Chemotherapy
Your questions answered
Foreword

This is one of a series of booklets written to provide information for patients and their relatives. It is impossible to include everything you may need to know. However, you need to feel you have been given enough information before you can make a decision about treatment. Throughout the booklet we suggest questions you may wish to ask your doctor about treatment. Your doctor or nurse will be able to answer specific questions.

This booklet has been prepared by The Royal Marsden specialist nurses, with input from doctors and patients.

Throughout the booklet, definitions of words written in **bold** can be found in the glossary at the back.

We hope you find it helpful and would welcome your comments so that the next edition can be improved further.
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What is cancer?

Our bodies are made up of tiny structures called cells, which are not visible to the naked eye. These cells are ‘building blocks’ and groups of cells form the tissues and organs of the body. The cells in these different tissues, such as the brain, liver, kidney and lungs, each have a unique function.

Normally cells reproduce themselves by dividing in a regular, controlled fashion so that growth and repair of the body tissues can take place.

Sometimes this control goes wrong and the cells divide in an uncontrolled way and are abnormal. This abnormal and excessive growth of tissue cells is called a tumour and can be benign or malignant.

Benign tumours are usually limited to a small area and are often slow growing. They only cause problems because of their size or where they happen to be situated. Once they’ve been treated, often by an operation, they don’t usually cause any further problems.

Cancers or malignant tumours also start within a limited area but they can spread to nearby organs or tissues. Cancer cells may be carried in the bloodstream to distant sites in the body. Here they may form new tumours called metastases or ‘secondaries’.

Cancer cells may also be carried in the lymphatic system which normally helps the body to fight infection. It is made up of a series of nodes (glands) and vessels (tubes) linked throughout the body.

Cancer is a word used to describe many different diseases which have different causes and which are treated in different ways. All cancers can be treated although not all can be cured.
**Who will plan my treatment?**

A multidisciplinary team will plan your treatment for you.

**The multidisciplinary team may include:**

- A surgeon who has a special interest in your type of cancer (if appropriate)
- A **medical oncologist** (a specialist doctor who treats cancer with drugs, including chemotherapy)
- A **clinical oncologist** or radiotherapist (a specialist doctor who treats cancer with radiotherapy and may also prescribe chemotherapy)
- A **haematologist** (a specialist doctor who diagnoses and treats blood disorders. These include cancers such as leukaemias [affecting the bone marrow], lymphomas [affecting lymphatic tissue] and myelomas [affecting the plasma cells in the bone marrow])
- A **pathologist** (a specialist doctor who examines body tissues and organs under the microscope)
- A **radiologist** with a special interest in your type of cancer (who specialises in the use of x-rays and other imaging to diagnose and treat disease)
- A **specialist nurse** who provides information and support.
- A **keyworker** who may be a specialist nurse or other healthcare professional involved in your care (acts as a point of contact for you and your carers throughout your treatment).

Other health care specialists (for example, pharmacist, dietician, and physiotherapist) may also be represented on this team and involved in your care.

It depends on the type of cancer you have as to which of the above healthcare professionals will be part of your multidisciplinary team. For example, not all cancers can be treated with surgery.
The multidisciplinary team meets regularly to discuss the best treatment options for each patient in their care. They will take into account the results of tests and your general health. Your doctor will then discuss the different treatment options available to you, taking into account your own preferences. You may find it helpful to take a family member or friend with you when your treatment is being planned.

Research

Research is an important part of developing better treatment for cancer. Many specialist centres take part in clinical trials or research studies. You may be asked if you would like to take part in a clinical trial, for example, looking at a new or improved way of using chemotherapy or other treatments. These trials may be assessing the effectiveness of a new treatment compared to standard treatment. If the new treatment produces better results or offers few side effects it may then be widely used for all people with a particular cancer.

The doctors and nurses will discuss any clinical trial in detail with you, if it is an appropriate treatment option in your case. Another Royal Marsden booklet, Clinical trials, provides more detailed information about clinical trials and what is involved.

Several national cancer information organisations can provide information on clinical trials currently in progress, or else provide website links to online clinical trial databases. You can find further details about these organisations on page 50–51.
Questions you may wish to ask your doctor about research studies or clinical trials

To help you think about what you want to ask your doctor, you may find the following questions helpful.

- What is the purpose of this study?
- Why have I been chosen?
- Do I have to take part?
- What will happen to me if I take part?
- What do I have to do?
- What is the drug or procedure that is being tested?
- How long will the study last?
- How could the trial help me?
- What are the side effects of taking part?
- What are the benefits and risk?
- Will I need any extra tests or investigations?
- What other treatments are available if I don’t take part in this trial?
- Who can I contact if I have any concerns or problems?
- What if new information becomes available?
- What happens when the research study stops?
- What if something goes wrong?
- Will my taking part in this study be kept confidential?
- What will happen to the results of the research study?
- Who is organising and funding the research?
**Consenting for treatment**

Before you can give your consent, your doctor will discuss with you what the treatment is likely to involve, the benefits and risks, and any available alternative treatments. You may also be given some written information about the treatment protocol that has been planned for you. It is important that you understand the information you have been given – ask questions if you don’t understand or if you want more information. To help you think about what you want to ask your doctor, you may find questions on page 8 and 9 helpful. You will then be asked to sign a consent form, which is a written record that you have agreed to the planned treatment and understand the benefits and risks involved. The main benefits and risks associated with the treatment will be written on the **consent form** before you sign it. You will then be given a copy of this.

**How long will I have to make up my mind about treatment?**

Ask your doctor how long you can take to make up your mind. The answer will depend on the type of cancer you have, as some cancers grow much faster than others. While it may be important to start treatment as soon as possible, many people will have plenty of time to make a decision.

Your treatment plan may involve one or more treatments, for example, surgery and / or chemotherapy and / or radiotherapy. You will be asked to sign consent forms for each treatment. Before each treatments starts, you will be asked to confirm that you agree to treatment.

It is important to remember that once you have made a decision about treatment, you can change your mind at any time, even after you have signed a consent form.
What is chemotherapy?

Chemotherapy means treatment with anti-cancer drugs and they are given to destroy or control cancer cells. There are over 50 different drugs. Drugs may be given on their own or several different drugs may be given together. This is called combination chemotherapy.

Why is chemotherapy given?

Many types of cancer can be treated with chemotherapy. The aim of the treatment will depend on the type of cancer you have and how advanced it is. Chemotherapy is given:

• To cure the cancer – chemotherapy is given to destroy all the cancer cells.

• To reduce the possibility of cancer coming back – chemotherapy may also be given to destroy any cancer cells that may be present in your body that are too small to detect. It is important to destroy these cells.

• To control the cancer – when the chemotherapy is unlikely to cure the cancer but may prevent it from growing for some time.

• To relieve symptoms – chemotherapy may be given to shrink a tumour if it is causing any symptoms.

How does chemotherapy work?

The drugs enter your bloodstream and therefore reach all parts of your body. This is called systemic treatment. Most radiotherapy and surgery are called local treatments. Chemotherapy drugs destroy cancer cells by damaging them so they can’t divide and grow.

The drugs can also affect normal cells which are growing and dividing quickly. Damage to normal cells may cause side effects (see page 23). These are usually temporary because healthy cells quickly grow back to normal. Permanent damage is rare with most chemotherapy regimens.
When is chemotherapy given?

Sometimes chemotherapy is used on its own in cancers that respond well to this treatment. However, chemotherapy is often used with other treatments.

- **Neo-adjuvant therapy** – chemotherapy may be given before surgery or radiotherapy to shrink the tumour.

- **Adjuvant therapy** – chemotherapy can also be used to help destroy any cancer cells that may remain after surgery or radiotherapy. The aim is to reduce the likelihood of your cancer returning in the future.

- **Peri-operative therapy** – chemotherapy may be given both before and after surgery.

- **Chemoradiation** – chemotherapy may be combined with radiotherapy.

