Living Well with Chronic Myeloid Leukaemia (CML)

A Guide for Patients

Leukaemia Care
YOUR Blood Cancer Charity
Introduction

Being diagnosed with chronic myeloid leukaemia (CML) can be a shock, particularly when you may never have heard of it. This booklet covers a range of topics to help you cope a little better.

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You may experience all of the emotions featured in this booklet, some of them, or maybe none at all. Experiences vary from person to person. This booklet, and the organisations listed in it, are available to help you whatever you may be facing.

Remember, you can also find tailored advice for your particular situation by talking to your GP or your medical team.

If you would like any information on the sources used for this booklet, please email communications@leukaemiacare.org.uk for a list of references.
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About Leukaemia Care

Leukaemia Care is a national charity dedicated to ensuring that people affected by blood cancer have access to the right information, advice and support.

Our services

Helpline
Our helpline is available 8:30am - 5:30pm Monday - Friday and 7:00pm - 10:00pm on Thursdays and Fridays. If you need someone to talk to, call 0808 801 0444.

Alternatively, you can send a message via WhatsApp on 07500068065 on weekdays 9:00am – 5:30pm.

Nurse service
We have two trained nurses on hand to answer your questions and offer advice and support, whether it be through emailing nurse@leukaemiacare.org.uk or over the phone on 08088 010 444.

Patient Information Booklets
We have a number of patient information booklets like this available to anyone who has been affected by a blood cancer. A full list of titles - both disease specific and general information titles - can be found on our website at www.leukaemiacare.org.uk/support-and-information/help-and-resources/information-booklets/

Support Groups
Our nationwide support groups are a chance to meet and talk to other people who are going through a similar experience. For more information about a support group local to your area, go to www.leukaemiacare.org.uk/support-and-information/support-for-you/find-a-support-group/

Buddy Support
We offer one-to-one phone support with volunteers who have had blood cancer themselves or been affected by it in some
way. You can speak to someone who knows what you are going through. For more information on how to get a buddy call 08088 010 444 or email support@leukaemiacare.org.uk

**Online Forum**

Our online forum, [www.healthunlocked.com/leukaemia-care](http://www.healthunlocked.com/leukaemia-care), is a place for people to ask questions anonymously or to join in the discussion with other people in a similar situation.

**Patient and carer conferences**

Our nationwide conferences provide an opportunity to ask questions and listen to patient speakers and medical professionals who can provide valuable information and support.

**Website**

You can access up-to-date information on our website, [www.leukaemiacare.org.uk](http://www.leukaemiacare.org.uk), as well as speak to one of our advisers on our online support service, LiveChat (9am-5pm weekdays).

**Campaigning and Advocacy**

Leukaemia Care is involved in campaigning for patient well-being, NHS funding and drug and treatment availability. If you would like an update on any of the work we are currently doing or want to know how to get involved, email advocacy@leukaemiacare.org.uk

**Patient magazine**

Our quarterly magazine includes inspirational patient and carer stories as well as informative articles by medical professionals: [www.leukaemiacare.org.uk/communication-preferences/](http://www.leukaemiacare.org.uk/communication-preferences/)
The monitoring of CML is important for assessing your response to treatment. Your response to TKI treatment will help the haematology team manage your CML, and allow appropriate changes in your CML treatment, if required. It is very important to take your tyrosine kinase inhibitor (TKI) medication as instructed by the haematology team. Missing TKI medication can be detrimental to your overall CML care. Attending hospital clinics as planned by the haematology team will aid doctors in charting your progress at important time points.

At the diagnosis of CML, the haematologist will arrange a full blood count. This test checks the haemoglobin, white cell count and platelet count. When CML is first diagnosed, the white cell count is elevated, and the platelet count can also be increased. The laboratory staff will look at your blood under the microscope (blood film). The blood film shows lots of mature white cells called neutrophils and also immature white cells. The hospital may also arrange a bone marrow aspiration and biopsy at diagnosis, and this looks more closely at the chromosomes of the dividing CML cells. The chromosomes contain long strands of DNA which control the behaviour and properties of the cells. In CML, chromosome 9 and chromosome 22 exchange material giving rise to the Philadelphia chromosome. A special technique called karyotyping can identify the abnormal Philadelphia chromosome. This technique can also identify other chromosome changes in addition to the Philadelphia chromosome abnormality.

Blood samples taken at the diagnosis of CML are sent to the laboratory to perform specialised tests. The quantitative reverse transcriptase polymerase chain reaction (RT-PCR) is a test that analyses a short sequence of DNA. In the case of CML, the test compares the ratio of abnormal BCR-ABL1 gene transcripts (associated with the Philadelphia...
chromosome) to the normal version of the gene. This baseline result enables monitoring of CML at specific time points during TKI therapy.

The European Leukaemia Net (ELN) 2013 guidelines are the standards by which the haematology team will make decisions on your response to TKI therapy. The ELN response criteria assesses your response to TKI therapy at three, six, 12 and 18 months. At three months, it is generally expected the full blood counts will normalise and the spleen will be of normal size. Every three months, a sample will be sent for RT-PCR to monitor the BCR-ABL1/ABL1 ratio. The haematologist will interpret the BCR-ABL1/ABL1 results and advise accordingly regarding any changes in medication, dependent on the guidelines. If the results show a failure of response, then your doctor will change you to another TKI therapy and check for mutations which could have potentially stopped the TKI from working.

The RT-PCR test is very sensitive and can detect low levels of disease. This test can detect 1 CML cell in 100,000. If there is a reduction in disease by hundred-fold, this is equivalent to a cytogenetic remission (no Philadelphia cells in the marrow). The ideal response to treatment is when the CML disease level falls by 1,000-fold, this is known as a major molecular response (MMR). The fall in the RT-PCR is not judged from the level at diagnosis of the individual patient, but to an International standard which is expressed as an International Scale (IS). The RT-PCR test can detect disease levels as low as a 100,000-fold reduction from diagnosis. This is important to bear in mind, as studies have shown CML patients with deep responses to treatment for a certain period of time can potentially stop TKI therapy after discussion with the haematology team. This is called Treatment Free Remission (TFR).
The tyrosine kinase inhibitors (TKI) are usually well-tolerated and rarely cause severe side effects (grade 3 or 4 toxicity). They are targeted drugs which means they eliminate the CML cells expressing BCR-ABL. It is important to take the medication as directed by the haematologist in charge of your care. Prior to starting the TKI therapy, you will be given information leaflets about the specific TKI your haematologist is proposing to start. A consent form listing the potential side effects of the specific TKI will usually be provided. Your doctor will explain the common, uncommon and rare side effects. If you experience symptoms which are affecting your quality of life when taking TKI therapy, then contact the haematology team and your CNS.

Viral testing for hepatitis B is always performed prior to TKI therapy. If you have had a previous hepatitis B infection, then the doctor will recommend screening to reduce the risk of the hepatitis virus from reactivating.

If you are a man with newly diagnosed CML, you will be offered the opportunity for sperm storage. If you are a woman, your doctor will reassure you that fertility is not recognised to be impaired. You can discuss this further with your doctor. For more information on relationships, sex and fertility, go to page 16.

The TKIs can cause general side effects such as tiredness and lethargy, abnormal liver function, skin rashes, muscle aches and joint pains, headaches, changes in bowel habit and a fall in blood counts. These side effects are mild and are measured as grade 1 or 2 side effects. If necessary, they can sometimes be treated with medication.

**Imatinib**

Imatinib is the most common TKI prescribed in the UK. Side effects usually occur in the first one to two months, though there are some long-term side effects associated with imatinib.

Fluid retention can be a
troublesome symptom in up to 2.5% of patients on imatinib. This can sometimes be treated with diuretics, and regular weight measurements can help guide treatment response. Fluid buildup around the eyes (periorbital oedema) can also be an issue, but, again, can sometimes be responsive to diuretics. If the fluid buildup around the eyes is difficult to control, the dose of TKI may be altered or another TKI may be required.

Muscle cramps, joint pains and general bone pain are also side effects of imatinib. Muscle cramps can be frustrating and affect your quality of life. The doctor will check your salt levels in the blood and restore deficiencies if present. Simple pain relief and tonic water or occasionally quinine can also be helpful to treat the cramps.

Symptoms affecting the eyes are common with imatinib. Imatinib can be associated with conjunctival haemorrhage (bleeding into the white section of the eye), eyelid swelling, dry eyes and blurred vision. It is important to discuss these with your doctor.

Imatinib is not thought to be associated with an increased risk of bacterial infections in the form of respiratory and urinary tract infections if the white cell count is not low. If you have fevers, it is important to contact your haematology department. Viral reactivations such as hepatitis B can also occur while on imatinib, particularly if there is a past history of hepatitis B. Responses to various vaccinations against influenza and bacterial infection (pneumococcus) can be less effective while on TKI therapy. Live vaccines are not advised, and if necessary should be discussed with your doctor.

Skin rashes can occur while on imatinib treatment. Skin rashes can be treated with steroid creams, antihistamines and oral steroids if necessary. If the rash is severe, the imatinib will be stopped temporarily. The doctor may decide to restart the imatinib
if the skin rash improves. Imatinib associated photosensitive skin reactions have been reported, but, regardless, avoiding direct sunlight and using high-factor sun protection is recommended.

During treatment with imatinib, the haematology team will monitor your full blood count. Imatinib can cause a reduction in the red cell count (haemoglobin), white cell count and platelet count. Your doctor will closely monitor the blood counts and the imatinib may need to be temporarily stopped if the blood counts fall below a certain level. Your doctor will then decide whether to restart at a lower dose.

Nausea, vomiting and abdominal discomfort are all recognised side effects of imatinib. Nausea can sometimes be reduced by taking imatinib on a full stomach or during the meal. Anti-sickness and anti-indigestion medication can also be helpful.

Imatinib is mainly metabolised (broken down) in the liver. It can cause abnormal liver function tests, and your doctor will be regularly monitoring the liver function. Imatinib rarely causes severe liver abnormalities but, if there are significant liver abnormalities, the liver tests usually recover once imatinib is stopped.

While on treatment for imatinib, your doctor may check your thyroid function if you have symptoms of low thyroid levels.

In female patients, imatinib may adversely affect a pregnancy and the foetus. Foetal malformations have unfortunately been documented with imatinib use. It is therefore vital to discuss with your doctor when it is safe to consider trying for a baby.

Safe and effective contraception is important while on imatinib. Imatinib will be stopped while you try to conceive. Imatinib is found in breastmilk, and it is therefore recommended not to breastfeed while on imatinib. Imatinib has been associated with reduced sperm counts in males but, more recently there has been less...
concern regarding male fertility, which your doctor can discuss with you.

**Nilotinib**

Nilotinib is associated with an increased risk of arterial blood clots in the lower limbs and coronary vessels and an increase in lipid and glucose levels. It is important to reduce the risk of arterial blood clots by prevention and early detection. It is advised to try and stop smoking, and good blood pressure control is encouraged. This is achieved by monitoring your blood glucose, cholesterol and blood pressure regularly. If angina develops while on nilotinib treatment, a cardiologist will help control the symptoms and optimise your heart medication and your haematologist may recommend that you switch TKIs unless there is another alternative. While on nilotinib, the blood sugars can increase. If you have pre-existing diabetes, it is important to monitor your blood sugars as your diabetic medication doses may need adjusting, and your doctor may decide to give you an alternative TKI.

Nilotinib can cause heart rhythm disturbances. Your doctor will arrange an ECG to be done prior to starting nilotinib to ensure this medication is safe for you, as certain heart rhythm changes may prevent you from taking nilotinib.

Liver abnormalities, including abnormal liver function tests, can occur while on nilotinib. Depending on the severity of the abnormal liver function tests, nilotinib may be temporarily withheld or stopped permanently. Rarely, nilotinib can also be associated with inflammation of the pancreas, especially if there is evidence of abdominal discomfort and high amylase levels.

Nilotinib, like imatinib, can rarely cause headaches, infections, nausea and change in bowel habit. Skin rashes and itching are common on nilotinib.
Coping with side effects of TKIs (cont.)

Dasatinib

Dasatinib can be associated with minor buildup of fluid in the lungs in up to 28% of patients. However, a small number of patients treated with dasatinib have continued gradual lung fluid buildup, called a pleural effusion, and can occur at any time after dasatinib treatment. The fluid buildup is usually not serious. However, in some people, it can cause dry cough, shortness of breath and reduced exercise tolerance. If the pleural fluid is causing symptoms, then the dasatinib will need to be stopped, and steroids (possibly diuretics) may be required. It is important to contact your doctor if you develop these symptoms. A chest X-ray can help to determine if you have a pleural effusion. It is important not to re-start the dasatinib until the fluid has resolved.

Dasatinib can also cause headaches, a fall in blood counts, skin rashes and infections (if the blood counts are low).

Bosutinib

Bosutinib can cause diarrhoea soon after taking the medication. Diarrhoea can still be problematic in some patients for some time, but loperamide is helpful to treat it from the outset.

Bosutinib can also cause a fall in blood counts, infections (if the blood counts are low) and skin rashes.

Ponatinib

Ponatinib can cause an increased risk of blood clots in the arteries (thrombosis) and veins. The risk is higher than with other TKIs. Thrombosis can develop in lower limbs and heart vessels. The risk of this can be reduced by controlling the blood pressure, stopping smoking, maintaining good diabetic control if you are diabetic, and treating high cholesterol levels.

Ponatinib has also been associated with inflammation of the pancreas. This can cause abdominal pain and your doctor
needs to be informed.

Ponatinib can cause a fall in blood counts, infections (if the blood counts are low), liver test abnormalities and skin rashes.
CML is generally treated with tyrosine kinase inhibitors which tend not to have a big impact on appearance, your body and your body image. However, for those who receive chemotherapy, there may be some noticeable changes.

Weight
Some people may experience weight gain as a side effect of treatment or a symptom of CML.

