An Introduction to Kidney Transplantation

Prevent, Detect, Support.
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An Introduction to Kidney Transplantation
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Disclaimer
Always consult your doctor before beginning any health treatment. This booklet is intended as a general introduction to the topic and should not be seen as a substitute for your doctor’s or other health professional’s advice. All care is taken to ensure that the information contained in this booklet is accurate at the time of publication.
Foreword

*An Introduction to Kidney Transplantation* is designed specifically for people who are considering treatment options for kidney disease.

If you are reading this for the first time it may be that you have just received news that you need to consider kidney disease treatment options, or you may be supporting a loved one in this situation.

*An Introduction to Kidney Transplantation* is designed to provide you with information about kidney transplantation as a kidney disease treatment option. Other booklets in this series cover home dialysis, haemodialysis, peritoneal dialysis, and supportive care. We encourage you to read all the booklets so you can make an informed decision about your treatment.

Kidney Health Australia is here to support you throughout your journey and our fully qualified team of nurses and allied health professionals are only a phone call away should you wish to speak to someone confidentially.

Welcome to Australia’s kidney community and please keep in touch so that we may be able to provide you with whatever support and assistance you require.

Mikaela Stafrace
CEO
Kidney Health Australia
What are kidneys?

Most people are born with two kidneys, each growing to the size of your fist. Your kidneys are bean shaped and are positioned near the middle of your back, on either side of your backbone (spine).

Your kidneys are part of the body’s urinary system.
What do healthy kidneys do?

Kidneys are the unsung heroes of the body.

The kidneys play a number of important roles in the body such as:
- Extracting excess fluid to make urine (wee)
- Controlling blood pressure
- Filtering waste products from the blood
- Controlling body fluids
- Stimulating the production of red blood cells
- Controlling potassium, calcium and phosphorus levels
- Regulating vitamin D
What happens when the kidneys don’t work properly?

Sometimes kidney function can change quickly. For example, your kidneys may stop working properly because of a sudden loss of large amounts of blood (e.g., during surgery) or as a result of an accident, illness or infection. A sudden change in kidney function is called acute kidney injury. This is often temporary but can occasionally lead to lasting kidney damage.

More often kidney function worsens over a number of years – it is a chronic, or ongoing, condition. Chronic kidney disease (also referred to as CKD) is called a silent disease as there are often no warning signs. Sometimes people lose up to 90 per cent of their kidney function before getting any symptoms.
Symptoms you may experience

• Retention of fluid causing breathlessness and swelling
• Tiredness
• Headaches
• Poor memory and concentration
• Irritability
• Sleep disturbances
• Feeling washed out
• Itchiness
• Nausea and loss of appetite
• Weight loss
• Altered sexual function

Dialysis is started when you no longer feel well enough to live your usual life, have excess fluid swelling, or if any of your blood results show levels of wastes that are becoming dangerous to your health.
You are not alone

In Australia:
- Over 2,000 people start dialysis or receive a transplant every year
- There are currently 21,000 people in Australia on dialysis or living with a transplant

Out of every 10 people needing treatment for kidney disease the causes are:

- Diabetes
- High blood pressure
- Nephritis
- Polycystic kidney disease
- Other
Treatment options

When you have end stage kidney disease your body cannot get rid of extra water and waste products. If this becomes dangerous to your health your kidney function will need to be replaced by either a kidney transplant or dialysis.

Some people choose not to have either dialysis or transplant and instead choose supportive care. With this option, medications and diet are used to manage the symptoms of end stage kidney disease.

A kidney transplant involves the transplantation of one kidney into your body from either a living or deceased donor.

Dialysis has a number of options, which include home dialysis (either peritoneal dialysis or home haemodialysis) and centre-based haemodialysis.

If you are suitable for a kidney transplant, this is the most effective treatment for kidney failure. Compared with other kidney failure treatment options, transplantation will increase your life expectancy and improve your quality of life.
RESOURCES

Kidney Health Australia fact sheets (translated versions available): Kidney Transplants, Treatment Options

Kidney Health Australia booklets: An Introduction to Haemodialysis, An Introduction to Peritoneal Dialysis; An Introduction to Home Dialysis, An Introduction to Supportive Care of Advanced Kidney Disease, Kidney Failure Treatment Options (translated versions available)

Kidney Health Australia book: *Living with Kidney Failure*


Kidney Health Information Service (KHIS) **1800 454 363**
Transplantation

A kidney transplant is treatment for kidney failure but it is not a cure. A transplant offers a more active life, without needing dialysis. However, your new kidney requires ongoing care. You will need to take medications to stop your body rejecting the kidney (anti-rejection) for as long as you have the transplanted kidney.

How long your transplanted kidney will work can depend on many factors. In Australia, around four in every ten kidney transplants are still working 15 years later. Some people have had kidney transplants that have lasted more than 30 years.

If a kidney transplant stops working, dialysis treatment will be necessary again. Another transplant may also be possible.

CALL TO ACTION

Talk to your health care team about whether a kidney transplant is a suitable option for you. Find out where your closest transplant hospital is.
Who can have a kidney transplant?
Kidneys can be donated by living donors or deceased donors. Having a transplant from a living kidney donor means you will have better outcomes. If you have a transplant from a living donor it usually means you will not have to wait as long for your transplant.

You can be given the option of transplantation if you are considered to be medically suitable by the kidney transplant team. Unfortunately a transplant is not suitable for everyone, as the risks are sometimes too high. The transplant team may decide that the risks of the operation or the medications may make your health worse.

However, it is everyone’s right to be considered for a transplant. You will need to speak with your nephrologist for further information about your circumstances if you are regarded as not being suitable for a kidney transplant.

Factors affecting your suitability for a kidney transplant include:
- Agreement with the idea of transplantation and acceptance of the risks involved
- Your general health, apart from kidney failure
- Willingness to go through with the tests and the transplant surgery
- Willingness to take the anti-rejection medications.

Most metropolitan cities have a hospital that performs kidney transplants. If you live in a regional or rural area of Australia or in a city that does not have a transplant hospital you will need to travel to the closest transplant hospital for your evaluation and tests, and for the transplant procedure itself. Your health care team will be able to discuss these travel arrangements with you.
Are kidney transplants successful?
Kidney transplants are very successful. On average, 95 per cent of transplants are working one year later. If the transplant works well for the first year, the chances are good that it will function very well for many years.

