STAGE 3 MELANOMA

What are the aims of this leaflet?

This leaflet provides some general information on melanoma skin cancer and the melanoma staging system. In particular, it tells you what stage 3 melanoma is and what will be the investigations/treatments. Details on where to find out more information are also provided at the end of the leaflet.

What is melanoma?

Melanoma is a type of skin cancer, which arises from the pigment cells (melanocytes) in the skin. One of the most important causes of melanoma is exposure to too much ultraviolet light in sunlight. The use of artificial sources of ultraviolet light, such as sunbeds, also increases the risk of getting a melanoma.

Melanocytes make a brown/black pigment (known as melanin), and often the first sign of a melanoma developing is a previous mole changing in colour or a new brown/black lesion developing on the skin. Most frequently there is darkening in colour but occasionally there is loss of pigmentation with pale areas or red areas developing. Occasionally melanomas have no pigment and appear just a pink/red colour. The development of this melanoma on the skin is known as the primary melanoma.

Melanoma is considered to be the most serious type of skin cancer because it is more likely to spread (metastasise) from the skin to other parts of the body than other types of skin cancer. If melanoma has spread to other parts of the body, those deposits are known as secondary melanoma (secondaries/metastases).
What is stage 3 melanoma?

Doctors use a staging system for melanoma to indicate both the likely outcome and the best treatment. The AJCC (American Joint Committee on Cancer) system is currently used in the UK to stage melanoma from 1 to 4. Stage 1 is the earliest melanoma and stage 4 is the most advanced.

The staging system takes into account the thickness of melanoma and if there has been any spread of melanoma from the skin to other parts of the body. Stage 1 and 2 melanomas are present in the skin only and have not spread elsewhere in the body. Stage 3 melanomas have generally spread towards or have reached the draining lymph glands (nodes). Stage 4 melanomas are those that have spread beyond the closest draining lymph glands to other parts of the body.

Lymph nodes, or lymph glands, are found in our lymphatic system, which is part of the body’s immune system. The lymphatics are tiny channels in the skin, which move lymph fluid around the body to the lymph nodes in the groin, under the arms (arm pits) and in the neck. The immune system may fail to destroy the melanoma cells and the cells can then grow within the lymph nodes. If secondary melanoma appears as a lump that can be felt (often about the size of an olive), then you will be diagnosed with stage 3 melanoma.

In some patients, a procedure called a sentinel lymph node biopsy is used to detect small numbers of microscopic cancer cells in the nodes, (which cannot be felt). If you have a positive sentinel node biopsy, you will also be diagnosed as having a stage 3 melanoma.

Stage 3 melanoma may also be diagnosed if there are tiny (microscopic) deposits of melanoma in the skin surrounding the primary melanoma (microsatellites), which cannot be seen with the eye but are found by the pathologist when the primary melanoma is removed from the skin. Sometimes these deposits can be seen in the skin near the primary melanoma and they are called in-transit secondaries or satellites. These deposits are considered to be ‘in-transit’ between the primary melanoma and the local lymph nodes. These deposits can be skin coloured or black/blue. They can sometimes be felt just under the skin and are sometimes associated with bruising.

Occasionally a primary melanoma is detected purely as a result of someone developing enlarged lymph nodes with no history of a skin lesion (an occult primary).
Stage 3 melanoma can be divided up into stage 3a, 3b, 3c and 3d. If you wish to know the details about your specific melanoma stage then you can discuss this with a member of your own hospital melanoma team.

Having a stage 3 melanoma means a greater chance that the cancer will come back later somewhere else in the body (stage 4 melanoma). However, this is by no means inevitable. So, developing stage 3 melanoma is a cause for concern but it is still potentially curable.

What tests may be performed when stage 3 melanoma is suspected?

**Fine needle biopsy**
This is one of the most common tests, and it sometimes termed a fine needle aspiration (FNA). A sample is taken from a suspicious lump using a small needle. The pathologist then examines the sample under the microscope to look for melanoma cells. This test is similar to a regular blood test and typically takes place in the clinic. When the lump is deeper in the tissues, a radiologist (the X-ray doctor) will use a scan to determine where to place the needle. This can be an ultrasound scan or a CT (computerised tomography) scan.