There are ways to help you maintain a baseline or good weight throughout. These include screening and assessing your weight and nutrition status at presentation of CML. If you experience extreme weight loss, you should be referred to a specialist dietician, where appropriate.

Physical pain
Bone and joint pain is more commonly reported in patients with CML as this is a side effect of some TKIs such as imatinib. Furthermore, certain procedures and treatments can be associated with pain, including bone marrow biopsies.

Chronic pain has also been found to be associated with a lack of appetite, fatigue, nausea and psychological distress.

There are different approaches to managing pain. If you experience pain, it can be treated with painkillers. Other approaches include complementary therapies, psychological interventions like mindfulness, meditation, nerve blocks and TENS (transcutaneous electrical nerve stimulation) used for blocking pain signals from the nerves to the brain.

Where pain is not adequately controlled, or there is no response from standard interventions, specialist palliative care teams can play a significant role in managing pain symptoms.

If you need any help in dealing with pain, speak to your medical team.

Hair loss
Hair loss or alopecia is a common and well-known side effect
experienced by people receiving high-dose chemotherapy; however, it is not a common side effect of TKIs. In fact, most TKIs can initially cause some hair thinning, but this normally resolves on its own within six months to a year.

You are entitled to a free wig if you are a hospital inpatient. For more information, go to the NHS website: [www.nhs.uk/NHSEngland/Healthcosts/Pages/Wigsandfabricsupports.aspx](http://www.nhs.uk/NHSEngland/Healthcosts/Pages/Wigsandfabricsupports.aspx)

### Infection control

The infection rate in CML patients is much the same as the general population. When you have an infection, it can take several doses of antibiotics to bring your temperature and symptoms under control, but antibiotics should offer an effective treatment.

Vaccinations are designed to protect people against infections by priming the immune system. Some vaccinations are live (they are capable of causing a milder version of the infection) while some are inactive (they contain dead particles of the infection and therefore incapable of causing an infection).

Live vaccines should not be given in general to blood cancer patients, but your doctor can discuss this in more detail. The seasonal influenza vaccine is recommended yearly.
Relationships, fertility and sex

Sex life and libido
The human and personal experience of sexuality varies from individual to individual. It is lived, sensed, experienced, expressed and communicated in different ways, via thoughts, fantasies, desires, beliefs, attitudes, values, practices and the way relationships are formed. It is related to the way human beings form emotional and physical bonds.

World Health Organisation states clearly that human sexuality has many dimensions and factors that include sex, gender, sexual identity, sexual orientation, eroticism, love/emotional connection and reproduction.

A great part of our sex life, libido, feelings and the way human beings form relationships is influenced by hormones that are released in our brains and travel through our bodies, having different effects in specific parts of it. Some of the hormones involved in sexuality are testosterone, dopamine, norepinephrine and serotonin. They are present in the body and brain and they can interfere in sexual impulses, libido and frequency of sexual intercourse.

There are also other factors that can influence and interfere with your sex life and libido on a permanent or temporary basis. The physical and emotional impact of having a cancer diagnosis and the side effects of treatment can have a huge influence on libido, wellbeing and relationships.

At diagnosis
The news of cancer may impact your wellbeing and create anxiety and fear which may lead to a loss of interest in sex and reduced libido. However, while CML is a form of cancer/neoplasm, most patients will have a normal life expectancy.

It is important that you communicate your feelings and emotions with your doctor, nurse and partner and not to isolate yourself. It’s important that you involve your partner as early as possible in the process. Having
an emotional support system whilst going through cancer is important, as well as maintaining connections and a bond with your partner through the process. Your partner may also wish to be involved in the consultations and, sometimes, may help you to envisage how to best cope with short, medium- and long-term strategies and plans for facing the future. It is important to have in mind that whenever there is a crisis in a person’s life such as cancer, having a good support system helps to overcome the difficulties of diagnosis, treatment and follow ups, as well as managing and maintaining a good quality of life. Bear in mind that having your partner in your life can be important, as they are there to help support and nurture you.

**During treatment**

Some treatments may put you off of having sex and reduce your libido and desire for sex. If you are feeling this at the moment, that’s perfectly normal and a common side effect. There are many ways to express your emotions and feelings. Remember that your first priority is your health, so you may find alternatives to sexual intercourse and ways to express your emotions which are as valuable as sex. It is important to involve your partner along the way.

It is okay if during treatment it takes some time to adjust to the new reality, and this may also affect your relationships. You might have less interest in sexual contact and desire to have intercourse for a number of reasons. Treatments may potentially affect your hormonal system in your body, and/or cause temporary or long-term changes in your body, which may interfere with your body image, your wellbeing, your self-image and sometimes your mental wellbeing. It is important that you talk to your partner and healthcare professionals as they will able to provide support.

Try to remember that you are not alone in what you are going through. Although it’s always
different from person to person and it is your own individual experience, there are other patients that somewhere along their cancer journey have felt similar to you. Your healthcare professionals can help you to get information and proper support for your particular struggle, if needed. In some situations, these problems may already be thought of and foreseen whilst professionals organised your treatment care plan. Reach out to your healthcare team so they can provide and/or direct you to the professionals that can help you.

Contraception

There is no contraindication on having sex whilst on treatment. However, if it makes you uncomfortable, you and your partner should respect each other’s decision. Exploration of your partner’s feelings and sensations can be done via other ways and can be as pleasant as sexual intercourse.

After you start your cancer treatment, your nurse/doctor may advise you to abstain from unprotected sexual relationships. TKIs will affect the development of a foetus therefore avoiding a pregnancy while on treatment is important; until you seek medical advice on this matter. You should always prevent a natural conception from occurring before you seek proper advice from your oncology/fertility team. Each treatment strength is different, so it is paramount that you ask for individual and specific advice before making a decision to naturally conceive. There isn’t a formula to calculate the exact date and time when you can naturally conceive, but you can discuss the timing with your doctor. Not waiting for the correct time to naturally conceive may put your baby at risk of malformation at birth, in the uterus or other complications through life. It is safer to confirm with your clinician and discuss this issue before attempting a pregnancy. Please talk to your doctor/nurse/fertility specialist
before you start planning a family.

**Pregnancy and breastfeeding**

Research studies show that cancer during pregnancy is relatively rare. Depending on the stage of the pregnancy when the diagnosis is disclosed, the way a pregnant woman with a diagnosis of CML is treated will vary. Every case is different and the whole team is involved as well as you and your partner. It is important that you seek information and ask questions about treatment, the impact on the baby and/or when is time to decide to go through a different route. It is important that you seek information and ask all the questions you want. The welfare of the child and its future will always be taken into consideration by the team, and decisions may need to be made, so it’s very important that you and your partner are comfortable with the explanation obtained, as well as feeling confident that you were given enough information in order to be able to make an informed decision leading up to the due date.

Experts tell us that TKI therapy during the first trimester of pregnancy is to be avoided as it can cause high genetic anomalies to the baby and could potentially impact the baby for life or result in a miscarriage or still birth.

Nevertheless, it’s always discretionary and up to your team of healthcare professionals to discuss the timing of CML treatment with you. In some situations, it may not be possible in this period due to side effects, so it is important that you always seek advice from your healthcare team.

If you are diagnosed in the last months of your pregnancy, your doctors will discuss treatment options with you, the period of pregnancy and the decision to start treatment before or after delivery. Seek advice from your consultant and medical team.
Breastfeeding

Milk is produced in the woman’s breast and released through the mammary ducts and expelled by the nipples.

Whether or not you can breastfeed your baby will depend on the treatment regime you are on, but it is not advised on TKI therapy. Other clinical situations may impose a challenge on breastfeeding, so seek a medical opinion from your nurses, midwives and doctors.

Breast milk is still the most complete food for babies, and it has several benefits which baby formulas cannot give. There are other components in breast milk which will also help the baby to develop their immune system and be more equipped to fight infections. In the first few days following delivery, you may be able to feed your baby with colostrum, the first form of milk from the mammary gland.

If you are on long term medication for CML, you potentially may not be able to breastfeed due to the medication you are taking. Always seek medical advice. There are several baby formulas in the market which will keep the baby fed and healthy.

Fertility after CML

When facing a diagnosis of CML, the priority is survivorship.

In the UK there are some restrictions to all types of fertility treatments. Funding for fertility treatments under the NHS follows strict rules and policies so it is important to speak to your General Practitioner and hospital consultant when you are thinking of applying for an assisted conception treatment, if you are struggling to naturally conceive.

As soon as possible, you should seek information from both oncologist and fertility specialists in order to make informed decisions about fertility and the next steps you should take.

The NHS funds the IVF (in-vitro fertilisation) and ICSI (intra cytoplasmic sperm injection) treatment for female partners with a different amount of
funding available in each region of the country. With this being said, for a woman that already has one child, or has reached a certain age, it is less likely that funding will be granted. In the event that you are not granted the funding, you may need to fund your treatment privately.

**Women and infertility**

Women are born with all the eggs they will ever have. After puberty, one egg is released each cycle/month, it travels from the ovary through a fallopian tube and reaches the uterus. If this egg is not fertilised by a sperm, the pregnancy will not occur. Ovarian reserve is the term used to describe the remaining egg supply of the ovaries.

When you are diagnosed with CML, your fertility is not thought to be at risk before TKI therapy, but your doctor can advise you further. However, once you have started and responded to treatment, there are options as to when and how you can stop treatment in order to have a family which you can discuss with your haematologist.

Adoption is also an alternative when other options are not possible.

**Men and infertility**

With some treatments, the cells responsible for producing sperm can be affected. Therefore, before undergoing TKI treatment, you should discuss the need for sperm freezing. This would give you the chance to try to father a biological child in the future via IVF or ICSI (in case your fertility is affected during TKI treatment). TKIs do not in general affect fertility and males can continue on treatment when they father a child without any ill effects.

For more information about sperm storage go to the HFEA (Human Fertilisation and Embryology Act) website at [www.hfea.gov.uk](http://www.hfea.gov.uk).

**Changes in relationships**

Following the news of a cancer diagnosis, its impact can vary from person to person as well as
in their partners and family. You should seek as much information as possible about your treatment as well as other factors as this may help to empower you to make the best decisions regarding your future.

Depending on your CML and duration of treatment, there will be a period where you will suffer from some physical symptoms. Cancer and treatment can make you feel physically ill, cause you pain, tiredness, decrease your libido, and cause you to feel all sorts of emotions. You may feel weak, unattractive, afraid, feel that your body is different because of the side effects or based on the fact that you may be losing or gaining weight. Sex can become the thing that you don’t want to do, or that is something that you don’t want to talk about. This can affect your relationship.

The libido can decrease as well as the frequency of intercourse.

There will be some changes in your life, when you are attending the hospital/clinics for your treatments.

It’s important that as soon as possible you involve close family and your partner in the process. This will give them the opportunity to be involved and be aware of the decisions that are being made.

An honest discussion is important. You may have loads of thoughts coming through your head about the disease, the future and about your partner and the relationship. It is okay to feel confused and sometimes to feel afraid and uncertain about it all. If you need to when it comes to breaking bad news to your partner, go to page 66 or seek help from your medical team.

Changes in your body can affect your self-esteem and your sense of feeling desired. Your libido may be affected also.

You may feel more emotional and you may even have difficulties in processing all the information that is given. You may need to ask the same questions many time. Feel free to enquire as many times as you need about information concerning you and
your treatment/future.

It is important to talk to your partner and seek support. You are not alone, and you shouldn’t be. There are other sources in the community that offer help. Charitable organisations and your medical/nursing team/GP can also guide you through other resources available in the community.
Emotional impact of CML

This chapter describes some of the more challenging emotional responses you may experience as you adjust to life following a diagnosis of CML. Everyone is different and what follows may or may not apply to you. There is no right way or wrong way to live with CML. As you gradually adjust to this major change in your life you will find what is most helpful and feels right for you.

Feelings
Living with CML can be described as feeling like being on an emotional rollercoaster ride. You have had to face the shock of being told about the diagnosis and then the challenge as you adjust to the CML treatment that is right for you. The knowledge that CML will always be a part of your life brings its own adjustment. You may experience a range of emotions that can catch you by surprise and may sometimes feel overwhelming. It is common to feel low, sad, angry or anxious. Feeling down or worried is a normal reaction and does not mean that you are weak or failing to cope. Your feelings may come and go. There may be times when you are not sure how you are feeling and other times when you feel okay. However, it is gratifying to know that for responding patients, the life-expectancy of CML patients is similar to that of the general population.

Hope and determination
Hope and determination will help you to face CML. Identify your reasons to have treatment and remind yourself of them when you feel that you are struggling. Notice what helps you to feel strong and draw on the support of others. We are very resilient beings and with time and good support, you can adjust to this enormous change in your life.

Anger
Anger is an emotion that we naturally feel when we are challenged and under threat and so it is not unusual for illness to provoke an angry response. You may feel angry with your body for becoming ill in the first place. Thoughts like, ‘Why me? It’s not fair, I don’t deserve this’
can make you feel angry. You may be annoyed with healthcare professionals and with the lack of control over your life. It may be frustrating that you have to live with the side effects of treatment such as fatigue or diarrhoea.

Anger makes us feel uncomfortable. Some of us shout and stomp around, but for others, feeling and expressing anger is a struggle. It can be a physical feeling of restlessness and frustration and of not being able to settle. It may leak out as irritability or impatience with the people around us.

It helps to acknowledge your anger at the time rather than hold on to it and let it build up, otherwise, it may burst out and leave you feeling helpless and out of control. Buy yourself some time to let the tension go by leaving the room, getting some fresh air, listening to some loud music, doing something physical or putting your angry thoughts down on paper. Try and identify what it is that is making you angry, so you can direct the energy that comes with anger in ways that help you to feel more in control of things. This could be setting some recovery goals for yourself, doing some exercise or physical activity to shift your pent-up tension or talking through your frustrations with someone who is a good listener.