- **Palliative chemotherapy** – if the cancer has spread to other parts of the body, the chemotherapy drugs carried in your bloodstream can reach these cancer cells. The aim is to help relieve symptoms and slow the growth of the cancer.

Questions you may wish to ask your doctor about treatment

Before you can make a decision about treatment, you need to feel you have been given enough information. All treatments carry risks as well as benefits and thinking through the possible options can help you to decide. You should be given a chance to ask questions, however it is not always easy to know what to ask.
To help you think about what you want to ask your doctor, you may find the following questions helpful:

- What are the main treatment options?
- What are the benefits of each of the options?
- What are the risks, if any, of each option?
- What are the risks if I decide to do nothing for the time being?
- How long can I take to decide about what treatment I have?
- Will this treatment cure me?
- When will I know if the treatment has worked?
- What will happen if this treatment doesn’t work?
**How will my chemotherapy treatment be planned?**

The chemotherapy treatment your doctor recommends will depend on several factors. These include:

- The type of cancer you have
- Where the cancer is in your body
- Where in the body it has spread to (if it has)
- Your general health

You will be offered the best standard treatment available that current research shows will be most helpful in your situation. This is why you may meet other people with the same cancer as you who are having different chemotherapy treatments.

**Questions you may wish to ask your doctor about chemotherapy**

It is important that you understand what will happen and why. You should be given a chance to ask questions. To help you think about what you want to ask your doctor, you may find the following questions helpful.

- What drug or drugs will I be given?
- How will the drugs be given?
- Will I have to spend any time in hospital?
- How often will I need treatment?
- How long will my treatment go on for?
- Can I still take other medicines (including any alternative therapies) during my chemotherapy treatment?
- How will I feel during treatment and are there any side effects I can expect?
- If there are any side effects, what can be done to help me cope with them?
• How long will it take for me to recover after I have finished the treatment?
• Are there any long-term side effects?
• Will I be able to continue with the same lifestyle that I’m leading now?
• Can I talk to someone who has had the same treatment?
• Who should I contact if I am worried about my diagnosis, treatment or prognosis?

Your doctor or nurse will explain your personal drug treatment to you. If you want more details or have any questions, please ask the staff caring for you.
How often will I have chemotherapy?

A course of chemotherapy is given according to a protocol (plan) and how often you have chemotherapy will depend on this. Each course usually consists of several cycles of chemotherapy with a rest period in between each cycle. The rest periods are often longer than the treatment periods so most of the time you’re not actually having chemotherapy. The rest periods allow your body to recover from any unwanted effects of the drug. As you have more cycles of treatment it is occasionally necessary to extend these rest periods. Don’t worry, this isn’t unusual.

If you’re having oral chemotherapy (tablets or capsules), you may take smaller doses daily for several weeks or months before having a rest period.

<table>
<thead>
<tr>
<th>Example 1</th>
<th>A course of chemotherapy = six cycles</th>
</tr>
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<tbody>
<tr>
<td>Cycle 1</td>
<td>Day 1  \text{Chemotherapy}</td>
</tr>
<tr>
<td></td>
<td>Days 2–21  \text{Rest period}</td>
</tr>
<tr>
<td>Cycle 2</td>
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</tr>
<tr>
<td>Cycle 3</td>
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</tr>
<tr>
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<td>Days 2–21  \text{Rest period}</td>
</tr>
<tr>
<td>Cycle 4</td>
<td>Day 1  \text{Chemotherapy}</td>
</tr>
<tr>
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<td>Day 1  \text{Chemotherapy}</td>
</tr>
<tr>
<td></td>
<td>Days 2–21  \text{Rest period}</td>
</tr>
</tbody>
</table>

= total of 18 weeks
### Example 2
**A course of chemotherapy = six cycles**

<table>
<thead>
<tr>
<th>Cycle</th>
<th>Day 1</th>
<th>Days 2–7</th>
<th>Day 8</th>
<th>Days 9–21</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cycle 1</td>
<td>Chemotherapy</td>
<td>Rest period</td>
<td>Chemotherapy</td>
<td>Rest period</td>
</tr>
<tr>
<td>Cycle 2</td>
<td>Day 1</td>
<td>Days 2–7</td>
<td>Day 8</td>
<td>Days 9–21</td>
</tr>
<tr>
<td>Cycle 3</td>
<td>Chemotherapy</td>
<td>Rest period</td>
<td>Chemotherapy</td>
<td>Rest period</td>
</tr>
<tr>
<td>Cycle 4</td>
<td>Day 1</td>
<td>Days 2–7</td>
<td>Day 8</td>
<td>Days 9–21</td>
</tr>
<tr>
<td>Cycle 5</td>
<td>Chemotherapy</td>
<td>Rest period</td>
<td>Chemotherapy</td>
<td>Rest period</td>
</tr>
<tr>
<td>Cycle 6</td>
<td>Day 1</td>
<td>Days 2–7</td>
<td>Day 8</td>
<td>Days 9–21</td>
</tr>
<tr>
<td></td>
<td>Chemotherapy</td>
<td>Rest period</td>
<td>Chemotherapy</td>
<td>Rest period</td>
</tr>
</tbody>
</table>

= total of 18 weeks

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**How long will my treatment last?**

The length of your treatment will depend on how well your disease responds to the drugs, for example, when your cancer starts to shrink. Your doctors may choose to give you treatment for several weeks and then check on your progress. Overall, it may take several months to complete your course of chemotherapy.

When adjuvant chemotherapy is given, the doctor may plan your treatment for a certain length of time known to be effective for people with similar tumours, for example, six months.

**How will I know if my chemotherapy is working?**

Your doctor will monitor your progress throughout treatment. There are several tests which may be repeated during your treatment. They may include scans, x-rays and blood tests. All the tests check on your health and the way chemotherapy is affecting you. They will vary from person to person so if you’re not sure why you’re having a particular test, ask your doctor or nurse.
Whether or not you have side effects from chemotherapy is not an indication of how well the treatment is working.

**Why has my chemotherapy plan been changed?**

Sometimes, as a result of the tests you have during treatment, your chemotherapy plan may be changed. For example, if the blood test shows that your ‘blood count’ (see page 25) is low, your doctor may decide to give you a longer rest period between cycles. Your doctor may reduce the dose of the drugs or you may be given medicine to ‘boost’ your blood count.

If the test results show that the chemotherapy treatment isn’t working well enough, then your doctor may change your treatment plan. Sometimes, different drugs may need to be tried to find out which ones are best for you.
How is chemotherapy given?

How chemotherapy is given will depend on the type of cancer you have and the drugs used.

Chemotherapy is most commonly given in one of two ways:

• By mouth (orally)
• By injection into a vein using a syringe or in an infusion (‘drip’)

Other – occasionally, a drug may be injected into a muscle or under the skin. Sometimes it can be injected into your spine, lungs or bladder if that is where treatment is needed.

You may have one drug, several drugs or different drugs given at different times. You may have tablets or injections or both.
Oral chemotherapy

Some chemotherapy drugs can be taken as tablets or capsules, which means you can continue your treatment at home. You will still need to visit the outpatient department for regular check-ups.

When should I take my tablets?

All bottles of tablets given out by the pharmacy department carry instructions about how and when to take them. You will also be told whether or not you have to take them with food and how to store them safely. Please follow these instructions carefully and if they’re not clear, ask your doctor, nurse or pharmacist. You may be given an extra information sheet with some medicines.

You will probably be provided with only enough tablets for one cycle of therapy. This is so your doctor can assess your progress before you start your second cycle. Sometimes, it may be necessary to change the dose of the tablets.

Your chemotherapy tablets have been prescribed specifically for you and you should not give them to anyone else. If you have chemotherapy tablets left when you have finished treatment, take them back to the hospital pharmacy. If you think you are running out of chemotherapy tablets and should have more, then contact your hospital doctor (not your GP).

What should I do if I am sick after taking my chemotherapy?

It is very important that you take your tablets regularly on the days specified. If you miss a dose for any reason, contact the ward for advice. Do not take a double dose.