**Scans**
Scans such as ultrasound, CT (computerised tomography), MRI (magnetic resonance imaging), PET (positron emission tomography) are usually done to check for any further spread of the melanoma in the body. Your melanoma team will explain why they have chosen a particular type of scan - each type has particular advantages. Further details on what these scans may involve can be found on the links provided at the end of the leaflet.

Routine scans can be considered for patients with stage 3 melanoma but the following advantages and disadvantages should be considered.

**Possible advantages and disadvantages of having regular scans**

<table>
<thead>
<tr>
<th>Possible advantages to having regular scans</th>
<th>Possible disadvantages to having regular scans</th>
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<tbody>
<tr>
<td>If the melanoma comes back (recurrent melanoma), it is more likely to be detected sooner. It is possible that this could lead to a better outcome by allowing treatment with</td>
<td>Although early drug treatment of recurrent melanoma might improve survival, there is currently limited evidence that earlier treatment definitely does this.</td>
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<td>drugs (such as immunotherapy drugs) to start earlier.</td>
<td>Some people find it reassuring to have regular scans</td>
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<td>Some people find that having regular scans increases their anxiety, waiting for the next scan, having the scan then waiting for the results.</td>
<td>Scans expose the body to radiation, which can increase the risk of other cancers in the future. Scans of the chest in particular cause a very small increase in the risk of thyroid cancer.</td>
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<tr>
<td>Scans of the brain and neck increase the risk of developing cataracts.</td>
<td>Scans may show abnormalities that are later found to be harmless, causing unnecessary investigations and anxiety.</td>
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**Genetic testing on your melanoma**

A sample of your melanoma tumour may be sent for genetic testing, in particular to look for a specific gene mutation (a change in the arrangement of the building blocks of your melanoma), known as a BRAF mutation. This gene mutation makes an altered BRAF protein, which makes the melanoma cells grow. Around 50% of melanomas will have a BRAF mutation. If you have changes in the BRAF gene, your melanoma is described as BRAF positive or mutant. If you don’t have changes, then your melanoma is BRAF negative or wild type. Although this information may not be used straightaway, it will help plan any future treatment with drug therapies if needed.

**What is the treatment for stage 3 melanoma?**

A multidisciplinary team (MDT) of experts will meet to discuss the best treatment option(s) for you. The MDT is made up of dermatologists, surgeons, pathologists, oncologists and specialist nurses A member of the MDT will explain your treatment options to you and you may also meet a melanoma/skin cancer clinical nurse specialist (CNS) who would be a point of contact for you and advise you accordingly.

Treatment options may include some of the following:
1. Surgery

**In-transit secondaries**
Surgery can be used to remove the in-transit or satellite lumps of melanoma in the skin near your primary melanoma scar, usually under a local anaesthetic (via an injection to numb the skin). This might not be possible when there are many deposits and so other treatments may be discussed with you, such as cryotherapy treatment, isolated limb infusion, or isolated limb perfusion, topical treatment with imiquimod or electro-chemotherapy (ECT). Further information can be found on the Macmillan Cancer Support website ([https://www.macmillan.org.uk/information-and-support/melanoma/recurrent-melanoma/treating/surgery/surgery-explained](https://www.macmillan.org.uk/information-and-support/melanoma/recurrent-melanoma/treating/surgery/surgery-explained)).

**Lymph node secondaries**
If you have spread of melanoma to your lymph nodes which can be felt, then all the lymph nodes in that area are typically removed (lymphadenectomy or complete lymph node dissection) for local control of the melanoma. The aim of the surgery is to remove the cancer to reduce the chance of it recurring in the same place. This is to avoid the melanoma from further growing in this site which may cause pain and lead to skin breakdown.

This surgery takes place while you are under a general anaesthetic (you are put to sleep) and will involve a stay in hospital. Usually during the surgery, drains (plastic tubes) are put in place with one end of the drain in the wound under the skin and the other end attached to a plastic bag. These tubes drain away any blood or lymph fluid from inside the wound and will be removed usually around 2 weeks after the operation.