**Feeling alone**

There are times when living with CML that you may feel alone. This can be despite having good support from your family, friends and healthcare team.

Worrying about how others might respond if you tell them what’s really on your mind, feeling that you need to hide your feelings to protect others or because they may not understand can leave you feeling isolated and alone. Feeling different from those around you whose lives are continuing as before can be a challenge.

**Loss, sadness and depression**

The experience of CML brings with it many losses. You might pine for the life that you had before the illness, and for the hopes and
Emotional impact of CML (cont.)

plans that have been changed for now, due to the loss of control over your life you feel because of treatment. You may feel a sense of loss for the healthy you and feel sad and vulnerable because of the changes to your body. It is normal to grieve for these losses and to feel sad when you think about the impact that CML has had on you and those close to you. Give yourself permission to allow these feelings of grief to surface. It is not self-indulgent or weak to feel sad as you adjust to your new reality.

Sadness can often make our mood feel low. People sometimes worry that feeling low can affect how they live with CML or respond to treatment. They can put themselves under pressure or are encouraged by others to remain ‘positive’ and to ‘fight’ the illness. This becomes a way of coping for some people and it works for them. It is important, though, to know that everyone copes in their own way and that you don’t have to be positive all the time. A hopeful and determined outlook will help you to get through the challenges but it is alright to feel low from time to time, particularly when your energy is low, or you experience setbacks.

Feeling low from time to time is normal and these feelings usually pass. But if your low mood persists over a couple of weeks, you feel depressed and it is affecting your day to day life, you may be suffering from depression. It can be hard to separate out the physical and psychological causes, but the feeling of depression is deeper, longer and more unpleasant than the short episodes of feeling low and unhappy that everyone sometimes experiences.

Some of the common symptoms of depression include:

- Feeling sad or low, most of the time
- Losing interest or enjoyment in people and things
- Finding it harder to concentrate or make decisions
• Feeling helpless, hopeless or worthless
• Feeling restless, agitated and irritable
• Feeling easily moved to tears
• Thinking negative thoughts about yourself, the world and the future
• Difficulty getting to sleep or waking up very early
• Having thoughts of self-harm or suicide

Depression does not mean that you cannot cope or are weak. It is a common medical condition and there is support available to help you get through it. You cannot simply ‘snap out of it’ or ‘pull yourself together’.

Talk to your GP or healthcare team if you think that you may be depressed. Depression can be treated with both medication and talking therapies such as cognitive behavioural therapy (CBT) and counselling. Your doctor may also recommend self-help books or an online course. Depression can make you feel less inclined to ask for help, so be guided by those close to you.

If you are having suicidal thoughts or feelings, you can call the Samaritan 24-hour helpline on 116 123. www.mind.org.uk and www.nhs.uk have further information and links to support.

**Anxiety and worrying about your health**

CML brings with it the challenge of living with uncertainty. What will happen to me? Will the treatment work? Will the CML stay in remission? It is natural to feel anxious when faced with fears for ourselves. Anxiety is like an internal alarm system going off. Being anxious will not affect the CML or harm your response to treatment. Do ask for support if it is making it difficult for you to have medical procedures.

We feel anxious when we feel threatened. The brain uses adrenaline to tell the body to run away from or to fight a danger...
and this is called the ‘fight-flight response’. It is helpful when we need to escape quickly from a physical threat such as a fire, but less helpful when the brain interprets stressful life events like illness in the same way. Because we can’t physically run away (although we may feel like it), those same fight-flight physical symptoms can persist and make you feel uncomfortable. It causes that familiar feeling of lying awake at night and the mind won’t switch off because it’s focused on the threat, and with it comes a racing heart, tight chest and tense muscles. When we feel constantly stressed, we struggle to relax, it is difficult to concentrate, and we can feel impatient, irritable and restless.

Worrying is something we do to try and cope with uncertainty. We try to predict and control future events by thinking about them, perhaps to try and prepare ourselves if the worst happens. Worry can be useful when it helps us to take action to sort out a difficult problem. But more often, these ‘What if …’ thoughts or images focus on things that could go wrong in the future. All that worry does is keep our attention focused on the threat and this keeps the anxiety going. It’s helpful to remember that our brain is responding to the threat of something that may happen in the future. It hasn’t happened yet, and it may never happen.

When we are faced with a threat like CML, our alarm system can become a bit too sensitive and lots of things can trigger our anxiety and keep it going. We are always on the look-out for threats. Each pain, niggle, clinic appointment or wait for test results can set off fearful thoughts about your health. This is normal and gradually eases as you adjust to living with CML. Your consultant and specialist nurse will be able to guide you about any physical symptoms that you need to watch out for.

If you notice that you are constantly checking for new symptoms, frequently seeking reassurance from others or that...
you avoid doing things, it may be a sign that anxiety is becoming a problem. If you feel that the uncertainty is making you feel so anxious and worried that it is affecting your daily life, you may find it helpful to talk to your healthcare team.


**Managing Stress**

You cannot remove the stress of CML completely but there are some things that you can do to help yourself manage when you are feeling stressed:

- **Exercise** is an excellent way to reduce the impact of stress. It could be any physical activity – walking, exercise or dance classes, yoga, gardening, housework etc. It helps us to use up some of that extra adrenaline that comes from worrying, helping us to feel calm. For more information about the potential benefits of exercise, go to page 40.

- **Relaxation and breathing exercises** can help to calm the mind and release muscle tension.

- **Doing an activity** that you enjoy can help you to stay in the present moment and enjoy life in the now rather than jumping ahead in your mind to predicting the future.

- **Talking about** what's on your mind to someone who is a good listener or writing worries or thoughts down on paper can make them feel less overwhelming. When we hear ourselves out loud, we are more able to rationalise and put worries into perspective. We can ask: "Is this a problem that I can do something about or is this an uncertainty that may never happen? Am I concentrating on negative fears for the future at the expense of hope and enjoying life again today?"

- **Mindfulness** is a great way to be in the present. It is simply
Emotional impact of CML (cont.)

- bringing our attention into the present moment and being fully aware of what we are experiencing - noticing what is going on using your senses rather than thinking about the past or worrying about the future. We are observers of our thoughts and feelings, allowing them to come and go without getting caught up in them. When we bring our attention into the present moment rather than our imagined future, we realise that we are coping, and getting on with life the best we can. If you have experienced trauma in the past, mindfulness may not be helpful and so seek advice. For more information about how mindfulness may be able to help you, go to page 86.

- Complementary therapies can help you relax and feel more in control, but make sure you are seeing a qualified therapist who has experience of treating people with cancer. If you would like to know about the types of complementary therapies available, go to page 75.

- Talking therapies such as counselling can help you to talk through your experience to try and make sense of it all. Talking to someone you don’t know and in confidence can be helpful to understand and process feelings as normal responses to a difficult situation. Counselling may be available through your doctor or local cancer support centre.

- Connect with others who have been affected by CML through a support group or online forum. Each person’s experience is unique, but it can help to hear from others who know what it is like to have CML.

- Do the things that work for you – maintain the activities and hobbies that were important to you before your diagnosis and try to keep these interests going if they are not too tiring and you are still enjoying them.

- See the people you want to see and talk about what you feel comfortable with. It is fine to change the subject if you don’t want to talk about something.
A new normal

A diagnosis of CML for most people will mean living with daily medication and its possible side effects. There will be regular blood tests and clinic appointments but much of life may carry on as before. With time and support, you can understand, manage and adjust to living with CML as one aspect amidst the rest of your life.

Having a good understanding of CML can help reduce the stress and uncertainty. Get as much information as you need from your clinical team and support organisations. Keeping a diary of your symptoms may be helpful to keep track of side effects to discuss with your team. Managing your condition day-to-day can help you feel that you are taking back control e.g. plan activities so that you can pace yourself to manage fatigue and follow a healthy diet.

Your life may have a very different outlook because of CML. Continue to seek support from those close to you. You might find that making decisions and plans with those around you helps you feel more prepared for what the future may hold. Perhaps you will discover that you or those around you are more resilient than you ever imagined.

A good response to treatment brings hope for the future, and life inevitably takes on a new meaning. People sometimes think about what is most important to them in life and make changes for the better. As you work through your process of adjustment, you can gradually incorporate all that you experience because of CML into your new view of yourself and your life.
Diet

Nutrition and wellbeing

With a diagnosis of CML, some aspects of your day to day life may change during the duration of your treatment. It is therefore important that you have information that will help you, or someone you care for, to stay as well as possible throughout their treatment.

What is neutropenia?

Treatment for some cancers acts to ‘kill off’ the harmful tumour cells in the body, and in doing so the treatment can also destroy healthy cells. When this happens with blood cancers, the protective white cells in our body (called neutrophils) which naturally fight infection, are reduced by both the disease and by the treatments.

Neutropenia occurs when the level of neutrophils falls below a certain level as a result of your illness. However, patients on long-term TKI therapy are rarely if ever neutropenic.

As your neutrophils are much lower than normal you are at greater risk of infection, including foodborne infections and food poisoning, and so one of the measures to prevent this is to ensure scrupulous food hygiene.

What is a healthy diet?

To help ensure your treatment is as effective as possible, a well-balanced nutritional intake is a good starting point. This can be difficult if you are not feeling well and some ideas to cope with these particular difficulties will be covered in this chapter.

Diet and nutritional intake is an area where there is a huge amount of information available to the public, and unfortunately not all of it is based on scientific evidence. It is advised that sensible healthy eating advice is followed for all patients and that no ‘special’ diets are used unless this is recommended by your consultant or a qualified dietitian, and forms part of your clinical treatment.

However, just knowing what makes up a well-balanced diet can lead to confusion. The Government’s Eatwell Guide shows that you should try to eat foods from all food groups every day and within certain proportions.
Try to have as many different food stuffs from all food groups as this will ensure you get all your necessary nutrients for good health.

**Carbohydrates**

These foods are often considered unhealthy. This is not true, and you should try to have some starchy carbohydrate at each main meal like porridge, cereals, potato, pasta, rice or couscous. These foods should make up at least a third of what you eat. Wholemeal and wholegrain varieties will increase your fibre intake, but white ones are also healthy foods. Nutritionally, all these foods are very similar and will provide you with a supply of energy throughout the day.

Particularly if you are being treated with imatinib, a carbohydrate full meal can also help with nausea.

**Fruit and vegetables**

Make sure you have your five portions of fruit and vegetables a day. This can be all kinds of fruit or vegetables: fresh, frozen, canned, dried or as juice or smoothies. Smoothies made with frozen fruit and added milk can make a high energy drink and are particularly helpful if your appetite is poor. Fruit and vegetables will also provide dietary fibre which helps regulate your bowels and prevent constipation which can be a side effect with your treatment.

**Dairy foods**

Milk, cheese and yoghurt are good sources of protein and calcium in the diet. If you have lost weight or your appetite is poor, these foods are very helpful as they tend to be relatively high in energy and can help to boost your overall calorie intake.

**Other protein foods (meat, poultry, eggs, pulses, peas, beans and lentils, nuts and meat alternatives such as Quorn or tofu)**

Protein is an important nutrient as it helps to maintain your muscle strength and enables you to go about your normal daily activities and reduces fatigue. It is recommended that you have at least two portions of protein foods...
from this group every day, but you may need to increase this if you have lost weight to enable you to replace lost muscle tissue.

**Oils and spreads**

The last very small group is for fats and spreads which are used in cooking. General healthy eating advice is to use less fat, and that the fat you do use comes from healthier unsaturated fats such as olive, sunflower, rapeseed or soya oils. If you need to gain weight it is okay to use a little more of these fats to increase overall energy intake but remember that the most important nutrient way to regain lost weight through illness is by increasing protein and overall energy intake.

Sitting outside the Eatwell Guide are treat foods such as cake, biscuits, crisps or sweets and chocolate. These foods are outside the main plate model as they are not essential to health. However, they can make an important contribution to your total energy intake and also can help boost appetite. A drink of ice-cold milk and a small slice of sponge cake makes a good high energy snack between meals.

In summary, the nutrients each food group provides are:

- **Bread and cereals** - provide energy, B-vitamins, iron and folic acid and dietary fibre.
- **Fruit and vegetables** - provide vitamins A, C and E, folic acid, dietary fibre and potassium.
- **Fish, eggs, Quorn, nuts, soya** - provide protein, iron, B-vitamins, thiamine and B12, zinc.
- **Milk and dairy products** - provide protein, Calcium, B-vitamins, riboflavin and B12.
- **Fats, oils and spreads** - provide energy, essential fatty acids and fat-soluble vitamins ADE and K.

**Neutropenia: what you need to be careful of**

If you have neutropenia, the following guidelines should always be considered when choosing and preparing food:

**Water**

Freshly drawn tap water and fizzy bottled water is safe to use as a drink and in cooking. This can
also be used to make ice at home.

**Milk, cheese and ice cream**
Milk must be pasteurised or UHT. This applies to all kinds of cows’ milk, sheep and goat’s milk, and milk alternatives such as soya milk.

Similarly, all cheese must be made with pasteurised milk. This includes fresh and vacuum-packed cheese as well as processed cheese spreads. You should avoid blue and veined cheeses, soft mould ripened cheese like camembert and feta cheese.

All commercially available ice cream maintained at the appropriate temperature is fine, so long as it is not allowed to thaw and refreeze. Do not use soft scoop ice cream or ice cream from open ice cream counters.

**Yoghurt**
Live yoghurt is not harmful and can be safely included in your diet. However, it is advised to avoid yoghurts and drinks containing prebiotics and probiotics.

**Honey**
Honey naturally contains yeasts and spore-forming bacteria. Despite its antibacterial properties, it is recommended that honey from the following sources is avoided during your treatment: raw honey, fresh honey comb, unpasteurised honey or strained honey. Pasteurised or filtered honey removes fine particles of dust, pollen or nectar and is fine to use. Where possible, use small jars or individual portions.

