Timing it right: the challenge of recipient selection for lung transplantation

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Abstract: Selection criteria for the referral and potential listing of patients for lung transplantation (LTx) have changed considerably over the last three decades but one key maxim prevails, the ultimate focus is to increase longevity and quality of life by careful utilization of a rare and precious resource, the donor organs. In this article, we review how the changes have developed and the outcomes of those changes, highlighting the impact of the lung allocation score (LAS) system. Major diseases, including interstitial lung disease (ILD), chronic obstructive pulmonary disease and pulmonary hypertension are considered in detail as well as the concept of retransplantation where appropriate. Results from bridging to LTx using extracorporeal membrane oxygenation (ECMO) are discussed and other potential contraindications evaluated such as advanced age, frailty and resistant infections. Given the multiplicity of risk factors it is a credit to those working in the field that such excellent and improving results are obtained with an ongoing dedication to achieving best practice.

Keywords: Recipient selection; lung transplantation (LTx); lung allocation score (LAS); cystic fibrosis (CF); chronic lung allograft dysfunction (CLAD)

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Introduction

Approximately 4,500 lung transplants are performed worldwide each year, but the number of potential recipients on the waitlist far outstrips this (1). Effective management of this precious resource relies in part on optimal recipient selection to ensure the best possible outcomes. Historically, this was interpreted as transplanting only the “perfect” candidate. Over time, and with a growing body of research we have come to understand that this approach led to many good candidates being turned away. With increased understanding of what represents an acceptable risk-benefit ratio, the focus has shifted to ensuring the best possible outcome for the largest number of recipients. This changing paradigm is clearly reflected in the gradual evolution of the selection criteria from the International Society for Heart and Lung Transplantation (ISHLT) over the past 20 years (2-4).

The lung allocation score (LAS) represents a major attempt to take this further. Implemented initially within the USA, and subsequently in Europe, the LAS attempts to fairly allocate or “match” a given donor with a particular recipient. In order to accomplish this, the LAS mathematically calculates the overall likelihood of post-transplant survival against the risk of mortality without transplant, computing the number of life days gained. This requires accurate and up to date information, with regular recalculation of scores, to provide a functional system (5). Not surprisingly, LAS programs have generated
much interest and numerous publications regarding their
development, promulgation and utility (5-23).

Careful assessment and evaluation of the potential lung transplant recipient is central to improving survival and quality of life via transplantation, and must be combined with judicious therapeutic intervention to increase the chance of surviving to transplant and to minimize the impact of co-morbidities peri-operatively and beyond. A number of indications and contraindications to recipient referral and selection for lung transplantation (LTx) have been discussed extensively in the literature. We will focus in particular on those major areas where the evidence base has been developed.

Age

The current ISHLT recommendation is that there should be no absolute upper age limit to LTx, with the caveat that increasing age often comes with a range of co-morbidities. The reality may be that biological, rather than chronological “age” is the important factor. A retrospective review of the United Network for Organ Sharing (UNOS) Standard Transplant Analysis and Research database for 15,844 adult lung recipients from 2005–2015 examined the effect of the donor-recipient age relationship on outcomes. Recipient age had a significant impact on post-transplant survival by multivariate analysis. Compared with recipients aged 30–39, who had the best overall survival, those >70 year had a mortality odds ratio of 1.81 (95% CI, 1.46–2.24, P<0.0001). While the overall trend demonstrated decreased survival with age (Kaplan Meier log rank P<0.0001), it must be mentioned that those aged 18–29 had an odds ratio of 1.52 (95% CI, 1.27–1.83, P<0.0001) (24).

Series that do not demonstrate an age-related survival advantage perhaps reflect a highly selected and well managed group of older patients. Complications following transplantation vary with age, with higher rates of drug toxicity and malignancy in older patients (25). Coronary vascular disease, cerebrovascular disease and frailty, which have an age-related association, are also independent risk factors for survival following transplant (26-30).

Weight

Body mass index (BMI) has been the primary tool utilized to assess an independent predictor of LTx outcome, however body composition is also important. Respiratory cachexia and emphysema, in particular, have been known to be associated with sarcopenia for over 20 years (31). Underweight is complex and the impact on recipient selection varies. A study of the ISHLT Registry on pediatric LTx for primary pediatric recipients (aged <18 years) between 1990–2008 found that underweight (BMI <18.5 kg/m²) patients did not have significantly worse outcomes following transplant (n=897) (32).