If you are sick immediately and can see the tablet, then repeat the dose one hour later. If you are sick later (even five minutes later), contact the ward for advice.

If, however, you are using liquid or opened capsules, or you were asked to crush or dissolve tablets, the dose should not be repeated.
How is chemotherapy given into a vein?

When chemotherapy is given intravenously (into a vein), it is given through a venous access device.

Types of venous access devices

There are several different types of devices available. These include:

**Cannula**

This is small tube which is put into a vein in your lower arm or the back of your hand. It is inserted at the beginning of each treatment and is removed after the chemotherapy has been given. Once the cannula has been taped to your arm, the chemotherapy drugs can be injected slowly into the vein through it. Sometimes the drugs are diluted and given through a ‘drip’ (intravenous infusion) into a vein in your arm.

Some treatment drugs cannot be given into the vein of the hand or the arm and can only be given through one of the central venous access devices described below.

**Skin-tunnelled catheter**

Sometimes it may be known by its brand name, for example a Hickman line. It is a fine tube which is inserted through your chest into a large vein leading to your heart. Because it is made of a non-irritant material, for example silicone, it can be left in place for several weeks or months. As well as using it to give you fluids and drugs, it may also be used to take blood samples.

**Peripherally Inserted Central Catheter (PICC)**

A PICC is inserted into a vein in your arm, above the bend in the elbow. It goes into the large vein leading to your heart. A PICC can be used to give you drugs and to take blood samples.

**Implanted port**

An implanted port is a device, which is inserted into your body under the skin. The usual position is on the chest. The port is made up of a portal body (or chamber) and this is connected via a
thin tube inserted into one of the body’s veins. A special needle is used to get into the chamber which can then be attached to a drip for injections, drips or blood tests.

You may be able to choose the type of device you have, however, your choice will depend on the type of therapy you are going to have, your physical condition and your life style. Sometimes a choice isn’t possible.

More information about skin-tunneled catheters, PICCs and implanted ports can be found in another Royal Marsden booklet, *Central venous access devices*.

**Will having chemotherapy hurt?**

Chemotherapy is generally no more painful than any other injection or blood test. However, occasionally veins become hardened or sore. Please tell your chemotherapy nurse or doctor if this happens, so that they can avoid the sore area.

Some drugs can cause damage if they leak out of the vein while they are being injected. If you feel *any stinging, ‘burning’ or pain* when you are receiving the drug, tell your nurse or doctor straight away. If your arm or hand, where the cannula was sited or the area around the catheter or port, is sore, painful or swollen following the injection, please contact your chemotherapy nurse or doctor. If this continues or we have difficulties with your veins, they may suggest you have a central venous access device inserted.

**How long does it take to give intravenous chemotherapy?**

Depending on the chemotherapy drug or combination of drugs, each cycle lasts from a few hours to a few days.

Your doctor may recommend that your chemotherapy is given continually at a lower dose, over weeks or months. This keeps a steady level of the drug in your bloodstream. You will receive the drug from a small infusion pump while you are at home. This is called *ambulatory chemotherapy*. 
What is intrathecal chemotherapy?

For some types of cancer, for example lymphomas and leukaemias, cancer cells can pass into the cerebrospinal fluid (CSF) which surrounds the brain and spinal cord. Your doctor may prescribe intrathecal chemotherapy. This means that anti-cancer drugs will be given through a needle into the space around the spinal cord in your back. This procedure is called a lumbar puncture. Injecting chemotherapy into the spinal fluid can destroy any cancer cells that may be there and prevent cancer cells growing.
Where will I have my chemotherapy treatment?

Most patients receive their chemotherapy as outpatients or day patients, visiting the hospital on the day of treatment. Your length of stay in hospital will depend on the types of tests, assessment, chemotherapy preparation and treatment you need. If you are a day patient, you are likely to be at the hospital between four and six hours, sometimes longer if the treatment is complex, or if you are taking part in a research study.

Before you are given your chemotherapy, you will usually have blood tests and see the doctor. As you may have to wait while your chemotherapy drugs are being prepared, you may want to bring someone with you. Or you could bring a book to read or something to occupy you while you wait.

A step-by-step guide to having chemotherapy

Your tests and assessment

When you arrive on the day unit or ward, you will:

• Have a blood test – to check your blood count is satisfactory before you receive your chemotherapy.

• See the doctor or nurse
  ○ to check you are well enough to be given chemotherapy
  ○ to check your chemotherapy prescription.

For some people this may happen the day before treatment is given.

Chemotherapy preparation and treatment

• Your chemotherapy will be prepared by pharmacy (if you have already had your blood tests and assessment your chemotherapy may have been prepared in advance)

• Your chemotherapy treatment will be given

• You will be given drugs to take home (if appropriate)

• You will be given your next appointment.
Sometimes you might need to stay in hospital overnight when treatment is first started or if several drugs and fluid infusions are to be given. In these circumstances, a 24–48 hour hospital stay is usually all that is needed.

There are special treatments which require longer admissions and, if necessary, they will be explained to you.

If your chemotherapy is given by continuous infusion at home, you will be asked to attend hospital for regular check-ups every few weeks.

**Can I still take other medicines while I am having chemotherapy?**

Tell your doctor about any other medicines you are taking or planning to take, including herbal medicines, vitamins, other dietary supplements and complementary therapies. Some drugs may interfere with your treatment.

If you are admitted to hospital, please bring all your current medicines with you. Show them to the doctor or ward nurse so they know what you are taking.

**Please ask your hospital doctor before taking any new medicines.**

**Will I be able to have vaccinations?**

During chemotherapy you will not be able to have live virus vaccines. You may be able to have flu vaccines but it is always important to consult your hospital doctors before having any vaccinations.
Will chemotherapy affect my everyday activities?

Many people continue with their usual activities while having chemotherapy, for example working between injections or cycles of tablets. However, most people do find that they are more tired than usual for the first few days after treatment. You may need to take life more slowly, working part time or cutting down on social activities. Take care not to overtire yourself, get enough rest and accept offers of help with everyday tasks.

The drugs, among other things, may cause you to feel a little ‘down’. It’s quite normal for this to happen at various times during your treatment. If you’d like to chat about this, contact your nurse or doctor or talk it over at your next appointment.

What if I have planned a holiday?

It may be possible to adapt your chemotherapy protocol to fit in with holidays or other special occasions. Please discuss this with your doctor in advance so your treatment can be planned around your arrangements.

It is important that you inform your hospital team of any travel plans while you are on chemotherapy treatment. There are a number of factors you will need to consider if you are thinking of travelling outside the UK including whether you can have some of the vaccinations you may need to visit certain countries.

There are risks associated with travelling outside the UK. If you need to use healthcare services in another country, the financial cost can quickly build up. Comprehensive travel insurance is strongly recommended. However, this is not always possible to arrange for pre-existing conditions and during treatment. It is also more difficult to arrange travel insurance to countries where healthcare is more expensive, for example the USA or Canada. You may need to consider if you are prepared to take the risk of travelling without insurance.
Airline travel is also associated with an increased risk of blood clots (see page 40). Longer flights present the greatest risk although blood clots can also form after long train or car rides. Some cancers and chemotherapy drugs increase this risk; discuss this with your doctor.

You must take care of your skin when you are exposed to the sun. Some chemotherapy drugs can make your skin more sensitive and you may burn easily (see page 33). Check with your doctor or specialist nurse if it is safe for you to swim.

Finally, many of the risks of chemotherapy including an infection developing while you have a low white blood cell count, may be increased if you do not have good access to medical care.

**Can I have a drink?**

Yes, it will usually be all right for you to have a small amount of alcohol, if you feel like it. There are one or two anti-cancer drugs which may interact with alcohol but you will be told about these.

Some people find that alcohol tastes unpleasant during chemotherapy treatment. Avoid drinking alcohol if you feel nauseous or if you have a sore mouth, it may make you feel worse.
Are there any side effects or complications?

