Anorectal Mucosal Melanoma
Information for patients and carers
**Introduction**

The information in this leaflet relates specifically to melanomas of the anorectal mucous membranes. The leaflet summarises a guideline [https://melanomafocus.com/activities/mucosal-guidelines/](https://melanomafocus.com/activities/mucosal-guidelines/) developed by experts in the field to advise cancer specialists who treat patients with this condition and is based upon the best evidence available. Skin cancers can also develop in the same areas of the body, in the skin rather than in the mucous membranes. These are known as cutaneous melanomas and are not covered by this guideline. If you have been diagnosed with a skin (cutaneous) melanoma please refer to the NICE guideline [www.nice.org.uk/guidance/ng14](http://www.nice.org.uk/guidance/ng14) and the other organisations listed at the end of this leaflet.

The number of medical terms has been kept to a minimum in this leaflet. If you come across a term you don't understand, please see the definitions and abbreviations section at the end of this document or ask your consultant or nurse.

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**What is it?**

Melanoma develops if there is uncontrolled growth of melanocytes, the cells responsible for pigments (darkening) the skin. Mucosal melanoma is a kind of melanoma that occurs in mucous membranes. These are the moist surfaces that line cavities within the body. Mucosal melanomas can occur in the eyes, mouth, nasal passages and larynx. They can also develop in and around the anus, vagina or penis. They are very rare in the Surveillance Epidemiology and End Results (SEER) database which registered melanomas over 22 years, only about 1 melanoma in 100 was a mucosal melanoma. The majority of melanomas grow within the skin. Mucosal melanomas can be more complicated to treat than skin melanomas as they are often diagnosed later. This is because they are in less visible places and because they are often not pigmented (darker), which makes them even more difficult to spot.

These melanomas are different in several ways from skin melanomas. For example, while the risk of getting skin melanoma is increased by too much exposure to the sun, there appears to be no link between sun and mucosal melanomas. No specific causes or links with lifestyle have been found for mucosal melanoma and as far as we know there is nothing you can do to prevent it. It also appears that mucosal melanoma tumours are different from skin melanoma tumours. This means that some treatments for skin melanomas may not be as effective for mucosal melanoma and the outlook, or prognosis (or natural history), for mucosal melanomas is not as good as for skin melanomas. The prognosis will be different depending on whether the tumour can be fully removed surgically and whether it has spread. If you would like more information about your individual situation, you should discuss this with your clinical team.
Questions you may wish to ask about your visits

How often will I have appointments at the cancer centre?
What sort of support will I need at these appointments?
How long will I be at the centre each time?
What further signs and symptoms should I look out for?
What symptoms should I report urgently?
How and who should I contact if I need to do so urgently?

What are the symptoms?
The symptoms for anorectal mucosal melanomas are similar to those for other cancers in the anus and rectum and for many other conditions. These include bleeding from the rectum (back passage), pain, swelling or a lump. There may be an urgent feeling to empty your bowels, although you may not be able to empty your bowels. As with skin melanomas, the area may be pigmented (darker) but this is not always the case. If you have any of these symptoms and your GP cannot find another cause, you should be referred quickly (normally within about two weeks) to a specialist with experience of this condition, usually a consultant colorectal surgeon but in some cases a dermatologist, for tests and further evaluation. It is most important that the team you see regularly treats patients with this condition. As it is so rare, you may need to travel to a specialist centre rather than attend your local hospital.

Going to the hospital

Tests and examination
At the hospital the consultant or a senior doctor on the team will examine you, including a rectal examination and proctoscopy. You may also have a sigmoidoscopy and an examination under anaesthesia (which means you will be given an anaesthetic to make you unconscious). This will let the consultant look closely at any lumps or other abnormalities in your rectum (back passage). You may also have an MRI scan of your pelvis. If the consultant identifies a suspicious lump, you will be offered an excision or a punch biopsy depending on your symptoms. The consultant may also carry out a core biopsy and/or a fine-needle aspiration to send off to be examined under a microscope. The guideline does not recommend that sentinel node biopsy is done routinely, but it may be performed in some cases to decide on future treatments or entering into clinical trials.