The most comprehensive assessment of the role of weight as a risk factor for LTx survival, was of 5,978 adults with cystic fibrosis (CF), chronic obstructive pulmonary disease (COPD), and diffuse parenchymal lung disease who underwent LTx in the United States between 1995–2003. Median follow-up time was 4.2 years. Multivariable-adjusted rates of death were 15% higher for underweight recipients (95% CI, 3–28%), 15% higher for overweight recipients (95% CI, 6–26%), and 22% higher for obese recipients (95% CI, 8–39%), irrespective of diagnosis. Both obesity and underweight were independent risk factors for death after LTx, contributing to up to 12% of deaths in the first year (33).

A single center study of 810 patients, of whom 403 (50%) were overweight and 109 (13%) obese by BMI criteria, demonstrated greater pre-transplant weight loss was associated with dose-response improvements in survival [hazard ratio (HR) 0.83, 95% CI, 0.72 to 0.97, P=0.018]. Modest (0% to 3%, HR 0.91), moderate (7% to 10%, HR 0.83), and high (>15%, HR 0.71) levels of weight loss conferring longer survival, independent of initial weight (P=0.533 for interaction). Weight loss was also associated with improved chronic lung allograft dysfunction (CLAD)-free survival [HR 0.84 (0.71 to 0.99), P=0.034] and shorter LOS (b=0.17, P<0.001). They concluded weight loss before transplantation was associated with improved short- and long-term clinical outcomes, independent of initial weight but acknowledged that the mechanisms by which weight loss improve clinical outcomes were unclear (34).

Infections

In the past, the inability to treat certain multi-resistant or tissue invasive organisms led to patients colonized or infected pre-transplant being excluded from candidacy. Much of the data comes from the CF community, who are the primary group involved. This is due in part to the selective pressure of long-term antibiotics as well as nosocomial transmission. As the body of research has expanded, there is increased recognition that many of these organisms should no longer be considered absolute
Carriage of pan-resistant bacteria poses a particular problem, both perioperatively as well as in long-term management. A cohort study of CF patients compared outcomes post-transplant out to 6 years of patients colonized pre-transplant with pan-resistant organisms (n=21) to those colonized with sensitive organisms (n=39). The pan-resistant group comprised of 21 patients with *Pseudomonas aeruginosa* and 6 with *Burkholderia cepacia*. The incidence of bacterial bronchitis (28% and 33%, respectively) and pneumonia (28% and 38%, respectively) did not differ between groups (P>0.2) at 6 months. 1-year survival was similar (81% vs. 83%) for both groups (P=0.2), though pan-resistant *B. cepacia* patients had a lower 1-year survival (50% versus 90%, P<0.05) compared with pan-resistant *P. aeruginosa* patients. The authors concluded that CF patients infected with pan-resistant *P. aeruginosa* have similar transplant outcomes as patients with sensitive bacteria and therefore should not be excluded from LTx (35). These results are supported by a study of 54 LTx patients transplanted with pan-resistant bacteria which found a 1-year survival of 92%, where 11/18 post-transplant deaths were in part related to infection (36).

Though active infection with *Mycobacterium tuberculosis* represents an absolute contraindication, there is growing evidence that not all non-tuberculous mycobacteria (NTM) need be considered in the same light. Effective treatment and even spontaneous clearance has been demonstrated post-transplant (37). In general, NTM colonization is not considered an absolute contraindication to LTx (38-40). *M abscessus* is the exception due to intrinsic resistance to antimicrobials and tendency to relapse even after prolonged therapy (40,41). Even there, the small number of cases reported makes it difficult to provide a confident statement regarding a definitive impact on outcome.

There remain some organisms that all units approach with extreme trepidation due to their perceived risk. The filamentous fungi *Scedosporium*, predominantly *S. apiosperum*, and *Lomentospora prolificans* may lead to severe disseminated infections after LTx (42,43). These are inherently resistant to all current antifungal drugs, though voriconazole and terbinafine have demonstrated successful suppression (44,45). Mortality rates from disseminated disease, including anastomotic dehiscence, empyema and intracerebral disease are unfortunately high (46,47). All attempts at reducing the fungal burden must be made before consideration of listing, and even in these instances, some units consider the risk too great.

