What is WM?

Non-Hodgkin lymphoma (“NHL”) is a cancer of the lymphatic system and lymphoplasmacytic lymphoma (“LPL”) is a rare type of NHL.

Jan Waldenström was the Swedish doctor who first described the disease now known as Waldenström’s macroglobulinaemia (“WM”) in 1944 and “macroglobulinaemia” was the word he used to describe the high levels of IgM “paraprotein” (meaning an abnormal version of the usually normal IgM protein) seen in the bloodstream of 95% of patients with LPL. WM is the term describing LPL characterised by production of IgM paraprotein.

The bone marrow is the source of all our blood cells, including red cells (which carry oxygen), white cells (which fight infections) and platelets (which help the blood to clot when needed). The blood cells made in the bone marrow are continuously released from the bone marrow into the bloodstream. The lymphatic system is part of the body’s immune system and helps us fight infection, and consists of organs such as the bone marrow, thymus, spleen, and the lymph nodes (or lymph glands). Lymph nodes are connected by a network of tiny lymphatic vessels that contain lymph fluid and are found in groups, particularly in the armpits, the neck and the groin. Some groups of lymph nodes are situated more deeply, in the chest and abdomen. There is also lymphatic tissue in other organs, such as the skin, lungs and stomach.

B-cells are a type of white blood cell that develop into “plasma cells”, which in turn make antibodies to help fight infections. Antibodies are made from a special type of protein called “immunoglobulin” and there are 5 types of immunoglobulins (Ig) in the body: IgG, IgA, IgM, IgD and IgE. Each of these antibodies has a different function and size. Usually, antibodies are made in response to infections in order to help the body fight against them and so develop immunity. IgM is the largest of these, as it circulates in groups of five. WM affects B cells when they are in the process of developing into plasma cells. These developing cells are called “lymphoplasmacytic cells”, because they have features of both lymphocytes and plasma cells, which is where the name “lymphoplasmacytic lymphoma” comes from.

In WM, the bone marrow produces abnormal lymphoplasmacytic cells. Although they are of no use to the body, these cells keep being made. As their numbers increase, they build up within the bone marrow, lymph nodes, spleen and other organs. In the bone marrow the result of this build-up is that the normal blood cells are ‘crowded out’ and this leads to a gradual reduction of normal blood counts. If the build-up occurs in the lymph nodes, spleen or even elsewhere, these tissues swell up and lumps sometimes form that can be visible or felt, although this is not as common a finding as in other lymphomas.
What is the difference between WM and LPL?
In most people with LPL the abnormal B cells produce IgM protein and the condition is then known as “Waldenström’s macroglobulinaemia”. The term ‘macroglobulinaemia’ just means that there is more of the large (or macro) IgM type of immunoglobulin in the blood (‘aemia’) than normal.

The B cells that build up in large numbers when someone has an LPL often produce large amounts of antibody. Unlike the antibodies we produce when we have an infection, however, the antibodies produced by the B cells in LPL have no useful function. In fact, some of them can have harmful effects on the body if they wrongly target the body’s own tissues or organs.

Large amounts of IgM in the blood can cause it to become thicker than normal. This is known as “hyperviscosity”. Sometimes, the IgM (being an antibody) may wrongly recognise the body’s tissues as foreign and attach on to them causing damage or inflammation. This may result in damage to nerves (‘neuropathy’) or destruction of blood cells (‘autoimmune haemolytic anaemia’).

In around 1 in 20 people with an LPL, either no abnormal immunoglobulin protein is produced or the abnormal B cells produce a different immunoglobulin type (the smaller IgG or IgA types, for example). When there is no immunoglobulin produced or the smaller immunoglobulins are produced, the condition is then called ‘lymphoplasmacytic lymphoma’ rather than WM and thickening of the blood does not occur.

Despite these technical differences in the terms used to describe LPL and WM, the tests and treatments for these conditions are the same. So, from now on we will just refer to all LPL in this article as ‘WM’.

Causes of Waldenström’s macroglobulinaemia
The causes of WM are unknown. Like other cancers, it is not infectious and cannot be passed on to other people. There does seem to be a familial tendency to have WM or other kind of lymphoma in immediate blood (“1st-degree”) relatives of WM patients. This tendency is between 5 and 20 times more common than in the normal population but is not enough to warrant screening of family members.

