

Radiotherapy consent form for sarcomas - bone and soft tissue tumours (extremity)

This form should only be used if the patient is over 16 years old and has capacity to give consent. If the patient does not legally have capacity please use an appropriate alternative consent form from your hospital.

Patient details				
Patient name: Patient unique identifier:		Date of birth: Name of hospital:		
responsible consultant c	oncologist of consultant therape	auto rautographier.		
Special requirements: eg, to	ransport, interpreter, assistance			
Details of radiothe	гару			
Radiotherapy type:	External beam radiotherapy	y		
Site and side:	☐ Specify site:	☐ Left ☐ Right		
Aim of treatment: (Tick as appropriate)	 Neo-adjuvant – treatment given before surgery Adjuvant – treatment given after surgery to reduce the risk of cancer coming back Definitive – without surgery Palliative – to improve your symptoms and/or help you live longer but not to cure your cancer 			
Concurrent systemic anti-cancer therapy (SACT), including chemotherapy: (Tick as appropriate)		be increased when receiving concurrent systemic anti-cancer therapy. r the side effects of this treatment.		
Contact details are provided	before starting, during or after y d here for any further queries, e to discuss your treatment further.	our radiotherapy.		

Possible early or short-term side-effects						
Start during radiotherapy, can peak shortly after completing radiotherapy and usually resolve within two to six months of finishing radiotherapy. Frequencies are approximate.						
	Expected 50%-100%	Common 10%–50%	Less common Less than 10%	Rare Less than 1%	Not applicable to you	
General radiotherapy risks						
Tiredness						
Skin soreness, itch or colour change in the treatment area – white/lighter skin: pink, red, darker than surrounding area; brown skin: maroon or darker than surrounding area; black skin: darker than surrounding area, yellow/purple/grey colour changes.						
Skin breakdown in the treatment area – oozing, weeping, scabbing and/or bleeding						
Hair thinning or loss in the treatment area						
Limb oedema – swelling in all or part of the limb due to fluid build-up						
Pain – may require pain killers / analgesics						
Post-surgical wound complications – which may require dressing, packing, drainage, antibiotics						
Post-surgical wound complications -requiring surgery and debridement						
Joint stiffness – reduced range of movement experienced over joints and/or muscles						
Joint immobility – which may affect how well the joint functions						
Tumour growth during pre-operative radiotherapy – which may make surgery more extensive or not possible. This is not a consequence of the radiotherapy.						
Specific risks which relate to the site of treatment						
Inflammation of the lungs – which may cause shortness of breath and/or cough						
Change in bowel habit – increased frequency (opening your bowels more often than usual), urgency (a sudden urge to open your bowels), incontinence, and/ or looser stool with more mucous or wind						
Urinary symptoms – increased frequency (passing more urine than usual), urgency (a sudden urge to pass urine), incontinence, discomfort on passing urine, and/or an inability to pass urine (you may require a urinary catheter)						

Patient unique identifier:

Continued on the next page

Patient name:

Patient name:		Patient unique identifier:			
Possible early or short-term side-	-effects co	ontinued			
	Expected 50%–100%	Common 10%–50%	Less common Less than 10%	Rare Less than 1%	Not applicable to you
Rectal and/or anal pain and/or a feeling of not completely emptying your bowels					
Vaginal soreness, itching, or discharge					
Other specific risks to you from your treatment					

I confirm that I have had the above side-effects explained.