**Nuts and dried fruit**
As long as these have undergone heat treatment, to kill off any potential bacteria, they are considered safe to use. Therefore, roasted nuts, nut butters, cooked dried fruit in tray bakes, flapjacks, cereal bars or scones are good sources of snacks.

**Smoothies and fruit juice**
If you purchase these, make sure they have been pasteurised. It is not recommended to have smoothies made at home with fresh fruit as it cannot be guaranteed that the bacteria has gone.

**Cooking**
Generally, all food should be...
Diet (cont.)

well cooked. This includes all meat, poultry, fish and eggs. Avoid any undercooked foods or anything traditionally served with uncooked ingredients such as sushi, fresh mayonnaise, homemade ice cream, dressings containing egg or any fresh pâté.

Do not reheat foods where possible and always follow standard food hygiene to store any leftovers in a refrigerator by covering with cling film or foil and using within 24 hours. Defrost frozen food in the fridge and once defrosted use at once following cooking instructions.

The one food you must not reheat is boiled rice. Cook, serve hot and eat immediately. Any leftovers must be thrown away.

Good food hygiene is important for everyone and simple things like washing your hands thoroughly before and after food preparation, before eating or after visiting the bathroom should always be followed.

Foodstuffs that interfere with TKI therapy

When being treated with TKIs, it is advised that you avoid certain fruits, particularly citrus fruits like grapefruits or Seville oranges, as well as pomegranate and star fruit as these can increase the drug levels in your blood.

For more information on how your food can interact with your treatment, you can order our Chronic Myeloid Leukaemia (CML) booklet. Alternatively, if you would like information regarding your specific TKI, check out our booklet entitled Chronic Myeloid Leukaemia (CML) – TKIs and TFR or speak to your doctor.

Remember: Mealtimes are a key time of day so try to eat with friends and family to help you maintain as normal a routine as possible. This is an enjoyable social activity and may help take your mind off your treatment for a short while, benefiting your overall health and recovery.
Side effects and their impact on diet

Nausea

If you can avoid preparing and cooking food, let someone else do it for you. The smell of food cooking can make you feel worse so try and stay away from the kitchen if possible. Also, fried, highly flavoured or fatty foods can make this seem worse so simple, chilled or cold meals are often tolerated better.

Sometimes sucking a boiled sweet, trying ginger flavoured chilled drinks or teas, nibbling a plain biscuit or drinking fizzy water can help. If you are struggling to drink, try eating fruit ice lollies or ice cubes to help you stay hydrated.

If nausea is so bad that it prevents you from eating and you are losing weight, anti-sickness medication can be prescribed alongside your treatment drugs to enable you to eat and feel better.

Poor Appetite

If you have a poor appetite, the thought of preparing and then eating a meal or snack can be difficult. Often the sight of a large amount of food is off-putting and so you can quickly find yourself in a situation where you are eating less and less every day. First of all, allow others to help if they offer. Someone else taking the decision of what to prepare means you do not have to think of it and when you get the meal you may find that you can manage a small amount after all. Also, small frequent meals are much better than three main meals a day so have little and often when possible. Don’t worry too much about healthy eating as it is better to have two puddings than no lunch at all. Also, to make sure every mouthful counts, do not routinely use low fat, low sugar foods and add extra spread, grated cheese, cream or preserves to help you increase calorie value of foods.

Constipation and Diarrhoea

Both of these bowel problems can be the result of oral medication, chemotherapy and also radiotherapy. You may be prescribed medication to help you deal with either condition alongside your CML treatment. However, remember that regular meals as part of a well-balanced diet, plenty of fruit and vegetables
Diet (cont.)

(but do not overdo) and fluids are key to managing both diarrhoea and constipation.

Also, regular physical activity, even a short walk for 10 minutes in the fresh air, will help regulate your bowels and make you feel better by being more active.

What about ‘special’ diets?

On a large population basis if scientists look at disease and dietary intake, there are links between dietary patterns in population and disease incidence. However, it is important that you do not follow any unnecessarily restrictive and limiting diets that cut out entire food groups - such as no dairy or no foods containing carbohydrates, foods that are gluten free or suddenly swapping to vegetarian or vegan diets if you previously ate animal products. There is no evidence that diet alone causes any type of cancer or that elimination of food stuffs from your diet will improve or cure a cancer over conventional medical care.

Another dietary consideration often asked about is the use of dietary supplements in the form of over the counter vitamins or minerals. There is no evidence that these components, that are normally required in very small amounts, will convey any additional benefit in the treatment of blood cancers. You may be given transfusions of red blood cells or prescriptions of iron tablets with some B-vitamins such as folic acid and these must be taken as advised.
Exercise

What are the benefits of exercise?

There is a huge amount of research and information available on the benefits of exercising with cancer. However, many people still feel unsure how this can help especially when you are going through treatment, feeling very unwell and tired.

You need to remember everyone is different. Treatment is specifically tailored to each individual and this should be the same with exercise. Side effects, coping mechanisms and general wellbeing effects are not the same for everyone and what works for one person isn’t necessarily going to work for you.

A good exercise programme can help to improve many cancer or treatment-related side effects.

Exercise is medicine, wonderful for our physical and emotional health. It is one thing you can do for yourself, which is very empowering. It can be carried out before, during and after treatment but you need to be mindful of how you are feeling.

It is possible that exercise lowers the risk of cancer relapse, although, whether it can prevent the recurrence of leukaemia, or any other type of blood cancer, is still unknown. Keeping your fitness up can reduce longer term risks such as heart disease which may be associated with some treatments.

When thinking about exercising, it is always good to speak to someone from your medical team to keep them informed as they may advise you of any precautions you need to take.

Some side effects are more common than others but, in general, if you are able to exercise at a level that is tailored to your physical needs, these side effects should not be made any worse and, in some cases, may actually improve.

Contrary to belief, exercise can help with tiredness and does not make it worse. An overemphasis on energy conservation can result in treatment-related losses.
of aerobic fitness and muscle strength. Exercise can help to decrease bone pain, improve sleep and appetite. If your weight has increased, gentle exercise may not stop you putting weight on, but it can help in reducing the amount gained.

Exercise can also have significant improvement on your wellbeing and quality of life, helping to relieve stress related to the disease and treatment.

**What sort of exercise is recommended?**

You need to ensure you choose something you enjoy which is also appropriate for your age and circumstance. There is no point doing something you have no interest in as most people find it hard enough getting motivated to exercise in the first place. If you don’t want to do it, without taking into account other factors that could come into the equation, you are likely to struggle.

Going through treatment is a worrying time but it is good to try and exercise gently. A lot will depend on your type of treatment and your current fitness level. You need to be aware of picking up infections, so gyms and swimming pools are not always the best options.

Always ensure you are sensible and listen to your body. Those who are already active should not look at increasing your fitness level if still on treatment but focus on maintaining what you have. Your fitness may drop but do not be discouraged, the fact that you are managing to still exercise is a great achievement. If you have never done exercise, it is still a good idea to try and do some gentle activity and walking or a home programme is usually the best prescription.

Walking enables you to go at your own pace and distance. The key is to keep it gentle and easy, so you can do little and often, even if it is a walk to the end of the garden.

Treatment can be tiring and some days you may struggle to get out of bed and feel completely exhausted. On these days, keep positive, rest and when you feel
better try to do a small amount of exercise. People who are exercise driven or determined to make a change need to be cautious. Pushing yourself on the days when you feel better can have a negative effect. Hard exercise could end up being too much and result in you spending the next few days completely exhausted, resulting in no exercise and leading to a negative effect. By being sensible, it means you can exercise the next day or every other day which results in you having more active days. You need to feel energised, not wiped out.

There are many variables that can determine which exercises or activities are effective and safe for your particular situation. Every day can bring new challenges and new accomplishments, so it is important to be able to modify your physical activity to suit your needs at any given time.

Things to consider:

- Exercise when energy levels are at their best.
- Customise your exercise each day to allow for any physical or psychological side effects you may be experiencing.

- A walking programme is a great way to start physical activity. If fatigue levels are high, start small and build up gradually. This could be as little as five minutes to begin with, adding on a few minutes each time.

Consult with a dietician (your doctor should be able to refer you to one, where appropriate) to address weight loss and focus on gentle strength exercises to improve muscle mass. It is important that you seek out tailored advice on what is appropriate for you as your personal experience can affect what exercises would be best.

Once treatment has finished and you are feeling better, you can start looking at increasing your activity level by basing your starting point on what you have currently been doing. If you were unable to do any activity during treatment now is the time to look at what you can do.
The guidelines for staying healthy are 150 mins of moderate aerobic activity a week plus two or more strength days. Moderate means getting slightly out of breath such as a brisk walk or cycling. You can still have a conversation, but you are aware of your increased rate of breathing and are feeling warm. This can be broken down into smaller bouts of exercise to suit you. For example, six lots of five minutes may seem more achievable to begin with, before slowly building up to the 30 minutes for one day.

Strength and resistance days are where you work your main muscle groups: legs, back, chest, hips, abdomen, shoulders and arms. Try to do this two to three times per week, ideally not on consecutive days. Improving muscular strength and stamina can help to improve your quality of life by making day to day activities easier and more enjoyable. Gaining strength can also reduce the chance of injury and empower you both physically and mentally.

While diet is often the most critical factor for weight loss, we cannot forget that the loss of lean muscle mass contributes towards this very significantly. When we do not use our muscles, we simply lose them.

For some people, this is far more than they have ever done or are capable of, so keep your goals achievable and realistic. There is no point setting the targets too high, keep them small so you are more likely to succeed, and it will be much easier, more positive and you can then look at your next step to take your fitness further.

Lifestyle plays a huge role in what we can do and there are many barriers which may need to be looked at. Not just side effects but motivation, finances, time, lifestyle, risk and fear. Exercising with a friend can help and there are lots of ways to exercise which are free, not just walking and cycling. Some local borough councils have set up exercise groups that are friendly, convenient and free of charge. Local exercise classes are a great way of meeting new people and can provide the support you need and give you some confidence.
Exercise (cont.)

If you prefer to exercise at home, there are a number of workout DVDs and home exercise programmes available.

Cancer charities and some centres often run classes specifically for those on and off treatment. This environment can feel safe and supportive for patients and far less overwhelming.

Exercise can be daunting, and walking is a good way to start. You can tailor it specifically to your needs in time, intensity and when it is convenient for you. Pedometers are a great way to increase your daily activity and can be picked up cheaply, or there are numerous phone apps which can help. It doesn’t matter how many steps you do a day, whether it’s a few hundred or more, you can build on this.

Try and aim to be consistent with your efforts by doing something every day. When energy levels are very low, perform some stretches or practice deep breathing techniques or balance movements.

Active Daily Living (ADL)

ADL is another great way to increase your fitness. These are jobs or tasks you naturally do during the day, whether at home or work, and building on them. They are ideal if you have a low fitness level or if you struggle for time as they can be built into your daily routine.

Jobs at home which get your heart rate up all count. These can include:

- Getting off the bus or tube a stop earlier
- Taking the stairs instead of the lift
- Iron a couple of items, putting them away, coming back and repeating
- Parking at the back of the supermarket car park
- Gardening

There are lots of things you can do. The aim is to get you thinking about little changes and then these little changes can then lead onto bigger ones. Progressing
slowly ensures you are being realistic and allows you to plan achievable goals which you can build on.

Before embarking on your chosen activity, you need to ensure you have an adequate warm up which can be incorporated into your exercise. It is important to start off gently and gradually increase your range of movement to get your body warmer and slowly build up your heart rate. For the cool down, bring the heart rate down slowly by gradually decreasing the intensity/speed of your exercise and not stopping immediately. If you have a slow progressive warm up, you may decide not to include stretches. However, it is really important to remember to stretch the muscles you have used after exercise. If you are unsure of which stretches to do or how to do them, speak to a physiotherapist or fitness advisor. Any areas of reduced flexibility should be worked on daily, ensuring you are gentle and work slowly without over stretching. Whilst there is no research to suggest yoga and tai chi can help prevent or treat cancer, some people find it makes them feel emotionally and physically better and can help with strength and range of movement. For more information about these complementary therapies and others, go to page 75.

Once you have established your exercise routine, ensure you build on it gradually. Increase the time before intensity until you are doing at least 20-30 minutes of moderate intensity exercise a few times a week before making it harder. Strength work which involves repetitions should be done moderately, working towards three sets of 12-15 repetitions (resting between each set). Ensure you can complete one set before continuing to two and then three sets and only then increase the weight.

Exercise is important for those receiving palliative care. Systematic reviews have shown exercise is feasible with important benefits in physical function, symptoms and quality of life. There can be improvement in fitness, functional ability, emotional wellbeing as well as symptoms such as fatigue,
Exercise (cont.)

dyspnea and anorexia. Quality of life can be helped by keeping you independent and able to carry out everyday tasks which are important to you. Depending where you are on your pathway, there are different options available. Keep in mind what would work best for you and speak to the team around you who can give you the help and support needed.

Once you have decided what you would like to do, you need to ensure you stay safe. If you are going through treatment and experience extreme fatigue, have a low blood count or are anaemic, you should not exercise. If you are unsure, your medical team will be able to advise you on this. After treatment, if you are experiencing extreme fatigue, balance problems, aching, heaviness, swelling, dizziness, shortness of breath for no reason and/or sharp pain in a specific area, you should not exercise and seek help from your GP/medical team.
Fatigue

Fatigue is one of the most common consequences of cancer and related treatments. Its cause, however, is not yet fully understood. Fatigue is thought to affect the majority of people who have had a cancer episode and may be defined as:

"A distressing, persistent, subjective sense of tiredness or exhaustion related to cancer or cancer treatment that is not proportional to recent activity and interferes with usual functioning."