Success rates are higher with living donor kidneys than for deceased donor kidneys.

<table>
<thead>
<tr>
<th>Percentage of people alive after five years who received a kidney from a:</th>
<th>Percentage of transplanted kidneys working after five years from a:</th>
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<tbody>
<tr>
<td>96%</td>
<td>90%</td>
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<tr>
<td>Living donor</td>
<td>Living donor</td>
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<tr>
<td>90%</td>
<td>81%</td>
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<tr>
<td>Deceased donor</td>
<td>Deceased donor</td>
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RESOURCES

Kidney donors

Kidneys for transplantation come from either living donors or deceased donors.

For every 10 kidney transplants performed in Australia:

three involve kidneys from living kidney donors

seven involve kidneys from deceased kidney donors

The surgery and pre-surgical care for you will be the same whether your transplanted kidney is from a living or deceased donor. It is recommended that you discuss your situation with your health care team, to make sure you understand your options.

CALL TO ACTION

Talk about kidney transplantation with your family and friends. Discuss the options of receiving a kidney from a living or deceased donor.
Living kidney donors

Living donors are usually relatives (parents, brothers, sisters, or adult children) of the person who needs the kidney transplant. If circumstances are suitable, people who are not related to you, such as your partner or friend, may also donate a kidney.

Another type of living kidney donation is called non-directed kidney donation or altruistic donation. An altruistic donor is a person who is willing to give the gift of life to someone who is not only unrelated to them by blood or marriage but a complete stranger. These donors are first matched in the Australian live kidney donor exchange program. With this program an altruistic donor starts a chain of kidney transplants. Instead of just one person benefiting from their donation, this donor can allow many incompatible pairs to be transplanted. See more about the Paired Kidney Exchange program on page 53.

The tests to determine whether a living donor will be a compatible donor for you are very complex and can take a long time. New techniques mean that transplants can often now be done in situations that were previously difficult. For example, if your potential living donor is not from a compatible blood group there are now ways to deal with this.

The main benefit of having a living donor is timing. It can sometimes take years before a kidney becomes available from a deceased donor. Having a living donor may mean you can have a transplant before you even need to start dialysis. This is called a pre-emptive transplantation.
Having a living donor also means that the transplant surgery can be organised and planned. This means that the time between when the kidney is removed from your donor and transplanted into you (called the **cold ischaemia time**) is shorter. A shorter cold ischaemic time improves your chances of a successful transplant surgery and contributes to longer kidney transplant survival.

Hospital admission and surgery can also be planned ahead of time, allowing you and your donor time to prepare for the surgery and recovery.

Another advantage of having a living kidney donor is that transplant success rates are higher with living donor kidneys than with deceased donor kidneys.

**RESOURCES**

- Kidney Health Australia fact sheets: Deciding to be a live donor, Live kidney donation, Organ and tissue donation and transplantation, Non-directed living kidney donation
- Kidney Health Information Service (KHIS) **1800 454 363**
Deceased kidney donors
Deceased donors are people who die with organs that are suitable for transplantation.

In Australia, most donations occur when a person is declared brain dead. This is called Donation after Brain Death (DBD). Brain death occurs when the brain is so badly damaged that it permanently stops functioning. This is usually because of bleeding in the brain, a stroke, infection or severe head injury.

Another method of organ donation is known as Donation after Cardiac Death or Donation after Circulatory Death (DCD). Potential DCD donors are patients in the Intensive Care Unit (ICU) who have suffered irreversible brain injury and may be near death, but do not meet formal brain death criteria. These patients have no chance of survival without ongoing artificial ventilation or life support. In these cases, the family has decided to stop care. When the patient’s heart stops beating, the organs are then removed in the operating room. These organs have had some loss of oxygen after the heart stops beating. This may make kidneys from this type of donor slow to start working.

Deceased donors may have recorded their decision to be an organ donor on the Organ Donor Register. Their family may make this decision after the person has died. This is a very special gift and the donor and their family are treated with the greatest respect throughout the decision and the surgery.

Unfortunately there are not many deceased donors each year. Organ donation requires special conditions and is only possible in less than 1 out of every 100 deaths that occur in hospital.
REMEMBER

Any person is around 10 times more likely to need an organ transplant than to become a deceased organ donor.

RESOURCES

Kidney Health Australia fact sheets: Organ and Tissue Donation and Transplantation
Australian Organ Donor Register www.donatelifegov.au
The kidney transplant process

**Assessment to have a kidney transplant**
Before you can be placed on the kidney transplant waiting list you will need to undergo many tests. These tests are to make sure that both your body and mind are healthy enough for you to have the kidney transplant surgery and take the medications after surgery. These tests are often called the transplant work-up.

Medical tests that are used to assess your health before a transplant include:
- Assessment of your overall health, including your weight
- Blood tests
- Dental check
- Pap smear and mammogram (females) or prostate tests (males)
- X-rays and other tests of your heart and lungs
- Tests involving your arteries, veins and bladder.

**Living Donor assessment**
If you have someone who wants to be your living kidney donor they will probably go through a similar work-up at the same time as you. They will also have tests to make sure their kidney is healthy enough to be donated. There will also be tests to check if the kidney from your living donor is compatible or *matched* with you. More information on how kidneys are matched can be found on page 29.

The process of assessment for both you and any potential living kidney donors can take between six months and a year depending on any issues that are faced.
Transplant waiting list
When you have been evaluated as being suitable for a kidney transplant your name will be placed on the transplant waiting list to receive a kidney from a deceased donor. You will be referred to as being active on the transplant waiting list.

You will only be activated on the transplant waiting list for a deceased donor once you have started dialysis. This does not mean you still can’t look into having a transplant from a living donor. If you have started dialysis you can go on the transplant waiting list while you are waiting to see if you have a compatible living kidney donor.

If your health gets worse while you are waiting for a kidney, you may be temporarily taken off the active list. This is because your transplant would have less chance of success if you are not well. Once you are well you will be active on the waiting list again.