If the consultant makes a diagnosis of anorectal mucosal melanoma, you would normally have a CT scan of your body to check whether the cancer has spread, and you may need a further excision. There are several specific genes which have been identified that may be involved in some but not all mucosal melanoma tumours. The pathologist should test the tissue sample from the biopsy to see if one of these identified genes is involved in your cancer as this will help decide the best treatment for you now and in the future.

Diagnosis, staging and prognosis
The cancer team can estimate the possible outcomes – your prognosis – based on the results of your tests and medical examination. If you are diagnosed with a mucosal melanoma, your cancer will be evaluated according to a process known as staging. The stage indicates whether your cancer has spread in your body and, if so, where. This staging process will help both you and the cancer team make decisions about your future treatment.

For the staging, the American Joint Committee on Cancer TNM staging system is used. In addition to other information the team should record in particular:
- Location of the primary (original) tumour
- Primary Tumour size and extent of tumours
- Lymph Node involvement (whether or not the cancer has spread to the nearby lymph nodes (also called glands)
- Presence or absence of distant Metastasis (whether or not the cancer has spread to distant areas of the body).

The cancer team will make a prediction of the outcome of your cancer – your prognosis - based on the results of your tests and examination and on what happened to others who have had similar cancers. If you and/or a relative or carer want to have a full discussion about your prognosis, make this clear to someone from the cancer team. It will mean that your appointments will be more informative and will enable you to be more involved in planning your treatment and other aspects of your life. On the other hand you may wish to take it ‘one step at a time’. Whatever your decision, you will be fully supported by the cancer team.
**What to expect from your care**

A specific specialist cancer team will look after you during your treatment. As this is a rare cancer which has similarities to both skin and colorectal cancers, there should be co-operation between the skin and colorectal experts. The guideline advises teams on specific ways in which they should work together.

When discussing your diagnosis and treatment you should be told:

- The name and contact details of your consultant and of your Cancer Nurse Specialist.
- Who your key worker is (this person is your first point of contact should you have questions or problems and is usually the Cancer Nurse Specialist; you need to make sure you are given the contact details of the person to contact if your keyworker isn’t available).
- How to make an appointment with the consultant quickly should you run into problems.
- About referral to support services (for example, palliative care services) should you need them at any point in your treatment.

**Treatment**

If the cancer hasn’t spread beyond the primary site the first line of treatment for an anorectal mucosal melanoma is surgery. This should be done by a surgeon who is very experienced in complex anorectal surgery and is based in a centre where there is support from an experienced skin cancer team.

In planning the operation, the consultant should discuss your current health and how the surgery may affect your continence (bowel control). If it is likely to make a difference, the consultant should discuss other treatment options with you to agree what is best for you. The aim of surgery is to remove a little of the surrounding healthy tissue (typically 1mm) to make sure that all of the cancer is removed. This is called having ‘clear margins’.

If the cancer hasn’t spread and the margins are clear, this may be the end of your treatment (see After Treatment, page 6) or, depending on the stage and tumour, you may be offered adjuvant therapy (see page 5).

If, after surgery, the margins of tissue aren’t clear of cancer, the surgery may be repeated. If another operation isn’t an option, the possibility of radiotherapy should be discussed with you.

If the tumour is large, or the cancer has spread, then a more complex operation may be performed. This may involve removing parts of the anus (back passage), rectum or colon and any affected lymph nodes. This may be an abdominoperineal resection (APR) or an anterior resection (see definitions). If this type of operation is being considered the need for a permanent or temporary stoma (see definition) should be discussed with you. You also may be offered further tests to help decide if surgery is the best option for you. If the cancer has spread (metastasised), you may be offered chemotherapy and/or radiotherapy instead of surgery (see page 7).

**Questions you may wish to ask about your treatment**

Why for why not is surgery the best option for me?

How should I expect to feel after my surgery?

What will the recovery time be?

Will I be able to carry on with life as normal?

What will be the effects of my surgery?

What are the risks that I will have some incontinence?

Will I need a stoma?

What symptoms should I look out for, indicating whether the surgery has worked or not?

What support will I need after surgery?

Will I need adjuvant therapy? If so, what is the reason for this and what will it involve?