How long does it last?
There are no hard and fast rules for how long fatigue will last. Many CML patients expect to feel fatigued during treatment with TKIs as they can exacerbate fatigue. However, as diagnosis and treatments improve we are beginning to learn that people living with and beyond cancer may experience significant fatigue which interferes with day to day life for several months and, in some instances, for years. The good news is that, over time, fatigue may subside, and a lot of people get back to their normal energy levels once blood counts have become controlled.

How can fatigue affect your daily life?
Your fatigue can negatively impact on all aspects of your day to day life. This may include disruption to your usual and taken-for-granted activities such as personal care, including showering and dressing; leisure and social activities, such as keeping fit or going out with family and friends; and others may experience disruption to
work and educational attendance and performance. For people who are working and trying to manage with compromised energy levels, there may be a tendency to use most of their available energy for work purposes. By the end of your working week, you may find there is little left in your ‘battery’ for leisure activities and socialising.

Being unable to do everything you need, want and have to do can prevent you from being in contact with other people as much as you would like. This can understandably lead to feelings of social isolation and sadness.

Additionally, people may experience a sense of ‘loss of former self’ akin to feelings of bereavement and might grieve for the normal life they seem to have left behind. Some patients also worry about the cognitive impact of fatigue which may lead to what some people describe as ‘brain fog’ or ‘chemo brain’. This commonly includes reduced concentration, memory issues and communication difficulties, all of which can be scary.

You may also have worries over the expectations others may have of you in terms of pre-illness roles, responsibilities and involvement in day to day activities and events. As fatigue is not always obvious to others, it might be that friends, family, or employers do not seem to understand the impact of fatigue on a person’s life or how best to support them. This may cause you to experience some feelings of frustration, anger or guilt if you are currently unable to do everything you normally do, or what others expect from you. For some, feelings of anxiety and a sense of sadness may lead to distress and low mood which can be difficult to cope with alone.

Relationships of an intimate nature may also be affected by the presence of fatigue and you may feel too tired to participate in sexual intercourse. Additionally, your partner may be cautious to do so and worry that they may make your fatigue symptoms worse. For more information about the impact a CML diagnosis can have on your relationship, go to page 16.

All of these affects are commonly reported by people living with and beyond cancer. However, there are strategies that you can employ to help you better manage these
Fatigue (cont.)

and feel more in control of your
day to day activities, roles and
responsibilities.

Steps to combat fatigue

The first step in self-managing
your fatigue symptoms is
acknowledging that you have
fatigue. Secondly, being self-
compassionate and not expecting
yourself to perform all the
activities you did before you
became ill will help if fatigue is an
issue.

Congratulate yourself on what you
do achieve rather than punishing
yourself for what you don’t,
regardless of how little it seems,
and do not compare yourself with
others or past performance.

If you accept that your available
energy is not currently what it
used to be, you can then begin to
think about how much energy you
have in your ‘battery’, how you are
using this and whether you would
like to use it differently. This
self-management approach will
hopefully help you to regain some
control over your daily activities,
roles and responsibilities.

Research evidence demonstrates
that activity management and
energy conservation may help.

Here are some top tips for
regaining control of your day to
day life:

• Identify activities, both physical
and cognitive, that drain your
battery/energy the most and
make a note of these.

• Identify activities that make you
feel you have topped up your
battery/energy levels. These
may include physical activity,
such as a paced walk, and do
not necessarily mean resting or
sleep. Make a note of these.

• Use a diary or calendar as a
reminder to note down what
drains your energy and what
tops you up.

• Pace the most draining
activities and use top up
activities to replenish your
battery in between. This may
help you get through to the end
of the day easier.

• Prioritise which activities you
need, want and have to do and
decide whether they really all
need to be done on the same
day. Avoid overdoing things
when you are having a good day.
• Plan activities in a diary and pace them across the week if possible, to spread out energy expenditure. Plan outings and events and incorporate top up points during the day.

• Getting back to, or slowly introducing, some form of physical activity into your daily routines may help to reduce cancer-related fatigue. There is also some research evidence to show this may also help you to return to pre-cancer levels of sexual activity. It doesn’t really matter what type of physical activity it is, so it is your opportunity to choose something you would enjoy. It may be through attending a gym or yoga class, or pursuing something less traditionally considered as physical activity, such as dancing, which has been shown in some research to reduce fatigue. For more information about exercise, go to page 40.

• Communicate your needs and share your knowledge of CML in a calm and articulate way with all those around you including family, friends, employers, colleagues and health professionals. Some people will need to be educated by you about the impact of fatigue on patients with cancer, as they are unfamiliar with this. If you would like more advice about telling others about your CML diagnosis, go to page 66.

• If you are struggling emotionally, try learning a relaxation exercise, mindfulness technique and/or seek medical help from your GP to discuss other support. This may be in the form of counselling or ‘talking therapies’ and/or include temporarily taking medications to raise your mood and reduce anxiety.

• Eat a well-balanced diet and have nutritious snacks throughout the day to top up your energy levels. If you would like more guidance about how to develop a healthy diet, go to page 32.

• Adopt a relaxing ‘wind down’ regime an hour or two before you go to bed at night in order to promote relaxation and aid restful sleep. Sleep hygiene strategies may be helpful and enjoyable.
Fatigue (cont.)

• Contact an organisation such as Relate [www.relate.org.uk](http://www.relate.org.uk) if you are struggling with communicating your needs to your partner, or intimate relationships.

• Access available information and local support groups and online forums (see [www.leukaemiacare.org.uk](http://www.leukaemiacare.org.uk)) as this may help reduce feelings of social isolation.

• If you have concerns linked to employment issues contact your local Citizens Advice Bureau or access the Advisory, Conciliation and Arbitration Service (ACAS) website [www.acas.org.uk](http://www.acas.org.uk). Reasonable Adjustments to your workplace and/or work role under the Equality Act 2010, can be explored to make things easier for you and your employer. More information about this can be found in the work and finances chapter of this booklet, starting on page 54.

• If you are a student, discuss the impact of fatigue with student disability services and request support from your organisation, such as extensions to hand-in dates for assignments if needed.

Remember the fatigue you are currently experiencing may well go away over time. In the meantime, self-managing your fatigue will help you feel more in control and add to your sense of wellbeing.
I’m still working - what are my rights?

A cancer diagnosis at any stage of your life is hard to digest and how you approach your diagnosis and treatment is very personal. If the diagnosis does occur while you are working, it can feel very daunting and scary. Some of the questions you may be concerned about are:

- How will I cope with treatment and work?
- How will the treatment affect my ability to complete my tasks at work?
- How much time will I need off for treatment and/or recovery time?
- What do I do if I need time off for treatment or medical appointments?
- Will I be assessed differently at work due to my cancer treatment?
- Will I be looked over for promotions because of the length of time I took off work for treatment or recovery from treatment?
- Will my bonus be impacted?
- How will my colleagues react to my cancer diagnosis and treatment?
- How will I come back to work after treatment?
- What if I feel I cannot come back to work after treatment?

This is by no means an exhaustive list of questions and, depending on your job, there may be many more.

Deciding to continue working during your treatment is a very personal decision and should be discussed with your manager or Human Resources (HR) department, so they can best assist you during this time.

Treatment plans and the side effects are as unique and complex as the individual undergoing treatment. Some people have little or no side effects and they find that continuing with their normal day to day life is the best way of dealing with their diagnosis.

Other people have severe side effects and can only continue with flexible working arrangements or cannot work at all. These
decisions can only be made once you have started treatment.

Although you may have decided to continue working for the time being, it does not mean that you cannot change your mind later and request more flexible working arrangements from your employer as a reasonable adjustment. You should note that your employer is required to assist you with reasonable adjustments while you are undergoing treatment and thereafter.

You may not be discriminated against for continuing to work but, if your employer does begin to treat you less favourably as a result of you having cancer at present or in the past, that is discrimination and is in contravention of the Equality Act 2010.

During treatment, your employer should allow you as much time off as you require to attend medical appointments or undergo treatment to aid your recovery.

Your employer should also allow you reasonable sick leave after an operation or invasive medical procedure. The time off required should be specified by your medical practitioner and communicated to your employer.

Once you are back at work, your manager or HR should keep in touch with you regularly to determine if you require any assistance with either your job, workload or if there is any other way that they can support you.

Going through cancer treatment can be very challenging but your rights at work are protected under the Equality Act. Try to be open and honest with your employer about your treatment plan and your needs, to enable you to negotiate your treatment and recovery as best as you can.

Not all employers are aware of their responsibilities and your rights under the Equality Act. It is important to make sure you are aware of your rights both as they relate to the Equality Act and your company procedures. If you encounter difficulties at work, speak to your manager and HR team first, in order to see if a solution can be obtained. If you are still not satisfied with the outcome, seek legal assistance from either a voluntary organisation, Citizens Advice Bureau (CAB) or an employment
The Equality Act and Cancer Patients

In England, Scotland and Wales, the Equality Act 2010 as well as the Disability Discrimination Act 1995 (DDA) protects employees. In Northern Ireland the extension, the Disability Discrimination Order of 2006, protects employees.

Under these acts, it is unlawful to treat an employee less favourably because of their disability. Cancer is legally classed as a disability and will also include a person who had cancer in the past, has gone through treatment and is now in remission. The employer cannot treat the employee less favourably because of his/her past cancer diagnosis.

The Equality Act and DDA, covers all areas of employment, including but not limited to:

- Recruitment
- The employee’s terms and conditions
- Benefits and bonuses
- Opportunities for promotion
- Training

The acts will even protect an employee when they no longer work for the employer.

In order for you to benefit from the protection of these acts, your employer must be aware of your diagnosis, except for Indirect Discrimination. It is advisable to discuss your diagnosis with either your Human Resources (HR) Department or your manager, as soon as possible, so that you can determine the best way forward.

There is no legal requirement for you to inform your employer of your diagnosis, but if they are not aware of your diagnosis, they cannot assist you during your treatment and/or recovery period. Regular absences or low performance when completing your tasks could be considered performance-related issues and will not be attributed to your treatment or period of recovery.

The Equality Act protects employees against Direct Discrimination; Indirect Discrimination; Harassment and Victimisation.

Discrimination can occur in various ways and may even be well intentioned. Some of
the following are examples of discrimination that you may encounter:

- You are not promoted or offered the same opportunities because of your cancer.
- Your employer does not make reasonable adjustments. These are suggested workplace adjustments that could assist you during your treatment and/or recovery.
- Giving you a warning because of persistent sick leave and not taking your cancer into consideration.
- Your employer may suggest that you do another role in the business which is a demotion from your current role and does so because you have cancer and without your consent.
- Your employer gives you a poor end of year appraisal and does not take your cancer into consideration.
- Your employer may give you a reduced or pro-rated bonus.
- Your employer may suggest that you resign from your employment or retire because you have cancer.
- Your employer may select you for redundancy or for reduced hours working because of your cancer.
- Not being offered a job because you have or have had cancer in the past.

**Direct Discrimination**

Direct Discrimination is when, due to your cancer diagnosis, you are treated less favourably than someone who does not have cancer. This can include not being offered a job or not promoted because you have or have had cancer.

**Indirect Discrimination**

Indirect Discrimination is when there is a "rule, policy or practice which applies to everyone" but it puts you at a disadvantage compared to the other employees who do not have the same disability. You can only challenge a practice, policy or rule which you think is indirectly discriminatory if it affects you personally.

For example, if your employer wants to make redundancies using the amount of sick leave...
taken as a selection criterion, that is indirectly discriminatory. An employee with cancer could be discriminated against because they have had to take time off work due to treatment and will be at a clear disadvantage compared with a person who has not had cancer. In these circumstances, you would need to make your employer aware that the policy is discriminatory because, although neutral in nature, it disadvantages you and others with cancer. With Indirect Discrimination, an employer may not be deliberately discriminating against anyone, but it is still discrimination and not knowing you have cancer is no defense.

It should be noted, Indirect Discrimination may be permitted if the employer can show that the aim of the policy is for a good reason or can be justified, that the aim of the organisation is fair, balanced and reasonable and that reasonable adjustments have been considered.

**Reasonable adjustments**

The Equality Act 2010 and the DDA, requires your employer to make reasonable adjustments.

Your employer is not required to make reasonable adjustments if they are unaware or have not been told that you have cancer, or they could not have reasonably known that you have cancer. However, "should have reasonably known," will be difficult for your employer to dispute once they start to see physical changes occurring due to your cancer treatment. For example, if your hair is falling out; you are looking ill; you are unable to do tasks that you previously could have done, or you are taking extended and/or frequent periods of sick leave.

Being required to make reasonable adjustments means that your employer is required to assist you to manage your work with either a change in how your employment is structured, by removing physical barriers, or by providing additional support.

There is no definitive description of what a reasonable adjustment is, but the following will be taken into consideration:

- The size of your employer
- How much will the adjustment
• How much will the adjustment benefit you
• Whether making the adjustment will affect the employer’s business service or financial situation

Here are some examples of reasonable adjustments:
• Having time off to go to medical appointments.
• Having time off to recover from treatment.
• Allowing you to tailor your hours to suit your needs. This is called a phased return to work and is very often used to ease employees back into work, after a period off sick.
• Allowing you to undertake lighter duties during or after treatment or allowing you more periods of rest.
• Changing your job description to remove tasks that could prove physically difficult or stressful. This would be done with your full co-operation and agreement.
• Sharing some of your tasks with another employee.
• Requesting physical assistance – changing the way your work station is set up.
• Requesting props to assist you at work, like a back rest or a cushion.
• Making sure that you can access your place of work – including adding a ramp, lift or hand rails to name a few.
• Requesting that your work location is adjusted – moving to a lower floor or working from home.