Yes, but these will depend on the type of chemotherapy you are given. Side effects are the secondary effects of treatment and can be acute or late. Acute (immediate) side effects occur during and immediately after treatment. Late (delayed) side effects develop after treatment has been going on for some time and may continue, at least for a while, after treatment is finished.

Chemotherapy drugs damage fast growing cells. As well as destroying cancer cells, they also cause damage to normal cells. It is this damage to normal cells that may cause side effects. Everyone reacts differently to chemotherapy and some people may have no side effects at all. The side effects you may experience with your chemotherapy protocol will be discussed with you. For example, not all chemotherapy drugs cause sickness or hair loss, so do check what is relevant for you.

We can offer help for most side effects, so please tell your doctor or nurse if you feel any different from normal. We want to maintain your confidence and well being as much as possible during treatment. Remember, most of the side effects of chemotherapy are temporary and will disappear after your treatment has finished.

There are some side effects that need to be treated quickly and it is important that you do not wait until the next morning or after the weekend. If you are unsure as to whether your symptoms need urgent treatment, contact the hospital or your GP for telephone advice.
You **must** contact your hospital team or GP **immediately** if you develop any of the following symptoms:

- a temperature of 38°C/100°F or higher
- shivering episodes
- flu-like symptoms
- gum / nose bleeds or unusual bleeding (if bleeding doesn’t stop after 10 minutes of pressure)
- mouth ulcers that stop you eating or drinking
- vomiting (that continues in spite of taking anti-sickness medication)
- diarrhoea (four or more bowel movements more than usual or diarrhoea at night)
- difficulty with breathing.

It is important that you tell your hospital doctor if you suffer from any side effects or anything else unpleasant that may have happened to you since your last visit. Your doctor can help you by giving you medication to reduce or stop you from experiencing these side effects in the future.

You may be given a card similar to the one below to carry around with you. The card lists the symptoms needing urgent treatment and tells you and a healthcare professional what to do.

Some more common effects which occur with several drugs are discussed here.
Your blood

Blood cells are made in the bone marrow, the spongy tissue found inside the hollow bones of hips, legs and arms. Your bone marrow makes red blood cells, white blood cells and platelets.

Chemotherapy temporarily reduces the rate at which blood cells are produced in your bone marrow. A blood test will be taken at the beginning of therapy and before each course of treatment. This is to make sure that your ‘blood count’ is satisfactory before you have your chemotherapy. Your doctor may also request a ‘blood count’ between treatments. The main effects of chemotherapy on your blood cells are:

- infection
- anaemia
- bruising or bleeding.

Infection

There are many types of white blood cells which make up the total white cell count. Their main function is to help your body to fight infection. If your white cell count is low (usually 7–14 days after the chemotherapy), you may become prone to infections and take longer to recover from them. In general, these arise from bacteria within our own bodies, and it is not necessary to avoid crowded places or isolate yourself from others. However, we advise you to keep away from people with serious infections, for example, chicken pox.

It is important to keep good personal hygiene. This includes taking daily baths or showers and washing clothes and bed linen regularly. Looking after your mouth (see page 29) will help prevent an infection from developing. Take care to wash your hands well when preparing food, before meals and after using the toilet.

During your treatment try to reduce the risk of developing an infection, for example take care not to graze or cut your skin when gardening, shaving or preparing food. If you do, clean the area with warm water and soap and cover it with a sterile dressing, for example an Elastoplast.
Tell your doctor or nurse if you develop a rash or diarrhoea, as both of these symptoms can be a sign of infection.

Your doctor may prescribe a course of antibiotics to prevent infection during your treatment. Another option is to give you a course of injections of growth factors, which are proteins that stimulate the production of blood cells. The commonly given growth factor to boost white cells is G-CSF.

The signs of infection (neutropenic sepsis) may be a **high temperature, shivering or flu-like symptoms or other signs of infection, such as a sore throat or cough**. If you feel unwell at any time or have a temperature (38°C/100°F or higher) you should contact the hospital immediately as you may need to be admitted for intravenous antibiotics.

<table>
<thead>
<tr>
<th>Possible side effects</th>
<th>Symptoms include:</th>
<th>What you should do</th>
</tr>
</thead>
</table>
| **Infection** due to low white blood cells – Also called neutropenic sepsis | • High temperature  
• Shivering or flu-like symptoms  
• Other signs of infection, such as a sore throat or cough  
• Rash  
• Diarrhoea | • Check your temperature if you feel unwell  
• Avoid grazing or cutting your skin.  
**Contact the hospital immediately** – refer to your alert card for contact details if you feel unwell at any time or have a temperature (38°C/100°F or higher) |

If you are on an intensive chemotherapy regimen, you may be given additional advice about diet. You may be advised to avoid certain foods that have been linked with food poisoning in the past, for example, soft cheese and under-cooked eggs.

**Anaemia**

The red blood cells contain a protein called haemoglobin (Hb), which carries oxygen around the body. If your haemoglobin is
low, you may become anaemic and begin to feel very tired and you may look pale. Anaemia may lead to shortness of breath when you exert yourself more than usual, such as when climbing stairs or doing housework.

If during your treatment you begin to feel more tired than normal or become breathless, tell your doctor. Eating a diet rich in iron, for example liver, red meat, fish, eggs and green leafy vegetables, may help to keep your haemoglobin up to its usual level.

If your haemoglobin is very low, you may need to be admitted for the day, or overnight, to receive a blood transfusion. Another way of treating anaemia is to stimulate the body to produce more red blood cells. You may be given erythropoietin or EPO, which is a naturally occurring growth factor, produced by the kidneys. It stimulates the bone marrow to produce red blood cells.

<table>
<thead>
<tr>
<th>Possible side effects</th>
<th>Symptoms include:</th>
<th>What you should do</th>
</tr>
</thead>
</table>
| Anaemia (low red blood cells / haemoglobin) | • Tiredness  
• Shortness of breath | • Eat a diet rich in iron.  
**Contact your hospital urgently (see page 55 for contact details) if you feel very unwell** |

**Bruising or bleeding**

Platelets help your blood to clot and prevent you from bleeding and bruising. Most types of chemotherapy do not normally affect the platelets in any significant way. However, with some treatment you may notice that you bruise easily or that you have a tendency to bleed from your nose or gums. Rarely, small groups of red-purple spots may appear on your skin. **All these could be signs of a low platelet count and you should contact your hospital doctor urgently.**
Don’t take drugs which could affect your platelets, such as aspirin. Ask your doctor if you’re not sure what to avoid. Use a soft toothbrush and an electric razor to prevent damage to your gums and skin. Take care not to cut yourself when using knives and wear thick gloves when gardening. Also take extra care if you are playing sports.

You may need to come into hospital for a platelet transfusion (which is like a blood transfusion but all the red cells and white cells have been removed) your platelet count can also be corrected by making the rest period between your courses of drugs longer, or by adjusting the doses of the drugs.

<table>
<thead>
<tr>
<th>Possible side effects</th>
<th>Symptoms include:</th>
<th>What you should do</th>
</tr>
</thead>
</table>
| **Bruising or bleeding** (due to low platelet count) | • Gum / nose bleed  
• Rarely small groups of red-purple spots on your skin | • If you are taking drugs such as aspirin or clopidogrel, ask your doctor if you should continue  
• Use a soft toothbrush and an electric razor to prevent damage to your gums and skin  
• Take care not to cut yourself  
Contact your hospital urgently (see page 55 for contact details) if you have bruising or bleeding or groups of red-purple spots appear on your skin |

**When symptoms may occur**

During the course of chemotherapy
Your digestive system

Some chemotherapy drugs can affect the lining of the digestive system and may cause the following problems:

- sore mouth
- taste changes
- nausea and vomiting
- changes in appetite
- diarrhoea or constipation.