**Adjuvant (Extra) therapy**

Depending on the results of the surgery and the cancer stage, you may be offered adjuvant therapy. The aim of adjuvant therapy is to reduce the chances that the cancer will come back after it has been removed entirely by surgery (a ‘belt and braces’ approach). Recently there have been great advances in the treatment of skin melanomas with immunotherapy, which stimulates your own body to fight the cancer. By understanding more about the molecular make-up of different tumours, treatments such as immunotherapy and targeted therapies may target unique aspects of the tumour. However, because mucosal melanomas are so rare, scientists are still uncertain whether these treatments work in the same way as with skin melanomas.

You may be asked if you would like to enter a clinical trial to help doctors answer this question. Or, as doctors learn more, you may be offered immunotherapy, particularly if the tumour contains one of the genes that can make such a treatment effective. (However keep in mind that some of these treatments are not currently funded by the NHS and you may have to pay for them yourself). Similarly, you may be offered adjuvant radiotherapy or be invited to enter a trial to learn more about the benefits of this type of treatment for people with mucosal melanoma.
After treatment

After your initial treatment, you should be offered regular follow-up appointments to check that no cancer has returned, either to the original site or elsewhere in your body.

You should be given information how to make an appointment quickly at any time after your treatment if you are concerned.

The schedule recommended in the guideline is for an appointment every three months during the first three years after your treatment. The procedure will be similar to the one you had when you first attended the hospital and should include a physical examination, a check of your anus and rectum and a proctoscopy/sigmoidoscopy under anaesthetic to check everything is still clear at the original site.

You should be offered a CT scan two or three months after surgery to establish a baseline and then every six months during the first three years in order to check that the cancer hasn’t spread to another site. You may also be offered an MRI of the rectum and surrounding lymph nodes. Very occasionally the cancer spreads to the brain, the consultant should discuss the risks and benefits of a brain scan with you.

If you would like to read more about the advantages and disadvantages of having scans refer to the NICE option grid set out in skin cancer guideline NG14: www.nice.org.uk/guidance/ng14/resources/followup-with-regular-ct-scans-yes-or-no-pdf-250598416

The risk of the cancer returning reduces with time. Therefore from three to five years after treatment you should be offered an appointment every six months and a CT scan once a year. From six to ten years after treatment you should be offered an appointment once a year or alternatively rapid access to a clinic if you are concerned.

After ten years you should not need to have appointments but can come back at any time if you are concerned.

Questions you may wish to ask about your treatment for metastatic disease

Is there an immunotherapy, targeted therapy or radiotherapy available in my situation?
If so, what is the likelihood that it will reduce my symptoms and/or extend my life?
Is it available on the NHS?
Are there side-effects with this treatment and, if so, what are they?
What clinical trials are available?
What is the aim of the clinical trial in question?
What support will I need while on this treatment?
How will I be able to tell whether it’s working or not?
If it is not working, how soon can I move on to something else?
Will this treatment prolong my life?
Will I have a reasonable quality of life while on this treatment?

If your cancer recurs or spreads (metastasis)

Surgery is again the first line of treatment if the tumour recurs. If the cancer has spread or if surgery is not an option a member of the cancer team should discuss the different treatments available. New treatments are being developed and your consultant should tell you about their success rates, as well as the risks and benefits in your particular case.

You may be offered immunotherapy treatment to slow or stop the growth of the tumour. Some anorectal melanomas have unique aspects that may be targeted with specific drugs. In this case the cancer team should discuss the options with you and the availability of any clinical trials you could enter. There is little evidence that the routine use of chemotherapy or bio-chemotherapy is effective in the treatment of metastatic mucosal melanoma and it not currently recommended by the guideline.

Depending on numerous factors relating to your tumour, you may be offered palliative radiotherapy. Your consultant will discuss these types of treatment with you.

Following treatment for a recurrence or metastasis, you should be offered regular appointments at two- to three-monthly intervals, depending on the kind of treatment you have had. These should include CT scans and an MRI or CT scan of the brain. If you remain well and the disease does not progress, after two to three years, appointments can be every six months. After five years, the interval can be extended to yearly.

If the disease progresses, there are other options to make you more comfortable, which are used for skin melanomas. General guidance is available in NICE guidance CSG4: www.nice.org.uk/guidance/csg4.

Sources of information and support

Since mucosal melanoma is so rare, there are no UK websites dedicated to information about this disease.