Reasonable adjustments are not easily or formally defined. They are usually quite specific to you and the assistance that you require during treatment or thereafter. Whether it is a change in your duties or hours, counselling, coaching or something more physical, always speak to your employer to determine whether they can assist you. Your employer is required to try and implement reasonable adjustments and cannot merely refuse. Reasonable adjustments can be either temporary or permanent. It is more difficult for an employer to
refuse a temporary adjustment and it is a good starting point when asking your employer to implement reasonable adjustments.

If you encounter difficulties having your reasonable adjustments agreed to, it is important to think about seeking legal assistance. There are also a number of free services available that can assist you, including the Disability Law Service or Working with Cancer.

**Victimisation**

Victimisation is when you, as a disabled person, are treated less favourably than another employee, for asserting your rights as a disabled person.

An example of this would be if you were to request time off for treatment and your manager does not grant your request. You decide to take the matter up with your HR department. They agree that you are entitled to the time off and instruct your manager to grant the time off. Your manager is now so annoyed that you went over their head to HR that they do not put you forward for training and you are given a negative appraisal.

The law will also protect your colleagues from Victimisation if they support you in making the complaint.

**Harassment**

Harassment is where certain behaviour is meant to or has the effect of either: violating your dignity, or creating an intimidating, hostile, degrading, humiliating or offensive environment.

Harassment would also include bullying in the workplace. According to the Workplace Bullying Institute, bullying at work means "repeated, health-harming mistreatment of a person by one or more perpetrators. It is abusive conduct that is: threatening, humiliating, or intimidating, or work interference or sabotage which prevents work from getting done, or verbal abuse."

**What do I do if I encounter discrimination in my workplace?**

If you feel that you have been discriminated against at work, speak to your manager or HR Department and see if the issue
can be resolved. Quite often issues arise due to miscommunication or the employer not being fully aware of your needs and/or your situation. Try to speak openly and update them about your situation throughout your treatment and after your return to work.

If you are unable to speak to your manager or HR department, speak to your Union if you belong to one or ask for help from an Occupational Health Adviser, which your employer can arrange for you. If you still feel like issues are not being addressed, seek assistance from the Citizens Advice Bureau or organisations like Working with Cancer or Macmillan Cancer Support which deal specifically with providing advice about work to cancer patients.

**What should I do?**

- Have a look at the relevant company procedures and policies. Your manager or HR department should be able to provide you with a copy.

- Approach your employer with solutions and suggestions. This will show that you are committed to your job and invested in resolving the situation.

- Make sure you know and understand your legal rights.

If the matter still stays unresolved despite you approaching your employer and speaking openly about it, you may be required to lodge a formal grievance.

Your employer will have their own grievance procedure, either in their employer handbook or stated in your contract of employment. If you are unable to find this, ask your manager or HR department for a copy.

To raise a formal grievance, you will need to write a letter to your manager detailing the incidents which have led you to believe that you have been discriminated against. It is suggested that you be specific and detailed when drafting this letter. Your employer will be required to investigate your grievance and deal with it fairly.

If you are not satisfied with how your employer is dealing with your grievance and you believe that it is not being dealt with fairly, or your employer is being
unreasonable, you can bring a complaint to an Employment Tribunal.

Before you bring a claim, you will have to notify ACAS (Advisory, Conciliation and Arbitration Service) of your intention to lodge a claim. ACAS will try and sort out disputes between the parties before the matter goes to an Employment Tribunal.

More information on how to lodge a claim with an Employment Tribunal can be found here: www.gov.uk/employment-tribunals

Alternatively, you should seek advice from an Employment Solicitor.

I’m unable to work: what benefits am I entitled to as a cancer patient?

There may be various benefits available to you, all of which will depend on your individual circumstances – there is no benefit for which you automatically qualify solely based on your cancer diagnosis.

Statutory Sick Pay
If you are an employee taking sick leave, then you are likely to be entitled to Statutory Sick Pay from your employer. You can have 28 weeks (about 6 months) of SSP before it runs out. This is usually straightforward to ‘claim’ because it is arranged and paid by your employer like your normal pay. If you (and your partner’s) income is low, then you might be able to top up your income with a benefit called Income Support.

Employment and Support Allowance
This benefit provides financial help to people who are unable to work following completion of the Statutory Sick Pay period (28 weeks). You may also be eligible if you can’t claim SSP because you’re self-employed, unemployed but not in receipt of Jobseeker’s allowance, you earn less that the National Insurance contribution threshold or are under State Pension age. ESA also provides personalised support to those who are able to work.

There are three different types of ESA, each with different eligibility criteria:

1. ‘New style’ ESA - to claim this you will need to have been
employed or self-employed and paid National Insurance contributions, usually within the last two to three years, and now have an illness or disability that affects your ability to work.

2. Contribution-based ESA – this follows the same type of criteria as ‘new style ESA’, but you also have to be in receipt of the severe disability premium or you should have received it within the last month and still be eligible for it.

3. Income-based ESA – this type of ESA is not related to your National Insurance contributions.

For 13 weeks after your claim you will receive the assessment rate and, if you are entitled to ESA, you will be placed in either the work-related activity group (up to £73.10 a week) or the support group (up to £111.65).

Tax Credits
Depending on your circumstances, a reduced income might entitle you to Working Tax Credit or Child Tax Credit or both. The eligibility criteria for Tax Credits can be complicated and will look at how many hours you’re working, how long you have been on sick leave, whether you are paying for childcare, and much more. Working Tax Credit is paid for people in full-time work and you can be treated as being in full-time work for the first 28 weeks of sickness if you were working sufficient hours beforehand. You have to be responsible for a child in order to get Child Tax Credit.

Personal Independence Payment
If you have care or mobility needs and you expect these needs to continue for another nine months or more, then you might be eligible for PIP. There is a fast-track system with guaranteed eligibility for the enhanced daily living component of PIP for claimants with a terminal illness who have a ‘DS1500 form’ from their doctor or specialist nurse. PIP is not means-tested, so you can claim it regardless of your employment status, income, and savings.

Housing Benefit and Council Tax Support
These two benefits can help with rent and council tax bills. You
cannot claim Housing Benefit for mortgage payments. These are both means-tested and will take account of your circumstances to determine how much help you are entitled to, such as non-dependents living with you and how many bedrooms you have. Housing Benefit is unlikely to cover the full cost of a privately rented property.

**Universal Credit**

This is a new benefit rolling out across the country which replaces Income-related ESA, Income-based JSA, Income Support, Housing Benefit, and Tax Credits and unifies them into one benefit. In some areas of the country, you must claim Universal Credit instead of the benefits it replaces. For sick or disabled people, Universal Credit will usually mean less weekly income than the benefits it replaces. If you live in an area that only offers and accepts new claims for Universal Credit, but you still receive the old benefits, you should seek specialist advice before making any new claims – it could trigger a switch to Universal Credit and leave you worse off.

This is only a brief overview of benefit entitlement for people too ill to work. There are many other benefits available, as well as further eligibility criteria for the benefits that are listed above. It is well worth seeking help from specialist advisors at Citizens Advice, Macmillan, or other benefits advice agencies locally. More information about benefits can be found at [www.gov.uk](http://www.gov.uk).

**Can those caring for me claim any benefits?**

If someone is caring for you then they should be able to claim Carer’s Allowance as long as:

- They spend at least 35 hours per week caring for you
- They don’t earn more than £116 per week
- They are over 16 and not a full-time student

In addition, they have to be caring for someone who receives a disability-related benefit such as Personal Independence Payment or Attendance Allowance. If the person being cared for does not receive such a benefit, then the carer cannot claim Carer’s Allowance.
If the carer does meet all the criteria, then they can claim Carer’s Allowance of (currently) £62.70 per week. If they have a low income, then they might be able to have their income topped up by another benefit called Income Support.

It is feasible that you could face a situation where your carer has to stop work to care for you but cannot claim Carer’s Allowance because you are not in receipt of Personal Independence Payment.

As a PIP claim can take months to reach an award, and often are unsuccessful, it could mean a long wait without any income for the carer. If your carer is on a low income they could be entitled to Income Support whilst you are waiting for your PIP or Attendance Allowance claim to be decided. However, they will not get paid beyond 26 weeks if your claim takes longer than that to decide. If you are living on your own and receiving a disability-related benefit in addition to other benefits, such as Employment and Support Allowance or Pension Credit, then seek advice before allowing a carer to claim Carer’s Allowance. Their claim could reduce your weekly income if you are receiving a severe disability premium.

If the person caring for you already receives state benefits or a State Pension, then they might not be able to receive any Carer’s Allowance payments. This is because other benefits ‘overlap’ Carer’s Allowance. Even if this is the case, the carer should still apply for Carer’s Allowance. Although they will not receive any payments directly, they should be awarded an ‘underlying entitlement to Carer’s Allowance’ that can be beneficial to them when calculating other benefits and credits them with National Insurance contributions.

In some circumstances, a carer under pension age can claim Carer’s Credit even if they aren’t entitled to Carer’s Allowance. This won’t give the carer any income, but it will credit them with National Insurance credits towards their State Pension.
Telling others about CML

Your relationships are unique to you and so there is no one way to tell others about CML. Our relationships can be a great source of support when we are ill and can bring people closer together. Relationships are often complicated, and the challenge of illness can open up difficulties as well. It generally helps when communication is as open and honest as possible. Here is some guidance to help you think about what feels right for you.

**Should I tell others about the CML?**

When you are first diagnosed, there is a lot of information to take in and you may be feeling shocked and overwhelmed by different emotions. You may need to give yourself some time to think about this enormous thing that is happening to you before you decide when and how you tell others about it. People may be anxious to know what is happening and this can make you feel under pressure to pass on information before you are ready. Let them know that you need some time to process the information yourself before you are ready to talk about it in more detail.

There will be some people that you want to know and others that you would prefer not to. It is up to you who you tell and how much you tell them but going through treatment for CML is not something that you can hide from the people closest to you. You may be concerned about how family and friends will react and be tempted to protect them from upset and worry by not telling them. It is usually harder for them to manage when they sense that something is wrong and try to guess what it is. Take some time to plan how you will tell children and elderly parents. It may help to sit down with someone else first and rehearse what you are going to say. Hearing yourself say the words out loud first will help you to feel calm when you come to talking about it.
How to tell a partner

Your partner may have been with you when you heard about the CML. If not, they will most likely want to know as soon as possible to be able to support you. It can be hard talking to them about it because they will have their own concerns and emotions and they will also need time to adjust to the shock of the diagnosis.

As you adjust to life with CML, your partner and family will be affected in their own way. They will also need the support of family and friends. They can also access support for themselves such as counselling and support for children through helplines and cancer support centres. Ask your healthcare team about what is available locally.

It can be helpful to have your partner with you for hospital appointments so that you are both able to hear the information being given and have the opportunity to ask questions. We often have different needs for information. Perhaps one prefers to have lots of information and ask lots of questions and the other may prefer to know less or to quietly think things through. One may like to talk about feelings and the other not. You may want to try and protect each other from upset or how you feel, and this can get in the way of talking to each other when you most need to. Finding ways to manage your different needs can be a challenge throughout your experience of CML.

Some couples describe having to go through the challenge of illness as an experience that brings them closer together. For others, the stresses can put a strain on the relationship and it can become difficult to find your way together. The more open and honest you are able to be with each other, the more you will be able to work together as a team to get through it.

When you are going through some treatments, it can be hard to find time for yourselves as a couple. When you are feeling relatively well, you may want to
set aside some time to catch up and give each other a chance to say how you are. Identify what is helping each of you get through this time and what is making it difficult. This could be on both an emotional and a practical level. It is also helpful to set aside some time when you can be a couple and do something "normal" like watch TV, have a trip out together or decide to have a CML-free day where you talk about other things.

**How to tell a friend or family member**

Telling people that you have a rare condition like CML can be hard to explain. Keep it simple and explain that CML is a type of blood cancer. It means that the bone marrow does not function properly, and this affects the blood cells it produces. The treatment will aim to get the CML under control.

If they have more questions, you may find it easier to direct them to information leaflets or our website for more information than to try and explain it all yourself.

People will have their own ideas and possible misunderstandings and it can create more uncertainty for you if you cannot answer all of their questions.

It is exhausting to have to keep repeating the same difficult conversation and you could ask someone else to tell people who are not so close to you. It can be easier over the phone or by letter or email as this gives you more control.

People will often want to do what they can to help and support you. Think about practical things that will help such as lifts to and from hospital appointments and help with shopping, housework or childcare. As treatment continues, you will have a better idea about what you need and what works for you.

Your family and friends will be keen to know how you are and for updates about your progress. Keeping in touch can be hard work when you are not feeling well, and you may want to consider having someone else as a point of contact to pass
Telling others about CML (cont.)

on information. This saves you from having to repeat similar conversations. You may not feel up for long conversations when you are feeling unwell but keeping in touch through texts can help you feel connected to others.

Some people send out a group email or use social media to pass on news. This can make it quick and easy to contact a lot of people at once. However, do consider who you want to see your posts, check your privacy settings and remember that information stays on the internet unless you remove it.

Coping with other people’s emotional reactions

You can be sure that there will be a wide range of reactions from people you talk to about your illness. Hopefully, they will be supportive and open to following your lead about what will be most helpful to you. Some may be upset and distressed by your news and you may find yourself needing to be strong and supportive for them. This is hard when you’re not feeling well yourself. People who care will want to help you as best they can, but they may first need time to adjust.

You may find that not everyone is sensitive to your needs and this can be upsetting and frustrating. If some people avoid you, it is usually because they don’t know what to say or are worried about saying the wrong thing. Often people make assumptions and say what they think is helpful. They might say that you look well, recount stories of others they know with a similar diagnosis or even a completely different type of cancer. They may encourage you to stay positive and to look ahead. They may have advice about particular diets or supplements you can take. This may not reflect how you are feeling or be what you want to hear. Where possible, try to let people know what you find helpful and unhelpful. In time, you will discover who is better at giving you the practical and emotional support that you need.