The kidney transplant waiting list is not just a queue where you slowly work your way to the front. Each time a deceased donor kidney becomes available a recipient is chosen taking into account a number of factors. A computer generates a score for these various factors, and the recipient with the highest score will be offered the kidney. See page 23 for more information about how deceased donor kidneys are matched and allocated.
Your waiting time is based on the day you started dialysis, regardless of when you were officially placed on the waiting list. If you come off the waiting list for any reason the same ‘dialysis start’ date is used when you become active on the list again. This means that you do not lose your accumulated waiting time.

If you are receiving a second, third or even fourth transplant, your waiting time is based on the date that dialysis was re-started after your transplanted kidney failed.

**REMEMBER**

The kidney transplant waiting list is not just a queue where you slowly work your way to the top. Your healthcare team can let you know if there are issues that may make your wait longer or shorter than average.
National and State allocation of deceased donor kidneys

There is a National Allocation Formula for allocating kidneys from deceased donors. This means a kidney may be sent interstate if that is where the recipient with the best match is.

If no one meets the criteria of the National Allocation Formula the State Allocation Formula is used. This means that the kidney is allocated to a recipient who lives in the same state as where the deceased donor is. In these cases, the kidneys are mostly allocated based on waiting time on dialysis.

Out of every 10 deceased donor kidneys, two are sent interstate and eight are allocated within the state.

Young children get priority for deceased donor kidneys. This is because dialysis impacts on their growth and education. As there are not many children waiting for a deceased donor kidney this policy has very little impact on the average waiting time.

Overseas visitors are not eligible for an Australian transplant.

Unfortunately it may take months or even years of being active on the kidney transplant waiting list before you receive a kidney from a deceased donor.

RESOURCES

Staying healthy while waiting for a kidney transplant
To give your transplant the best chance of success, it is important that you look after yourself while you are on the transplant waiting list.

You will also need to provide routine tissue typing and antibody blood tests every one to two months. This allows current samples to be used in the cross match tests. Small amounts of these samples are sent around Australia in case a donor matches with you in another state. If you do not do these blood tests, the laboratory will not be able to assess your compatibility with a donor and you may miss an opportunity. You will also have regular reviews where you can discuss any concerns about your transplant preparation.

Stay healthy while waiting for your kidney transplant by:
- Keeping to your regular dialysis schedule
- Following your recommended diet and fluid restrictions
- Taking your medications as prescribed
- Controlling your weight with diet and physical activity
- Visiting your dentist regularly to watch for any infections
- Protecting your skin, which will help to prevent skin cancer after your transplant
- Being a non-smoker
- Keeping your vaccinations up to date
- Avoiding blood transfusions if possible.
RESOURCES

Kidney Health Australia fact sheet: Common kidney disease symptom management options, Depression and chronic kidney disease, Looking after yourself with chronic kidney disease (translated version available)

Kidney Health Australia book: Living with Kidney Failure

Kidney Health Information Service (KHIS) 1800 454 363
What happens when a deceased donor kidney becomes available?
When a kidney from a deceased donor becomes available for transplantation a number of laboratory tests are done. These tests include tissue typing and cross matching. The tests are done to work out who would be the best match for the kidneys.

Once the laboratory tests have been completed the Organ Donor Coordinator will receive a list of potential recipients. They will then contact the Transplant Unit responsible for the first potential recipient on the list.

If you are selected to receive a kidney from a deceased donor the hospital will contact you by telephone.

**REMEMBER**

Make sure the hospital has your current telephone numbers as you may receive the phone call at any time – day or night. If you cannot be located and do not contact the hospital within a certain time (usually around 30 minutes), the kidney will need to be given to the next person on the waiting list. It is important that you are always prepared to answer your phone.
You will also need to be able to get to the hospital very quickly as deceased donor kidneys are usually transplanted within 24 hours of their removal from the donor. This is to make sure that there is limited damage to the kidney from the loss of oxygen and blood supply. The kidney will be put on ice until it can be transplanted into your body.

CALL TO ACTION

Pack a hospital bag so you are ready at any time. Include pyjamas, warm socks, toiletries, pen and paper, small amount of change, essential phone numbers, any necessary medical documents.
Deciding whether to accept the kidney

Once you have received the phone call that there may be a kidney available for you, there may be further issues to think about. Your doctors will need to assess the compatibility, the safety, as well as the quality (likely survival) of the kidney that is being offered for donation.

Some of these issues may involve possible risks to your health. For example, if it is discovered that the donor had suffered from cancer or other infections.

In recent years there are more kidneys coming from elderly donors who may have had illnesses such as high blood pressure, diabetes or a history of smoking. In some cases, donors may have had a history of taking illegal drugs.

All these factors need to be taken into account by your doctors and by yourself. The quality of the kidney being offered may also influence the decision to go ahead with the transplant.

You may have to decide whether it’s best to accept an immediately available, but less-than-ideal deceased donor kidney or take the chance and wait for a healthier one in the future.

CALL TO ACTION

Take some time to think about these issues before you receive your call that a kidney is available. Talk to your health care team and your family about your decisions.
How are kidneys matched and allocated?

Kidney donor and recipient matching can be divided into several areas. These areas include blood group matching, tissue type matching and cross matching. Soon Australia will also be considering survival matching. This involves the assessment of how long a certain kidney is likely to last. It also involves allocating kidneys partly based on how long different recipients are likely to survive.

For these reasons, allocation and acceptance of a kidney offer needs to consider compatibility, your waiting time and the quality of the kidney being offered and its likely benefit to you.

A computer can manage some of these factors. Your doctors will also need to consider other issues and possibly discuss these with you if needed.

Each area of compatibility is important for donor and recipient matching for both living and deceased donors.

If you are receiving a kidney from a living donor it may be possible to get around some of these matching issues. For example, it may be possible to prepare your immune system so that you are able to receive a kidney that would have been incompatible.

It may also be possible for you and your living kidney donor to participate in the Australian Paired Kidney Exchange Program (see page 53).
Donor and recipient matching

1. Blood group
If your transplanted kidney comes from a deceased donor, the ordinary blood groups (A, B, AB, O) and red blood cells of the donor must be compatible with yours. This is similar to matching blood groups for a blood transfusion. We all have antibodies to blood groups different to ours.

