Living-donor lung transplantation for end-stage lung disease

Interventional procedures guidance
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www.nice.org.uk/guidance/ipg170

Your responsibility

This guidance represents the view of NICE, arrived at after careful consideration of the evidence available. When exercising their judgement, healthcare professionals are expected to take this guidance fully into account, and specifically any special arrangements relating to the introduction of new interventional procedures. The guidance does not override the individual responsibility of healthcare professionals to make decisions appropriate to the circumstances of the individual patient, in consultation with the patient and/or guardian or carer.

All problems (adverse events) related to a medicine or medical device used for treatment or in a procedure should be reported to the Medicines and Healthcare products Regulatory Agency using the Yellow Card Scheme.

Commissioners and/or providers have a responsibility to implement the guidance, in their local context, in light of their duties to have due regard to the need to eliminate unlawful
discrimination, advance equality of opportunity, and foster good relations. Nothing in this
guidance should be interpreted in a way that would be inconsistent with compliance with
those duties. Providers should ensure that governance structures are in place to review,
authorise and monitor the introduction of new devices and procedures.

Commissioners and providers have a responsibility to promote an environmentally
sustainable health and care system and should assess and reduce the environmental
impact of implementing NICE recommendations wherever possible.

1 Guidance

1.1 Current evidence on the efficacy of living-donor lung transplantation for
end-stage lung disease and its safety profile for suitable recipients
appears adequate to support the use of this procedure.

1.2 The procedure should only be used in selected patients who would
otherwise die.

1.3 However, limited evidence suggests that living-donor lung
transplantation for end-stage lung disease carries a significant risk of
morbidity for donors. Therefore clinicians wishing to undertake this
procedure should take the following actions.

- Inform the clinical governance leads in their Trusts.

- Ensure that donors receive thorough physical and psychological screening, and
counselling about the morbidity associated with this procedure. They should
also be provided with clear written information. In addition, use of NICE's
information for the public is recommended.

- Audit and review clinical outcomes of all people donating lungs for
transplantation.

1.4 Living-donor lung transplantation for end-stage lung disease should only
be performed in specialist centres in the context of a multidisciplinary
team. Donor lungs should be harvested by specialist thoracic surgeons.

1.5 Clinicians should enter all donors and recipients into the UK National
Audit of Intrathoracic Transplantation.
2 The procedure

2.1 Indications

2.1.1 Lung transplants are performed in patients with non-malignant pulmonary disease that is unresponsive or minimally responsive to treatment and who have a life expectancy of less than a year. The underlying causes include cystic fibrosis, severe pulmonary fibrosis, pulmonary hypertension and obliterative bronchiolitis.

2.1.2 The majority of live-donor lung recipients are patients with cystic fibrosis. The majority of lung donors are first-degree relatives who are compatible in terms of size and ABO blood group.

2.1.3 Living donation is an alternative to cadaveric organ donation. Living donation is an option for patients for whom cadaveric transplantation is unavailable, or who are deteriorating clinically to the point of transplant ineligibility while waiting for a cadaveric donor. Living donation may also be an option for critically ill children, as there is a particular shortage of suitable cadaveric donors for this age group.

2.2 Outline of the procedure

2.2.1 Living-donor lung transplantation requires three operations: two donor lobectomies (one donor providing the right lower lobe and another, the left), and the recipient bilateral pneumonectomy and lung implant.

2.2.2 The recipient operation is performed through a chest incision. Procedures are performed on cardiopulmonary bypass. Once the pneumonectomies have been completed the harvested lobes are implanted sequentially.

2.3 Efficacy

2.3.1 In a study of 123 adult and paediatric patients who had undergone living lung transplantation, 1-, 3- and 5-year survival was 70%, 54% and 45%,
respectively. Infection was the main cause of death (33/63; 52%), followed by obliterative bronchiolitis (8/63; 13%).

2.3.2 In a non-randomised study from the same centre outcomes were compared between living (n = 59) and cadaveric (n = 43) lung recipients who had survived more than 3 months after transplant. The study found no significant differences between the groups with respect to survival; 1-, 3- and 5-year survival was 83%, 64% and 62%, respectively, in the living lung group compared with 83%, 81% and 75% in the cadaveric lung group. A true comparative analysis is difficult, however, because those receiving living lung transplants often have poorer outcomes by nature of eligibility criteria (for example, underlying lung disease and preoperative severity of illness).

2.3.3 Where pulmonary function was measured in the studies it was reported that patients who had undergone living lung transplantation had improved function compared with preoperative values. For more details, refer to the 'Sources of evidence' section.

2.3.4 Some Specialist Advisors expressed uncertainties about the long-term outcomes of recipients following living lung transplantation and the incidence of obliterative bronchiolitis compared with those undergoing cadaveric lung transplantation.

2.4 Safety

2.4.1 There was limited information reported on the complications in recipients following living lung transplantation. In the studies that included both adult and paediatric patients, the incidence of acute rejection ranged from 0.8 to 1.5 episodes per patient. In a small study of 30 patients, complications following living lung transplantation included pulmonary oedema in 20% (6/30), haemorrhage necessitating rethoracotomy in 7% (2/30) and cardiac tamponade in 7% (2/30). Tracheostomy was required in 15 patients (50%), and reintubation in seven patients (23%).

2.4.2 There were no reports of donor mortality following lobectomy. In one study it was reported that 20% (50/253) of donors had one or more perioperative complications following lobectomy. The most common
complication was the need for a thoracostomy tube in 30% (15/50), either for persistent drainage or for air leaks. The most significant complication was pulmonary artery thrombosis, which occurred in two patients (1%). Eight patients (3%) also required reoperation because of bleeding (1.2%), bronchopulmonary fistula (0.4%), unresponsive pericarditis (0.4%), loculated pleural effusion (0.4%), a sterile empyema (0.4%) and a retained sponge (0.4%). In a study following 253 donor lobectomies it was reported that donors who could be contacted at 1 and 2 years had reduced pulmonary function compared with preoperative values. For more details, refer to the ‘Sources of evidence’ section.

2.4.3 The Specialist Advisors considered the main complications in recipients to be rejection and hyperexpansion of the transplants leading to significant lung injury and subsequent failure. With respect to donors, the Specialist Advisors listed potential complications following donor lobectomy as prolonged air leak, bleeding, pleural sepsis and pulmonary embolism. The Specialist Advisors also stated that donors were likely to experience loss of lung function following lobectomy.

3 Further information

Sources of evidence

The evidence considered by the Interventional Procedures Advisory Committee is described in the interventional procedure overview of living donor lung transplant for end-stage lung disease.

Information for patients

NICE has produced information on this procedure for the public. It explains the nature of the procedure and the guidance issued by NICE, and has been written with patient consent in mind.

Update information

Minor changes since publication
January 2012: minor maintenance.


Endorsing organisation

This guidance has been endorsed by Healthcare Improvement Scotland.