Following surgery, there is sometimes a fluid collection under the wound known as a seroma, which usually resolves over time. However, one of the possible permanent side effects of a complete lymph node dissection is swelling of the limb known as lymphoedema. This can be mild but may be more troublesome especially in the leg after surgery to the groin, but even then, is usually manageable with the use of compression bandaging/stockings. Other complications of a complete lymph node dissection include scarring, infection, numbness, blood clots particularly if the groin nodes have been removed and stiffness. You will have physiotherapy to help you regain movement. It will help your recovery if you persevere with this.
If you have been found to have stage 3 melanoma following a positive sentinel node biopsy, in the past patients would also have been offered a complete lymph node dissection. However new evidence suggests that this further procedure does not improve survival from melanoma. This is therefore no longer routinely recommended after a positive sentinel node biopsy result.

2. **Talimogene laherparepvec (T-VEC)**

Talimogene laherparepvec (or T-VEC) is a genetically modified virus that is designed to infect and kill melanoma cells. It has to be injected directly into tumours, so is only suitable for people who have lumps that can be felt (e.g. skin lumps or lymph nodes) with no disease in internal organs, and it is considered when surgery is not an option. Ultrasound guidance may be needed to make the injection safer. Side-effects of T-VEC include pain at the injection site and rashes. You should keep the injection site covered for 48 hours after treatment, and discard dressings in the bin provided by the hospital.

3. **Adjuvant treatments/Clinical trials**

Adjuvant therapy refers to any additional treatment that is given after a cancer is surgically removed and when there is no other disease visible clinically or on radiological scans. The aim of adjuvant treatments is to stop or slow the growth of any potentially remaining cancer cells (reducing the risk of secondaries). This treatment will be discussed with you by your doctor and may be either treatment with a) targeted therapy or b) immunotherapy.

a) **Targeted Therapy**

If your melanoma is shown to have a BRAF mutation when it is sent for genetic testing, then targeted therapy with a BRAF inhibitor drug will be suitable for you and works by inhibiting the growth of any melanoma cells which may be present in the body. These are tablets or capsules which are taken twice a day. BRAF inhibitor drugs do not work unless you have a BRAF mutation. A BRAF inhibitor (e.g. dabrafenib) is usually given in combination with a MEK (another building block) inhibitor drug (e.g. trametinib). Side effects of these drugs include rashes, joint pain, high temperature (fever), ECG changes, high blood pressure, changes in eyesight, and weakness of the heart muscle. The adjuvant treatment is for 1 year.
b) Immunotherapy

Immunotherapy is treatment that encourages this action of the body to fight against the melanoma.

*Nivolumab* and *pembrolizumab* are drugs which target a protein called a PD1 receptor on the surface of T cells. Blocking of the PD1 receptor activates the T cells to produce an immune response against melanoma cells. It is given as an intravenous infusion every 2 weeks (nivolumab) or every 3 weeks (pembrolizumab). Side effects include tiredness, diarrhoea, feeling sick, skin reactions, changes to liver/kidney function, changes in the function of glands such as the thyroid and adrenals, and eye inflammation. The adjuvant treatment is for 1 year.

**What is the follow up for patients with stage 3 melanoma?**

Patients with stage 3 melanoma will be followed up regularly in the outpatient clinic. In the UK, after a diagnosis of stage 3 melanoma, patients are generally seen about every 3 months for 3 years, then every 6 months to 5 years. At your clinic appointment, you will be examined by either a doctor or a CNS for any further spread of melanoma. It is not unusual for your follow up to be shared between the different doctors, and/or a CNS, involved in your care.

Certainly, it is important to be aware and familiar with your body so that you can recognise a new lump (or lesion) if one appears. If in doubt call your specialist nurse and arrange a sooner appointment with your medical team if concerned.

**Self care (What can I do?)**

Normally, your CNS will discuss the following with you after treatment:

**You can examine your skin**

Most people do not develop further melanomas; however, some do and they may also develop other forms of skin cancer. The best way to detect skin cancer is to check all your skin every month (please see Patient Information Leaflet on early detection). Essentially, you are looking for changes in the size, shape or colour of any moles, a new mole, or a mole that looks different to the others. There are patient information web-packages, which outline how to look after your moles (please see the web links at the end of this...
You can also look for recurrence of the melanoma by checking for any new lumps near your primary melanoma scar.