*Burkholderia cepacia complex* (BCC) presents similar challenges. A retrospective cohort study of 22/247 patients with CF infected with BCC found that early mortality rates were higher in the BCC group (3 month survival: 85 vs. 95%, P=0.05) and that the subset of 8/22 patients infected with *B. cenocepacia* (genomovar III) faced significantly higher risks of death than the 14 patients infected with other BCC (HR 3.2, 94% CI, 1.1–5.9, P=0.04). Long-term outcomes of the non-genomovar III organisms were similar. These findings were reflected in a large UK cohort, and *B. cenocepacia* is generally considered a contraindication to transplant (48,49). As with many of these organisms, the data only carries us so far. Some units will consider non-genomovar III BCC for transplant while others quote a history of poor outcomes. Graft infection and loss of function as well as disseminated infection are not uncommon, and it remains the responsibility of the individual units to counsel individual patients regarding the likely risk.

**Extracorporeal membrane oxygenation (ECMO)**

While we strive to perfect the time of listing for transplant, some patients suffer a catastrophic event and deteriorate quickly to the point they require ECMO. Some of these were already on the waitlist who have deteriorated faster than expected, others were previously healthy. Prolonged periods of ECMO are associated with worse outcomes and high mortality (50), so the decision to transplant off ECMO is a critical one and must be made early (51).

Among those with previously healthy lungs, the decision must balance the long-term risks of a transplant against the possibility of recovery. There are case reports of recovery following long periods of ECMO up to 56 days (52). Against this, there are patients who develop fibrotic end-stage lung with adult respiratory distress syndrome from influenza and secondary infection (53,54). Experienced units have employed ECMO bridging for almost 2 decades and the results appear acceptable in selected cases (53). A study of 38 patients (median age 30.1 years, range, 13–66 years) who underwent ECMO support with intention to bridge to primary LTx 1998–2011 demonstrates this point well. Median bridging time was 5.5 days (range, 1–63 days) days and four died prior to transplant. Of the 34 who received an LTx, 26 survived to discharge and returned to “normal”
life. The 1-, 3- and 5-year survival for all transplanted patients was 60%, 60%, and 48%, respectively. Long-term survival outcomes for those who survived beyond 3 months were similar (55). Subsequent work has demonstrated that outcomes for transplant off ECMO tend to be better in higher volume centers (56).

**Specific diseases**

**CF (19,57-62)**

Patients with CF have some of the best survival outcomes following LTx, with a median survival of 9 years for those alive 1 year following transplantation (63). Despite this, CF continues to be one of the more challenging diseases to determine the best time for referral and listing for LTx. In part, this is because the outcomes for CF patients continue to improve, such that the median predicted survival for a child born in 2019 with CF is 43.6 years. Conversely, that single statistic means half of all CF patients will die before age 43.6 years and this is a large potential target group. In historical studies, the decision to proceed to LTx may have been made to early, such that two studies found that in certain subgroups of CF, the non-transplanted comparators had longer survival (64,65).

Central to the challenge of timing is the issue that no single factor has been demonstrated to be overwhelmingly predictive of early mortality in CF. This may be confounded by the recognition that CF survival data include patients who would not otherwise be suitable for LTx (3). One major marker of disease severity frequently utilized as an indication for referral, is an FEV1 that has fallen to less than 30% predicted, however on multiple occasions this has been found not to correlate with a median survival less than 50% at 2 years (64,66-68). One study which constructed ROC curves utilized data from 14,572 patients in the Cystic Fibrosis Foundation National Patient Registry and found that an FEV1 <30% predicted had a sensitivity of 42% and a specificity of 95% for 2-year mortality. While the low FEV1 is a marker of disease severity, many patients remain stable at this point for many years. This “stability” may work in the patient’s favor as it may allow time for the CF patient to be fully appraised of the information about LTx and thereby to make a truly informed decision. Unfortunately a multiple logistic regression model that also incorporated, among other things, the number of exacerbations, infection with *Pseudomonas* or *Burkholderia cepacia*, sex and age proved no more accurate (67). From a more practical perspective, the use of FEV1 <30% as a benchmark for referral helps ensure that there is sufficient time to complete the transplant workup and allow sufficient time for a donor organ to become available for patients who are listed.

While technically more challenging to model, the greater risk perhaps lies with those in whom the obstruction is not just severe, but rapidly progressive. A study of 635 patients with CF, found that while the median survival of the 61 patients with an FEV1 <30% was 3.9 years (95% CI, 2.6-4.1years), those who died before the median survival had a significantly more rapid decline in lung function, with an average rate of change of -1.8% predicted/year against 0.73% predicted/year (Cox proportional hazards model P=0.0001). Further, the rate of change had been elevated in the 5 years prior to death (68). Again, there is conflict within the evidence. A larger modeling study of 5-year survival found that rate of decline was not a significant predictor (66). Rate of change has been excluded from some other studies due to the complexity of accurate modeling (67).