If your transplanted kidney comes from a living donor it may be possible to have the transplant even if you and your donor are not from a compatible blood group. This is called an ABO incompatible transplant. Extra treatments will be needed to remove the antibodies before the transplant can go ahead. ABO incompatible live donor transplants are being performed in most Australian transplant units with excellent results.

2. Tissue type matching
Your immune system is designed to keep you safe from foreign invaders such as viruses and bacteria. Everyone has special proteins on their cells called HLA proteins. Your combination is unique to you and is recognised by your immune system so it will not attack your own cells.

Your immune system will recognise a combination of foreign proteins not belonging to you and will attack them. It will then develop cytotoxic antibodies against these HLA proteins. This is how your immune system defends your body against infections. This is also the underlying cause of your body rejecting a transplant and the reason you need to take anti-rejection medications.
Cytotoxic antibodies are called *donor specific antibodies* if they are shown to be against your potential donor’s HLA proteins. These can make it hard for you to be compatible with other donors. These antibodies may have developed because you have previously been exposed to other people’s HLA proteins through a blood transfusion, through pregnancy, or from a previous transplant. Sometimes there is no clear reason found for the presence of these HLA-antibodies.

Your immune system remembers this exposure and is ready to attack cells with similar HLA-proteins on them. The *cross match* test also checks for this.

**HLA proteins and cytotoxic antibodies**

You have a combination of A, B and DR HLA proteins inherited from your parents. When you are exposed to other HLA proteins your immune system attacks them. Your body then develops antibodies against these HLA proteins. If you have already have antibodies against a donor cell HLA protein (B6 in the diagram above) then your body may attack those donor cells and reject the donor kidney.
When you are tissue typed, a blood test is taken to determine your HLA proteins and cytotoxic antibodies. This is usually one of the first steps in your transplant work-up. Your HLA proteins and cytotoxic antibodies can also change over time, so these tests will be repeated every one to two months.

You will share some HLA proteins with other people and the more you share the more likely your body is to accept their kidney. How similar your tissue typing is with someone else’s refers to your ‘HLA-match’. This is usually given as a number out of six as there were six groups of proteins identified many years ago that were found to be important.

For many people, HLA-matching is less important than it was a few decades ago. This is because the anti-rejection medications have improved. We also understand more about tissue typing than we used to, and this has led to longer survival of transplanted kidneys.
REMEMBER

Most kidney transplants that are performed do not have a high level of HLA protein matching and still have excellent results. Many people would wait a long time or never get transplanted if they waited for a very good HLA-match.

What is more important is whether you already have anti-HLA or cytotoxic antibodies in your blood against any foreign HLA proteins in the potential transplant kidney.

3. Cross match
In the cross match test, your blood cells are added to blood cells taken from the donor. A positive reaction means that there is a high likelihood of severe rejection of the donor kidney. In the case of a deceased donor, the transplant will not go ahead if there is a positive cross match.
What happens during a kidney transplant?

**Before surgery**
You will be admitted to the hospital where the transplant surgery will take place. You will have a physical examination, and you may need to have a dialysis session. If you have any active infections or other significant medical problems the transplant surgery may need to be cancelled.

You will be given anti-rejection medications before and after the surgery to control your immune system and stop your body from rejecting the new kidney.

**During surgery**
Surgery for a kidney transplant takes about two to three hours. A cut is made in your lower abdomen, on the right or left side. The new kidney is placed in your pelvis. The renal **artery** and **vein** of the transplant kidney are connected to an artery and vein in your pelvis (usually the iliac artery and vein that lead down towards the leg). The **ureter** of the transplanted kidney is connected to your bladder so urine can flow.

Many people are surprised to learn that their own failed kidneys are not removed. They are left in your body to continue to provide whatever amount of function they may still have. Sometimes it may be necessary to remove your failed kidneys if they are very large (if you have polycystic kidney disease) or in the case of chronic infection. If this is needed, your failed kidneys will be removed in a separate surgical procedure before your transplant surgery.
Kidney transplant anatomy

Diseased kidneys

Transplanted kidney

Transplanted ureter
After surgery
After the transplant surgery it is normal to feel some pain around your wound. You will be given medication to help with this.

Your transplanted kidney may start to make urine immediately, or you may need dialysis for a few days. If you need dialysis it does not mean that your transplanted kidney will not work. It just may need a bit of time to recover from the transplant procedure.

A catheter will be placed in your bladder for around five days to drain your urine into a bag. This helps your health care team to check that your kidney is working well. Often there are tubes from your wound draining away extra fluid. These are usually removed after a few days.

You will have blood tests every day to check the function of your transplanted kidney, to measure medication levels and to detect any problems early so they can be treated quickly.

The blood tests for creatinine and eGFR measure how well your kidney is working. These may already be familiar to you. In many cases your creatinine and eGFR will go back to what they were before you first got kidney disease.

The amount of fluid you take in is important. The nurses may be asking you regularly how much you have had to drink. If your kidney is working you may find yourself having to drink lots of fluid.
A physiotherapist may assist you with an exercise plan. You will probably be able to sit out of bed on the first day after your surgery. You should be able to go for a walk within a couple of days.

Reducing the risk of an infection is very important. For this reason, you may be looked after in a separate room or ward of the hospital, your visitor numbers may be limited, and you may not be allowed to receive flowers from visitors.

The length of time that you will need to stay in hospital after the transplant surgery will depend on how well your body responds to the new kidney and whether you have any complications. Most people are in hospital between six and ten days. You may feel better immediately after your surgery or you may take longer to adjust.

**REMEMBER**

It is important to do coughing, breathing and leg exercises while you are restricted to bed rest.

This helps to:
- Reduce the risk of a chest infection
- Reduce the chance of clots in your legs
- Keep your airways clear
- Encourage good blood flow.
Going home with a successful transplant

Caring for yourself after your kidney transplant is very important. The first few months following your transplant are often when things will be the most unstable and you may be at risk of developing problems. It is important that you report any changes in your health to your health care team as soon as possible.

You will have regular check-ups to monitor the function of your new kidney. These will be daily at first, then weekly, then monthly.