See the people that you want to
see and talk about what you feel comfortable with. Ideally, people will give you the opportunity to talk about how you are feeling if that is what you want or respect your wishes if you prefer not to talk about it. Some of us are better listeners than others. It is fine to change the subject if you don’t want to talk about something. You could say that you are giving yourself a break and having a day without talking about your illness. Asking others about themselves and what they are doing can steer the conversation away from illness.

Going through CML is an enormous challenge and you and those close to you are likely to feel stressed at times. It’s natural to feel anxious, angry or frightened when faced with CML and it is often those closest to us who may feel the brunt of these strong emotions. We often take out our feelings on those close to us. This can put a strain on relationships. The more open and honest we can be during these times, the easier it is to work through any difficulties or misunderstandings.

**Telling children or vulnerable people**

Our instinct is to protect children from upset and you may be tempted to put off telling them about your diagnosis. You may worry about how they will cope and how you will support them, especially when you have so much to deal with yourself. We know from research and the experience of others that children cope better when they are told about an illness. Children are very good at sensing when something is wrong, and it can be more distressing for them to not know what this is. Not telling them can leave them feeling frightened when they don’t understand what is happening. Talking about the illness with openness and honesty sets the tone for the way ahead.

Telling your children means that you can be more open about what you say at home and not have to worry about them hearing something at school or from someone else. It gives them the opportunity to ask questions and
to say how they are feeling. It can help you all to feel closer and to work through things together.

Telling children about the CML in a way that they can understand will vary according to their age. If you have more than one child, telling them together or as closely together as possible will avoid one carrying the burden of knowing or wondering why they were told last.

You know your child best and so trust your instincts when it comes to supporting them. You might like to think about what you will say in advance and choose a time to avoid interruptions. Try not to put pressure on yourself to have the perfect conversation. The aim is to convey enough information to begin an on-going process with many more opportunities to talk about what is going on. You may be surprised by their questions or reactions. It is fine to say if you can’t answer all of their questions but can try to find out.

Younger children will respond to small chunks of information at a time and may be keen for reassurance about how it will impact on them. Let them know that they have not caused the CML and that they cannot catch it from you. They may carry on playing as if they haven’t understood but they will be processing the information in their own way. They may then surprise you with a very direct question when you least expect it.

Teenagers are likely to want more information but will also need time to work through their feelings. They might be reluctant to talk about it with you and may want to spend time alone or will seek support from their friends or other family members. Letting your child’s school know will mean that they can also offer support.

As time goes on, help them to be prepared for any changes in your appearance such as hair loss and weight changes. Explain that fatigue may make it harder for you to do the same activities with them. If they are able to visit you in hospital, explain what they can expect to see. Playing with hospital toy sets, books and imaginary games can
help younger children express themselves and feel connected to what is happening. Reassure them that their routine and activities will stay the same as much as possible. Explain that family and friends may be more involved in activities such as picking them up from school and be clear about what the plan for them is. Maintaining boundaries and limits for all ages will help them to feel contained and more able to know where they stand.

You may have vulnerable people in your life, perhaps a frail parent or relative or friend who is unwell that you would like to tell. Again, plan how you will tell them as appropriate to their circumstances and explain that you may not be able to visit them as frequently whilst you are going through treatment. They may want to feel that they too are able to support you by thinking of you.

Our easy read document, All About Leukaemia, may be a useful resource to help tell your loved ones about your diagnosis. To order a copy, call the helpline on 08088 010 444

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**Telling work about the diagnosis**

Being diagnosed with CML will mean that you need to start treatment straightaway or in a matter of weeks if it has been discovered very early. You or someone else will need to let your employer know about your situation. Your medical team will be able to advise you about timescales. You may need to negotiate a reduction in working hours or arrange with your employer for times when you are in hospital or for when you are not well enough to go into work.

Your consultant or your GP can arrange letters to your employer to confirm your diagnosis and the
Telling others about CML (cont.)

effects it may have on your work life. It is often worth taking time to explain CML to your employer, as it is likely they will never have heard of the disease.

Hopefully, your employer will be supportive. You may worry that you could be given the sack or made redundant if you cannot work. This should not happen because people with any form of cancer are covered in law by the Equality Act. This means that legally your employer cannot discriminate against you and must make reasonable arrangements relating to your disease.

You may also be wondering about what you tell your colleagues. You may want to keep your diagnosis to yourself and it is important that your privacy is respected. Telling those you are closest to gives them a chance to support you and may help you feel connected to work. It can also make it easier when you do go into work if they know that you are at risk of infection or if fatigue or poor concentration is a problem.

Returning to work after your initial diagnosis or after the initial side effects of your treatment have subsided can feel like another big challenge. You may wonder if people will want to talk about it too much or not at all. If some people avoid you, it is usually because they don’t know what to say or are worried about saying the wrong thing. It can help if you or your colleagues let people know if you do or don’t want to talk about it. Plan beforehand what you might say if someone asks how you are. It is up to you what you say and in how much detail. At other times, it can seem that people have forgotten what you are going through and you need to gently remind them that you are still coping with side effects and that your recovery is ongoing. As with all of your conversations about CML, being honest and saying what you need to say is generally the most helpful approach and makes it easier for you both.
Complementary therapies have been used in a supportive capacity both within the NHS and by charities for the enhanced wellbeing of patients and clients within haematology for over 30 years. Aromatherapy and reflexology are generally the most popular and available within the NHS due to their effectiveness; assisting with overall wellbeing and relaxation, as well as addressing pain, nausea and other side effects from treatment and medication as part of a package of care. Consultants have commented that the more relaxed the body and mind is, the better accepting of their treatment it is, even taking bloods.

In the next section, there is some information about a variety of complementary therapies and exercise that may be helpful at different stages; before, during and after your treatment.

The guidance is to enable you to have an overview of the benefits of different therapies and also includes contacts and information to enable you to find an appropriate therapist privately.

During your treatment and hospital visits you may be offered a choice of treatments from a complementary therapy team; this service would usually be free and provided by experienced therapists who are accustomed to CML and the treatment you are receiving.

**Complementary Therapist**

A complementary therapist will offer supportive therapy as part of an integrated practice, alongside standard medical treatment for CML, not an alternative treatment.

When looking for a therapist it is important to ensure your therapist has appropriate training for your needs. The therapist will generally be a member of a representing body for their therapies and must have the appropriate insurance.

The Complimentary and Natural Healthcare Council (CNHC) is a representing body for yourself as the client. All therapists on this
register will have a certain level of qualifications and insurance. Most NHS therapists will be members. When approaching a therapist do check that they have taken part in some additional training or continued professional development (CPD) or have experience with haematology-oncology clients and understand the appropriate guidelines that apply.

When treating patients with CML, all hands-on massage therapists are advised to work with light touch techniques and massage due to the possibility of low platelet counts and the risk of bruising. This doesn’t affect the quality of treatment but ensures you receive massage or reflexology with the appropriate lightness of touch, and therefore the best outcomes.

Before starting on a course of treatments the therapist should take a detailed case study on your overall health and wellbeing in order to tailor the treatment to you, accommodating your needs.

**Types of complementary and supportive therapies**

**Acupuncture**

Acupuncture is a form of Traditional Chinese Medicine (TCM). This treatment uses the application of fine needles in points in the body which have been identified as helpful for various symptoms and side effects that are present at the time. Many people find acupuncture very relaxing and useful for health problems such as fatigue, sleep problems, nausea and pain. It can also support emotional wellbeing.

Health professionals and doctors also train in a method of acupuncture called Western Medical Acupuncture which is often used as part of an integrated service. NICE guidelines recommend acupuncture for headaches, whilst it has also been reported as beneficial for nausea and vomiting. The WMA approach is different to TCM, but they are used for similar concerns.
For more information on the two approaches – including how they differ, and which may be more suitable for you – speak to a professional acupuncturist or your medical team.

Aromatherapy

Aromatherapy is the use of essential oils which are aromatic volatile oils extracted from plants. Aromatherapy as a therapy is most often used with massage but can be used in bath and skin preparations, inhalation, compresses and other methods. The chemical constituents or make-up of each oil assists the aromatherapist in choosing the most appropriate oil or combination of essential oils for their client’s needs. These therapeutic properties of essential oils may assist with psychological, emotional and physical concerns and, importantly, relaxation. Clinical aromatherapists would also choose essential oils to assist with side effects of standard medical treatments which might include nausea, poor sleep and pain.

The use of touch and essential oils are documented with positive results. For example, lavender has been found to be beneficial for stress and anxiety due to the amount of the chemical esters in this oil which are known to be calming and soothing. When buying essential oils for home use, good quality oils should be used especially when they have contact with the skin. Essential oils should always be diluted, the exceptions being lavender and tea tree which are generally acknowledged as being suitable to apply directly to the skin. However, if you are unsure, a patch test should be carried out with the oils first. Oils used in an aromatherapy fan or ceramic burner are using the olfactory senses and so, the aroma must be pleasurable. Only a few drops are needed, and, if using in the bath, always follow guidelines. If you have nothing to hand, the fat content of a couple of tablespoons of milk will disperse the oil into water. Coconut oil is also suitable.

For a massage, the aromatherapist will blend a dilution of essential oil in a
vegetable oil, often known as the carrier oil. A very weak dilution, usually of around one percent, would be made up for those with CML who have undergone treatment, or those who are less well. An aromatherapy massage should be light touch and leave you with a feeling of deep relaxation. Relaxation is an important part of recovery. Massage can be full body or directed to specific areas of discomfort. A hand massage can be extremely relaxing.

In addition to massage the aromatherapist may make a personal inhaler to assist with specific problems. Inhalers can be useful to carry with you for nausea, using a blend that might include Spearmint. Inhalers can also assist with deepening the breath. Frankincense may be used to help when we are anxious, and our breathing becomes shallow. Blends of essential oils can also be diluted for use in a roll-on bottle which can be used like a perfume on pulse points. Suitable blends can be extremely helpful assisting with stress and anxiety, or the feel-good factor. An aromatherapist may give the balance of an oil blend in this way for home use and a lovely reminder of the benefits of the treatment received.

Nourishing blends of carrier oils can feed the body through the skin, providing a variety of nutritional benefits when absorbed, as well as excellent skin care.

**Bowen technique**

The Bowen technique is a gentle therapy which aims to rebalance the body’s energy flow to assist the body to heal itself. This treatment utilises a light rolling movement of the thumb and fingers over the skin. This system uses subtle and precise mobilisations called "Bowen moves" which affect the muscles, tendons, nerves and fascia (connective tissue) by application of this gentle treatment with non-invasive pressure. These procedures follow prescribed sequences to affect specific body systems. The therapist will take frequent pauses during the
treatment to allow the body to adjust and absorb the treatment.

Energy therapies
Bach flower remedies and flower essences

Flower essences are preparations which utilise flowering plants and the sun’s energy. Bach flower remedies were created from British plants. Other examples of essences are made from varieties of rose or other countries’ indigenous plants such as Australian Bush plant remedies.

Flower essences are considered by some to assist with emotional imbalance, often assisting with emotions, mood, mental state, shock and sometimes trauma. These are often seen as precursors to physical illness. The essence from the flower of the plant is usually stabilised in alcohol, although special preparations are available for those who wish to avoid any alcohol content. Just a few drops are added to water or dropped under the tongue.

Each plant has a unique signature addressing different emotions. A blend of essences is common as a rescue remedy, as in Bach flower remedies, and thought to assist with stressful situations such as hospital visits and exams.

Flower essences are available from many chemists and health shops with accompanying information to assist in correct selection.

Therapists who specialise in flower essences will take a case history and then blend a group of essences to address their client’s emotional needs.

Energy therapies; healing, reiki, therapeutic touch

These therapies are frequently used in supportive and palliative care. Although the word ‘healing’ is used as part of the treatment, this is not referring to cure and likewise ‘spiritual’ is not associated with any religion. These modalities are often used with the intent to assist with pain relief, promote relaxation, improve sleep, to contribute to a
Complementary therapies (cont.)

sense of well-being or to reduce side effects of chemotherapy and radiotherapy.

Healing

Healing, sometimes referred to as spiritual healing, is described by The National Federation of Spiritual Healers as “a process that promotes better health, through the channeling of healing energies through healer to receiver”. Healing should not be confused with curing cancer.

There is some evidence that suggests this supportive therapy may quicken the body’s own healing process. The practitioner will use light touch or no touch as appropriate for the patient who will be clothed.

Reiki

Reiki is a method of healing which promotes the aim of homeostasis through balancing the body’s own Ki (energy) with Rei (universal energy). This healing modality originated in Japan with its origin in self-healing. Reiki treatments are received in a similar way to healing with the recipient fully-clothed.

Therapeutic Touch

Therapeutic touch is a modern interpretation of some traditional healing approaches. It is described as an exchange of energy between client and practitioner, a healing meditation. During the treatment, the practitioner will place their hands a little way above the client’s body and rhythmically move their hands from head towards the feet with the aim of creating a sense of deep relaxation for the client with the potential of reduced anxiety.

Hypnotherapy

Clinical hypnotherapists believe that creating a more positive relaxed state of mind helps the body to repair and heal.

Hypnotherapy is a way of reducing the stress response. During a first hypnotherapy session, the hypnotherapist will take their client through a progressive relaxation, relaxing first the body and then the mind. The therapist may provide clients with relaxation techniques and recordings to help them relax at home and, after a little practice,
these may assist a reduction in stress levels with an improved sense of relaxation, with the aim of feeling much better as a result.

Hypnotherapy aims to create powerful and positive changes. Unlike other therapies that work over a prolonged period, hypnotherapy will help you to move on as quickly as possible, leaving unwanted anxiety, stress, fear, phobia or habit behind, often after only two or three sessions.

There is significant evidence for the use of hypnosis within integrative care in reducing different side effects to treatment such as nausea or anxiety or overcoming phobias. Some medical practitioners receive training in these techniques.

