Renal Transplantation

GUIDELINES ON RENAL TRANSPLANTATION


Introduction

The number of patients registered as starting end-stage renal disease (ESRD) therapy per year has increased in all countries. The most common cause of ESRD is diabetes (affecting at least 16% of patients with diabetes) closely followed by high blood pressure, Glomerulonephritis, Polycystic Kidney Disease and Interstitial Nephritis.

Renal transplantation is now widely considered the treatment of choice for patients with ESRD due to improved short- and long-term survival benefits over dialysis treatment (aided by effective immunosuppressants such as cyclosporine A, mycophenolate mofetil (MMT), tacrolimus (TAC) and Sirolimus.

Within Europe attitudes and practices concerning renal transplantation differ significantly. This text can only provide general practice guidelines rather than taking national legislation into account.

Evaluation Criteria

Absolute contraindications are: uncontrolled hypertension, diabetes mellitus, Proteinuria (>300mg/24 h), abnormal glo-
mular filtration rate compared to normal range for age, microscopic haematuria, high risk of tromboembolism, medically significant illnesses (chronic lung disease, heart disease, etc), history of bilateral kidney stones.

- Assess potential donor for HIV-1 and -2, hepatitis C virus (HCV) and hepatitis B surface antigen (HBsAg), hepatitis D (HDV)-positive serology, acute hepatitis, cytomegalovirus (CMV), Epstein-Barr virus (EBV) (only in paediatric recipients), viral infection, sepsis, tuberculosis, infection of unknown aetiology, family history of (or clinical signs that may be caused by) Creutzfeldt-Jakob disease, and active syphilis.

- Transplant from an HCV-positive donor to an HCV-positive recipient is permissible, but only life-saving transplants should be performed in an HCV-negative recipient (after informed consent). Kidney transplant is permissible in an HBsAg-positive recipient, or an HBsAg-negative recipient with anti-HBsAg antibody titre < 10 mIU/ml, but only life-saving transplants should be performed in an HBsAg-negative recipient with no anti-HBsAg antibody (after informed consent).

- Reject donors with cancer or histories of breast carcinoma, melanoma, leukaemia, or lymphoma as well as donors who have had died from brain haemorrhage of unknown aetiology. However, non-melanoma, low-grade skin cancer and some CNS tumours may be acceptable. For a complete list consult the full guideline text.

- Tumours that do not contraindicate donation include basal cell carcinoma, non-metastatic spinocellular carcinoma of the skin, cervical carcinoma in situ, and carcinoma in situ of the vocal cords.
Donors affected by selected low-grade (grades 1 and 2) brain tumours are suitable for kidney donation. However, ventriculo-peritoneal shunting is an absolute contraindication for donors affected by brain tumours of any grade. Potential recipients affected by selected high-grade (grades 3 and 4) tumours should be transplanted only when clinically urgent.

**Kidney donation**

*Living donor* transplantation has a higher success rate than cadaveric donation. The donor must be medically and psychologically suitable and the donated organ healthy. The lower limit of calculated creatinine clearance suitable for a single kidney transplant is > 60 ml/min. A ‘marginal’ kidney has a calculated creatinine clearance of 50-60 ml/min. Kidneys with creatinine clearance < 50 ml/min are only suitable for dual transplant. Any brain-dead comatose subject (i.e., heart-beating kidney donors) is a potential organ donor. The physical condition of the donor and of the organ to be donated is more important than donor age.

**Non-Heart beating donors (NHBD):** Novel techniques for in-situ perfusion of recently dead bodies should enable greater use of NHBD. Legal implications in many countries inhibit a more wider use of this option.