At these visits your wound will be assessed to make sure it is healing well, and you will be checked for any signs of rejection of the transplanted kidney. Slow wound healing can be caused by some medications, diabetes and obesity. If you are at increased risk you will be monitored closely.

You will need to think about where you will stay after you are discharged from hospital. If you live a long distance from the transplant unit you will need to find short-term accommodation close to the hospital. Travel and accommodation assistance schemes are available to help with this (see What Support is Available on page 62).

CALL TO ACTION

Think about where you will stay after you are discharged from hospital. If you need to arrange accommodation it is best to do this sooner rather than later.
Living with your new kidney
You should be able to return to normal activities and work within three to six months of your transplant. You will need to allow time for your wound to heal and for your stomach muscles to get strong again. Regular exercise is an important part of staying healthy. It is recommended that you start with gentle exercise first.

It is recommended that you do not drive a car until six weeks after your transplant. You should also avoid any heavy lifting for six weeks.

You will need to manage your medications carefully. It is important that you take your medications exactly as prescribed by your doctor.

It is important your kidney has enough fluid to work properly. You need to make sure you drink enough water or fluids to stop you feeling thirsty, and to keep your urine a pale yellow colour.
CALL TO ACTION

Check the colour of your urine regularly. If the colour is too dark drink some water. If the colour is unusual, red or the urine is frothy speak to your health care team.
More tips on staying healthy

Avoid infections
- Limit contact with people with colds or viruses (especially small children)
- Wash your hands well
- Treat scratches
- Keep your vaccinations up to date

Reduce your risk of skin cancer
- Follow the **slip, slop, slap, seek, slide** advice provided by the Cancer Council of Australia (slip on a shirt, slop on sunscreen, slap on a heat, seek shade, and slide on sunglasses)
- Have regular checks for other cancers such as breast cancer (women) and prostate cancer (men)

Be a non-smoker
- Having a transplant increases your risk of having health problems caused by smoking
- This includes lung cancer, stroke, and heart attack
- Your chances of a long-term, successful transplant may be reduced if you smoke

Let your health care team know if you get sick
- If your illness is treated quickly your risk of damaging your transplant is reduced
Stay at a healthy weight
- Your anti-rejection medications may increase your appetite
- Talk to your doctor or renal dietitian about how to stay at a healthy weight

Avoid food poisoning
- Wash your hands well
- Keep cooking utensils, chopping boards, and knives separate for raw and cooked meats
- Cook foods thoroughly – no pink left in cooked meats such as mince, sausages and chicken
- Avoid eating foods that have a higher risk of food poisoning – soft cheeses (such as brie and camembert), pate, salami, and raw seafood

CALL TO ACTION
Visit www.foodauthority.nsw.gov.au > Food safety & you > Keeping food safe > Key tips for tips on how to avoid food poisoning.
Sexuality and fertility after a transplant

Sexual activity will not harm your transplanted kidney. It will also not increase your risk of infection. It is recommended that you wait about four weeks after your transplant before having sexual intercourse.

As your health improves after your kidney transplant your fertility may increase. This is true for both men and women.

If you are a woman and would like to have a baby it is usually suggested that you wait at least 1 year after your transplant before trying to get pregnant. This allows time for you to become stable on your anti-rejection medication.

If you become pregnant some of your medications may need to be changed and you will be carefully monitored to make pregnancy safer for you and your baby.

REMEMBER
Speak to your doctor if you are thinking about becoming pregnant after your kidney transplant.

RESOURCES
Kidney Health Australia fact sheet: Sexuality, fertility and kidney disease
Kidney Health Australia book: Living with Kidney Failure
Emotional well-being
Having a transplant should have a positive effect on your life. However it is still a major life event. You may have mood swings and feel stressed or depressed as you adjust to your transplant, and as your body responds to your anti-rejection medications. It is not unusual to have these feelings. The earlier you talk to someone the quicker you can get support.

You may also:
- Have to come to terms with having another person’s organ inside you. Some people worry that they may develop the donor’s personality but this is not possible
- Have conflicting feelings about receiving a kidney from a deceased donor. You may feel relief and happiness at having a new kidney, but upset because the donor’s family and friends are grieving
- Be overwhelmed by feelings of being normal again
- Find that your relationships change as your family and friends adjust to you becoming more active
- Experience a roller-coaster of emotions about the possibility of rejection. You may feel anxious about attending medical appointments and then relief after being told that your transplant kidney is functioning well
- Be overwhelmed at the thought of taking many medications with their possible side effects
- Feel guilty about your transplant because there are other people still on dialysis.
Many people who receive a deceased donor kidney may wonder who the kidney came from. This information is not released but Donate Life offer a number of ways of recognising donors. This includes annual services that involve both donors and transplant recipients.

REMEMBER

People deal with a transplant in different ways. Your emotions affect your health so it is important to talk about your thoughts and feelings with your family, friends and health care team.

CALL TO ACTION

Many transplant centres also hold information workshops for people with a kidney transplant. These can add to the discussions that you have with your health care team.

RESOURCES

www.donatelife.gov.au > Donor family support services
Transplant rejection

You must take anti-rejection medications for as long as your transplanted kidney is working. These medications partially block the activity of your body’s immune system. This prevents your body from attacking your transplanted kidney.

There are many different medications which are used to prevent rejection. For each of your medications find out:

- When and how to take them
- What to do if you miss a dose
- Possible dietary restrictions
- Interactions with other medications
- What side effects you may experience
- How to store your medications.

In the early period after your transplant, you may have some rejection episodes. These may only be picked up by your regular blood tests. These episodes can usually be managed with changes to your medications.

Some rejection episodes may require extra treatments such as plasma exchange (to remove antibodies in your blood) or special infusions.

**REMEMBER**

Most rejection episodes can be managed successfully. However, rejection can take place at any time after your transplant so it is very important that you follow any medical instructions and receive regular monitoring.
Types of rejection episodes

Hyperacute rejection
This can occur minutes or hours after the transplant. This type of rejection is very rare. It is untreatable and the kidney is removed immediately.