Hypnotherapists will often tape record sessions for use at home in the same way as a meditation, with mindfulness techniques often incorporated into it. During a hypnotherapy session, the patient or client will be guided into a trance state, through suggestion of voice. This trance is a state of awareness where the client is in touch with the practitioner’s voice but in a deep sense of relaxation. The therapist will make positive suggestions which may assist in a change in mental attitude towards a problem or life concern, either physical or emotional, according to the desire of the patient. Hypnotherapists also have reported success with addictions.

Massage

This touch therapy is the most well-known, but sometimes met with reluctance. A clinical/professional therapist will have completed extensive training and qualifications in anatomy and physiology in addition to extensive massage techniques. Therapeutic Massage is a combination of different touch techniques manipulating the soft tissue using the hands and perhaps forearms. Each treatment will vary according to the client’s requirements and health state. CML patients and others recovering from ill health should receive a lighter touch as they may be at risk of bruising. Pressure will vary dependent on
the sensitivity of the body and will ideally be a gentle rhythmical touch. Oil, or sometimes powder, is used to ensure a smooth and comfortable treatment. The therapist will offer a full body massage or massage to specific areas of discomfort such as neck and shoulders. In cancer and palliative care, several studies have taken place identifying assistance with anxiety, pain and a variety of reduced physical and emotional symptoms.

**M Technique**

Touch M has been widely used in hospital and hospice care. This very light touch manual treatment utilises a formula where each treatment will be given in the same sequence and same level of light touch throughout, making it accessible to anyone, however unwell. Treatment can be full body or just to specific areas. Training is available to anyone who wishes to give touch to friends and family, also for health professionals and therapists. M technique is extremely relaxing and invaluable to anyone with low energy or high stress levels or where deeper massage is inappropriate.

**Reflexology**

A reflexologist will apply an intermittent pressure using their thumb and fingers to areas of the feet or hands which they believe will access all body systems with the aim of bringing balance, or homeostasis, to their client. A reflexologist does not diagnose but, with reference of a foot chart, which relates areas of the feet to areas of the body, they can identify potential areas of imbalance through an altered state in the feet in different areas. Clients may also feel areas of discomfort in their feet or sensation in their body. If this is the case, the therapist and client together will identify areas of imbalance in the body. This holistic treatment is found by many clients to offer deep relaxation, boost their immune system and assist with maintaining good health. Some studies and client feedback have suggested that reflexology can assist with anxiety, pain, nausea, sleep and digestive problems.

**Reflexology Lymphatic Drainage (RLD)**

RLD is additional training available to qualified
reflexologists which focusses specifically on the lymphatic system. Early indications from the founder suggest RLD is useful in reducing volume for lymphoedema and assisting with oedema as well as inflammatory and auto-immune disorders. This treatment of feet or hands is deeply relaxing.

Movement, mindfulness and meditation

Mindfulness and meditation

Meditation, or mindfulness meditation, is often offered by yoga teachers as a separate class or as part of a regular yoga class. The aim of meditation is to bring your attention and awareness more fully to the present moment, being aware of thoughts, physical sensations and emotions, as they arise, experiencing them as they are without trying to change anything. This simple act of bringing attention and awareness to our experience from moment to moment allows us to find better ways of responding to situations in our lives rather than our automatic habitual reactions. This can lead us to reduce physical and emotional distress and live a happier life no matter what we are facing.

A candle or flower or other object may be used to focus the mind to help build concentration. The instructor or teacher will use the patterns of your breathing and may also use guided meditation as well as other techniques for self-practice.

Mindfulness classes and courses in healthcare have become increasingly popular, including the following programmes: Mindfulness-based stress reduction, Mindfulness Based Cognitive Therapy and Mindful Self-Compassion.

Mindfulness invites you to pay attention to your present moment experience with kindness and curiosity; thus, changing your relationship with your experience. Proven benefits include reduced stress symptoms, improved quality of sleep and enhanced quality of life.

Yoga Nidra

Yoga Nidra is a form of guided visualisation bringing attention to different parts of the body. This practice requires the
participant to lie or sit quietly while their mind is guided through a structured sequence of visualisations. Yoga Nidra may be offered as a one to one session or in a class. This practice is deeply relaxing and can also be taught for self-practice.

**Tai Chi**

Tai Chi could be described as a moving meditation, using a sequence of gentle flowing movements that combine breathing, movement and awareness exercises with meditation.

There are many forms of Tai Chi and, practiced regularly, may help reduce symptoms or side effects including stress and anxiety, increase flexibility and balance and improve overall physical and emotional wellbeing.

**Tripudio Movement Systems (TMS)**

Tripudio Movement was developed to provide a short practical sequence of movements that can be taught for home practice, encouraging a healthy, flowing lymphatic and circulatory system utilising a mixture of gentle mobilisation and stretches which can be done seated or standing.

This Tripudio Flow sequence is also used at the beginning and the end of a one-hour class which uses a mixture of seated and standing gentle stretching, mobilisation, strengthening and low impact aerobic exercise all set to music. There is no floor work involved and TMS is suitable for any age or ability. Classes are often on offer at cancer centres, hospices and hospital clinics, as well as in the community.

**Yoga**

There are many different styles of yoga, some very vigorous and others slow or relaxing. Gentle classes which focus on a mix of stretching/postures, breathing and some meditation or relaxation would be most appropriate. To support your wellbeing, find a class that offers a gentle, relaxing approach or a class that is specifically designed for people who are affected by cancer.

Yoga aims to bring balance to the whole (mind, body and spirit) through awareness of your breath and movement. Classes usually last between one to one and a half hours and require loose or comfortable stretch clothing and bare feet.
Mindfulness

What is mindfulness?

Mindfulness is a way of living our lives that can help us become more resilient and cope more effectively with whatever we are facing, including living with a diagnosis of leukaemia. Mindfulness is a natural human tendency that we can all develop. Through the practice of mindfulness, we become more aware of the experience of what is happening from moment to moment and therefore engage in an effective way of responding to it. The present moment is the only time that we can change anything, including our automatic reactions and responses to any given situation.

With regular practice, we can cultivate mindfulness; we learn to rest back into the awareness of the moment to moment comings and goings of physical sensations, thoughts, emotions, sounds, sights, smells and tastes. We can bring curiosity to this rather than judgement. This can help us to become less reactive and we can become more effective in our responses to whatever we are experiencing. This helps us become more resilient and cope much better.

What are the benefits of mindfulness?

Mindfulness can help us learn to respond more effectively to whatever is happening in our lives. Whilst we cannot prevent difficulties from happening, through mindful awareness we can learn to cope better with situations.

Increasing our awareness of what is happening from moment to moment enables us to notice our automatic reactions and learn to respond more effectively. We can respond better to our constant stream of thoughts and worries. Research in people affected by cancer has shown this to be helpful with pain, stress, fatigue, anxiety and depression, and generally helpful with general wellbeing. Essentially, mindfulness can help us to live happier and healthier lives, whatever we are facing.

How to be more mindful

We can incorporate mindfulness into our lives in two ways.
One is the formal practice of mindfulness, setting time aside to bring awareness to sensations occurring during our breathing, when we move, when we are lying down and when we are walking. These practices are often done slowly, giving the attention time to become fully absorbed. These practices can be done for anything from a few minutes to longer practices of up to an hour. Research tells us that it is the regular daily practice of mindfulness which is so beneficial and anything from 10-30 minutes has been effective in its cultivation.

We can also be mindful in the activities of everyday life. We can turn off the phone, TV or music and just pay full attention to sensations when we are, for example, standing and walking, sitting down or lying, chopping up the vegetables, taking a shower or getting dressed. Maybe walking outside in the park and feeling the air and sun or rain on our faces and the ground underneath our feet.

There are eight-week courses available in mindfulness provided by trained teachers called Mindfulness Based Stress Reduction and Mindfulness-Based Cognitive Therapy. Courses can be found on www.bemindful.co.uk

**Mindfulness Practices**

These come in various forms including the mindful body scan, mindful walking, mindful movement, and mindful sitting practice. There are podcasts available that can help you and have been especially designed for people affected by blood cancers http://www.abbvie.co.uk/mindfulness.html

**Mindfulness of the soles of the feet**

While you are standing or sitting, gently bring your attention to the soles of the feet and start to notice the sensations - whatever is happening from moment to moment and remembering that there is no right or wrong way to experience the present moment so whatever is happening is completely fine. Maybe there is heat, coolness, moisture, dryness, tingling or numbness or maybe other sensations. The point is to just notice whatever is there from moment to moment. Each time that you find your mind has
Mindfulness (cont.)

wandered away, which all our minds do, just bring the attention back to the soles of the feet again.

You can practice this for a few seconds or a few minutes. Try it for yourself.

**Mindfulness of breathing**

Find a comfortable position, sitting up or lying down, allowing your body to settle. Become aware of your body in the position that you have chosen and take a couple of breaths. Let go of any tension in the body as you breathe out. Now bring your attention to the movement of the breath in the body and become aware of where you notice it most. Allow your attention to rest on the movement there as you breathe, gently coming back to that area each time your mind wanders away, as it surely will. You can practice this as long or short a time as you have available.

**Is mindfulness helpful for everybody?**

Many people find mindfulness helpful and find ways to apply it in their lives which are beneficial, especially when facing the uncertainty of a leukaemia diagnosis. Like anything, some people may find it less helpful than others, but generally speaking, many people find that there is something helpful that they can take from the practices and use to support them in daily life.

If you are suffering from a mental health condition, you should check with your doctor that this is appropriate for you right now.
Glossary

Allopecia
Hair loss.

Alternative therapies
These are treatments used instead of conventional western medicine.

Antibiotics
These are used to treat or prevent bacterial infections. They work by killing bacteria or preventing them from reproducing and spreading.

Blood cells
Blood cells have many different functions including transporting oxygen and nutrients to the lungs and tissues, forming blood clots to prevent excess blood loss and carrying cells and antibodies that fight infection.

Chemotherapy
A form of cancer treatment that uses one or more anticancer drugs as part of a standardised treatment regime.

Complementary therapies
These are treatments used alongside conventional western medicine.

Fatigue
Extreme tiredness, which is not alleviated by sleep or rest. Fatigue can be acute and come on suddenly or chronic and persist.

Full Blood Count (FBC)
A blood test that counts the number of different blood cells.

Holistic Needs Assessment (HNA)
This can help identify some of the physical and psychological needs of patients and ensure patients’ needs and management plans are met in a timely and appropriate way.

Mindfulness
Mindfulness is a way of approaching life that can help with resilience and coping more effectively in the present moment.

Mucositis
Chemotherapy-induced mucositis causes the mucosal lining of the mouth and gut to waste away and break down forming ulcers, leading to a sore mouth and diarrhoea.
Glossary (cont.)

Neuropathy
Also known as parathesia, but more commonly called pins and needles, this is indicated by tingling or numbness on the skin.

Neutropenia
A condition in which the number of neutrophils (a type of white blood cell) in the bloodstream is decreased.

Neutrophil
A type of white blood cell that helps fight infection.

Red blood cell
The blood cell that carries oxygen. Red cells contain haemoglobin, which permits them to transport oxygen (and carbon dioxide).

Platelet
A disc-shaped element in the blood that assists in blood clotting. During normal blood clotting, the platelets clump together (aggregate).

White blood cell (leukocytes)
One of the cells the body makes to help fight infections. There are several types of white blood cells. The two most common types are the lymphocytes and neutrophils.

If you would like more definitions of terms you may come across during your blood cancer journey, you can order our A - Z of Leukaemia by calling Patient Services on 08088 010 444

Tell us what you think!
If you would like to give us some feedback about this patient information booklet, please hover over the code to the right using your phone or tablet’s camera. Click the link as it appears and this will take you to a short web form to fill in.

Suitable for Android, iPhone 7 and above.
Useful contacts and further support

There are a number of helpful sources to support you during your diagnosis, treatment and beyond, including:

- Your haematologist and healthcare team
- Your family and friends
- Your psychologist (ask your haematologist or CNS for a referral)
- Reliable online sources, such as Leukaemia Care
- Charitable organisations

There are a number of organisations, including ourselves, who provide expert advice and information.

**Leukaemia Care**

We are a charity dedicated to supporting anyone affected by the diagnosis of any blood cancer. We provide emotional support through a range of support services including a helpline, patient and carer conferences, support group, informative website, one-to-one buddy service and high-quality patient information. We also have a nurse on our helpline for any medical queries relating to your diagnosis.

Helpline: **08088 010 444**

www.leukaemiacare.org.uk

support@leukaemiacare.org.uk

**Bloodwise**

Bloodwise is the leading charity into the research of blood cancers. They offer support to patients, their family and friends through patient services.

**020 7504 2200**

www.bloodwise.org.uk

**Cancer Research UK**

Cancer Research UK is a leading charity dedicated to cancer research.

**0808 800 4040**

www.cancerresearchuk.org

**Macmillan**

Macmillan provides free practical, medical and financial support for people facing cancer.

**0808 808 0000**

www.macmillan.org.uk

**Maggie’s Centres**

Maggie’s offers free practical, emotional and social support to people with cancer and their families and friends.

**0300 123 1801**

www.maggiescentres.org

**Citizens Advice Bureau (CAB)**

Offers advice on benefits and financial assistance.

**08444 111 444**

www.adviceguide.org.uk
Leukaemia Care is a national charity dedicated to providing information, advice and support to anyone affected by a blood cancer.

Around 34,000 new cases of blood cancer are diagnosed in the UK each year. We are here to support you, whether you’re a patient, carer or family member.

Want to talk?

Helpline: 08088 010 444
(free from landlines and all major mobile networks)

Office Line: 01905 755977

www.leukaemiacare.org.uk
support@leukaemiacare.org.uk

Leukaemia Care, One Birch Court, Blackpole East, Worcester, WR3 8SG
Registered charity 259483 and SC039207

Leukaemia Care
YOUR Blood Cancer Charity