In most cases, WM is preceded by a condition known as “monoclonal gammopathy of uncertain significance” (“MGUS”) which is how WM itself is thought to start. This is the very early stage, when there are very few LPL cells in the body (often undetectable in the tissues even if they are sampled by a biopsy) but there is a detectable amount of abnormal IgM (usually a low level). This may be picked up on a blood sample done for an unconnected reason, and at this stage people are typically feeling normal and have no symptoms. The cause of MGUS (and hence WM) is not known, but it is commoner as people get older.

Over time (usually years), these cells may gradually build up and accumulate. If they accumulate enough to affect the functioning of the body, symptoms such as fatigue, weight loss, sweats, fevers or infections (due to an under functioning immune system) may develop and WM is eventually diagnosed.
MGUS and WM are at the opposite ends of a spectrum that moves forwards over time. The rate of progression is variable and typically spans many years. The number of persons progressing from MGUS to WM builds up over time. By 5 years, 10% progress; by 10 years, 18% progress and by 15 years nearly a quarter progress to WM.

The symptoms of Waldenström’s macroglobulinaemia

WM often develops over a long period of time and many people have no symptoms at all. This means that the condition is sometimes found by chance while having investigations for another condition or on a routine blood test. About a quarter of people with WM are diagnosed by chance like this. Most people with WM, however, gradually develop symptoms due to the disease. Symptoms develop for two main reasons. The first is that abnormal B cells fill up the bone marrow or collect in the lymph nodes or the spleen (and, rarely, in other places in the body). The second reason for developing symptoms in WM is the presence of large amounts of IgM protein circulating in the blood.

Symptoms caused by a build-up of lymphoma cells

When abnormal B cells accumulate and fill up the bone marrow, it is not able to make as many normal blood cells as usual. This can cause:

- tiredness, weakness and breathlessness, due to a lack of red blood cells (anaemia)
- a tendency to develop infections, due to a lack of the white blood cells that help fight infections
- a tendency to bruise or bleed easily, due to a lack of platelets.

Unlike people with other types of non-Hodgkin lymphoma, people with WM do not often have swelling of their lymph nodes or spleen. If lymph nodes do enlarge you might notice swollen glands and if the spleen is swollen this can be uncomfortable or painful.

People with WM can also experience fevers, night sweats and weight loss. These are symptoms that are a particular feature of lymphomas and you might hear them referred to as “B symptoms”. They reflect the additional metabolic activity (the production and use of energy) of the LPL cells in the body. If such symptoms become marked, such as to require (for example) a change of nightclothes on a regular basis or significant weight loss, they should be mentioned to the doctor and may indicate the need for treatment. While all these symptoms can also be caused by other conditions, they should always be checked out by a doctor.

Rarely, WM lymphoma cells can build up in other parts of the body, forming masses or tumours. These can occur almost anywhere in the body -
they have been described in the spine, limbs, lungs, around the eye socket and palate, in the gut and the skin. These masses are usually slow-growing but they can cause symptoms if they press on surrounding organs, nerves or blood vessels. Very rarely, people may experience symptoms affecting the central nervous system, such as fits, weakness of the facial muscles, double vision. These symptoms may be a sign of the WM affecting the brain and are called "Bing-Neel Syndrome". Special tests are needed to identify these cells. This is a very rare complication of WM, which can be treated if recognised promptly.

**Symptoms caused by the IgM protein**

If there is a large amount of IgM protein in the bloodstream this can make your blood thicker and more slow-flowing than normal. This is called **hyperviscosity**. Hyperviscosity develops in up to 30% of people with WM. This can cause symptoms such as:

- nosebleeds
- blurring or loss of vision
- dizziness or headaches
- drowsiness, poor concentration or confusion
- shortness of breath due to heart failure or lung congestion.

Hyperviscosity can cause changes in the back of the eyes (the retinas) due to pressure in the retinal blood vessels. These changes can be seen readily by a hand-held instrument called an ophthalmoscope, which is found in GP surgeries and hospital outpatients. It provides the doctor with a simple tool for assessing whether the hyperviscosity needs to be treated urgently (if there is bleeding in the back of the eyes). A blood test called **plasma or serum viscosity** can be done to measure the degree of thickness of the blood, but it is not available in all hospitals.

Sometimes people develop numbness or tingling in their hands and feet, or problems with their balance. This may be due to nerve damage in the extremities ("**peripheral neuropathy**") caused by the abnormal IgM in the blood. It is important to mention such symptoms to the doctor, especially if they are getting worse over time. It may be necessary to carry out special tests to examine the nervous system in more detail, for example: a scan of the brain or spinal cord, a lumbar puncture to look for signs of inflammation in the spinal fluid, nerve conduction studies to see how well the nerves are conducting electrical impulses or even a nerve biopsy (a procedure done under a local anaesthetic).