**Top sun safety tips:**

- Protect your skin with clothing, and don’t forget to wear a hat that protects your face, neck and ears, and a pair of UV protective sunglasses.
- Spend time in the shade between 11am and 3pm when it’s sunny. Step out of the sun before your skin has a chance to redden or burn.
- When choosing a sunscreen look for a high protection SPF (SPF 30 or more) to protect against UVB, and the UVA circle logo and/or 4 or 5 UVA stars to protect against UVA. Apply plenty of sunscreen 15 to 30 minutes before going out in the sun, and reapply every two hours and straight after swimming and towel-drying.
- Keep babies and young children out of direct sunlight.
- The British Association of Dermatologists recommends that you tell your doctor about any changes to a mole or patch of skin. If your GP is concerned about your skin, make sure you see a Consultant Dermatologist – an expert in diagnosing skin cancer. Your doctor can refer you for free through the NHS.
- Sunscreens should not be used as an alternative to clothing and shade, rather they offer additional protection. Sunscreens do not provide 100% protection.

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**Vitamin D advice**

The evidence relating to the health effects of Vitamin D blood levels, sunlight exposure and Vitamin D intake remains inconclusive. Avoiding all sunlight exposure if you suffer from light sensitivity, or to reduce the risk of melanoma and other skin cancers, may be associated with Vitamin D deficiency.

Patients diagnosed with melanoma are advised to have their Vitamin D level checked by their hospital team. If levels are low they are advised to consider taking vitamin D3, 10-20 micrograms (400-800IU) per day, and increasing their intake of foods high in Vitamin D such as oily fish, eggs, meat, fortified margarines and cereals. Vitamin D3 supplements are widely available from supermarkets and chemists. Specific advice will be given by your doctor depending on your level of Vitamin D.
Having had a melanoma may impact on future applications for life or health insurance, particularly for the first five years after diagnosis. Equally you may be able to make a claim against any critical illness insurance, you may have taken out prior to the diagnosis of melanoma. If you have particular concerns about this, you should seek financial advice.

**Where can I get more advice, support & information about melanoma?**

When you have been diagnosed with melanoma you might experience a range of emotions including worry, confusion, or even feeling unable to cope. It will probably help if you discuss and share your thoughts and feelings with someone close. This might be a family member or friend. It could also be your doctor, specialist nurse or another member of the team looking after you.

When you are diagnosed with melanoma, you will be given a lot of information. All this information at once can be hard to take in. If you are not clear about anything during your treatment, please don’t be afraid to ask.

**Web links to detailed leaflets:**

**British Association of Dermatologists**
- Information on early detection and prevention of melanoma  
- Information on sun-safety  
- Information on Vitamin D  
  [http://www.bad.org.uk/vitaminD](http://www.bad.org.uk/vitaminD)

**Cancer Research UK (CRUK)**
- Information on melanoma staging  

**GenoMEL: The Melanoma Genetics Consortium**
- Information on Vitamin D
http://genomel.org/info-for-patients/%EF%BF%BCsun-protection-and-vitamin-d-after-a-diagnosis/

**Melanoma Focus**
- **Patient Decision Aid**  
  https://pda.melanomafocus.com/
- **Sentinel Node Consensus**  
- **Video Immunotherapy Treatment for Patients**  
  https://vimeo.com/252355507/00d629311d

**Macmillan Cancer Support**
- **http://www.macmillan.org.uk/Cancerinformation/Cancertypes/Melanoma/Melanoma.aspx**
- **Financial support**  
  http://www.macmillan.org.uk/HowWeCanHelp/FinancialSupport/Financialguidance/Financialguidance.aspx

**National Cancer Action Team (NCAT)/ NHS Choices**
- **Information on Vitamin D**  
  www.nhs.uk/Conditions/vitamins-minerals/Pages/Vitamin-D.aspx

**Links to patient support groups:**

- **Melanoma UK**  
  Web: https://www.melanomauk.org.uk/

- **Melanoma Action and Support Scotland (MASScot)**  
  17 Cairnhill Road  
  Bearsden  
  East Dunbartonshire  
  Glasgow, G61 1AU  
  Tel: 0773 823 1260  
  Email: leigh@masscot.org.uk  
  Web: www.masscot.org.uk

For details of source materials used please contact the Clinical Standards Unit (clinicalstandards@bad.org.uk).

**This leaflet aims to provide accurate information about the subject and is a consensus of the views held by representatives of the British Association of Dermatologists: individual patient circumstances may**
differ, which might alter both the advice and course of therapy given to you by your doctor.

*This leaflet has been assessed for readability by the British Association of Dermatologists’ Patient Information Lay Review Panel*