Sore mouth

During chemotherapy the lining of your mouth may become sore and prone to infection. This is usually happens around 7–10 days after chemotherapy. It’s important that you keep your mouth clean and healthy and the following suggestions may help:

- Drink plenty of fluids, to keep your mouth moist and fresh.
- Use a mouthwash regularly. Ask your doctor, nurse or oral hygienist which one is best.
- Clean your teeth regularly using fluoride toothpaste and a soft toothbrush. If someone needs to clean your teeth for you, they may find a child’s toothbrush easier to use.
- Clean your dentures after meals as well as at night.
- Keep your lips moist using a lip salve available from a chemist.

Tell your doctor or nurse if your mouth becomes sore. It may be possible to prescribe some tablets to prevent this in future. If you need further advice, ask to see an oral hygienist. Depending on the cause of your sore mouth it may be possible to provide you with medication to relieve the discomfort.

During chemotherapy, you may be more prone to infection and bleeding. Always consult your doctor before having any dental work done. Your blood count may need to be checked before the dentist can decide whether or not to proceed with dental treatment.
Taste and smell changes

Some people experience a change in their sense of taste or smell. You may find that food may taste more salty, bitter or metallic. This is only temporary. Normal taste and smell usually returns two or three months after the end of treatment. If an unpleasant taste occurs during an injection, it can be disguised by a strong-flavoured sweet. Occasionally, some people may experience food cravings. All these things which can affect your mouth, and appetite, are covered in more detail in The Royal Marsden booklet *Eating well when you have cancer.*

<table>
<thead>
<tr>
<th>Possible side effects</th>
<th>Symptoms include:</th>
<th>What you should do</th>
</tr>
</thead>
<tbody>
<tr>
<td>Taste changes</td>
<td>Changes in the way food and drink taste</td>
<td>Suck a strong flavoured sweet if you have an unpleasant taste during an injection</td>
</tr>
<tr>
<td>Changes in smell</td>
<td>Increased awareness of smells that may now become unpleasant, for example cooking and perfumes</td>
<td>Avoid areas where unpleasant smells are present</td>
</tr>
</tbody>
</table>

When symptoms may occur

During your course of chemotherapy
Nausea and vomiting

Nausea and vomiting aren’t side effects of every chemotherapy drug although many people believe they are. Some drugs or protocols cause a greater reaction than others. Also everyone is individual – a similar drug combination may cause nausea in one person but not in another.

If you do feel nauseous or even vomit following your treatment, there are many anti-sickness (anti-emetic) drugs to help overcome this problem. These come as tablets, injections or suppositories. You will be given anti-sickness injections before the chemotherapy and tablets to take home with you. You should take these regularly as prescribed, even if you are not feeling sick, as some anti-sickness drugs are better at preventing than stopping sickness. Some people find taking ginger helpful. Should you still have nausea or vomiting, please tell your doctor or nurse and you can be given different anti-sickness drugs that may be more effective. If you experience vomiting at home then anti-sickness tablets may not be effective and you should ask your doctor for suppositories with your next course.

You may also find relaxation or other therapy helpful. Another Royal Marsden booklet, *Coping with nausea and vomiting*, gives advice on managing the effects of nausea and vomiting.

<table>
<thead>
<tr>
<th>Possible side effects</th>
<th>Symptoms include:</th>
<th>What you should do</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nausea and vomiting</td>
<td>• Feeling sick (nausea)</td>
<td>• You should take anti-sickness tablets or suppositories regularly as prescribed</td>
</tr>
<tr>
<td></td>
<td>• Being sick (vomiting)</td>
<td><strong>Contact your hospital</strong> (see page 55 for contact details) if you continue to have nausea or vomiting. You can be given different anti-sickness drugs that may be more effective</td>
</tr>
</tbody>
</table>

When symptoms may occur

7–14 days after chemotherapy
Changes in appetite

If you experience any tastes changes or lose your appetite, you may not feel like drinking or eating. However, you should try to drink plenty of fluids, about 10 glasses or cups each day, and eat well.

Try sipping clear, cold fluids, such as water and soft drinks, slowly through a straw. Fizzy drinks like soda water and ginger ale are quite refreshing, as are lemon or herbal teas. Avoid coffee, which has a strong taste and may also make you more thirsty.

You may need to change your meal times and have small, frequent meals or snacks of whatever you fancy. Eat slowly and chew your food well. After a meal, relax, in a sitting or slightly reclined position, instead of lying down.

Eat light meals on the day of your treatment.

Another Royal Marsden booklet that gives advice on problems associated with eating is Eating well when you have cancer. If you are worried at all about diet, please ask to see the dietitian.

<table>
<thead>
<tr>
<th>Possible side effects</th>
<th>Symptoms include:</th>
<th>What you should do</th>
</tr>
</thead>
<tbody>
<tr>
<td>Changes in appetite</td>
<td>• You may not feel like drinking or eating</td>
<td>• Drink plenty of fluids</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Have small frequent meals or snacks.</td>
</tr>
<tr>
<td>When symptoms may occur</td>
<td></td>
<td>Ask to see the dietitian if you are worried about your diet</td>
</tr>
<tr>
<td>7–14 days after chemotherapy</td>
<td></td>
<td></td>
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</tbody>
</table>

Diarrhoea or constipation

Diarrhoea or constipation may occur with a few chemotherapy drugs. They can often be managed by tablets, medicines or a change of diet. Please tell your doctor or nurse if you have any problems.
Possible side effects | Symptoms include: | What you should do
---|---|---
Diarrhoea or constipation | • Frequent or loose bowel movements
• Not able to control bowel action
• Irregular or hard bowel movement | • Drink plenty of fluids
Contact your GP or hospital (see page 55 for contact details) if you have diarrhoea (four or more bowel movements more than usual or diarrhoea at night)

### When symptoms may occur

7–14 days after chemotherapy

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### Your skin

Some chemotherapy drugs may cause sore, dry hands and feet. This problem can be relieved with medication so please tell your doctor if it happens.

With some drugs, your nails may become darker than usual and they may develop ridges or white lines. These changes usually grow out over a few months after the treatment has finished.

Certain drugs may discolour the skin causing dark lines along the veins or where there is friction, for example due to tight clothing. Others may cause a local red reaction at the injection site or along the vein. This fades a short time after treatment.

Any rash should always be reported to your doctor. Some drugs may cause patches of red skin, particularly on the palms of your hands and soles of your feet. This skin may become dry and cracked. Using a good hand cream may help to prevent this.

During treatment, and for several months afterwards, your skin may be more sensitive to the sun and you may burn more easily. A moisturiser, such as aqueous cream, will help prevent dryness.

Try to stay in the shade between 10am and 3pm, and wear a wide brimmed hat, wear sunglasses that block out 100% of the ultraviolet (UV) rays and a T-shirt or other loose clothing.

Whenever you spend any length of time in the sun apply a sunscreen with an SPF (sun protection factor) of 15 or more that
blocks both UVB and UVA light. Put it on at least half an hour before exposure to sun. Remember vulnerable areas such as your ears, neck, back of the hands, and feet.

After your treatment has finished, you should protect your skin from extremes of temperature and continue using sunscreen (factor 15 or above).

<table>
<thead>
<tr>
<th>Possible side effects</th>
<th>Symptoms include:</th>
<th>What you should do</th>
</tr>
</thead>
<tbody>
<tr>
<td>Skin changes</td>
<td>• Sore, dry hands and feet</td>
<td>• Use a moisturiser, such as aqueous cream on dry skin</td>
</tr>
<tr>
<td>When symptoms may occur</td>
<td>• Nails become darker than usual</td>
<td>• Protect your skin in the sun</td>
</tr>
<tr>
<td>During your course of chemotherapy</td>
<td>• Nails may develop ridges and white lines</td>
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</tr>
</tbody>
</table>

**Your hair**

Hair loss (alopecia) is common side effect of chemotherapy. Not all drugs cause hair loss and some drugs only cause thinning of the hair or cause it to become brittle.

Hair loss can be very distressing, however it is always temporary and your hair will grow again when treatment finishes. Occasionally, hair will start to grow back before the end of the chemotherapy. Sometimes, hair may grow back a different colour or texture.