In some people with WM the IgM has a tendency to cause the red blood cells to stick together in the cooler parts of the body such as the hands and feet, the tip of the nose or the ear lobes. This can cause poor circulation in these areas, especially when it is cold, which may cause colour changes or ulceration in these areas. If you notice this symptom you should mention it to your doctor because you might need to have a special blood test to look for a protein called a **cryoglobulin**.

In some patients, the antibody properties of the IgM may cause the red cells to clump together in the cooler parts of the body (the fingers, toes) and be broken down by the immune system resulting in so-called **Cold...**
Agglutinin Disease (CAD). This can result in the sudden onset of anaemia with symptoms such as fatigue and shortness of breath and the presence of dark brown urine due to the spill over of red cell pigment (haemoglobin) in the urine (haemoglobinuria). This may occur in the setting of MGUS or WM and needs prompt treatment. A transfusion of blood may be needed in the short term.

Diagnosis of Waldenström’s macroglobulinaemia

Your doctor will ask about your symptoms and general health, and will carry out a full physical examination. If the doctor suspects that you have hyperviscosity you will need blood tests, a bone marrow biopsy and scans to quantify the WM and treatment is likely to be commenced urgently.

Testing for IgM

Blood consists of blood cells (red blood cells, white blood cells and platelets), floating in a liquid called “plasma”, which contains a vast array of proteins with different functions. When a blood sample is taken, it is collected in one of several types of blood tubes. Some tubes have an additive to prevent the blood from clotting so that the blood cells can be analysed (e.g. a blood count); others have an additive that is designed to clot the blood cells, so removing all the clotting factors which bind the blood cells together, leaving just serum (which contains the remaining proteins, including immunoglobulins). Finally, some tubes contain additives which prevent clotting of the blood; the tube is spun so that the cells are separated from the rest of the plasma and analysis is carried out on the plasma itself (e.g. to check the clotting system).

In WM, the excessive amount of IgM is measured by a technique called “serum protein electrophoresis” (SPE). This test is carried out on patient’s serum and separates all proteins in the blood according to their electrical charge by running an electric current through the serum sample, either in a tube or in a gel.

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As IgM is such a large molecule, it is often difficult to measure the amount of protein accurately. The higher the level, the less likely it is that the result will be accurate. Some laboratories just report the total IgM level (in other words, the abnormal IgM resulting from WM as well as normal IgM that is still present), while others report the M-protein or ‘paraprotein’, which is the abnormal IgM on its own. It does not really matter which method is used, as long as it is consistently applied when following up an individual patient.

Other immunoglobulin levels (e.g. of IgA and IgG) are sometimes low and it is thought that this might be the reason why some people with WM are prone to sinus and bronchial infections.

A simple blood sample will show if the levels of red blood cells, white blood cells or platelets are low. Your blood tests will also show how well your liver and kidneys are functioning.

**Bone marrow biopsy**

Since WM is a disease of the bone marrow, it is essential to sample the marrow in order to confirm that LPL cells are present, how plentiful they are, and how much normally functioning bone marrow is remaining. This procedure is called “bone marrow biopsy” (BMB).

The samples are usually taken from the back of your hip bone (pelvis). You will be given an injection of local anaesthetic to numb the area. The doctor or nurse will then pass a needle through the skin into the bone and draw a small sample of liquid marrow into a syringe (bone marrow aspirate). After this, they will take a small core of marrow from the bone (a trephine biopsy). Both samples will then be looked at under a microscope.

A BMB can be done on the ward or in the outpatients department and no prior fasting is required unless sedation is used. The whole procedure takes about 15-20 minutes. It may be uncomfortable as the liquid marrow is drawn into the syringe or as the bone biopsy is taken with a needle, but any discomfort should only last for a few seconds. A small dressing or plaster is placed on the skin after the procedure and you will be free to move around straight afterwards.
You may feel bruised and have an ache for a few days after the test, but this can be eased with mild painkillers if necessary. The results of the BMB usually take around 7 to 10 days to come through because of the processing involved.

Increasingly, new ways of analysing the bone marrow in more detail are becoming available, including some genetic tests that are carried out on the LPL cells. Recently discovered genetic abnormalities called MYD88 and CXCR4 have been investigated by researchers. In some hospitals, a test is now carried out on the bone marrow to see if these abnormalities are present. Their precise significance in terms of prognosis and treatment is still the subject of concentrated research.

**Scans**

**CT (computed tomography) scan**

A CT scan is a special type of x-ray. It is used to find out if WM has affected lymph nodes, or organs such as the liver or spleen. A number of pictures are taken from different angles and fed into a computer which shows detailed pictures of the inside of the body. The process involves lying still for 30-45 minutes.