Several factors have been associated with increased rates of lung function decline and survival. A prospective cohort study of 446 adult CF patients found that those experiencing more than 2 exacerbations per year had an increased risk of experiencing a sustained 5% decline in FEV1 (HR 1.55, 95% CI, 1.10–2.18, P=0.01). They were also at increased risk of transplantation or death (60). Similarly, colonizing organisms, and in particular *Burkholderia cepacia* have been associated with increased rate of decline in lung function and also survival (62). These results have proven less statistically robust, often losing significance in multivariate analysis, however they contribute to the assessment of the patient as a whole, and should still be taken into consideration (67,69).

While valid concerns remain that some CF patients might be listed too early, the implementation of the LAS has brought new insights. A cohort study of 1,437 CF patients found that those with LAS <50 had a 1-year mortality of 12%, whereas those whose LAS was 50 or greater had a 1-year mortality of 16.1% (58). By multivariate analysis there was an increased risk out to 2 years (HR 1.38, 95% CI, 1.04–1.83, P=0.03). There is little argument that patients with chronic respiratory failure or pulmonary hypertension secondary to CF should be listed for transplantation, given the risks of deterioration and failure to survive to achieve LTx.

**Interstitial lung disease (ILD)**

ILD, and more specifically idiopathic pulmonary fibrosis (IPF) are now the most common indication for LTx (63).
The incidence increases with age, and peaks among those aged greater than 75 years (70,71). The associated rise in comorbidities, including weight and frailty, works against successful LTx in many of these patients (25,72). Despite this, the proportion of LTx for IPF increased from 16% to 33% of all LTx from 1990 to 2014 (63). The median survival for IPF is variously quoted as 2–5 years and due to the potential for an unpredictable and rapid progression, it is recommended all patients should be referred to a transplant center at the time of diagnosis unless there is a definite contraindication to LTx. Waiting list mortality rates have historically been in the order of 28–47% (3,73,74). The implementation of the LAS has been associated with a reduction in waiting list mortality rates to around 11%. The corollary to this is that a higher LAS appears to be associated with a higher mortality at 1-year post LTx, specifically 2% for every additional 1 point (75). While this has raised concern in certain quarters, it represents the reality that transplanting more unwell patients will lead to a higher risk.

Declining lung function has been determined to be a strong predictor of mortality, and therefore a trigger for listing. A study of 1,099 patients with IPF found on multivariate analysis that declines in % predicted FVC of >10% and % predicted DLCO of >15% over 24 weeks were associated with increased mortality [HR 3.65 (95% CI, 2.03–6.57, P<0.001) and HR 2.41 (95% CI, 1.19–4.87, P=0.015)] respectively (73). Similarly, exercise-induced hypoxemia has been demonstrated to be associated with mortality amongst those with both biopsy proven usual interstitial pneumonia (UIP) and biopsy proven nonspecific interstitial pneumonia (NSIP) (76). A cohort study of 105 patients found that desaturation below 88% during a 6-minute walk test (6MWT) was associated with an increased hazard of death (HR 4.2; 95% CI, 1.4–12.445; P=0.01) amongst those with UIP. Other predictors of mortality include hospitalization for exacerbations, and the development of pulmonary hypertension hence these have been included in the criteria for listing (3). These criteria should not be taken in isolation of the patient as a whole. There remains a significant mortality in patients with preserved lung function. Some novel markers exist. Telomere shortening shows promise in predicting mortality and early requirement for transplantation (77).

LTx for connective tissue disease (CTD)-ILD has long been a topic of debate with strong opinions for and against. Some centers do not offer LTx for this indication and those that do usually exclude patients with ongoing evidence of disease activity despite appropriate immunosuppressive medication, due to the fear that disease flares will contribute to mortality. There is a paucity of data on which to base a robust opinion, however the current consensus opinion of the ISHLT has been to support LTx in those with well controlled CTD-ILD using the same criteria as IPF. A recent study found that cumulative 1-year survival was 80% post-transplant for selected patients with CTD-ILD compared with 60% for IPF (n=62). Even after age matching, outcomes remained comparable (78). A larger retrospective cohort study that excluded systemic sclerosis (SSc) found no significant differences between 275 CTD-ILD patients and 6,346 IPF patients out to 10 years in terms of survival, rates of acute cellular rejection or CLAD, predominantly bronchiolitis obliterans syndrome (BOS) (79). SSc is considered a particularly difficult indication for LTx. Concerns remain that aspiration, secondary to esophageal dysmotility and gastroparesis may damage the transplanted lung. For this reason, some centers still consider SSc an absolute contraindication to LTx (3). This risk may be overstated. A retrospective case-control study of 69 patients at a center that did not exclude patients with severe gastro-intestinal involvement found comparable outcomes out to 5 years between SSc and non CTD-ILD patients (80). These findings were echoed in a recent international multicenter cohort study of 90 patients (81).

