome in patients who were transplanted with EVLP lungs and controls. One-year retransplantation-free survival was close to 90%. There is no statistically significant difference between groups in our study with regard to long-term survival or freedom from retransplantation; however, numbers at risk are very low beyond 3 years after transplantation.

In our experience, although short-term outcome was inferior in the group transplanted with lungs after EVLP compared to conventional lungs, the medium-term follow-up showed similar results between groups. EVLP seems to be a safe method of increasing the availability of transplantable organs.

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Conflict of interest: Göran Dellgren has research grants from Astellas A/S for a study in immunosuppression after lung transplantation (the ScanCLAD study) and from Abbott/St Jude regarding a destination therapy study on LVAD (the SweVAD study), none of which are relevant for the content of this study. Michael Perch has a research grant from Roche for a study in Pirfenidone for chronic rejection after lung transplantation (the EPOS study). All other authors declared no conflict of interest.

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