Hair loss does not always happen straight away; it usually starts within a few weeks of beginning treatment. Sometimes it starts within a few days. It can occur on all parts of the body, including the hands, face, arms, legs, underarms and pubic area. If you lose hair around your eyes or in your nose, you may experience a runny nose and weepy eyes because of this.
Chemotherapy

Cold cap

For some types of chemotherapy, cooling the scalp with a cold cap as the drug is given can prevent hair loss. This works by reducing the blood flow to the scalp so that less of the drug reaches the hair follicles on your head. However, the cold cap doesn’t work for everyone. It only blocks certain drugs and isn’t suitable to use with all types of cancer. Your doctor or nurse will be able to tell you if this is appropriate in your case. If you have a cold cap you will need to allow extra time at the hospital for your treatment.

Wigs, available on the NHS and privately, can be obtained in advance if you are likely to lose your hair.

Chemotherapy can cause your hair to become dry and brittle, so take good care of it. Use a neutral pH shampoo and conditioner (baby shampoo is too alkaline). Cut down on the number of times you wash your hair each week.

Use a wide toothed comb to avoid pulling at your hair and hair roots. Avoid harsh chemicals, such as hair dyes and perms, and excessive heat from heated rollers and hair dryers during chemotherapy and for a few months following completion of chemotherapy.

<table>
<thead>
<tr>
<th>Possible side effects</th>
<th>Symptoms include:</th>
<th>What you should do</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hair loss</td>
<td>• Total hair loss  or  • Thinning of hair or  • Brittle hair</td>
<td>• Use a neutral pH shampoo and conditioner • Cut down on the number of times you wash your hair each week. • Order a wig in advance if you are likely to lose your hair • Avoid harsh chemicals, such as hair dyes and perms</td>
</tr>
<tr>
<td>When symptoms may occur</td>
<td>Within a few weeks of beginning chemotherapy</td>
<td></td>
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</tbody>
</table>
Your fertility

Some chemotherapy drugs can damage the ovary or testis, leading to an increased risk of infertility (inability to have a child) and also, in women, early menopause. If this is an important issue for you, make sure you discuss it fully with your doctor before treatment starts.

Chemotherapy may affect sexual organs or functions in various ways. You may experience changes in your desires or desired level of sexual activity, or you may notice no difference. Loss of libido (sex drive) is not uncommon in both women and men. However, chemotherapy in itself doesn’t usually affect sexual performance or cause impotence. The stress of your illness or the treatment schedule may make you feel more tired than usual. If fatigue is a problem you may want to set aside time for physical intimacy after a period of rest.

Although your sexual needs and desires are highly individual, the following advice may be helpful:

- Try to find out as much as possible about how your treatment may affect you
- Share your worries and feelings with people who care for you

Feel free to discuss any concerns you may have with your doctor or nurse. Even if they are unable to help you, they can refer you to someone who can.

For women

Chemotherapy may affect your ability to conceive a child which may be temporary or permanent, depending on your age and your treatment. If you are concerned about this you may want to discuss it with your doctor before starting treatment.

Women having certain chemotherapy protocols may notice changes in their menstrual cycle. If you are still having periods then it is quite possible that these will gradually stop while you are on chemotherapy. This is less likely if you are in your 20’s or 30’s. If you are in your 40’s, your periods may not return and menopausal symptoms may begin.
Hormonal changes may cause hot flushes and vaginal dryness. If you suffer from any menopausal symptoms, talk to your doctor who may prescribe something to help relieve them.

If you are concerned about preserving your fertility, your specialist cancer doctor may want to refer you to a fertility clinic where you can discuss what options may be available. However, not all of the following options will be suitable for every woman and it may also depend on whether you can safely delay treatment. The options include freezing fertilised or unfertilised eggs, or ovarian tissue:

- Freezing embryos (In Vitro Fertilisation or IVF) – collecting and freezing embryos for later implantation.
- Freezing unfertilised eggs (oocytes) – a procedure that may be considered by women who do not currently have a partner and do not wish to use a sperm donor. This is a fairly new and experimental technique.
- Freezing ovarian tissue – which contains hundreds of immature eggs that could potentially be saved and used to start a future pregnancy. This is still at a very early and experimental stage.
- Drugs to protect the ovaries. There is some evidence that drugs which induce a temporary menopause, for example, Zoladex, may protect the ovaries during treatment from long term chemotherapy damage. This approach is experimental.

**Pregnancy** – It may be possible to become pregnant during the time you are having chemotherapy, but it isn’t advisable to do so. Some chemotherapy drugs can damage an unborn child. During treatment and for about one year afterwards, sperm and eggs may not be formed normally, if they are produced at all. Your doctor will be happy to discuss this further with you.

You or your partner should use a barrier method of contraception during treatment not only to prevent pregnancy but also to prevent any possible contamination with chemotherapy. Non-barrier methods should be fine after treatment is finished but they need to be used for one year after completion of chemotherapy. If you know you are pregnant before starting treatment or become pregnant during treatment, you must tell your doctor immediately.
We are aware that some women may be diagnosed with cancer during pregnancy. If you are in this situation, your specialist doctor will discuss with you, the benefits and risks of having chemotherapy.

**Breast feeding** – There may be a risk of harm to a child who is being breastfed since the drug may be concentrated in the milk. It is very important that you check with your doctor first, if you wish to breast feed while receiving chemotherapy.

**For men**

Some chemotherapy drugs can damage the testis and this may affect your ability to father a child. Although sterility (failure of sperm production) isn’t associated with many drugs, chemotherapy may reduce the number of sperm or their motility (movement). After treatment, some men remain infertile while in others, the sperm count returns to normal.

If sterility is likely to be a permanent side affect, you may be offered the opportunity to bank sperm before starting treatment (sperm is frozen for artificial insemination at a future date). Before sperm banking takes place, you will be asked to have a blood test for Human Immunodeficiency Virus (HIV) antibodies, Hepatitis B and Hepatitis C. This is routine practice to ensure that healthy sperm are banked.

During treatment and for about one year afterwards, it isn’t advisable for you to father a child because sperm may not be formed normally. You or your partner should use affective contraception.
Possible side effects

<table>
<thead>
<tr>
<th>Changes in your fertility</th>
<th>Symptoms include:</th>
<th>What you should do</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>• Early menopause (for women)</td>
<td>Men can consider • Sperm banking (if appropriate)</td>
</tr>
<tr>
<td></td>
<td>• Inability to conceive or father a child (men and women)</td>
<td>Women can consider • Freezing embryos</td>
</tr>
<tr>
<td></td>
<td>• Some drugs can damage the DNA development of a foetus</td>
<td>• Freezing unfertilised eggs (oocytes)</td>
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<td>• Freezing ovarian tissue</td>
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<td></td>
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<td>• Drugs to protect the ovaries</td>
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<td></td>
<td></td>
<td>Men and women • Use effective contraception</td>
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</table>

**When symptoms may occur**

During and following treatment

**Other effects**

Some chemotherapy drugs may, more rarely, cause the following effects. Your doctor will tell you if any of your chemotherapy drugs are likely to cause any of these:

**Allergic reactions**

Occasionally, a small number of patients may experience a severe, occasionally life-threatening, drug related reaction. Symptoms may include a feeling of swelling in the tongue or throat, a cough and breathlessness. You will be monitored closely during your treatment and if any of these symptoms occur, you will be treated appropriately.

**Bladder symptoms**

Some drugs may cause discolouration of your urine after chemotherapy. For example, your urine may turn pink or red up to 24 hours following each treatment. Your chemotherapy nurse will warn you if this is likely to happen. If you think you notice blood in your urine, you should report this to your doctor.
Rarely, a drug can cause irritation to the bladder (cystitis) since it is excreted in the urine. You should drink about an extra pint (half a litre) of fluid within the 24 hours after each injection.