Acute rejection
Acute rejection appears as an inflammation in the kidney. This usually needs quick diagnosis and action. It usually occurs in the early weeks following a transplant. This is why you will have frequent blood tests in the first weeks after your transplant. Mild acute rejection is very common in the first year after a kidney transplant.

Acute rejection sometimes causes pain and fever but usually has no symptoms. An increasing eGFR or creatinine is usually the first sign of acute rejection. A biopsy of the kidney transplant is often used to diagnosis rejection and to decide on the best treatment.

Chronic rejection
Chronic rejection refers to a gradual process, which leads to scarring and damage in the transplanted kidney. This usually occurs over several years and can be very difficult to treat. Often this is due to anti-HLA antibodies that are present (see page 31). During this time the kidney is gradually losing its function and may eventually fail, leading to the need for dialysis and another kidney transplant.
Anti-rejection medication side effects

Anti-rejection medications have a number of possible side effects. These side effects are usually able to be managed by changing the dose of your medications.

Some of the most common side effects include:

- Increased risk of infections
- Puffiness of your face and stomach
- Unwanted hair growth or hair loss
- Increased appetite and weight gain
- Mood swings
- Increased risk of high blood pressure, diabetes and cancer (particularly skin cancer)
- Pimples
- Muscle weakness and shakiness in your hands
- Stomach problems such as indigestion and diarrhoea

If you become worried about any possible side effects, you should discuss them with your health care team.

If you need treatment from other health professionals (such as a dentist), let them know that you have had a kidney transplant.
REMEMBER

Check with your nephrologist before taking any medication prescribed by another health professional. Check before taking any medication that you can purchase over-the-counter from the supermarket or chemist, as these may affect your anti-rejection medications. Do not stop taking your medication unless you are told to do so by your health care team.
The live kidney donation process

Kidney transplants from live donors now make up around three out of every ten kidney transplants in Australia each year.

To be a living kidney donor, the potential donor must have normal kidney function, overall good health, and be emotionally suitable for living donation.

The evaluation process includes blood tests and other investigations, as well as a psychological assessment.

Medical assessment
If someone is considering being a living kidney donor they will need a thorough medical check by a doctor. These tests will check that donating a kidney will be safe for the donor both in the short term and the long term. The transplantation team will also decide whether the donor is fit for surgery and has a suitable kidney to donate.
A number of tests are needed to make sure that the living donor is suitable to donate their kidney to you. These may include:

- Tissue typing
- Blood group
- Testing the recipient for anti-HLA antibodies
- Blood tests and scans to check their kidneys are functioning well and to make sure there are no signs of kidney damage
- Tests to find any viruses that may lead to an infection
- Tests for heart disease, diabetes, lung problems and cancer.

During these tests donors may find out that they have a health problem that they did not know about. If this happens they will receive support and referral to a specialist.

**REMEMBER**

Not everyone is able to be a living kidney donor. Some conditions that may prevent someone from being a living kidney donor include:

- Having cancer (or a history of cancer), hepatitis or AIDS
- Having diabetes, or having an increased risk of developing diabetes
- Having heart or lung problems (including high blood pressure)
- Being overweight
- Being a smoker
- Psychological concerns.
**Psychological assessment**
A psychological assessment is performed to make sure the living kidney donor is certain about donating and making a voluntary and informed choice.

Living kidney donors must be fully aware of the risks to their own health. The transplant team helps the donor to weigh up the risks and benefits and understand the medical procedure. This is called informed consent.

A psychological assessment also helps to make sure that the donor:
- Is comfortable with the idea of donation
- Is not being forced or paid for the donation
- Has a good understanding of the physical and emotional outcomes of kidney donation.

**REMEMBER**
The living kidney donor can change their decision to donate at any time; right up to the time that the surgery takes place.

**RESOURCES**
- Kidney Health Australia fact sheets: Deciding to be a live donor, Live kidney donation, Non-directed living kidney donation
- Kidney Health Information Service (KHIS) 1800 454 363
Australian Paired Kidney Exchange Program (AKX Program)

If you have a living kidney donor that is not a match for you it may be possible to enrol in the AKX Program. The AKX Program identifies matches between transplant recipients and living donors.

A computer program searches the entire database of registered recipient and living kidney donor pairs. The program looks for combinations where the donor in an incompatible pair can be matched to a recipient in another pair. If the computer finds a compatible match, two or more transplants can occur by swapping the living kidney donors.

More than 170 transplants have been successfully performed in Australia as part of the AKX program.

RESOURCES

www.donatelife.gov.au > About us > Kidney exchange programme
What happens during a living kidney donor transplant?

**Before surgery**
In the week before surgery, you and your living kidney donor will be re-tested to make sure you are both healthy and emotionally ready for the surgery. Another serum cross match test is done. The day before surgery, you and your donor go to hospital. You (the kidney recipient) may have a dialysis session or other special treatments.

**During surgery**
There are two ways of removing the kidney from a living donor; **laparoscopic nephrectomy** or **open nephrectomy**. The transplant team will discuss these procedures in detail with your living kidney donor. Most live donation procedures now occur through laparoscopy.

The procedure for transplanting the living donor’s kidney into your body is the same as the procedure for a deceased donor kidney (see page 34).
Laparoscopic nephrectomy
This is also called keyhole surgery. The surgeon inserts special instruments into the living donor’s abdomen through three small cuts (also called incisions). A thin tube with a video camera on the tip (called a laparoscope) is inserted into the donor’s abdomen. This allows the surgeon to see inside the living donor’s body.

The kidney is collected in one of two ways:
• Endo catch – a drawstring bag is used to catch the kidney and draw it out of their body through a small cut about 10–12cm long
• Handport – the surgeon’s hand is used to remove the kidney.
Open nephrectomy
The surgeon makes a cut about 15–25cm long on the side or front of the living donor’s abdomen. The cut is along the bottom of their lower rib to a point just above their belly button. The cut will be on their left or right side depending on which kidney is being donated. The vessels connecting the kidney to its blood supply are clamped and cut. Their ureter (which is connected to their bladder) is disconnected.
**After surgery**

After the transplant surgery your living kidney donor will feel some pain around their wound. They will be given medication to help with this.

The length of time that your living donor will need to stay in hospital may be around four to ten days.