You may be asked not to eat or drink anything for at least four hours before your appointment. Most people who have a CT scan are given a drink beforehand or an injection into a vein in the arm at the time of the scan. This injection of a substance called “contrast” allows particular areas of the inside of the body to be seen more clearly and may make you feel hot all over for a few minutes afterwards. If you have kidney problems, the use of contrast should be avoided as this can make the kidneys worse.

**PET (Positron Emission Tomography)-CT scan**

A PET-CT scan is a body scan in which an injection of labelled glucose is injected in to the vein and a scan performed to see whether the labelled glucose is taken up by the body’s tissues. Such uptake occurs when the tissues are actively metabolising. This kind of scan—which involves more x-rays than a CT scan—does not usually add further...
information in the setting of WM, unless there is a suspicion that the condition is behaving more aggressively than before.

**How is Waldenström’s macroglobulinaemia treated?**

Although WM is not curable, it is very treatable and most people live with this disease for many years. Some people who are diagnosed with WM do not need any treatment at first. WM often develops slowly over years and the term often used to describe this gradual behaviour is ‘indolent’. This means that some people may not need treatment for months or, very often, years. If this is the approach your doctor recommends you will have regular check-ups in the outpatients’ clinic. This active monitoring is often called ‘watch and wait’ or ‘watchful waiting’ (see below).

**Watch and wait**

If the doctor decides that no treatment is needed, you will have regular check-ups to assess how you are feeling and to take blood tests to measure your blood cell counts and IgM levels. This kind of follow-up, with check-ups but without treatment, is quite common in people with a low-grade non-Hodgkin lymphoma. If you have no symptoms of WM you will typically be seen in the clinic every 3–6 months for clinical review and blood tests.

Even though this approach is only taken because it is in your best interests medically, it can be hard to wait for symptoms to develop or for things to become worse before anything is done. It can make you feel anxious and unable to enjoy your relative good health.

**When does treatment start?**

Your doctor will consider starting treatment if:

- you begin to get increasing symptoms attributable to WM
- the level of IgM protein in your blood is increasing or hyperviscosity develops
- your blood count changes, such as developing low levels of red blood cells (anaemia)
- you develop complications such as a progressive neuropathy that is felt to be due to the effects of the abnormal IgM

The treatment you will be given will depend on your particular circumstances and the medical team will prescribe the most suitable drugs for you on the basis of:

- the results of all the tests
- your symptoms – for example how severe they are and whether or not you have neuropathy
- your age and general health.

Treatment is aimed at improving your quality of life and keeping you well for as long as possible, with the least possible side effects. The main treatment for WM has for many years been chemotherapy.
However as a result of research and clinical trials, many novel therapies are becoming available to treat WM patients — these are the so-called “biological therapies”. Other treatments such as blood transfusions, growth factor injections or plasma exchange may also be used to improve particular symptoms. These additional treatments are called ‘supportive treatments’.

If your doctor thinks that you need treatment, you might have one or more of these treatments:

- chemotherapy drugs
- steroids
- monoclonal antibodies
- biological treatments
- stem cell transplant.

**Chemotherapy**

Chemotherapy (chemo) is the use of anti-cancer (cytotoxic) drugs to destroy cancer cells, and can be given: as tablets; into a vein (‘intravenously’); or as an injection under the skin (‘subcutaneously’). You may be given just one type of chemotherapy drug or you may be given two or more such drugs together (combination chemotherapy). The selection of treatment is determined by the nature of the problems you are facing as a result of your WM and your own state of health, which will impact on your ability to tolerate different treatments. Some treatments can affect the stem cells in your bone marrow and should be avoided if there is a chance of needing a stem cell transplant in the future (see section on stem cell transplants for what this involves). You may be offered the chance to participate in a clinical trial.

Your specialist will explain which treatment is appropriate for you, but if ever you feel unsure of what is being offered or why, you can consider seeking a second opinion. It is important to feel comfortable with the options that have been recommended and to have a good understanding of what is to follow.

If you are treated with just one chemotherapy drug, any side effects are likely to be mild. If you are having treatment with a combination of drugs you may have more side effects. However most treatments for WM are given in outpatients and do not require admission to hospital unless complications such as infections occur. Your doctor or specialist nurse can tell you what to expect. You should always tell them about any side effects you have and ask them questions about your concerns. Some very effective medicines are available to reduce side effects if necessary. Treatment for WM usually spans 4 to 6 months. Sometimes the condition seems to be slow to respond but patience is important as a response is forthcoming in most cases,
given adequate time, and it is important not to switch to a new therapy too soon.