**Kidney recipients**

Pre-transplant therapy:

In most cases, pre-existing pathologies that adversely affect graft survival are to be corrected before transplantation. These include:

- congenital malformation or functional disorder of the lower
Renal Transplantation

urinary tract (pre-transplant urodynamics assessment being the key-investigation)

- low-compliance bladders (avoid ureteral implantation in a fibrotic, thickened bladder wall due to the high risk of transplant loss)

- absent bladder or sphincter insufficiency (in low-compliance bladders with intact sphincters, both bladder augmentation and continent pouches with umbilical stoma are alternatives)

- autosomal dominant polycystic kidney disease (ADPKD), uni- or bilateral nephrectomy is necessary when there is insufficient space for the transplant kidney or due to complications.

- chronically infected kidneys or suspected renal or urothelial cancer.

Preoperative cardiovascular work-up is mandatory to improve organ and patient survival in the post-transplant period. Co-morbid conditions, such as diabetes mellitus and cardiovascular disease, have a major impact on morbidity and mortality of kidney transplant patients. All potential transplant candidates should be seen by an ear, nose and throat specialist, dentist dermatologist, urologist and gynaecologist to exclude infectious foci.

Matching donors and recipients

The ABO blood group and the HLA-A,-B and -DR phenotypes should be determined for all candidates awaiting kidney transplantation.

To avoid hyperacute rejection of kidney transplants cross-
Histocompatibility (HLA) matching
Transplant outcome correlates with the number of HLA mismatches. HLA-A, HLA-B and HLA-DR phenotypes should be tested in all potential recipient patients and donors. Kidneys from cadaveric donors should be allocated to potential recipients with the lowest number of mismatches at HLA loci. HLA-compatibility is less important in graft outcome in living donation, possibly because other risk factors for rejection can be minimized in living donation.

Cross matching
A cross match test must be performed before each transplant to avoid hyper-acute rejection (HAR) of transplant T-cells. Routinely a lymphocytotoxicity assays is used and T- and B-cell cross matches performed (B-cell cross matches are more sensitive for class II antigens (HLA-DR antigens)). Especially in patients with autoimmune disease false-positive results may occur. A flow cytometry cross match may be used to confirm positive results (especially in recipients at high risk of HAR, including children and sensitized patients with pre-existing circulating antibodies).

Pre-existing HLA-antibodies
In highly sensitized (PRA > 80%) patients, analysis of HLA antibody specificities (e.g. flow cytometry and enzyme-linked immunoabsorbent assay (ELISA-based methods) should be carried out to select acceptable HLA patterns in the potential donor (matched antigens and acceptable mismatches), which should result in negative cross match tests.

ABO blood group matching
Matching of ABO blood group antigens is critical in preventing HAR. Kidneys from potential donors with blood group O can theoretically be used for transplantation in A, B or AB recipients. To avoid an imbalance between demand and supply in cadaveric organs, for O recipients, ABO identity is mandatory. In living donor transplantation, ABO compatibility is as acceptable as ABO identity.

- **Viral disease**
  - Cytomegalovirus (CMV)
    Testing of cytomegalovirus infection status is necessary to define the risk of developing CMV-disease in the recipient and to plan prophylactic treatment.
    In CMV IgG antibody-negative recipients who have received a transplant from a CMV-positive donor, adequate prophylaxis with gangcyclovir is strongly recommended as the risk of primary CMV is very high.
  - HBV and HCV infection
    Whether or not HCV-positive recipients can receive HCV-positive organs is still a matter of debate because of concerns about long-term morbidity and mortality.
    Transplant recipients with HBsAg-positive infection should be monitored very closely after renal transplantation, using liver function testing and the measurement of viral replication by HBV DNA, the same applies to Hepatitis C-positive patients;

**Complications**
Renal transplant practitioners must be continuously alert for acute rejection, particularly during the first 6 months post transplant. Complication may be:

**Immunological**: Immunological rejection is the commonest
cause of early and late transplant dysfunction. HAR is a rare complication, usually occurring within minutes or hours of surgery. Delayed HAR may occur within a week of transplant. HAR is recognized by acute anuria, fever and a swollen graft and treated by graft nephrectomy.