**Emphysema**

COPD, principally emphysema, was formerly the most common indication for LTx. There are numerous reasons for this. Patients are usually slow to deteriorate on the waitlist, facilitating adequate opportunity for transplant. Further, there is a perception that the surgical transplant procedure is less complex, thereby carrying a lower morbidity and mortality. The introduction of the LAS has changed this somewhat. A multi-center retrospective cohort study of 341 patients found that recipient diagnoses changed following implementation of the LAS, with an increase in IPF and a decrease in COPD and CF (P<0.002). There was a decrease in waiting time from 680.9±528.3 days to 445.6±516.9 days following implementation of the LAS (P<0.001). Hospital mortality and 1-year survival were the same between groups (5.3% vs. 5.3% and 90% vs. 89%) (7).

Determining the optimal timing of transplantation is challenging as survival in COPD is not easy to predict. The development of the body-mass index, airflow obstruction, dyspnea, and exercise capacity (BODE) index, a simple...
multidimensional grading system, which was superior to the FEV1 at predicting the risk of death from any cause and from respiratory causes among patients with COPD has been utilized as a useful tool by which to stratify patients being assessed for LTx (82-83). Longitudinal changes in a modified BODE index of more than 1 point from baseline to 6, 12, and 24 months of follow-up was predictive of subsequent mortality in National Emphysema Treatment Trial (NETT) patients (84). Comparing the BODE index to the Global Obstructive Lung Disease (GOLD) 2011 revision ABCD categories the BODE index was found to be superior in predicting survival (85). Further analysis has provided insights into how best to use these tools, identifying the importance of age and physiological parameters, specifically DLCO, as independent predictors of survival (86).

While it may be ideal to defer transplant until after lung volume reduction (LVRS), either bronchoscopic or surgical, has been considered, many potential candidates do not meet criteria for these (87). This is an evolving landscape and the advent of new, minimally invasive techniques may allow more patients to defer transplant (88).

**Pulmonary Hypertension**

Compared with the prevailing options when LTx was first developed there are many new therapeutic options for pulmonary arterial hypertension (PAH) can stabilize many patients so they do not need referral for transplant. When interpreting the data regarding the optimal timing for referral and listing for transplant in PAH it is important to factor in the historical context in which they were made. Despite these developments, LTx remains an important salvage option (89-92). Introduction of the LAS in particular has led to an improved likelihood of LTx for listed patients with PAH (93). Unfortunately, wait list mortality remains comparatively high based on a study of 7,952 adults listed for LTx 2002–2008 (8). Under the LAS system, patients with PAH were less likely to be transplanted than patients with IPF (HR, 0.53; P<0.001) or CF (HR, 0.49; P<0.001) and at greater risk of death on the waiting list than patients with COPD (HR, 3.09; P<0.001) or CF (HR, 1.83; P=0.025) after adjustment for demographics and transplant type.

One large single center analysed outcomes of 316 patients with PAH referred for consideration of LTx including idiopathic PAH (n=123), associated with congenital heart disease (n=77), CTD (n=102), or chronic thromboembolic disease (n=14) (94). Of the 100 patients listed for LTx, 57 underwent bilateral LTx, 22 underwent heart-LTx, 18 died while waiting, and 3 were still waiting. The waiting list mortality was the greatest for patients with CTD-PAH (34% vs. 11% in the remaining patients, P=0.005). After LTx, the 30-day mortality decreased from 24% in the 1997–2004 group to 6% in the 2005–2010 group (P=0.007). The 10-year survival was worse for those with idiopathic PAH (42% vs. 70% for the remaining patients, P=0.01). The long-term survival reached 69% at 10 years in the patients with CTD PAH who survived to transplant.

