HORIZONS



Understanding the impact of cancer diagnosis and treatment on everyday life

## BASELINE CERVICAL CANCER CRF

## FOR STAFF USE ONLY

## **CRF Completion Instructions**

- This CRF is for completion by members of site staff NOT study participants
- Please complete the CRF when a patient has been recruited to the study
- Please complete as much of the CRF as possible
- If you have any queries, please contact the HORIZONS Coordinating Centre, email address <u>HORIZONS@soton.ac.uk</u>
- Please tick boxes when appropriate
- When you have completed the CRF, please keep a copy for your own records and return a copy to us, by post, fax or email along with the completed return cover sheet

Participant's Study ID				
Participant's date of birth	d d m m	уууу	У	
Participant's weight k	rg Pa	ırticipant's he	eight	cms
Participant's blood pressure which they were measured)	⊇ (Please give the most re	ecently reported	d figures and	d the date on
Systolic	mmHg	D	ate meası	ured
Diastolic	mmHg	d d	m m	у у у у
Participant's tumour type (p	please tick one box)			
Туре	Sub-type			
Cervical	Squamous cell carcir	noma		
	Adenosquamous car	cinoma		
	Adenocarcinoma			
	Clear cell carcinoma			
	Other (please descri	be on line be	low)	
	Not currently known	1		
Date of participant's curren	_			
(date that histological diagnosis	was reported)	d d m	n m y	у у у

Participant	's Study ID		/			
•	's FIGO stage (pl		box OR tick	the box ir	ndicating the F	IGO
Stage 1	Stage 1A2					
	Stage 1B1					
	Stage 1B2					
Stage 2	Stage 2A1					
	Stage 2A2					
	Stage 2B					
Stage 3	Stage 3A					
	Stage 3B					
	not currently kr		ne box)			
Grade 1/lo	w grade/well di	fferentiated				
Grade 2/m	oderate/interm	ediate grade				
Grade 3/hi	gh-grade/poorl	y differentiate	d			
Grade not	currently know	า				
Participant	's pre-treatmen	t ECOG status	(please tick	one box)		
ECOG 0 (the	e patient has no sy	mptoms)				
ECOG 1 (the	e patient has symp	otoms but is am	oulatory)			
ECOG 2 (the	e patient is bedrid	den less than ha	lf the day)			
ECOG 3 (the	e patient is bedrid	den half the day	or longer)			

ECOG 4 (the patient is chronically bedridden and requires assis-

tance with the activities of daily living)

Participant's Study ID	
Is the participant pre or	post menopause? (please tick one box)
Pre menopause	
Post menopause	
Unknown	
Has the participant had a pre-	vious diagnosis of cancer (please tick one box)
Yes No	Unknown
	above question, please provide some information
about the patient's previous	cancer(s) by completing the box(es) below
PREVIOUS DIAGNOSIS 1	
Type of cancer	
Date of diagnosis	
Treatment received	
Date treatment ended	
PREVIOUS DIAGNOSIS 2	
Type of cancer	
Date of diagnosis	
Treatment received	
Date treatment ended	

Participant's Study ID			
Has the participant had (please tick one box)	any genetic tests fo	r inherited cancers?	
Yes Yes	No	Unknown	

If you answered "Yes" to the above question, please provide some information about the participant's other genetic test(s) by completing the table(s) below

Name of genetic test for cancer (1)	Result of genetic test
	Positive
	Negative
	Ambiguous/uncertain
	Awaiting result
	Unknown

Name of genetic test for cancer (2)	Result of genetic test
	Positive
	Negative
	Ambiguous/uncertain
	Awaiting result
	Unknown

Has a first degree relative of the participant (parent, sibling or child) been diagnosed with cancer? (please tick one box)						
Yes		No	Unkno	wn		
-	If you answered "yes" to the above question, what type of cancer and when was it diagnosed? (Please complete the table below)					
	Type of cancer	A	ge at diagnosis	Date of diag	nosis	
Relative 1						
Relative 2						
Relative 3						
Does the participant have any of the following co-morbidities (please tick all that apply)  Myocardial infarct						
Angina/coronary artery disease  Congestive Heart Failure						
Cardiac Arrythmia	S					
Hypertension						
Venous Disease (PE/DVT)						
Peripheral Arterial Disease						
Restrictive Lung Disease or COPD (chronic bronchitis, emphysema, asthma etc.)						
Liver Disease (portal hypertension, chronic/acute hepatitis, cirrhosis etc.)						
Stomach Ulcers or Inflammatory Bowel Disease						
Acute or Chronic Pancreatitis						
End-stage Renal Disease (chronic renal insufficiency, dialysis etc.)						
Thyroid problems	Thyroid problems (hyperthyroidism, hypothyroidism etc.)					

Participant's Study ID / / /

Participant's co-morbiditi	es continued (please tick all that apply)	
Diabetes Mellitus Type 1		
Diabetes Mellitus Type 2		
Stroke/TIA		
Dementia		
Paralysis (paraplegia or hemip	olegia)	
Neuromuscular Condition chronic neuromuscular disorde	(multiple sclerosis, Parkinson's, myasthenia gravis, other r)	
Clinical diagnosis of anxiet	У	
Clinical diagnosis of depres	ssion	
Other psychiatric Diagnosis	S (schizophrenia, bipolar disorder etc.)	
Osteoarthritis		
Rheumatoid Arthritis		
Other Rheumatological Dismyositis, rheumatic polymyosit	ease (systemic lupus, mixed connective tissue disorder, polyis, scleroderma etc.)	
HIV/AIDS		
Alcohol Abuse (or history of, plications)	must be accompanied by social, behavioural or medical com-	
Drug/Substance Abuse (or medical complications)	history of, must be accompanied by social, behavioural or	
Morbid Obesity		
Other (please give details)		
What is the participant's for cervical cancer)	proposed treatment start date (main first-line treatment date)	ent
Please add your name and	d signature and the date that you completed this C	RF
Name	Signature	
Date	d d / m m / y y y	