Melanoma Focus (melanomafocus.com) is a charity that commissions and funds melanoma research, while providing support and information for patients, carers and healthcare professionals. It funded this leaflet and the associated guideline for health professionals, on which this leaflet is based. See https://melanomafocus.com/activities/mucosal-guidelines/, where there are links to a Resources page giving access to the various guideline documents.

Macmillan (macmillan.org.uk) provides support for people who have cancer.

Although not specific to mucosal melanoma, Cancer Research UK has a great deal of information about anal and rectal cancer, including treatments and living with cancer, which is relevant to anorectal mucosal melanoma.

NHS Choices has information on anal and rectal cancer treatments and other aspects of care.

The American Joint Committee on Cancer (AJCC), which publishes the staging manual used by doctors, has a patient leaflet explaining staging: https://cancerstaging.org/CSE/Registrar/Documents/needtoknow.pdf.

See https://melanomafocus.com/activities/mucosal-guidelines/ where there are links to a Resources page giving access to the various guideline documents.

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Definitions and abbreviations

**Abdominoperineal resection (APR)**

In APR surgery the anus, rectum and the lower part of the colon are removed. Depending on the circumstances this will be done either as open surgery with a large incision (cut) in the abdomen or laparoscopically with a series of small incisions where cameras and instruments are inserted. Because the rectum and anus are removed, a new exit point for bowels (faeces) will need to be created from the abdomen (belly) into a bag. This may with be a permanent (colostomy) or temporary stoma (usually an ileostomy).

**Adjunct therapy**

Adjunct therapy is treatment in addition to that used to treat the initial cancer. The aim is to reduce the possibility that the cancer will come back after it has been removed entirely by surgery.

**Anterior resection**

Surgery where the rectum and anus are removed, depending on the circumstances this will be done either as open surgery with a large incision (cut) in the abdomen or laparoscopically with a series of small incisions where cameras and instruments are inserted. Because the rectum and anus are removed, a new exit point for bowels (faeces) will need to be created from the abdomen (belly) into a bag. This may with be a permanent (colostomy) or temporary stoma (usually an ileostomy).

**Bowel Cancer**

Beating bowel cancer

For more information see: www.beatingbowelcancer.org/understanding-bowel-cancer/treatment/surgery/understanding-stomas.

**Biopsy techniques**

A biopsy is when a piece of the abnormal tissue is removed so the cells can be examined under a microscope to determine whether it is cancer and if so what type.

There are many ways of taking a biopsy. The ones mentioned in this leaflet are listed below.

See NHS Choices, from which this information was taken, for more detail. www.nhs.uk/conditions/biopsy.

If a core biopsy, the area is numbed with a local anaesthetic, a wide hollow needle is inserted and a sample of the cells taken.

**Excision biopsy**

An excision biopsy aims to remove the lesion or lump entirely. It will be examined to determine whether more treatment is needed.

**Fine needle aspiration biopsy**

A fine-needle aspiration (FNA) biopsy is similar to a core biopsy but isn’t as deep and may not need a local anaesthetic. When the needle is in position, it will cut out a small sample of tissue.

**Punch biopsy**

For a punch biopsy, the area is usually numbed with a local anaesthetic, and a small hole is made in the skin and samples of the top layers of tissue are taken.

**Sentinel node biopsy**

Sentinel node biopsy is a test to determine whether there are cancer cells in the lymph nodes around your groin. The consultant will carry out a small operation to remove some tissue to be tested. There is more information here: www.cancerresearchuk.org/about-cancer/penile-cancer/getting-diagnosed/tests-stage/sentinel-lymph-node-biopsy.

**Ileostomy**

An ileostomy is a stoma formed by bringing the end or a loop of the small bowel (ileum) out on to the surface of your abdomen (belly). The food waste passes out of the ileostomy and is collected in an external pouch (generally known as an ileostomy bag). Ileostomies can let the area rest and heal after surgery.

**Ileum**

The small bowel that lies below the caecum (appendix) and the ascending colon. The ileum is usually numbed with a local anaesthetic, and a small hole is made in the skin and samples of the top layers of tissue are taken.

**Lymph nodes**

Lymph nodes are often the first place the cancer spreads.