Early reports of LTx for idiopathic PAH may have comprised a number of subtypes. Recent diagnostic tools and an increased interest in the sub-specialty of PAH has led to an understanding of the importance of discriminating idiopathic pulmonary venous hypertension and especially pulmonary capillary haemangiomatosis (PCH) as subtypes of pulmonary venous occlusive disease (PVOD) due to the rate of progression and failure to respond to standard PAH therapies (95,96). Some patients with PVOD may be bridged to LTx with intravenous epoprostenol, however pulmonary edema may develop (97). Patients with PCH should be referred immediately to a LTx unit with expertise in PAH so that a rapid work-up can be undertaken to facilitate urgent listing.

**CLAD**

CLAD, specifically the two main phenotypes, BOS and the restrictive allograft syndrome (RAS) have been the subject of recent consensus documents of the ISHLT and are now well defined, supplanting previous discussion papers (98-102). Retransplantation of the lung has been variably seen as an acute solution to primary graft dysfunction or as salvage for severe CLAD with respiratory failure. Building experience has documented the futility of the former approach and the qualified benefits which might accrue from the latter when applied appropriately. Hall et al. reviewed the UNOS database to identify 542 patients undergoing LTx at their institution 1995–2014 of whom 87 were retransplants (103). Predictors of worse survival included recipient age 50–60 years (relative risk, 4.3; P=0.02) or older than 60 years (relative risk, 10.2; P<0.001), and time to retransplant of less than 2 years (relative risk, 3.8; P=0.01). Retransplant for BOS had longer median survival than for RAS (2.7 vs. 0.9 years; P=0.055). They opined that lung retransplantation was associated with significantly worse survival than primary LTx but may be appropriate in well-selected patients with certain diagnoses.
However Scully et al. reported that in well selected paediatric cases, graft survival in patients who underwent re-LTx greater than 1 year after primary transplant was not statistically different than for primary LTx patients (P=0.21; graft half-life 2.8 vs. 4.0 years), and if re-LTx greater than 1 year posttransplant occurred in patients who were not ventilator dependent, survival was further improved (P=0.68; graft half-life 4.7 vs. 4.0 years) (104). To complement these single center studies, Osho et al. evaluated 9,270 primary LTx and 456 re-LTx recipients since LAS implementation, based on UNOS data (105). They concluded late lung retransplantation appears to be as beneficial as primary transplantation in propensity-matched patients. However, survival was severely reduced in those retransplanted less than 90 days after primary transplantation.

Modifications in surgical technique including retransplantation of the lung via sternum-sparing anterolateral thoracotomies off-pump has been reported to be associated with improved survival outcomes in the era starting in April 2010 at 30 days (98% vs. 76.3%, P=0.002) as well as at 1 year (80.6% vs. 63.2%; P=0.01) (106).

Notwithstanding these improvements, practical and ethical considerations remain regarding the practice of retransplantation that need to be addressed by each unit according to local policies, organ availability and utility (107).

Recent experience demonstrates the importance of CLAD phenotype in assessing the risk of retransplantation (108). A retrospective analysis of 143 patients who underwent re-LTx for CLAD [94 BOS (66%), 49 RAS (34%)] in four LTx centers 2003–2013 demonstrated unadjusted and adjusted survival after re-LTx for RAS was worse compared to BOS (HR 2.60, 1.59–4.24; P<0.0001 and HR 2.61, 1.51–4.51; P=0.0006, respectively). Patients waiting at home prior to re-LTx experienced better survival compared to hospitalized patients (HR 0.40; 0.23–0.72; P=0.0022). Patients with RAS redeveloped CLAD earlier and were more likely to redevelop RAS. The authors advised re-LTx for RAS should be critically discussed, particularly when additional peri-operative risk factors were present.

**Summary and conclusions**

Selection criteria for LTx have changed significantly over the last three decades with the effect of permitting access to groups of patients who may not have been considered in the past. The increasing development of medical technologies and wide experience from high volume centers in particular has allowed successful therapy of many of the co-morbidities previously considered as contraindications to transplant. LAS data demonstrate just how much the focus has shifted towards the sickest patients particularly within the USA and Germany. Short term survival seems comparable to previous cohorts even if resource utilization may be greater. The balance ever was risk versus benefit for an individual patient with an overriding consideration of the implications of best use of a precious resource being mindful of the debt that all who work in the field of transplantation hold to the generosity of the donor community.

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**Footnote**

Conflicts of Interest: The authors have no conflicts of interest to declare.

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