One of the most common side effects of chemotherapy is being more prone to infections. Always let your doctor or nurse know if you have any signs of an infection, such as a cough, fevers, shivering or shaking so it can be treated straight away.

The chemotherapy drugs most commonly used to treat WM are listed below.

**Chlorambucil (Leukeran®)**
Chlorambucil is taken as a tablet and is usually given for 7 to 10 consecutive days per month for the time recommended by your specialist. It should be kept in the fridge. It may be given with a steroid called prednisolone and with the monoclonal antibody, rituximab (see later section). Treatment is given monthly for 6 months. Chlorambucil should be avoided in anyone in whom a stem cell transplant is being considered as it damages stem cells and makes it difficult to collect them later.

**Fludarabine (Fludara®)**
Fludarabine is usually taken as tablets, but it may be given as a ‘drip’ into a vein (intravenous infusion) for up to 5 days per month. It may be given with other agents such as cyclophosphamide and rituximab.

Usually 6 months of treatment are required. Fludarabine should be avoided in anyone in whom a stem cell transplant is being considered as it damages stem cells and makes it difficult to collect them later.

**Cladribine (Leustat®)**
Cladribine is a similar type of drug to fludarabine. It may be given as a ‘drip’ into a vein (intravenous infusion) or as an injection just under the skin (subcutaneous injection). It may be given with other agents such as rituximab. Each treatment cycle consists of 5 daily doses of cladribine as a subcutaneous injection, plus 4 weekly intravenous infusions of rituximab. This is typically repeated just once after 2 or 3 months of a rest period. Cladribine should be avoided in anyone in whom a stem cell transplant is being considered as it damages stem cells and makes it difficult to collect them later.

**Cyclophosphamide**
Cyclophosphamide may be taken as tablets or given into a vein (intravenously). It is usually given in combination with other agents, such as in the DRC regimen, which consists also of dexamethasone (a steroid), rituximab, or the CHOP regimen, which is made up of cyclophosphamide, hydroxydaunorubicin, vincristine (or oncovin), and prednisolone. These regimens are not toxic to stem cells and can be used even if a transplant is planned in the future.
**Bendamustine (Levact ®)**

Bendamustine is a newer treatment for WM and is given as a ‘drip’ into a vein. It is usually given on day 1 and 2 of a 4-week cycle with rituximab, which is given only on day 1 of the cycle. Although not currently approved by the National Institute for Health and Care Excellence (NICE), bendamustine is available and funded through the government-backed National Cancer Drugs Fund. Bendamustine can be used safely in patients who may need a stem cell transplant in the future.

**Steroids**

Steroids are often used as part of your treatment as they can help the other drugs to destroy the abnormal B-cells and make chemotherapy more effective. They're usually given as tablets, but may also be given as an injection into a vein (intravenously). Steroids are usually given for 1 to 5 days per month depending on the chemotherapy schedule. The side effects of steroids—including fluid retention, weight gain, restlessness, agitation and sleep disturbance, a tendency to high blood sugar and high blood pressure—are temporary and usually go away when treatment finishes.

**Monoclonal antibody therapy**

Monoclonal antibodies are drugs that recognise, target and stick to particular proteins on the surface of cancer cells. They can stimulate the body’s immune system to destroy these cells.

A monoclonal antibody called rituximab (Mabthera ®) is used to treat B cell lymphomas like WM. It targets the protein CD20 found on B-lymphocytes and is given as a drip into a vein (infusion). Rituximab may be given with chemotherapy and/or steroids. It may cause temporary allergic reactions at the time of the infusion (and rarely afterwards) and the first infusion is given over 6 hours to try and limit this reaction. Subsequent infusions can usually be given over 90 minutes.

Ofatumumab (Arzerra®) and obinutuzumab (GA101) are newer anti-CD20 monoclonal antibodies that also appear effective in indolent lymphomas including WM.