**Colostomy**

A colostomy is a stoma formed by bringing part of your colon (large bowel) out on to the surface of your abdomen (belly). The waste from a colostomy is usually more formed than from an ileostomy, as it has had some of the water removed on its way around the bowel. The waste passes out of the colostomy and is collected in an external pouch (generally known as a colostomy bag).

**Wide local excision (WLE)**

In a wide local excision (WLE) the surgeon cuts out the tumour (cancerous tissue) along with a small margin of around 3 millimetres of healthy tissue to increase the likelihood that all of the tumour has been removed.

**Palliative care support services**

More information on what forms of home care are available can be found at: www.nhs.uk/conditions/social-care-and-support/home-care and www.cancerresearchuk.org/about-cancer/melanoma/advanced-melanoma/support-home-for-you-your-family.

**Immunotherapy**

Also called biologic therapy or bio-chemotherapy, stimulates the patient’s own immune system to fight cancer.

**Lymphoma**

An area of tissue that isn’t normal. This term can be used when it is uncertain whether the area is cancer (malignant) or not (benign).

**Sentinel**

An area of tissue that isn’t normal. This term can be used when it is uncertain whether the area is cancer (malignant) or not (benign).

**Palliative**

Palliative care support services

For more information see: www.nhs.uk/conditions/oncology/Palliative-care-support-services.

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For more information see: www.nhs.uk/conditions/oncology/Palliative-care-support-services.

**Immunotherapy**

Also called biologic therapy or bio-chemotherapy, stimulates the patient’s own immune system to fight cancer.

**Chemotherapy**

A type of treatment that uses drugs to kill cancer cells.

**Sentinel lymph node biopsy (SNB)**

A procedure using a tiny device that transmits radioactivity, The tube is fitted with a light and camera so the clinician can look at the inside of the bowel very clearly on a television screen. Because it only looks at the lower bowel, it is much quicker than a colonoscopy and an anaesthetic isn’t usually needed.


**Magnetic resonance imaging (MRI)**

A procedure using a magnetic field and radio waves. These show up soft tissues and organs. The new, metastatic tumour is the same type of cancer as the primary tumour. For example, if mucosal melanomas spread to the liver, the cancer cells in the liver are mucosal melanoma cancer cells, not liver cancer cells.

**PET-CT**

A procedure using a PET scan (see above) with a Positron Emission Tomography (PET) scan. A mildly radioactive substance is injected, which shows up parts of the body where cells are more active. That is, where a cancer might be growing. For more information see the Cancer Research UK website link above and www.nhs.uk/conditions/pet-scan.

**Sigmoidoscopy**

A procedure using a sigmoidoscope, a narrow tube about as wide as a finger, which is inserted into the back passage. The tube is fitted with a light and camera so the clinician can look at the inside of the bowel very clearly on a television screen. Because it only looks at the lower bowel, it is much quicker than a colonoscopy and an anaesthetic isn’t usually needed.

**Colostomy**

A stoma is an opening made on your abdomen (belly) which allows food waste to pass out of the body into a bag. This may be temporary (ileostomy) or permanent (colostomy).

**Palliative care support services**

For more information see: www.nhs.uk/conditions/oncology/Palliative-care-support-services.

**STAGING**

For more information on the tests mentioned in this leaflet see the Cancer Research UK website link: www.cancerresearchuk.org/about-cancer/bowel-cancer/getting-diagnosed/tests-stage.

**CT scan**

A computerised tomography (CT) scan is taken up to 1½ hours. For more information see: www.cancerresearchuk.org/about-cancer/melanoma/stages-types and https://cancersupport.org.uk/CS/Refer/al/docs/CN/needletoscan.pdf.

**Stoma**

A stoma is an opening made on your abdomen (belly) which allows food waste to pass out of the body into a bag. This may be temporary (ileostomy) or permanent (colostomy).

**Abbreviations**

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<th>APR</th>
<th>Abdominoperineal resection</th>
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<tr>
<td>AUG</td>
<td>Ano-uro-genital</td>
</tr>
<tr>
<td>CNS</td>
<td>Clinical Nurse Specialist</td>
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<tr>
<td>CT</td>
<td>Computed tomography</td>
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<tr>
<td>MRI</td>
<td>Magnetic resonance imaging</td>
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<tr>
<td>PET</td>
<td>Position emission tomography</td>
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