They will then need up to six weeks of recovery time at home. It is recommended that your living kidney donor **does not drive a car** until six weeks after the surgery. They should also avoid any **heavy lifting** for six weeks.

Living donors who have laproscopic (keyhole) surgery often recover quicker than living donors who have an open nephrectomy. Everyone’s experience is different so it is important to talk about recovery with the health care team, family and friends.
What are the risks to living kidney donors?

Physical risks
Being a living kidney donor means having major surgery. This always carries a risk of serious complications including death.

Complications that may occur during surgery include:
- Punctured lung (pneumothorax)
- Lung infection (pneumonia)
- Nerve damage
- Blood clots (thromboembolism)
- Bowel perforation
- Bleeding.

The transplant surgeon will talk to your living kidney donor about these risks in detail.

Living kidney donors are unlikely to develop kidney problems in the future. The removal of one kidney triggers the other kidney to increase in size and function. The remaining kidney can provide up to 75 per cent of normal kidney function rather than the expected 50 per cent. Any decline in overall kidney function is usually mild. This has no impact on the life span of your living kidney donor.
However, living kidney donors may be at increased risk of kidney disease if their remaining kidney is injured or a disease develops unexpectedly. There is also a small risk of increased blood pressure and protein in their urine as they get older.

An annual check-up including a general health review, blood pressure measurement, and blood and urine tests are recommended for all living kidney donors.

RESOURCES

Kidney Health Australia fact sheet: How to look after your kidneys, Life with a single kidney
**Emotional risks**

Most living kidney donors say they feel very satisfied with the donation experience. Even if the transplant is not successful, many donors say they feel positive about their decision.

There is a small chance for living kidney donors to experience mixed feelings after the donation. These feelings may be more likely if the transplant surgery has not gone as well as expected.

It is also common for living kidney donors to experience depression and anxiety after surgery. This is normal, and happens after many different types of surgery. Your donor’s health care team can suggest ways of helping to manage this.

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**CALL TO ACTION**

*beyondblue* (beyondblue.org.au or call 1300 22 4636) provide a confidential telephone information and advice helpline if you are feeling anxious or depressed.
Combined kidney and pancreas transplantation

If you have developed kidney failure due to the complications of diabetes it may be possible for you to be considered for a combined kidney and pancreas transplant.

If you are medically and emotionally suitable for a kidney transplant you may also be suitable for a combined kidney and pancreas transplant if you:
- Have type 1 diabetes and are dependent on insulin
- Are ideally aged less than 50 years
- Have no history of heart disease.

A number of extra tests will be performed to check what impact diabetes has had on your blood vessels, eyes, nerves and kidneys. These tests are needed to make sure that the benefits of the transplant surgery are greater than any risks.

Outcomes of combined kidney and pancreas transplants vary from person to person. If you would like to consider a combined kidney and pancreas transplant your health care team can discuss this with you.
What support is available?

There is support available for you if you are considering a kidney transplant or living with a transplant. Your hospital will also have staff available to help. This may include an accommodation officer, a patient liaison officer, and a social work department.

RESOURCES

Kidney Health Australia has a range of fact sheets on this topic. See the Patient & carer support page at kidney.org.au
Accommodation assistance
If you live in a rural or regional area you may need to find short-term accommodation close to your transplant hospital. Kidney Health Australia has secure, comfortable quality accommodation where you can rest and recover from your transplant surgery. Kidney Transplant Houses are currently available in Perth and Melbourne, and construction is underway in Adelaide. More information is available at kidney.org.au > your-kidneys > support > patient-and-carer-support > kidney-transplant-houses.

Some hospitals have an accommodation liaison officer to help patients and their families to find suitable accommodation close to the hospital. In hospitals that do not have an accommodation officer, the patient liaison officer or social work department will be able to help.

Transport assistance
If you live in a rural or regional area you may be eligible for financial support from your state government to help with the costs of travelling to and from your transplant hospital. More information is available at kidney.org.au > your-kidneys > support > patient-and-carer-support > country-travel.

Travelling after a transplant
Travelling for work or pleasure within Australia and even overseas is possible after receiving a kidney transplant. However, you will require special attention to make sure that both you and your transplanted kidney stay healthy.

Kidney Health Australia has a fact sheet that provides details and useful tips about travelling with a kidney transplant available here kidney.org.au > about-us > resources-library > fact-sheets.
Financial assistance
If chronic kidney disease is causing you and your family financial hardship, there is a range of assistance available.

Start by speaking with the patient liaison officer or the social work department at your hospital. More information on the type of financial assistance is available at kidney.org.au > your-kidneys > support > patient-and-carer-support > finance.

There is also support available for living kidney donors. If your living kidney donor is employed they might be eligible for the Supporting Leave for Living Organ Donors Programme. This scheme provides reimbursement to their employer for up to 9 weeks of leave (based on a 38 hour week) of leave, at an amount up to the National Minimum Wage. For more information or to register see the Department of Health page www.health.gov.au/internet/main/publishing.nsf/Content/Leave-for-living-organ-donors or call (02) 6289 5055.
How do I make the choice?

The choice between a kidney transplant, dialysis and supportive care depends on factors such as your age, health and lifestyle.

Education is the first step to choosing what type of treatment you will have. Take advantage of the nurses and social workers who offer group and individual education sessions. Use booklets (like this one), good quality websites, videos, and consider talking to other people in the same position. Your health care team can offer advice on any possible reasons why a particular treatment may not be possible due to current or previous health issues.

My Kidneys, My Choice is a useful decision aid that will help you to make your choice. It contains a check-list of issues that you should consider when making your choice. The decision aid is available from the Treatment for kidney disease > Choosing your treatment page at kidney.org.au.
RESOURCES

- Kidney Health Australia fact sheets (translated versions available): Kidney Transplants, Treatment Options
- Kidney Health Australia booklets: An Introduction to Haemodialysis, An Introduction to Peritoneal Dialysis; An Introduction to Home Dialysis, An Introduction to Supportive Care of Advanced Kidney Disease, Kidney Failure Treatment Options (translated versions available)
- Kidney Health Australia book: Living with Kidney Failure
- [www.kidney.org.au](http://www.kidney.org.au) > your-kidneys > support
- [www.kidney.org.au](http://www.kidney.org.au) > your-kidneys > support > kidney-disease > treatment > choosing-your-treatment
- Kidney Health Information Service (KHIS) 1800 454 363
What if I am struggling to cope?