**Biological therapies**

A great deal of international collaboration and effort in recent years means that there are novel therapies becoming available for patients with WM. This includes agents which target chemical pathways within the LPL cells and also affect the way LPL cells collaborate with the environment in which they live (the bone marrow microenvironment). Such agents include bortezomib (Velcade®), carfilzomib (Kyprolis®), ibrutinib (Imbruvica®) and idelalisib (GS1101). Others include the so-called immunomodulatory drugs that are related to thalidomide, including lenalidomide (Revlimid®) and pomalidomide (Imnovid®).
**Bortezomib (Velcade®)**

Bortezomib is a new type of treatment for WM called a ‘proteasome inhibitor’ and appears to kill LPL cells as well as hamper their support networks within the bone marrow. It has shown encouraging results in trials and appears to be especially effective at lowering a high level of IgM and may also be useful in the setting of high blood viscosity. It is given subcutaneously either twice a week for 2 weeks and then a week’s break; or weekly for 4 weeks followed by a week’s break. It is currently the subject of a clinical trial in the UK (R2W), which is testing it in combination with cyclophosphamide and rituximab against fludarabine, cyclophosphamide and rituximab.

The main side effects of bortezomib are tingling, numbness or pain in the hands and feet, but this seems to be less of a problem now that it is given subcutaneously. It is important to highlight any symptoms of peripheral neuropathy that are present at the outset, in order that a close eye is kept to avoid worsening of this problem with the use of Velcade.

**Carfilzomib (Kyprolis®)**

Carfilzomib is a ‘next-generation’ proteasome inhibitor related to bortezomib. It has been analysed in conjunction with rituximab and dexamethasone in 31 patients with WM, of whom nearly 90% were previously untreated. The overall response rate in this trial was 81%, with 65% achieving more than a partial response. It was well tolerated and, after 9 months, two-thirds of the patients treated remain stable. Studies are ongoing in the US, and it is not yet possible to prescribe carfilzomib for WM outside of a trial.

**Ibrutinib (Imbruvica®)**

Ibrutinib is an investigational agent designed to specifically target and inhibit a signaling protein in cells called Bruton’s tyrosine kinase (BTK). BTK is a key mediator of B-cell survival, meaning that, through multiple signaling systems within cells, BTK regulation helps to direct malignant B-cells to lymphoid tissues, thus allowing such abnormal cells access to a special environment that encourages their survival. As mentioned previously, more than 90% of patients with WM have the MYD88 mutation, which is intimately connected to BTK. Through deactivation of BTK, ibrutinib has been found to abolish binding of MYD88 to BTK in MYD88-expressing WM cells.

Ibrutinib comes in the form of a tablet, taken daily. The pivotal study was done at the Dana Farber Cancer Institute by Steven Treon and colleagues, who treated 63 patients with relapsed or refractory WM with ibrutinib for up to 2 years. The number of treatment cycles ranged from 1 to 18, but after a median of 9 cycles, the serum IgM level fell from 36 g/l to 12.6 g/l and the haemoglobin rose from 10.6 g/dl to 13.4 g/dl, while the amount of WM in the bone marrow fell from 70% to 36.7%. Two-thirds of patients showed a partial response or better.
Another mutation has been identified that seems to provide further insight into which patients are more likely to respond to ibrutinib. The C-X-C chemokine receptor type 4 (CXCR4) plays a crucial role in modulating the biology of B-cell lymphoproliferative disorders (see section on Future Directions for more information).

The USA’s Food and Drug Administration (FDA) granted ibrutinib ‘Breakthrough Therapy Designation’ for use as a single agent in previously treated or resistant WM. A European trial is in development and is planned to start in 2014/5 with a view to UK participation.

**Idelalisib (GS-1101)**

Idelalisib is another investigational drug, also a tablet. It is an inhibitor of a molecule known as phosphoinositide 3-kinase (PI3K) delta. Signalling in cells by this PI3K delta pathway is critical for the growth, development, survival and movement of B lymphocytes and is overactive in many B-cell lymphomas including WM. Idelalisib acts by blocking this pathway and is being developed both as a single agent and in combination with approved and investigational therapies. Idelalisib is only available within a clinical trial, the Gilead 0125 Study, which is currently open in the UK.

**Thalidomide (Thalomid ®) and Lenalidomide (Revlimid ®)**

Thalidomide and lenalidomide are two related drugs, known as ‘immunomodulatory drugs’. Both have been trialed in WM patients with limited success, due to problematic side effects that were noted (peripheral nerve damage in the case of thalidomide and marked anaemia in the case of lenalidomide). These difficulties, as well as the advent of other, newer, agents, have led to a cessation of trials of these agents at this time.