Patient initials

May happen many months or years after radiotherapy and may be permanent. Frequencies are approximate.					
	Expected 50%-100%	Common 10%–50%	Less common Less than 10%	Rare Less than 1%	Not applicable to you
General radiotherapy risks					
Tiredness – chronic fatigue					
Permanent skin colour change – usually lighter or darker for any skin tone and/or texture change (thicker or thinner skin)					
Increased sensitivity of skin to sun					
Permanent hair loss in and around treatment area – if hair starts to regrow, it may be patchy					
Small visible blood vessels which look like spidery marks in the treatment area					
Long-term pain – may require pain killers					
Chronic muscle wasting – decrease in size and wasting of muscle tissue					
Lymphoedema – swelling in all or part of the limb due to fluid (lymph) build-up					
Soft tissue fibrosis – scarring or hardening of the skin, muscle, fat, fibrous tissue					
Joint stiffness – reduced range of movement experienced over joints and/or muscles					
Joint immobility – which may affect how well the joint functions					
Delayed wound healing					
Insufficiency fracture – a fracture that develops in the bone within the treatment area					
Bone infarction/osteonecrosis – death of bone caused by poor blood supply within the treatment area					
Nerve damage – which may cause pain, numbness, or weakness in the limb					
Radiation-induced necrosis/ulceration - radiation exposure kills or leads to the death of the soft tissue over the treatment area / tissue is unable to heal leaving an open sore					
Secondary amputation – surgery to remove the limb					
Increased risk of a different cancer in the treatment area					

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Patient unique identifier:

Patient name:

Possible late or long-term side-effects

Patient name:	Patient unique identifier:
Patient name:	Patient unique identifier:

Possible late o	r long-term	cida_affacts	continued
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	Expected 50%–100%	Common 10%–50%	Less common Less than 10%	Rare Less than 1%	Not applicable to you
Specific risks which relate to the site of treatment					
Lung fibrosis (scarring) – which may cause breathlessness and/or chronic cough					
Change in bowel habit – increased frequency (opening your bowels more often than usual), urgency (a sudden urge to open your bowels), incontinence, and/ or looser stool with more mucous or wind					
Rectal or anal pain – this may cause pain when opening your bowels and may affect your sex life if you receive anal sex					
Bleeding from your bladder or bowel					
Urinary symptoms – increased frequency (passing more urine than usual), urgency (a sudden urge to pass urine), incontinence, discomfort on passing urine, and/or an inability to pass urine (you may require a urinary catheter)					
Vaginal dryness, shortening and narrowing – this may impact vaginal intercourse. You may be advised to use vaginal dilators after treatment to reduce risks					
Infertility – unable to produce viable egg/ sperm, or for uterus to be unable to carry a fetus					
Early menopause – symptoms of this may start during or shortly after radiotherapy. Egg and hormone production will stop					
Low testosterone levels					
Erectile dysfunction					
Other specific risks to you from your treatment					

from your treatment

I confirm that I have had the above side-effects explained.

Patient initials

Patient name:	Patient unique identifier:		
Statement of health professional	to be filled in by health professional with appropriate knowledge of proposed procedure)		
 I have discussed what the treatment is likely to involve, the ir I have also discussed the benefits and risks of any available a I have discussed any particular concerns of this patient. 			
Patient information leaflet provided: Yes / No – Details:			
Copy of consent form accepted by patient: Yes / No	_		
Signature:	Date:		
Name:	Job title:		
Statement of patient		Statement of:	
 I have had the aims and possible side effects of treatment opportunity to discuss alternative treatment and I agree to 	interpreter witness (where appropriate)		
 described on this form. I understand that a guarantee cannot be given that a partic radiotherapy. The person will, however, have appropriate e I have been told about additional procedures which are not to treatment or may become necessary during my treatme include permanent skin marks and photographs to help wit planning and identification. I agree that information collected during my treatment, increcords may be used for education, audit and research. All I am aware I can withdraw consent at anytime. 	 I have interpreted the information contained in this form to the patient to the best of my ability and in a way in which I believe they can understand. or I confirm that the patient is unable to sign but has indicated their consent. 		
Tick if relevant		Signature:	
☐ I confirm that there is no risk that I could be pregnant. ☐ I understand that I should not become pregnant during treating.	atment.		
Note: if there is any possibility of you being pregnant you must tell your hospital doctor/h can cause significant harm to an unborn fetus. Testosterone and other hormone treatmer			
	Name:		
I understand that if I were to continue to smoke it could have side-effects I experience and the efficacy of my treatment.	ve a significant impact on the	Data	
I do not have a pacemaker and/or implantable cardioverter	Date:		
☐ I have a pacemaker and/or implantable cardioverter defibring risks associated with this explained to me.	Patient confirmation of consent		
Signature:	gnature:		
Patient name:	Date:	I confirm that I have no further questions and wish to go ahead with treatment.	
		Patient initials	
		Date:	