**Blood clots**

Some cancers can increase the risk of developing blood clots. Some chemotherapy drugs (and some other anti-cancer drugs) can also increase the risk of developing blood clots. The type of blood clot seen most often occurs in the veins causing problems like deep vein thrombosis (DVT). This usually refers to a clot in one of the leg veins, causing swelling and/or pain in the leg. If a part of the clot breaks free, it may travel to the lungs, causing shortness of breath or chest pain. This is called a pulmonary embolus (PE). Blood clots can be life threatening and treatment with blood-thinning drugs (anti-coagulants) is usually given to help dissolve the clot and prevent further problems.

Blood clots in the arteries are rare but there is an increased risk with some anti-cancer drugs. For example, drugs which affect tumour blood vessels (anti-angiogenic drugs). Blood clots in the arteries may lead to a heart attack, stroke or prevent the blood supply reaching a limb. Please inform your doctor immediately if you are worried you may have a blood clot.

**Eye symptoms**

Your eyes may become dry, irritable or weepy (conjunctivitis). Please mention this to your doctor or nurse if you get any of these symptoms so that you can be given advice on how to relieve them.

**Fluid retention**

You may notice swelling of your ankles (ankle oedema) or breathlessness. Usually, this is mild and goes away when treatment ends. Tell your doctor if you experience either of these symptoms.

**Flu-like conditions (fever, headaches, pain)**

This reaction does not last very long and can be treated with simple paracetamol.
Hearing symptoms
Some chemotherapy drugs can cause tinnitus (a continual buzzing noise in the ears) or high frequency hearing loss. Tell your doctor if you are worried about changes in your hearing.

Kidney function
There may be changes in the way your kidneys work. You will be monitored closely if you are receiving a chemotherapy drug that could cause any damage. These are usually temporary effects and are unlikely to cause you any symptoms.

Liver function
There may be changes in the way your liver works. You will be monitored closely if you are receiving a chemotherapy drug that could cause any damage. These are usually temporary effects and are unlikely to cause you any symptoms.

Memory and concentration
Some patients find that their short-term memory and concentration are not as good while they are undergoing chemotherapy. This usually resolves soon after treatment has finished, although it can rarely be a long term effect. This may affect your ability to work during treatment if your job involves a great deal of mental effort.

Nervous system symptoms
Some chemotherapy drugs can cause problems with the nerves in the body. This is called peripheral neuropathy. It can cause sensations like tingling, burning, numbness or pins and needles in the hands and / or feet. You may have difficulty with doing up buttons and other fine movements. These sensations may worsen on exposure to cold and can also affect the nose and throat causing swallowing and breathing problems. It may help to wear gloves, warm socks and a scarf. If your mouth is affected avoid cold drinks. Tell your doctor if you get any of these sensations, so they can be monitored and if necessary changes can be made to the dose or the type of drug you receive. Most of the time, these symptoms will get better after your treatment ends, although it may take some time.
If the symptoms become severe, the nerves could be damaged permanently and your chemotherapy drugs may be changed.

**Steroids**

Steroids are natural substances that are made in the adrenal glands, which lie just above the kidneys in the body, and in the reproductive organs. There are also man-made steroids.

Steroids, such as dexamethasone and prednisolone, can be given for different reasons. For some types of cancer, steroids have a direct anti-tumour effect. You may be given dexamethasone with your chemotherapy to help prevent sickness. Sometimes it is given to help prevent other side-effects or to reduce an allergic-type reaction.

The side effects you may experience with steroids will depend on the dose and the length of time you are taking them. Discuss this with your doctor, nurse or pharmacist. The more common side effects include:

- Irritation of the lining of the stomach causing indigestion or heartburn. You may be given drugs to prevent this particularly with higher doses or longer courses of steroids.
- An increased appetite and weight gain.
- Fluid retention. You may notice swelling of your ankles (ankle oedema). Usually, this is mild and goes away when treatment ends.
- Mood changes or difficulty in sleeping. You may find it helpful to discuss with your doctor, which time of day is best to take your steroids. It is usually best to avoid taking them in the evening if possible.

**Tissue damage (extravasation)**

Some drugs can cause damage if they leak out of the vein while they are being injected. This may result in some pain and take some time to heal. If you feel any stinging, burning or pain when you are receiving the drug, or develop any problems at the site, such as redness or blistering once you are home, tell your nurse or doctor straight away.
Long term risks

Developing a second cancer

With some drugs, there is a small possibility of developing a second cancer or serious blood disorder including leukaemia (cancer of the blood cells). However, the benefits of having chemotherapy generally outweigh the risk of developing a second cancer. Your doctor will discuss this with you if your chemotherapy drugs include this risk.

Damage to the heart muscle

This is very rare but can occasionally happen with prolonged usage of some chemotherapy drugs, especially doxorubicin and epirubicin. You should inform your doctor if you have had any pre-existing heart problems and he/she may arrange for you to have a test to see how well your heart is working before you start treatment, if this is likely to be a concern. You should tell your doctor if you notice chest pain or breathlessness.
How will I feel during my chemotherapy?

The fact that you need to have chemotherapy, and the effect it has on your life, can prompt a range of feelings. Fear, anxiety and depression are common to many people with cancer and are entirely normal. This may be in addition to how you feel physically.

**Emotional wellbeing**

Some people find it helps to gather information about their disease and treatment because this lessens the fear of the unknown. Find out as much as you want to know and don’t be afraid to ask questions. Your emotional wellbeing is as important as your physical health.

Loss of memory and concentration may be side effects of chemotherapy and usually resolve soon after treatment has finished. This can affect your ability to listen and remember things which are explained at your treatment visits. Don’t feel embarrassed to ask for information to be repeated as often as you need it. You may find this affects your ability to work during treatment if your job involves a great deal of mental effort.

You may find it helpful to bring a friend, partner or relative to hospital visits, and to make a note of any questions you have beforehand. The doctors and nurses caring for you will be happy for you to do this. There is a blank section on pages 52–54 for you to make notes or write down questions.

Talking with an understanding friend, relative, other patient or one of the organisations listed on pages 50–51 may be helpful.

Many people don’t understand about cancer or its treatment and may avoid you because they’re not sure what to say or how to help. Try to be open in talking with others about your illness, treatment, needs and feelings.

You might like to try new methods to help you cope with treatment and its side effects, such as meditation, relaxation exercises, distraction, visualisation (imagery) or hypnosis. You may want to ask what is available within the hospital or locally in the community.
Remember everyone needs some support during difficult times. Please don’t hesitate to ask for help from your doctor or chemotherapy nurse, during or after your treatment. They are keen to make sure you get the support you need.

**Fatigue**

During your chemotherapy you may start to feel tired or listless. This may be general fatigue or it may be that you become tired more easily after normal activities. This is quite normal and usually occurs with all types of chemotherapy.

Once you know what makes you more tired and when this happens, you can plan ahead. Try to plan your day so that light activities are spaced between more energetic activities. Gentle exercise can be helpful. Make sure you get enough rest and only do what you feel you can cope with.

People will often be willing to lend their support. If you get tired easily, limit your activities and only do the things that mean the most to you. Ask your family and friends to help with household chores, cooking and so on. Work part-time or see if you can work more flexible hours. Conserve your energy for the important events in your life.

Tiredness can also be a sign of anaemia, so do tell the nurse or doctor if you are worried about how you feel.

**What help is available during my treatment?**

Some people may experience problems as a result of their cancer and / or treatment that have an effect on different areas of their life. There is a wide range of services to help you recover and lead an active and productive life. Some services can help you with physical difficulties, for example, physiotherapists and occupational therapists. They may be able to teach you new ways of doing certain activities or help you achieve and maintain independence in other areas of your daily life. If you are finding it difficult to eat well, dieticians can provide advice whether it is during or after treatment.
Other services may be able to help you cope with social and emotional worries, for example social workers. They are experienced in working with people who have to adjust to change or crisis in their lives. A social worker can discuss with you any help you may need at home and give you information and advice on welfare benefits and coping with financial difficulties.

What happens when chemotherapy treatment is finished?