**Pomalidomide (Imnovid ®)**

Pomalidomide is a third-generation immunomodulatory drug. Like thalidomide and lenalidomide, pomalidomide can stop cancer cells from growing abnormally. Pomalidomide may also stimulate the immune system to fight the cancer cells and possibly improve the effectiveness of the steroid dexamethasone and the monoclonal antibody rituximab to fight WM cells. This drug, which is taken in tablet form, has been used experimentally in a related condition called multiple myeloma and information from these other research studies suggests that pomalidomide may help to reduce or prevent the growth of cancer cells. Clinical trials are underway which show promising results in WM without the kind of side effects noted with thalidomide and lenalidomide.
Stem cell transplantation

Some people with WM may have treatment involving the use of their own stem cells (autologous stem cell transplant, ASCT) or stem cells from a donor (allogeneic stem cell transplant, allo-SCT). Stem cells (in this context a specific type of stem cells called ‘haematopoietic cells’ or ‘HPCs’, i.e. “blood-producing cells”) are primitive cells found in the bone marrow that can develop into mature blood cells. They are found in the bone marrow and can be collected from the patient or donor before high dose chemotherapy is given to the patient. ASCT is sometimes called “high-dose chemotherapy with stem cell support”.

There are serious side effects associated with these treatments. They are not suitable for everyone and are not done routinely and doctors take into account a person's general health and fitness before recommending them. This often means that the risks of carrying out stem cell transplants in older persons who have other health problems are too high to recommend this approach.

Stem cell transplants are only performed after chemotherapy has been given to reduce the burden of the disease and put it into a remission and serve to consolidate that remission.

Autologous stem cell transplant

Patients due to undergo ASCT have some of their own stem cells collected and stored in advance of receiving a course of high-dose chemotherapy to kill any remaining lymphoma cells. This allows them to have higher doses of chemotherapy to destroy the lymphoma cells before the bone marrow is repopulated by the returned stem cells. The stem cells make their way to the bone marrow, where they form new blood cells to restore the bone marrow to normal function. While the stem cells are making their way to the bone marrow and becoming re-established there, the patient is particularly vulnerable to infection and requires a period of inpatient treatment and monitoring. This form of stem cell transplant is not curative, but it can lead to a long-lasting remission; in other words, the disease can stay at a very low level for quite a long time (typically a number of years) before further treatment is needed.

Figure 10. A schema showing the steps involved in autologous stem cell transplantation.
**Allogeneic stem cell transplant**

In this kind of transplant the stem cells come from another person. The donor might be a close relative, such as a brother or sister, or may be someone unrelated who has a matching tissue type. After the patient receives high dose chemotherapy, the donor’s stem cells are infused into the bloodstream via a cannula and within 2 or 3 weeks, produce donor blood cells in the patient’s bone marrow. These new cells resupply the patient with blood cells which can also directly fight against any leftover lymphoma cells. In this kind of stem cell transplant, the donor’s immune system is used as the weapon within the patient. There is an ongoing risk that the donor immune system may react against the patient’s healthy tissues as well (graft-versus-host disease), causing a variety of complications after the transplant. Special treatment is needed to suppress the patient’s immune system for a period of time to allow it to accept the donor’s cells (even though they are a tissue match), and this inevitably leads to the risk of unusual and dangerous infections. As with ASCT, a period of hospitalisation is inevitable.

While this form of transplant can offer the possibility of cure for some people with WM, it is a more hazardous procedure than an autologous transplant and the patient’s general health has to be good before you would be considered for it. The risks and benefits need to be weighed very carefully before embarking on this form of treatment.

![Figure 11. The steps involved in obtaining donor stem cells and their eventual use in the patient](image)

**Supportive treatments**

Supportive treatments are designed to counteract some of the symptoms of the lymphoma and the side effects of the treatments. In WM, these supportive treatments include antibiotics that are given to prevent infections (which may occur during chemotherapy cycles), blood transfusions and plasmapheresis.

**Blood transfusions**

Your red cell or platelet counts can decrease as a result of the WM itself or because chemotherapy is affecting your bone marrow as a side effect. If the counts fall to levels that cause troublesome symptoms, the medical team will consider giving you red cell or
platelet transfusions. Transfusions are given through a cannula (a thin flexible tube) into the vein and this can be done either as a day case or as an inpatient. If you have to have several blood transfusions over time, your blood will need to be checked regularly to check the cell counts and ferritin level. The ferritin level is an indication of your body's store of iron and it is important that this doesn't get too high.

White blood cells cannot be given by transfusion; rather growth factor injections can be used to boost the white blood count if it is low and increasing the risk of infection.

**Plasma Exchange**

If the IgM protein in the blood is causing symptoms, especially if it is causing heart or breathing problems, the blood can be thinned by a procedure called ‘plasmapheresis’. This is alternatively called 'plasma exchange' and it takes 1–3 hours.