It is normal to feel anger, despair, fear and other negative emotions during the journey of kidney disease diagnosis and treatment. These feelings usually improve with time. If the impact of kidney disease treatments and the changes to your lifestyle are overwhelming you, then it is important that you talk to someone.

The nurses, social worker or doctor at your renal unit are the best place to start. Social workers are employed in renal units to provide professional counselling to patients and their family members. They are very familiar with the issues you are experiencing. There is no need to feel embarrassed or ashamed or to suffer in silence.

You can also see your GP who can arrange a referral to a psychologist or social worker in your community. These services can be accessed through Medicare.

RESOURCES

Organisations such as beyondblue (beyondblue.org.au or call 1300 22 4636) provide a confidential telephone information and advice helpline

If you find yourself feeling low, please ask for help.
Where can I get more information?

This booklet contains information about kidney transplantation. Other booklets in this series cover home dialysis, haemodialysis, peritoneal dialysis, and supportive care.

We encourage you to read all the booklets so you can make an informed decision about your treatment.
RESOURCES

Living with Kidney Failure booklet 8th edition, Kidney Health Australia
Community newsletter

www.kidney.org.au

Connect with us on Facebook, Instagram and Twitter

Social groups – Kidney Club

Kidney Health Information Service 1800 454 363
TelEconnect 1800 454 363
ABO incompatible transplant
A type of transplant where your donor’s blood type and your blood type are not compatible. With this type of transplant you receive medical treatment before and after your kidney transplant to lower antibody levels in your blood and reduce the risk of antibodies rejecting the donor kidney.

Acute kidney injury:
A loss of kidney function that happens quickly which may or may not be permanent.

Acute rejection
Where your body quickly begins to reject your transplanted kidney in the first few weeks after your transplant.

Altruistic donation
Where someone anonymously donates a kidney to a recipient on the transplant waiting list.

Anti-rejection medications
Medication to reduce the risk of your body rejecting your new kidney.

Artery
The large blood vessel that takes blood from the heart to other parts of the body.

Artificial ventilation
Where a special machine called a ventilator is used to allow you to breathe.
**B**

**Blood group**
Refers to the red blood cell groups (O, B, A or AB) that determine the compatibility for blood transfusion and kidney transplantation.

**Biopsy**
A small piece of tissue is removed for testing and examination under a microscope.

**C**

**Catheter**
A plastic tube that is used to take fluid in or out of your body.

**Chronic kidney disease**
A term used to describe kidney damage or reduced kidney function that lasts for more than 3 months.

**Chronic rejection**
Where your body gradually rejects your transplanted kidney in the months and years after your transplant.

**Cold ischaemia time**
The time between when the kidney is removed from your donor and transplanted into your body.

**Compatibility**
An assessment of whether your body will accept the transplanted kidney.

**Creatinine**
Waste that is produced by your muscles. It is usually removed from your blood by your kidneys and passes out in your urine. When your kidneys aren’t working very well, the creatinine stays in your blood.
**Cytotoxic antibodies**
Antibodies that your body develops to prevent infections.

**D**

**Diabetes**
A chronic disease caused by problems with the production and/or action of insulin in the body which helps control blood sugar levels.

**E**

**eGFR**
Where a mathematical formula is used to estimate your glomerular filtration rate. (See Glomerular Filtration Rate)

**G**

**Glomerular filtration rate (GFR)**
The best measure of kidney function and helps to determine your stage of kidney disease. It shows how well your kidneys are cleaning your blood. Your GFR is usually worked out from your creatinine blood test. (See eGFR)

**H**

**Haemodialysis**
A treatment for kidney failure. Your blood is pumped through special tubing to a haemodialysis machine. The machine acts like a kidney, filtering waste products from your blood before returning it to your body.
Health care team
Living with kidney disease means you will have contact with a great many health professionals. The members of your health care team will change depending on your circumstances. Your health care team may include these health professionals: nephrologist, surgeon, nurse, transplant coordinator, GP, social worker, psychologist, dietitian, physiotherapist, pharmacist. Your health care team are here to help you – ask questions and keep them up to date with how you are feeling.

HLA (Human Leukocyte Antigen) proteins
Special proteins on your cells that are unique to you, and are responsible for controlling your immune system.

Hyperacute rejection
Rejection that occur minutes or hours after your transplant. This is very rare.

K
Keyhole surgery
Surgery that uses very small incisions and special instruments for the surgeon to see inside your body.

L
Laparoscopic nephrectomy
Where a kidney is removed from a donor using a number of small incisions rather than one large incision (compare open nephrectomy).
N

Nephrologist
A doctor who specialises in kidney function.

Non-directed kidney donation
Where someone anonymously donates a kidney to a recipient on the transplant waiting list.

Nephrectomy
Surgical removal of one or both kidneys.

O

Open nephrectomy
Where a kidney is removed from a donor using one large incision (compare laparoscopic nephrectomy).

Organ Donor Register
The Australian Organ Donor Register is the only national register for people to record their decision about becoming an organ and tissue donor for transplantation after death. Joining the register is voluntary and you can elect which organs and tissues you are willing to donate. For more information visit: www.donatelife.gov.au/decide

P

Peritoneal dialysis
Treatment for kidney failure during which dialysis fluid is moved in and out of your peritoneal cavity to remove wastes and fluid from your blood.

Plasma exchange
A treatment for kidney rejection that removes antibodies in your blood.
Polycystic kidney disease
An inherited kidney disease that produces fluid-filled cysts in your kidneys.

Pre-emptive transplant
When someone receives a kidney transplant before starting dialysis.

R
Rejection
Where your immune system attacks your transplanted kidney.

T
Terminal
An illness that is expected to lead to death.

U
Urine
The name for excess fluid and waste products that are removed from your body by your kidneys. Commonly called wee.

V
Vein
A blood vessel that returns blood to your heart.
Connect with us
www.kidney.org.au
Freecall 1800 454 363