**Acute Cellular rejection (ACR):** Patients with ACR should be tested as soon as possible for anti-HLA IgG antibodies reactive with the graft by CDC cross matching. Patients with ACR should be treated with parenteral methylprednisolone (500 mg to 1 g), given intravenously in three, daily, pulses.

**Chronic allograft rejection (CAR, humoral or unknown pathogenesis):** During the years of follow-up after renal transplantation, transplant practitioners must regularly monitor urinary protein secretion, serum creatinine and creatinine clearance. Changes in these parameters over time should trigger hospital admission for renal biopsy. If CAR is confirmed, appropriate medical treatment (e.g. control of hypertension and acidosis with ACE inhibitors) should be started.

**Malignancy:** The incidence of neoplasia in transplanted patients is higher than in the general population and is an important cause of morbidity and mortality in transplanted patients due to:
- A prior malignancy in the recipient: known or latent
- Transmission of a donor neoplasia to the recipient
- Development of a new tumour in the recipient

Immunosuppression following transplantation may stimulate growth of dormant metastases; tumours treated more than 5 years prior to transplantation may recur. The length of the waiting period varies according to tumour type.
**Immunosuppression**

The principle underlying successful immunosuppression is ‘the balance of survival’, i.e., practitioners have to prescribe a sufficient dosage of drug to suppress rejection, without at the same time endangering the life and health of the recipient. Prophylactic immunosuppression should be continued indefinitely.

Our understanding of the mechanisms involved in immune rejection has allowed the development of safer modern immunosuppressives, which are aimed at specifically suppressing sensitized lymphocyte activity against the kidney transplant.

**Cyclosporine A**

Modern therapy is based on cyclosporine A, used together with more recent drugs, such as mycophenolate mofetil (MMF) instead of azathioprine. Prednisolone is still regarded by the majority of practitioners as a fundamental adjunct to primary immunosuppression, although prednisolone withdrawal has been possible.

Cyclosporine is nephrotoxic in the majority of patients, and its long-term use may be a cause of chronic allograft nephropathy. It also causes hypercholesterolaemia, hypertension, gum hypertrophy, hirsutism and acne.

**Tacrolimus**

Tacrolimus is a more powerful immunosuppressive, but is associated with diabetes, neurological and electrolyte abnormalities, and nephrotoxicity.

It is a calcineurin inhibitor like cyclosporine A, and is therefore also associated with nephrotoxicity, though less common-
ly and to a lesser extent than cyclosporine. Blood monitoring levels of these two drugs are therefore mandatory to prevent both overdosing, leading to nephrotoxicity, and underdosing, which may lead to rejection.

**MMF**

There is well-documented evidence that MMF reduces the incidence of biopsy-proven acute rejection after transplantation. MMF is now routinely used as a primary- or second-line therapy in place of Azathioprine in many units. Nowadays, Azathioprine is usually reserved only for those patients who cannot tolerate MMF. Recently published data indicates that co-administration of MMF with cyclosporine, with or without prednisolone, allows a reduction or cessation of macrolide dosage.

**Sirolimus**

Sirolimus was licenced for clinical use in 1999 by the FDA and as an adjunct to cyclosporine therapy in Europe in 2002. The drug, a non-nephrotoxic, broadly reactive anti-proliferative for rejection, has been found to act synergistically with, and be equipotent to, cyclosporine. It shows dose-dependent, reversible thrombocytopenia and hypercholesterolaemia. Long-term follow up has not yet been reported with Sirolimus.

**Conclusion**

Over the past decade, due to a growing number of regional and national registries for treated ESRD, more information has become available concerning incidence rates, prevalence and the differences in the outcomes of the various treatment modalities.
An improved short- and long term survival of grafts, renal transplantation has become the treatment of choice for ESRD, further promoted by the introduction of new immunological suppressants.

This short booklet is based on the more comprehensive EAU guidelines (ISBN 90-806179-8-9), available to all members of the European Association of Urology at their website - www.uroweb.org.