In this procedure a cannula is placed into a vein in each arm. Blood is slowly removed from one arm and the blood is passed through a special machine that separates the liquid part of the blood—the ‘plasma’, which contains the IgM protein—from the blood cells. The blood cells are then passed back, together with an artificial plasma substitute, into the other arm. This might only need to be done once—before the chemotherapy has taken effect for example—but it might have to be done several times.

**Taking part in a clinical research trial**

You might be asked if you would like to take part in a clinical trial, a research study that tests new medical treatments. Clinical trials are very important in improving future treatments for people with your type of lymphoma. Also, some of the newer treatments are only available for people who are taking part in trials.

There is currently a multi-centre trial that is recruiting people with WM in the UK called the **R2W Trial**. This trial is comparing the combination of bortezomib, cyclophosphamide and rituximab (BCR) with a combination of fludarabine, cyclophosphamide and rituximab (FCR) for initial therapy of WM.

There is also a study in the UK for patients with relapsed WM known as the **Gilead 0125 Study**. It is a phase 3, randomized, double-
blind, placebo-controlled study evaluating the efficacy and safety of idelalisib (GS-1101) in combination with bendamustine and rituximab for previously treated indolent non-Hodgkin lymphomas. Participants need to have enlarged lymph nodes visible on a CT scan to be eligible for this trial.

Not all hospitals take part in clinical trials and there might not be a trial that is suitable for you when you are diagnosed, but this is something that you might like to discuss with your specialist when planning treatment. You do not have to take part in a clinical trial and can always opt to have the standard treatment instead.

**How will I be followed up?**

WM is an indolent lymphoma that patients can have for many years. With current therapies it is not possible to eradicate every last abnormal cell from the body and thus WM is very likely to relapse at some point after treatment. Consequently, all patients need to be followed up regularly in the outpatient department even when in remission, so as to detect a return of the disease and plan the next steps in a timely way so as to avoid a significant fall in wellbeing.

At each visit, blood tests are taken to check the level of the IgM protein and the blood counts to make sure that the WM is stable. If new symptoms develop—the same ones at diagnosis or new ones—or if there is a rise in the IgM protein or a fall in the blood counts, another bone marrow biopsy or CT scan will be advised to reassess the disease status.

In addition any new symptoms that arise between appointments should prompt contact with your medical team to discuss the symptoms. If necessary the next appointment could then be brought forward.

**What happens when WM comes back?**

When the WM comes back it can be treated again. The treatment will depend on how well prior treatments have been tolerated, how long it is since the last course of treatment and your state of general health.

The same treatment can be used again if a year or more has passed since its initial use. If the WM relapses more quickly than this, a different drug or a combination of drugs or a stem cell transplant might be considered.

In a small number of people WM turns in to a faster-growing type of lymphoma. If this happens it usually causes new symptoms and would be detected by tests (such as a lymph node biopsy). This is called 'transformation' and, although it sounds worrying, it can be
treated using drugs that are normally used for high-grade lymphomas.

**Future directions**

Recently identified genetic abnormalities found in patients with WM are being further explored to examine their significance for prognosis and treatment.

The MYD88 mutation is found in more than 90% of patients with WM and in up to 80% of persons with IgM MGUS. It predicts for progression from IgM MGUS to WM, but its significance in WM is still being analysed. The presence of MYD88 does not appear to impact on the effectiveness of ibrutinib.

The C-X-C chemokine receptor type 4 (CXCR4) plays a crucial role in modulating the biology of B-cell lymphoproliferative disorders. In nearly one-third of patients with WM this specific genetic mutation in the CXCR4 gene is present and, in experiments with laboratory cell samples (‘cell lines’), the researchers found that WM tumour cells with the CXCR4 mutation proliferated more quickly than those without the mutation. In mice, WM cells spread further and faster if they carried the CXCR4 mutation than if they did not: to the animals' liver, bone marrow, lymph nodes, kidney, and lungs. When researchers in the USA re-examined WM cells from human patients, they found that cells from patients with the most aggressive disease were the most likely to have CXCR4 mutations. It is hypothesised that the CXCR4 mutation drives the disease, in other words that it spurs WM cells to grow, divide, and spread. When researchers treated WM-carrying mice with an antibody that targets the mutation, progression of the disease halted, providing hope of a novel therapy to target this mutation in the future.

As previously mentioned, there are a large number of novel therapies being tested in WM with promising results. This fact and ongoing developments in scientific research mean that the future for people affected by WM is brighter than ever.

It is crucial to report any concerns to your medical team when they arise, be well informed and keep as fit and active as possible throughout the course of the illness.

*Dr Shirley D'Sa, University College Hospital, London 2014 - Review April 2015*