Once your chemotherapy treatment has finished, most side effects gradually disappear. Everyone is an individual and will recover in their own time. You may find that it takes a while for your energy levels to recover. You may also find that it will take some time for your emotions to settle down. As well as feeling relieved that your chemotherapy has finished, you may miss this close support of the hospital you had during treatment. You may find The Royal Marsden booklet After treatment helpful.

You will be given an appointment for a follow-up check. Ask who you should contact if you have any problems or concerns before your appointment.
## Glossary

These are some of the terms you may come across during your chemotherapy treatment.

<table>
<thead>
<tr>
<th>Term</th>
<th>Definition</th>
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</thead>
<tbody>
<tr>
<td>Adjuvant therapy</td>
<td>Treatment given in addition to the initial therapy.</td>
</tr>
<tr>
<td>Ambulatory chemotherapy</td>
<td>Chemotherapy given through a small infusion pump while you are at home.</td>
</tr>
<tr>
<td>Anaemia</td>
<td>Having too few red blood cells, which may cause symptoms including tiredness, weakness and shortness of breath.</td>
</tr>
<tr>
<td>Benign tumour</td>
<td>A tumour or growth that is not cancerous.</td>
</tr>
<tr>
<td>Blood count</td>
<td>The number of three different kinds of blood cells in the body. These are white blood cells, red blood cells and platelets.</td>
</tr>
<tr>
<td>Cannula</td>
<td>A small tube which is put into a vein in the arm or the back of the hand through which drugs are given.</td>
</tr>
<tr>
<td>Chemoradiation</td>
<td>Chemotherapy combined with radiotherapy</td>
</tr>
<tr>
<td>Chemotherapy</td>
<td>Treatment with drugs which destroy cancer cells.</td>
</tr>
<tr>
<td>Clinical oncologist</td>
<td>A cancer specialist who treats cancer with radiotherapy treatments (high energy x-rays) most clinical oncologists will also prescribe chemotherapy (anti-cancer drugs) and hormone therapies.</td>
</tr>
<tr>
<td>Clinical trial</td>
<td>Research aimed at improving cancer care.</td>
</tr>
<tr>
<td>Combination chemotherapy</td>
<td>Several different drugs given together.</td>
</tr>
<tr>
<td>Consent form</td>
<td>A written record that you have agreed to the planned treatment.</td>
</tr>
<tr>
<td>Term</td>
<td>Definition</td>
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<tr>
<td>Haematologist</td>
<td>A specialist doctor who diagnoses and treats blood disorders.</td>
</tr>
<tr>
<td>Implanted port</td>
<td>A device, which is inserted into your body under the skin. It is used to give fluid and drugs, and it can also be used for taking samples.</td>
</tr>
<tr>
<td>Intrathecal chemotherapy</td>
<td>Chemotherapy drugs given through a needle into the spinal fluid around the spinal cord in the back.</td>
</tr>
<tr>
<td>Keyworker</td>
<td>A specialist nurse or other healthcare professional involved in your care who is a point of contact for you throughout all or part of your care.</td>
</tr>
<tr>
<td>Malignant tumour</td>
<td>A tumour or growth that is made up of cancer cells.</td>
</tr>
<tr>
<td>Medical oncologist</td>
<td>A cancer specialist who treats cancer with drugs. These include chemotherapy (anti-cancer drugs), biological therapies and hormone therapies.</td>
</tr>
<tr>
<td>Metastases or ‘secondaries’</td>
<td>The spread of cancer cells to other parts of the body.</td>
</tr>
<tr>
<td>Neo-adjuvant therapy</td>
<td>Treatment given before surgery or radiotherapy to shrink the tumour.</td>
</tr>
<tr>
<td>Pathologist</td>
<td>A specialist doctor who examines body tissues and organs under the microscope.</td>
</tr>
<tr>
<td>Peripheral neuropathy</td>
<td>Sensations like tingling, burning, numbness or pins and needles in the hands and / or the feet.</td>
</tr>
<tr>
<td>PICC (a peripherally inserted central catheter)</td>
<td>A fine flexible tube inserted into the vein in the arm, above the bend in the elbow. It is moved up into the large vein leading to the heart. It is used to give fluids and drugs, and it can also be used for taking blood samples.</td>
</tr>
<tr>
<td>Term</td>
<td>Definition</td>
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<tr>
<td>Prognosis</td>
<td>The predicted outcome of a disease or the life expectancy.</td>
</tr>
<tr>
<td>Radiologist</td>
<td>A specialist doctor in the use of x-rays and other imaging to diagnose and treat disease.</td>
</tr>
<tr>
<td>Specialist nurse</td>
<td>Provides information and support.</td>
</tr>
<tr>
<td>Systemic treatment</td>
<td>Treatment that affects the whole body.</td>
</tr>
</tbody>
</table>
Sources of information and support

Macmillan Cancer Support
89 Albert Embankment
London SE1 7UQ
Tel: 020 7840 7840
Macmillan Cancerline: Freephone 0808 808 0000
Website: www.macmillan.org.uk

Provides free information and emotional support for people living with cancer and information about UK cancer support groups and organisations.

Offers free confidential information about cancer types, treatments and what to expect.

Other useful organisations

National Institute for Health and Clinical Excellence (NICE)
MidCity Place
71 High Holborn
London WC1V 6NA
Tel: 0845 003 7780
Website: www.nice.org.uk

NICE provides guidance for healthcare professionals, and patients and their carers that will help to inform their decisions about treatment and healthcare.

Cancer Research UK
PO Box 123
London WC2A 3PX
Cancer Information Service available on
Tel: 020 7061 8355 or 0800 226237 (freephone)
Email: cancer.info@cancer.org.uk
Website: www.cancerresearch.org.uk or patient information
Website: www.cancerhelp.org.uk
Trained cancer nurses can give information and support relating to cancer and its treatments. Publications are available and their patient information website, www.cancerhelp.org.uk has information on specific cancers. Click on the ‘Clinical Trials’ icon on this website home page to access up-to-date details on UK current clinical trials.

**UK National electronic library for health**  
*www.library.nhs.uk*

National UK health information site – covers all aspects of health, illness and treatments.

**National Cancer Institute (USA)**  
*www.nci.nih.gov*

Provides comprehensive information on cancer and its treatments. (Please note that not all the information will necessarily relate to treatment in the UK.)

**American Cancer Society**  
*www.cancer.org*

American Cancer Society gives detailed information on specific cancers and coping with cancer. Please note that not all the information will necessary relate to treatment in the UK.

**Maggie’s Centres**  
*www.maggiescentres.org*

Offers psychosocial support, through a network of cancer caring centres across the UK and online, to anyone affected by cancer.

**Further reading**


Provides information for people diagnosed with cancer on getting information, treatment options, choices and self help.
Notes/Questions

You may like to use this space to make notes or write questions as they occur to you, to discuss with your specialist nurse or doctor.
Notes/Questions
How you can contact us?

These are the contact details if you have any queries about your illness or treatment or experience any problems mentioned in this booklet:

Name of consultant (chemotherapy)

Names of clinical team including key worker

at ____________________________ Hospital

Telephone number

Name of ward / day unit

Telephone number

Other useful contact details:
The Royal Marsden publishes a number of booklets and leaflets about cancer care. Here is a list of information available to you.

Diagnosis
- A beginner's guide to the BRCA1 and BRCA2 genes
- CT scan
- MRI scan
- Ultrasound scan

Treatment
- Central venous access devices
- Chemotherapy
- Clinical trials
- Radiotherapy
- Radionuclide therapy
- Your operation and anaesthetic

Supportive Care
- After treatment
- Coping with nausea and vomiting
- Eating well when you have cancer
- Lymphoedema
- Reducing the risk of healthcare associated infection
- Support at home
- Your guide to support, practical help and complimentary therapies

Your hospital experience
- Help Centre for PALS and patient information
- How to raise a concern or make a complaint
- Your comments please
- Your health information, your confidentiality